

Singular

A Computer Algebra System for Polynomial Computations

Manual

Version 4-2-0, 2020

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Short Contents

1	Preface	1
2	Introduction,General concepts,Preface,Top	4
3	General concepts	15
4	Data types	72
5	Functions and system variables	154
6	Tricks and pitfalls	304
7	Non-commutative subsystem	312
	Appendix A Examples	691
	Appendix B Polynomial data	759
	Appendix C Mathematical background	766
	Appendix D SINGULAR libraries	785
8	Release Notes	944
9	Index	956

1 Preface

SINGULAR version 4-2-0
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For information on how to cite SINGULAR see

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You can also support SINGULAR by informing us about your result obtained by using SINGULAR.

Availability

The latest information regarding the status of SINGULAR is always available from <https://www.singular.uni-kl.de>. The program SINGULAR and the above mentioned parts are available via anonymous ftp through the following addresses:

GMP, libreadline

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<https://gmplib.org>

NTL

© Victor Shoup
<http://www.shoup.net/ntl>

cdd (C implementation of the Double Description Method of Motzkin et al)

© Komei Fukuda

http://www-oldurls.inf.ethz.ch/personal/fukudak/cdd_home/

FLINT © Bill Hart, Sebastian Pancratz, Fredrik Johansson

<http://www.flintlib.org>

gfanlib © Anders Jensen

<https://users-math.au.dk/~jensen/software/gfan/gfan.html>

Singular-Factory

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<https://www.mathematik.uni-kl.de/ftp/pub/Math/Singular/Factory>

Singular-libfac

© Messollen, University of Saarbrücken:

<ftp://jim.mathematik.uni-kl.de/pub/Math/Singular/Libfac/>

SINGULAR binaries and sources

<ftp://jim.mathematik.uni-kl.de/pub/Math/Singular/> or via a WWW browser
from <http://www.mathematik.uni-kl.de/ftp/pub/Math/Singular/>

Cygwin <https://www.cygwin.com/>

Xemacs <https://www.xemacs.org>

Some external programs are optional:

4ti2 (used by `sing4ti2.lib`, see Section D.4.33 [`sing4ti2.lib`], page 833)

<https://4ti2.github.io>

gfan (used by `tropical.lib`, see Section D.13.6 [`tropical.lib`], page 909)

<https://users-math.au.dk/~jensen/software/gfan/gfan.html>

graphviz (used by `resgraph.lib`, see Section D.5.11 [`resgraph.lib`], page 846)

<https://www.graphviz.org/>

normaliz (used by `normaliz.lib`, see Section D.4.24 [`normaliz.lib`], page 825)

© Winfried Bruns and Bogdan Ichim

<https://www.normaliz.uni-osnabrueck.de>

polymake (used by `polymake.lib`, see Section D.13.4 [`polymake.lib`], page 906)

© Ewgenij Gawrilow and Michael Joswig

<https://polymake.org/>

surf (used by `surf.lib`, see Section D.9.3 [`surf.lib`], page 885)

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<http://surf.sf.net>

surfer (used by `surf.lib`, see Section D.9.3 [`surf.lib`], page 885)

<https://imaginary.org/program/surfer>

surfex (used by `surfex.lib`, see Section D.9.4 [`surfex.lib`], page 885)

© Oliver Labs (2001-2008), Stephan Holzer (2004-2005)

<https://github.com/Singular/Singular/tree/spielwiese/Singular/LIB/surfex>

TOPCOM (used by `polymake.lib`, see Section D.13.4 [`polymake.lib`], page 906)

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<http://www.rambau.wm.uni-bayreuth.de/TOPCOM/>

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2 Introduction,General concepts,Preface,Top

2.1 Background

SINGULAR is a Computer Algebra system for polynomial computations with emphasis on the special needs of commutative algebra, algebraic geometry, and singularity theory.

SINGULAR's main computational objects are ideals and modules over a large variety of baserings. The baserings are polynomial rings or localizations thereof over a field (e.g., finite fields, the rationals, floats, algebraic extensions, transcendental extensions) or over a limited set of rings, or over quotient rings with respect to an ideal.

SINGULAR features one of the fastest and most general implementations of various algorithms for computing Groebner resp. standard bases. The implementation includes Buchberger's algorithm (if the ordering is a wellordering) and Mora's algorithm (if the ordering is a tangent cone ordering) as special cases. Furthermore, it provides polynomial factorization, resultant, characteristic set and gcd computations, syzygy and free-resolution computations, and many more related functionalities.

Based on an easy-to-use interactive shell and a C-like programming language, SINGULAR's internal functionality is augmented and user-extendible by libraries written in the SINGULAR programming language. A general and efficient implementation of communication links allows SINGULAR to make its functionality available to other programs.

SINGULAR's development started in 1984 with an implementation of Mora's Tangent Cone algorithm in Modula-2 on an Atari computer (K.P. Neuendorf, G. Pfister, H. Schönemann; Humboldt-Universität zu Berlin). The need for a new system arose from the investigation of mathematical problems coming from singularity theory which none of the existing systems was able to handle.

In the early 1990s SINGULAR's "home-town" moved to Kaiserslautern, a general standard basis algorithm was implemented in C and SINGULAR was ported to Unix, MS-DOS, Windows NT, and MacOS.

Continuous extensions (like polynomial factorization, gcd computations, links) and refinements led in 1997 to the release of SINGULAR version 1.0 and in 1998 to the release of version 1.2 (with a much faster standard and Groebner bases computation based on Hilbert series and on an improved implementation of the core algorithms, libraries for primary decomposition, ring normalization, etc.)

For the highlights of the new SINGULAR version 4-2-0, see Section 8.1 [News and changes], page 944.

2.2 How to use this manual

For the impatient user

In Section 2.3 [Getting started], page 6, some simple examples explain how to use SINGULAR in a step-by-step manner.

Appendix A [Examples], page 691 should come next for real learning-by-doing or to quickly solve some given mathematical problem without dwelling too deeply into SINGULAR. This chapter contains a lot of real-life examples and detailed instructions and explanations on how to solve mathematical problems using SINGULAR.

For the systematic user

In Chapter 3 [General concepts], page 15, all basic concepts which are important to use and to understand SINGULAR are developed. But even for users preferring the systematic approach it will be helpful to take a look at the examples in Section 2.3 [Getting started], page 6, every now and then. The topics in the chapter are organized more or less in the natural order in which the novice user is expected to have to deal with them.

- In Section 3.1 [Interactive use], page 15, and its subsections there are some words on entering and exiting SINGULAR, followed by a number of other aspects concerning the interactive user-interface.
- To do anything more than trivial integer computations, one needs to define a basering in SINGULAR. This is explained in detail in Section 3.3 [Rings and orderings], page 30.
- An overview of the algorithms implemented in the kernel of SINGULAR is given in Section 3.4 [Implemented algorithms], page 36.
- In Section 3.5 [The SINGULAR language], page 40, language specific concepts are introduced, such as the notions of names and objects, data types and conversion between them, etc.
- In Section 3.6 [Input and output], page 48, SINGULAR's mechanisms to store and retrieve data are discussed.
- The more complex concepts of procedures and libraries as well as tools for debugging them are considered in the following sections: Section 3.7 [Procedures], page 50, Section 3.8 [Libraries], page 54, and Section 3.9 [Debugging tools], page 67.

Chapter 4 [Data types], page 72, is a complete treatment of SINGULAR's data types in alphabetical order, where each section corresponds to one data type. For each data type, its purpose is explained, the syntax of its declaration is given, related operations and functions are listed, and one or more examples illustrate its usage.

Chapter 5 [Functions and system variables], page 154, is an alphabetically ordered reference list of all of SINGULAR's functions, control structures, and system variables. Each entry includes a description of the syntax and semantics of the item being explained as well as one or more examples on how to use it.

Miscellaneous

Chapter 6 [Tricks and pitfalls], page 304, is a loose collection of limitations and features which may be unexpected by those who expect the SINGULAR language to be an exact copy of the C programming language or of some other Computer Algebra system's language. Additionally, some mathematical hints are collected there.

Appendix C [Mathematical background], page 766, introduces some of the mathematical notions and definitions used throughout this manual. For example, if in doubt what exactly a "negative degree reverse lexicographical ordering" is in SINGULAR, one should refer to this chapter.

Appendix D [SINGULAR libraries], page 785, lists the libraries which come with SINGULAR, and all functions contained in them.

Typographical conventions

Throughout this manual, the following typographical conventions are adopted:

- text in **typewriter** denotes SINGULAR input and output as well as reserved names:
The basering can, e.g., be set using the command **setring**.
- the arrow \mapsto denotes SINGULAR output:

```
poly p=x+y+z;
p*p;
↳ x2+2xy+y2+2xz+2yz+z2
```

- square brackets are used to denote parts of syntax descriptions which are optional:

```
[optional_text] required_text
```

- keys are denoted using typewriter, for example:

```
N (press the key N to get to the next node in help mode)
```

```
RETURN (press RETURN to finish an input line)
```

```
CTRL-P (press the control key together with the key P to get the previous input line)
```

2.3 Getting started

SINGULAR is a special purpose system for polynomial computations. Hence, most of the powerful computations in SINGULAR require the prior definition of a ring. Most important rings are polynomial rings over a field, localizations thereof, or quotient rings of such rings modulo an ideal. However, some simple computations with integers (machine integers of limited size) and manipulations of strings can be carried out without the prior definition of a ring.

2.3.1 First steps

Once SINGULAR is started, it awaits an input after the prompt `>`. Every statement has to be terminated by `;`.

```
37+5;
↳ 42
```

All objects have a type, e.g., integer variables are defined by the word `int`. An assignment is made using the symbol `=`.

```
int k = 2;
```

Test for equality resp. inequality is done using `==` resp. `!=` (or `<>`), where 0 represents the boolean value FALSE, and any other value represents TRUE.

```
k == 2;
↳ 1
k != 2;
↳ 0
```

The value of an object is displayed by simply typing its name.

```
k;
↳ 2
```

On the other hand, the output is suppressed if an assignment is made.

```
int j;
j = k+1;
```

The last displayed (!) result can be retrieved via the special symbol `_`.

```
2*_; // the value from k displayed above
↳ 4
```

Text starting with `//` denotes a comment and is ignored in calculations, as seen in the previous example. Furthermore SINGULAR maintains a history of the previous lines of input, which may be accessed by CTRL-P (previous) and CTRL-N (next) or the arrows on the keyboard.

The whole manual is available online by typing the command `help;`. Documentation on single topics, e.g., on `intmat`, which defines a matrix of integers, is obtained by


```
help intmat;
```

This will display the text of Section 4.7 [intmat], page 88, in the printed manual.

Next, we define a 3×3 matrix of integers and initialize it with some values, row by row from left to right:

```
intmat m[3][3] = 1,2,3,4,5,6,7,8,9;
m;
```

A single matrix entry may be selected and changed using square brackets [and].

```
m[1,2]=0;
m;
↳ 1,0,3,
↳ 4,5,6,
↳ 7,8,9
```

To calculate the trace of this matrix, we use a `for` loop. The curly brackets { and } denote the beginning resp. end of a block. If you define a variable without giving an initial value, as the variable `tr` in the example below, SINGULAR assigns a default value for the specific type. In this case, the default value for integers is 0. Note that the integer variable `j` has already been defined above.

```
int tr;
for ( j=1; j <= 3; j++ ) { tr=tr + m[j,j]; }
tr;
↳ 15
```

Variables of type string can also be defined and used without having an active ring. Strings are delimited by " (double quotes). They may be used to comment the output of a computation or to give it a nice format. If a string contains valid SINGULAR commands, it can be executed using the function `execute`. The result is the same as if the commands would have been written on the command line. This feature is especially useful to define new rings inside procedures.

```
"example for strings:";
↳ example for strings:
string s="The element of m ";
s = s + "at position [2,3] is:"; // concatenation of strings by +
s , m[2,3] , ".";
↳ The element of m at position [2,3] is: 6 .
s="m[2,1]=0; m;";
execute(s);
↳ 1,0,3,
↳ 0,5,6,
↳ 7,8,9
```

This example shows that expressions can be separated by , (comma) giving a list of expressions. SINGULAR evaluates each expression in this list and prints all results separated by spaces.

2.3.2 Rings and standard bases

In order to compute with objects such as ideals, matrices, modules, and polynomial vectors, a ring has to be defined first.

```
ring r = 0, (x,y,z), dp;
```

The definition of a ring consists of three parts: the first part determines the ground field, the second part determines the names of the ring variables, and the third part determines the monomial ordering to be used. Thus, the above example declares a polynomial ring called `r` with a ground

field of characteristic 0 (i.e., the rational numbers) and ring variables called x , y , and z . The `dp` at the end determines that the degree reverse lexicographical ordering will be used.

Other ring declarations:

```
ring r1=32003, (x,y,z), dp;
    characteristic 32003, variables x, y, and z and ordering dp.

ring r2=32003, (a,b,c,d), lp;
    characteristic 32003, variable names a, b, c, d and lexicographical ordering.

ring r3=7, (x(1..10)), ds;
    characteristic 7, variable names x(1), . . . , x(10), negative degree reverse lexicographical
    ordering (ds).

ring r4=(0,a), (mu,nu), lp;
    transcendental extension of  $Q$  by  $a$ , variable names mu and nu, lexicographical ordering.

ring r5=real, (a,b), lp;
    floating point numbers (single machine precision), variable names a and b.

ring r6=(real,50), (a,b), lp;
    floating point numbers with precision extended to 50 digits, variable names a and b.

ring r7=(complex,50,i), (a,b), lp;
    complex floating point numbers with precision extended to 50 digits and imaginary
    unit i, variable names a and b.

ring r8=integer, (a,b), lp;
    the ring of integers (see Section 3.3.4 [Coefficient rings], page 36), variable names a and
    b.

ring r9=(integer, 60), (a,b), lp;
    the ring of integers modulo 60 (see Section 3.3.4 [Coefficient rings], page 36), variable
    names a and b.

ring r10=(integer, 2, 10), (a,b), lp;
    the ring of integers modulo  $2^10$  (see Section 3.3.4 [Coefficient rings], page 36), variable
    names a and b.
```

Typing the name of a ring prints its definition. The example below shows that the default ring in SINGULAR is $Z/32003[x, y, z]$ with degree reverse lexicographical ordering:

```
ring r11;
r11;
⇒ // coefficients: ZZ/32003
⇒ // number of vars : 3
⇒ //      block 1 : ordering dp
⇒ //      : names x y z
⇒ //      block 2 : ordering C
```

Defining a ring makes this ring the current active basering, so each ring definition above switches to a new basering. The concept of rings in SINGULAR is discussed in detail in Section 3.3 [Rings and orderings], page 30.

The basering is now `r11`. Since we want to calculate in the ring `r`, which we defined first, we need to switch back to it. This can be done using the function `setring`:

```
setring r;
```

Once a ring is active, we can define polynomials. A monomial, say x^3 , may be entered in two ways: either using the power operator `^`, writing `x^3`, or in short-hand notation without operator,

writing `x3`. Note that the short-hand notation is forbidden if a name of the ring variable(s) consists of more than one character (see Section 6.4 [Miscellaneous oddities], page 308 for details). Note, that SINGULAR always expands brackets and automatically sorts the terms with respect to the monomial ordering of the basering.

```
poly f = x3+y3+(x-y)*x2y2+z2;
f;
↳ x3y2-x2y3+x3+y3+z2
```

The command `size` retrieves in general the number of entries in an object. In particular, for polynomials, `size` returns the number of monomials.

```
size(f);
↳ 5
```

A natural question is to ask if a point, e.g., $(x,y,z)=(1,2,0)$, lies on the variety defined by the polynomials `f` and `g`. For this we define an ideal generated by both polynomials, substitute the coordinates of the point for the ring variables, and check if the result is zero:

```
poly g = f^2 *(2x-y);
ideal I = f,g;
ideal J = subst(I,var(1),1);
J = subst(J,var(2),2);
J = subst(J,var(3),0);
J;
↳ J[1]=5
↳ J[2]=0
```

Since the result is not zero, the point $(1,2,0)$ does not lie on the variety $V(f,g)$.

Another question is to decide whether some function vanishes on a variety, or in algebraic terms, if a polynomial is contained in a given ideal. For this we calculate a standard basis using the command `groebner` and afterwards reduce the polynomial with respect to this standard basis.

```
ideal sI = groebner(f);
reduce(g,sI);
↳ 0
```

As the result is 0 the polynomial `g` belongs to the ideal defined by `f`.

The function `groebner`, like many other functions in SINGULAR, prints a protocol during calculations, if desired. The command `option(prot);` enables protocolling whereas `option(noprot);` turns it off. Section 5.1.110 [option], page 230, explains the meaning of the different symbols printed during calculations.

The command `kbase` calculates a basis of the polynomial ring modulo an ideal, if the quotient ring is finite dimensional. As an example we calculate the Milnor number of a hypersurface singularity in the global and local case. This is the vector space dimension of the polynomial ring modulo the Jacobian ideal in the global case resp. of the power series ring modulo the Jacobian ideal in the local case. See Section A.4.2 [Critical points], page 728, for a detailed explanation.

The Jacobian ideal is obtained with the command `jacob`.

```
ideal J = jacob(f);
↳ // ** redefining J **
J;
↳ J[1]=3x2y2-2xy3+3x2
↳ J[2]=2x3y-3x2y2+3y2
↳ J[3]=2z
```

SINGULAR prints the line `// ** redefining J **`. This indicates that we had previously defined a variable with name `J` of type ideal (see above).

To obtain a representing set of the quotient vector space we first calculate a standard basis, and then apply the function `kbase` to this standard basis.

```
J = groebner(J);
ideal K = kbase(J);
K;
↳ K[1]=y4
↳ K[2]=xy3
↳ K[3]=y3
↳ K[4]=xy2
↳ K[5]=y2
↳ K[6]=x2y
↳ K[7]=xy
↳ K[8]=y
↳ K[9]=x3
↳ K[10]=x2
↳ K[11]=x
↳ K[12]=1
```

Then

```
size(K);
↳ 12
```

gives the desired vector space dimension $K[x, y, z]/\text{jacob}(f)$. As in SINGULAR the functions may take the input directly from earlier calculations, the whole sequence of commands may be written in one single statement.

```
size(kbase(groebner(jacob(f))));
↳ 12
```

When we are not interested in a basis of the quotient vector space, but only in the resulting dimension we may even use the command `vdim` and write:

```
vdim(groebner(jacob(f)));
↳ 12
```

2.3.3 Procedures and libraries

SINGULAR offers a comfortable programming language, with a syntax close to C. So it is possible to define procedures which bind a sequence of several commands in a new one. Procedures are defined using the keyword `proc` followed by a name and an optional parameter list with specified types. Finally, a procedure may return a value using the command `return`.

We may e.g. define the following procedure called `Milnor`: (Here the parameter list is `(poly h)` meaning that `Milnor` must be called with one argument which can be assigned to the type `poly` and is referred to by the name `h`.)

```
proc Milnor (poly h)
{
  return(vdim(groebner(jacob(h))));
}
```

Note: if you have entered the first line of the procedure and pressed `RETURN`, SINGULAR prints the prompt `.` (dot) instead of the usual prompt `>`. This shows that the input is incomplete and SINGULAR expects more lines. After typing the closing curly bracket, SINGULAR prints the usual prompt indicating that the input is now complete.

Then we can call the procedure:

```
Milnor(f);
↳ 12
```

Note that the result may depend on the basering as we will see in the next chapter.

The distribution of SINGULAR contains several libraries, each of which is a collection of useful procedures based on the kernel commands, which extend the functionality of SINGULAR. The command `listvar(package)`; list all currently loaded libraries. The command `LIB "all.lib"`; loads all libraries.

One of these libraries is `sing.lib` which already contains a procedure called `milnor` to calculate the Milnor number not only for hypersurfaces but more generally for complete intersection singularities. Libraries are loaded using the command `LIB`. Some additional information during the process of loading is displayed on the screen, which we omit here.

```
LIB "sing.lib";
```

As all input in SINGULAR is case sensitive, there is no conflict with the previously defined procedure `Milnor`, but the result is the same.

```
milnor(f);
↳ 12
```

The procedures in a library have a help part which is displayed by typing

```
help milnor;
```

as well as some examples, which are executed by

```
example milnor;
```

Likewise, the library itself has a help part, to show a list of all the functions available for the user which are contained in the library.

```
help sing.lib;
```

The output of the help commands is omitted here.

2.3.4 Change of rings

To calculate the local Milnor number we have to do the calculation with the same commands in a ring with local ordering. We can define the localization of the polynomial ring at the origin (see Appendix B [Polynomial data], page 759, and Appendix C [Mathematical background], page 766).

```
ring r1 = 0, (x,y,z), ds;
```

The ordering directly affects the standard basis which will be calculated. Fetching the polynomial defined in the ring `r` into this new ring, helps us to avoid retyping previous input.

```
poly f = fetch(r,f);
f;
↳ z2+x3+y3+x3y2-x2y3
```

Instead of `fetch` we can use the function `imap` which is more general but less efficient. The most general way to fetch data from one ring to another is to use maps, this will be explained in Section 4.11 [map], page 103.

In this ring the terms are ordered by increasing exponents. The local Milnor number is now

```
Milnor(f);
↳ 4
```

This shows that `f` has outside the origin in affine 3-space singularities with local Milnor number adding up to $12 - 4 = 8$. Using global and local orderings as above is a convenient way to check whether a variety has singularities outside the origin.

The command `jacob` applied twice gives the Hessian of `f`, in our example a 3x3 - matrix.

```

matrix H = jacob(jacob(f));
H;
↪ H[1,1]=6x+6xy2-2y3
↪ H[1,2]=6x2y-6xy2
↪ H[1,3]=0
↪ H[2,1]=6x2y-6xy2
↪ H[2,2]=6y+2x3-6x2y
↪ H[2,3]=0
↪ H[3,1]=0
↪ H[3,2]=0
↪ H[3,3]=2

```

The `print` command displays the matrix in a nicer format.

```

print(H);
↪ 6x+6xy2-2y3, 6x2y-6xy2, 0,
↪ 6x2y-6xy2, 6y+2x3-6x2y, 0,
↪ 0, 0, 2

```

We may calculate the determinant and (the ideal generated by all) minors of a given size.

```

det(H);
↪ 72xy+24x4-72x3y+72xy3-24y4-48x4y2+64x3y3-48x2y4
minor(H,1); // the 1x1 - minors
↪ _[1]=2
↪ _[2]=6y+2x3-6x2y
↪ _[3]=6x2y-6xy2
↪ _[4]=6x2y-6xy2
↪ _[5]=6x+6xy2-2y3

```

The algorithm of the standard basis computation may be affected by the command `option`. For example, a reduced standard basis of the ideal generated by the 1×1 -minors of H is obtained in the following way:

```

option(redSB);
groebner(minor(H,1));
↪ _[1]=1

```

This shows that 1 is contained in the ideal of the 1×1 -minors, hence the corresponding variety is empty.

2.3.5 Modules and their annihilator

Now we shall give three more advanced examples.

SINGULAR is able to handle modules over all the rings, which can be defined as a basering. A free module of rank n is defined as follows:

```

ring rr;
int n = 4;
freemodule(4);
↪ _[1]=gen(1)
↪ _[2]=gen(2)
↪ _[3]=gen(3)
↪ _[4]=gen(4)
typeof(_);
↪ module
print(freemodule(4));
↪ 1,0,0,0,
↪ 0,1,0,0,

```

```

↳ 0,0,1,0,
↳ 0,0,0,1

```

To define a module, we provide a list of vectors generating a submodule of a free module. Then this set of vectors may be identified with the columns of a matrix. For that reason in SINGULAR matrices and modules may be interchanged. However, the representation is different (modules may be considered as sparse matrices).

```

ring r =0,(x,y,z),dp;
module MD = [x,0,x],[y,z,-y],[0,z,-2y];
matrix MM = MD;
print(MM);
↳ x,y,0,
↳ 0,z,z,
↳ x,-y,-2y

```

However the submodule MD may also be considered as the module of relations of the factor module r^3/MD . In this way, SINGULAR can treat arbitrary finitely generated modules over the basering (see Section B.1 [Representation of mathematical objects], page 759).

In order to get the module of relations of MD , we use the command `syz`.

```

syz(MD);
↳ _[1]=x*gen(3)-x*gen(2)+y*gen(1)

```

We want to calculate, as an application, the annihilator of a given module. Let $M = r^3/U$, where U is our defining module of relations for the module M .

```

module U = [z3,xy2,x3],[yz2,1,xy5z+z3],[y2z,0,x3],[xyz+x2,y2,0],[xyz,x2y,1];

```

Then, by definition, the annihilator of M is the ideal $\text{ann}(M) = \{a \mid aM = 0\}$ which is, by definition of M , the same as $\{a \mid ar^3 \in U\}$. Hence we have to calculate the quotient $U:r^3$. The rank of the free module is determined by the choice of U and is the number of rows of the corresponding matrix. This may be retrieved by the function `nrows`. All we have to do now is the following:

```

quotient(U, freemodule(nrows(U)));

```

The result is too big to be shown here.

2.3.6 Resolution

There are several commands in SINGULAR for computing free resolutions. The most general command is `res(...,n)` which determines heuristically what method to use for the given problem. It computes the free resolution up to the length n , where $n = 0$ corresponds to the full resolution.

Here we use the possibility to inspect the calculation process using the option `prot`.

```

ring R;          // the default ring in char 32003
R;
↳ //   characteristic : 32003
↳ //   number of vars : 3
↳ //           block  1 : ordering dp
↳ //           : names  x y z
↳ //           block  2 : ordering C
ideal I = x4+x3y+x2yz,x2y2+xy2z+y2z2,x2z2+2xz3,2x2z2+xyz2;
option(prot);
resolution rs = res(I,0);
↳ using lres
↳ 4(m0)4(m1).5(m1)g.g6(m1)...6(m2)..

```

Disable this protocol with

```
option(noprot);
```

When we enter the name of the calculated resolution, we get a pictorial description of the minimized resolution where the exponents denote the rank of the free modules. Note that the calculated resolution itself may not yet be minimal.

```
rs;
↳ 1      4      5      2      0
↳R <-- R <-- R <-- R <-- R
↳
↳0      1      2      3      4
print(betti(rs),"betti");
↳
↳      0      1      2      3
↳ -----
↳ 0:    1      -      -      -
↳ 1:    -      -      -      -
↳ 2:    -      -      -      -
↳ 3:    -      4      1      -
↳ 4:    -      -      1      -
↳ 5:    -      -      3      2
↳ -----
↳ total: 1      4      5      2
```

In order to minimize the resolution, that is to calculate the maps of the minimal free resolution, we use the command `minres`:

```
rs=minres(rs);
```

A single module in this resolution is obtained (as usual) with the brackets `[` and `]`. The `print` command can be used to display a module in a more readable format:

```
print(rs[3]);
↳ z3,   -xyz-y2z-4xz2+16z3,
↳ 0,    -y2,
↳ -y+4z,48z,
↳ x+2z, 48z,
↳ 0,    x+y-z
```

In this case, the output is to be interpreted as follows: the 3rd syzygy module of R/I , `rs[3]`, is the rank-2-submodule of R^5 generated by the vectors $(z^3, 0, -y + 4z, x + 2z, 0)$ and $(-xyz - y^2z - 4xz^2 + 16z^3, -y^2, 48z, 48z, x + y - z)$.

3 General concepts

3.1 Interactive use

In this section, aspects of interactive use are discussed. This includes how to enter and exit SINGULAR, how to interpret its prompt, how to get online help, and so on.

There are a few important notes which one should not forget:

- every command has to be terminated by a ; (semicolon) followed by a `(RETURN)`
- the online help is accessible by means of the `help` function

3.1.1 How to enter and exit

SINGULAR can either be run in an ASCII-terminal or within Emacs.

To start SINGULAR in its ASCII-terminal user interface, enter `Singular` at the system prompt. The SINGULAR banner appears which, among other data, reports the version and the compilation date.

To start SINGULAR in its Emacs user interface, either enter `ESingular` at the system prompt, or type `M-x singular` within a running Emacs (provided you have loaded the file `singular.el` in your running Emacs, see Section 3.2.2 [Running SINGULAR under Emacs], page 25 for details).

Generally, we recommend to use SINGULAR in its Emacs interface, since this offers many more features and is more convenient to use than the ASCII-terminal interface (see Section 3.2 [Emacs user interface], page 22).

To exit SINGULAR type `quit;`, `exit;` or `$` (or, when running within Emacs preferably type `C-c $`).

SINGULAR and ESingular may also be started with command line options and with filenames as arguments. More generally, the startup syntax is

```
Singular [options] [file1 [file2 ...]]
ESingular [options] [file1 [file2 ...]]
```

See Section 3.1.6 [Command line options], page 19, Section 3.1.7 [Startup sequence], page 22, Section 3.2.2 [Running SINGULAR under Emacs], page 25.

3.1.2 The SINGULAR prompt

The SINGULAR prompt `>` (larger than) asks the user for input of commands. The “continuation” prompt `.` (period) asks the user for input of missing parts of a command (e.g. the semicolon at the end of every command).

SINGULAR does not interpret the semicolon as the end of a command if it occurs inside a string. Also, SINGULAR waits for blocks (sequences of commands enclosed in curly brackets) to be closed before prompting with `>` for more commands. Thus, if SINGULAR does not respond with its regular prompt after typing a semicolon it may wait for a `"` or a `}` first.

Additional semicolons will not harm SINGULAR since they are interpreted as empty statements.

3.1.3 The online help system

The online help system is invoked by the `help` command. `?` may be used as a synonym for `help`. Simply typing `help;` displays the “top” of the help system (i.e., the title page of the SINGULAR manual) which offers a short table of contents. Typing `help topic;` shows the available documentation on the respective topic. Here, `topic` may be either a function name or, more generally, any

index entry of the SINGULAR manual. Furthermore, topic may contain wildcard characters. See Section 5.1.54 [help], page 191, for more information.

Online help information can be displayed in various help browsers. The following table lists a summary of the browsers which are always present. Usually, external browsers are much more convenient: A complete, customizable list can be found in the file `LIB/help.cnf`.

Browser	Platform	Description
html	Windows	displays a html version of the manual in your default html browser
builtin	all	simply outputs the help information in plain ASCII format
emacs	Unix, Windows	when running SINGULAR within (X)emacs, displays help inside the (X)emacs info buffer.
dummy	all	displays an error message due to the non-availability of a help browser

External browsers depend on your system and the contents of `LIB/help.cnf`, the default includes:

`htmlview` (displays HTML help pages via `htmlview`),
`mac` (displays HTML help pages via `open`),
`mac-net` (displays HTML help pages via `open`),
`mozilla` (displays HTML help pages via `mozilla`),
`firefox` (displays HTML help pages via `firefox`),
`konqueror` (displays HTML help pages via `konqueror`),
`galeon` (displays HTML help pages via `galeon`),
`netscape` (displays HTML help pages via `netscape`),
`safari` (displays HTML help pages on MacOSX via `safari`),
`tkinfo` (displays INFO help pages via `tkinfo`),
`xinfo` (displays INFO help pages via `info`),
`info` (displays INFO help pages via `info`),
`lynx` (displays HTML help pages via `lynx`).

The browser which is used to display the help information, can be either set at startup time with the command line option (see Section 3.1.6 [Command line options], page 19)

```
--browser=<browser>
```

or with the SINGULAR command (see Section 5.1.153 [system], page 270)

```
system("--browser", "<browser>");
```

The SINGULAR command

```
system("browsers");
```

lists all available browsers and the command

```
system("--browser");
```

returns the currently used browser.

If no browser is explicitly set by the user, then the first available browser (w.r.t. the order of the browsers in the file `LIB/help.cnf`) is chosen.

The `.singularrc` (see Section 3.1.7 [Startup sequence], page 22) file is a good place to set your default browser. Recall that if a file `$HOME/.singularrc` exists on your system, then the content of this file is executed before the first user input. Hence, putting

```

if (! system("--emacs"))
{
  // only set help browser if not running within emacs
  system("--browser", "info");
}
// if help browser is later on set to a web browser,
// allow it to fetch HTML pages from the net
system("--allow-net", 1);

```

in your file `$HOME/.singularrc` sets your default browser to `info`, unless `SINGULAR` is run within `emacs` (in which case the default browser is automatically set to `emacs`).

Obviously, certain external files and programs are required for the `SINGULAR` help system to work correctly. If something is not available or goes wrong, here are some tips for troubleshooting the help system:

- Under Unix, the environment variable `DISPLAY` has to be set for all X11 browsers to work.
- The help browsers are only available if the respective programs are installed on your system (for `xinfo`, the programs `xterm` and `info` are necessary). You can explicitly specify which program to use, by changing the entry in `LIB/help.cnf`
- If the help browser cannot find the local html pages of the `SINGULAR` manual (which it will look for at `$RootDir/html` – see Section 3.8.11 [Loading a library], page 66 for more info on `$RootDir`) and the (command-line) option `--allow-net` has *explicitly* been set (see Section 3.1.6 [Command line options], page 19 and Section 5.1.153 [system], page 270 for more info on setting values of command-line options), then it dispatches the html pages from <https://www.singular.uni-kl.de/Manual>. (Note that the non-local net-access of HTML pages is disabled, by default.)

An alternative location of a local directory where the html pages reside can be specified by setting the environment variable `SINGULAR_HTML_DIR`.

- The `info` based help browsers `tkinfo`, `xinfo`, `info`, and `builtin` need the (info) file `singular.hlp` which will be looked for at `$RootDir/info/singular.hlp` (see Section 3.8.11 [Loading a library], page 66 for more info on `$RootDir`). An alternative location of the info file of the manual can be specified by setting the environment variable `SINGULAR_INFO_FILE`.

Section 3.1.6 [Command line options], page 19

Info help browsers

The help browsers `tkinfo`, `xinfo` and `info` (so-called info help browsers) are based on the `info` program from the GNU `texinfo` package. See section “Getting started” in *The Info Manual*, for more information.

For info help browsers, the online manual is decomposed into “nodes” of information, closely related to the division of the printed manual into sections and subsections. A node contains text describing a specific topic at a specific level of detail. The top line of a node is its “header”. The node’s header tells the name of the current node (`Node:`), the name of the next node (`Next:`), the name of the previous node (`Prev:`), and the name of the upper node (`Up:`).

To move within `info`, type commands consisting of single characters. Do not type `RETURN`. Do not use cursor keys, either. Using some of the cursor keys by accident might pop to some totally different node. Type `1` to return to the original node. Some of the `info` commands read input from the command line at the bottom. The `TAB` key may be used to complete partially entered input.

The most important commands are:

`q` leaves the online help system

n	goes to the next node
p	goes to the previous node
u	goes to the upper node
m	picks a menu item specified by name
f	follows a cross reference
l	goes to the previously visited node
b	goes to the beginning of the current node
e	goes to the end of the current node
SPACE	scrolls forward a page
DEL	scrolls backward a page
h	invokes info tutorial (use l to return to the manual or CTRL-X 0 to remove extra window)
CTRL-H	shows a short overview over the online help system (use l to return to the manual or CTRL-X 0 to remove extra window)
s	searches through the manual for a specific string, and selects the node in which the next occurrence is found
1, . . . , 9	picks i-th subtopic from a menu

3.1.4 Interrupting SINGULAR

On Unix-like operating systems and on Windows NT, typing **CTRL-C** (or, alternatively **C-c C-c**, when running within Emacs), interrupts SINGULAR. SINGULAR prints the current command and the current line and prompts for further action. The following choices are available:

a	returns to the top level after finishing the current (kernel) command. Notice that commands of the SINGULAR kernel (like std) cannot be aborted, i.e. (a)bort only happens whenever the interpreter is active.
c	continues
q	quits SINGULAR

3.1.5 Editing input

The following keys can be used for editing the input and retrieving previous input lines:

TAB	provides command line completion for function names and file names
CTRL-B	moves cursor to the left
CTRL-F	moves cursor to the right
CTRL-A	moves cursor to the beginning of the line
CTRL-E	moves cursor to the end of the line
CTRL-D	deletes the character under the cursor Warning: on an empty line, CTRL-D is interpreted as the EOF character which immediately terminates SINGULAR.

BACKSPACE	
DELETE	
CTRL-H	deletes the character before the cursor
CTRL-K	kills from cursor to the end of the line
CTRL-U	kills from cursor to the beginning of the line
CTRL-N	saves the current line to history and gives the next line
CTRL-P	saves the current line to history and gives the previous line
RETURN	saves the current line to the history and sends it to the SINGULAR parser for interpretation

When run under a Unix-like operating system and in its ASCII-terminal user interface, SINGULAR tries to dynamically link at runtime with the GNU Readline library. See section “Command Line Editing” in *The GNU Readline Library Manual*, for more information. If a shared version of this library can be found on your machine, then additional command-line editing features like history completion are available. In particular, if SINGULAR is able to load that library and if the environment variable SINGULARHIST is set and has a name of a valid file as value, then the input history is stored across sessions using this file. Otherwise, i.e., if the environment variable SINGULARHIST is not set, then the history of the last inputs is only available for previous commands of the current session.

3.1.6 Command line options

The startup syntax is

```
Singular [options] [file1 [file2 ...]]
ESingular [options] [file1 [file2 ...]]
```

Options can be given in both their long and short format. The following options control the general behaviour of SINGULAR:

- d, --sdb Enable the use of the source code debugger. See Section 3.9.3 [Source code debugger], page 68.
- e, --echo[=VAL]
Set value of variable echo to VAL (integer in the range 0, . . . , 9). Without an argument, echo is set to 1, which echoes all input coming from a file. By default, the value of echo is 0. See Section 5.3.2 [echo], page 298.
- h, --help
Print a one-line description of each command line option and exit.
- allow-net
Allow the help browsers based on a web browser to fetch HTML manual pages over the net from the WWW home-site of SINGULAR. See Section 3.1.3 [The online help system], page 15, for more info.
- browser="VAL"
Use VAL as browser for the SINGULAR online manual.
VAL may be one of the browsers mentioned in LIB/help.cnf, for example html (Windows only), mozilla, firefox, konqueror, galeon, netscape, safari (OsX only), xinfo, tkinfo, info, builtin, or emacs. Depending on your platform and local installation, only some browsers might be available. The default browser is html for Windows and one based on a web browser for Unix platforms. See Section 3.1.3 [The online help system], page 15, for more info.

- `--no-rc` Do not execute the `.singularrc` file on start-up. By default, this file is executed on start-up. See Section 3.1.7 [Startup sequence], page 22.
- `--no-stdlib` Do not load the library `standard.lib` on start-up. By default, this library is loaded on start-up. See Section 3.1.7 [Startup sequence], page 22.
- `--no-warn` Do not display warning messages.
- `--no-out` Suppress display of all output.
- `--no-shell` Runs Singular in restricted mode to disallow shell escape commands. Objects of type `link` will also be unable to use.
- `-t, --no-tty` Do not redefine the characteristics of the terminal. This option should be used for batch processes.
- `-q, --quiet` Do not print the start-up banner and messages when loading libraries. Furthermore, redirect `stderr` (all error messages) to `stdout` (normal output channel). This option should be used if SINGULAR's output is redirected to a file.
- `-v` Print extended information about the version and configuration of SINGULAR (used optional parts, compilation date, start of random generator etc.). This information should be included if a user reports an error to the authors.
It also list all the used directories/files (see Section 8.5 [Used environment variables], page 953).

The following command line options allow manipulations of the timer and the pseudo random generator and enable the passing of commands and strings to SINGULAR:

- `-c, --execute=STRING`
Execute `STRING` as (a sequence of) SINGULAR commands on start-up after the `.singularrc` file is executed, but prior to executing the files given on the command line. E.g., `Singular -c "help all.lib; quit;"` shows the help for the library `all.lib` and exits.
- `-u, --user-option=STRING`
Returns `STRING` on `system("--user-option")`. This is useful for passing arbitrary arguments from the command line to the SINGULAR interpreter. E.g., `Singular -u "xxx.dump" -c 'getdump(system("--user-option"))'` reads the file `xxx.dump` at start-up and allows the user to start working with all the objects defined in a previous session.
- `-r, --random=SEED`
Seed (i.e., set the initial value of) the pseudo random generator with integer `SEED`. If this option is not given, then the random generator is seeded with a time-based `SEED` (the number of seconds since January, 1, 1970, on Unix-like operating systems, to be precise).
- `--min-time=SECS`
If the `timer` (see Section 5.3.8 [timer], page 300), resp. `rtimer` (see Section 5.3.10 [rtimer], page 303), variable is set, report only times larger than `SECS` seconds (`SECS` needs to be a floating point number greater than 0). By default, this value is set to 0.5

(i.e., half a second). E.g., the option `--min-time=0.01` forces SINGULAR to report all times larger than 1/100 of a second.

`--ticks-per-sec=TICKS`

Set unit of timer to TICKS ticks per second (i.e., the value reported by the `timer` and `rtimer` variable divided by TICKS gives the time in seconds). By default, this value is 1.

`--cpus=CPUs`

set the maximal number of CPUs to use.

`--cntrlc=C`

set the default answer for interrupt signals to C which should be a for abort, c for continue or q for quit.

The next three options are of interest for the use with ssi links:

`-b, --batch`

Run in batch mode. Opens a TCP/IP connection with host specified by `--MPhost` at the port specified by `--MPport`. Input is read from and output is written to this connection in the format given by `--link`. See Section 4.9.5 [Ssi links], page 96.

`--MPport=PORT`

Use PORT as default port number for connections (whenever not further specified). This option is mandatory when the `--batch` option is given. See Section 4.9.5 [Ssi links], page 96.

`--MPhost=HOST`

Use HOST as default host for connections (whenever not further specified). This option is mandatory when the `--batch` option is given. See Section 4.9.5 [Ssi links], page 96.

Finally, the following options are only available when running ESingular (see Section 3.2.2 [Running SINGULAR under Emacs], page 25 for details).

`--emacs=EMACS`

Use EMACS as Emacs program to run the SINGULAR Emacs interface, where EMACS may e.g. be emacs or xemacs.

`--emacs-dir=DIR`

Set the singular-emacs-home-directory, which is the directory where singular.el can be found, to DIR.

`--emacs-load=FILE`

Load FILE on Emacs start-up, instead of the default load file.

`--singular=PROG`

Start PROG as SINGULAR program within Emacs

The value of options given to SINGULAR (resp. their default values, if an option was not given), can be checked with the command `system("--long_option_name")`. See Section 5.1.153 [system], page 270.

```
system("--quiet"); // if ‘quiet’ 1, otherwise 0
↳ 1
system("--min-time"); // minimal reported time
↳ 0.5
system("--random"); // seed of the random generator
↳ 12345678
```

Furthermore, the value of options (e.g., `--browser`) can be re-defined while SINGULAR is running using the command `system("--long_option_name_string ", expression)`. See Section 5.1.153 [system], page 270.

```
system("--browser", "builtin"); // sets browser to 'builtin'
system("--ticks-per-sec", 100); // sets timer resolution to 100
```

3.1.7 Startup sequence

On start-up, SINGULAR

1. loads the library `standard.lib` (provided the `--no-stdlib` option was not given),
2. searches the current directory and then the home directory of the user, and then all directories contained in the library `SearchPath` (see Section 3.8.11 [Loading a library], page 66 for more info on `SearchPath`) for a file named `.singularrc` and executes it, if found (provided the `--no-rc` option was not given),
3. executes the string specified with the `--execute` command line option,
4. executes the files `file1`, `file2` ... (given on the command line) in that order.

Note: `.singularrc` file(s) are an appropriate place for setting some default values of (command-line) options.

For example, a system administrator might remove the locally installed HTML version of the manual and put a `.singularrc` file with the following content

```
if (system("version") >= 1306) // assure backwards-compatibility
{
  system("--allow-net", 1);
}; // the last semicolon is important: otherwise no ">", but "." prompt
```

in the directory containing the SINGULAR libraries, thereby allowing to fetch the HTML on-line help from the WWW home-site of SINGULAR.

On the other hand, a single user might put a `.singularrc` with the following content

```
if (system("version") >= 1306) // assure backwards-compatibility
{
  if (! system("--emacs"))
  {
    // set default browser to info, unless we run within emacs
    system("--browser", "info");
  }
}; // the last semicolon is important: otherwise no ">", but "." prompt
```

in his home directory, which sets the default help browser to `info` (unless SINGULAR is run within emacs) and thereby prevents the execution of the "global" `.singularrc` file installed by the system administrator (since the `.singularrc` file of the user is found before the "global" `.singularrc` file installed by the system administrator).

3.2 Emacs user interface

Besides running SINGULAR in an ASCII-terminal, SINGULAR might also be run within Emacs. Emacs (or, XEmacs which is very similar) is a powerful and freely available text editor, which, among others, provides a framework for the implementation of interactive user interfaces. Starting from version 1.3.6, SINGULAR provides such an implementation, the so-called SINGULAR Emacs mode, or Emacs user interface.

Generally, we recommend to use the Emacs interface, instead of the ASCII-terminal interface: The Emacs interface does not only provide everything the ASCII-terminal interface provides, but offers much more. Among others, it offers

- color highlighting
- truncation of long lines
- folding of input and output
- TAB-completion for help topics
- highlighting of matching parentheses
- key-bindings and interactive menus for most user interface commands and for basic SINGULAR commands (such as loading of libraries and files)
- a mode for running interactive SINGULAR demonstrations
- convenient ways to edit SINGULAR input files
- interactive customization of nearly all aspects of the user-interface.

In order to use the SINGULAR-Emacs interface you need to have Emacs version 20 or higher, or XEmacs version 20.3 or higher installed on your system. These editors can be downloaded for most hard- and software platforms, sources from either <http://www.gnu.org/software/emacs/emacs.html> (Emacs), or from <http://www.xemacs.org> (XEmacs). (Download of binaries depend on your OS). The differences between Emacs and XEmacs w.r.t. the SINGULAR-Emacs interface are marginal – which editor to use is mainly a matter of personal preferences.

The simplest way to start-up SINGULAR in its Emacs interface is by running the program `ESingular` which is contained in the Singular distribution. Alternatively, SINGULAR can be started within an already running Emacs – see Section 3.2.2 [Running SINGULAR under Emacs], page 25 for details.

The next section gives a tutorial-like introduction to Emacs. This introductory section is followed by sections which explain the functionality of various aspects of the Emacs user interface in more detail: how to start/restart/kill SINGULAR within Emacs, how to run an interactive demonstration, how to customize the Emacs user interface, etc. Finally, the 20 most important commands of the Emacs interface together with their key bindings are listed.

3.2.1 A quick guide to Emacs

This section gives a tutorial-like introduction to Emacs. Especially to users who are not familiar with Emacs, we recommend that they go through this section and try out the described features.

Emacs commands generally involve the `CONTROL` key (sometimes labeled `CTRL` or `CTL`) or the `META` key. On some keyboards, the `META` key is labeled `ALT` or `EDIT` or something else (for example, on Sun keyboards, the diamond key to the left of the space-bar is `META`). If there is no `META` key, the `ESC` key can be used, instead. Rather than writing out `META` or `CONTROL` each time we want to prefix a character, we will use the following abbreviations:

`C-<chr>` means hold the `<CONTROL>` key while typing the character `<chr>`. Thus, `C-f` would be: hold the `<CONTROL>` key and type `f`.
`M-<chr>` means hold the `<META>` key down while typing `<chr>`. If there is no `<META>` key, type `<ESC>`, release it, then type the character `<chr>`.

For users new to Emacs, we highly recommend that they go through the interactive Emacs tutorial: type `C-h t` to start it.

For others, it is important to understand the following Emacs concepts:

window In Emacs terminology, a window refers to separate panes within the same window of the window system, and not to overlapping, separate windows. When using SINGULAR

within Emacs, extra windows may appear which display help or output from certain commands. The most important window commands are:

<code>C-x 1</code>	<code>File->Un-Split</code>	Un-Split window (i.e., kill other windows)
<code>C-x o</code>		Goto other window, i.e. move cursor into other window.

cursor and point

The location of the cursor in the text is also called "point". To paraphrase, the cursor shows on the screen where point is located in the text. Here is a summary of simple cursor-moving operations:

<code>C-f</code>	Move forward a character
<code>C-b</code>	Move backward a character
<code>M-f</code>	Move forward a word
<code>M-b</code>	Move backward a word
<code>C-a</code>	Move to the beginning of line
<code>C-e</code>	Move to the end of line

buffer

Any text you see in an Emacs window is always part of some buffer. For example, each file you are editing with Emacs is stored inside a buffer, but also SINGULAR is running inside an Emacs buffer. Each buffer has a name: for example, the buffer of a file you edit usually has the same name as the file, SINGULAR is running in a buffer which has the name `*singular*` (or, `*singular<2>*`, `*singular<3>*`, etc., if you have multiple SINGULAR sessions within the same Emacs).

When you are asked for input to an Emacs command, the cursor moves to the bottom line of Emacs, i.e., to a special buffer, called the "minibuffer". Typing `(RETURN)` within the minibuffer, ends the input, typing `(SPACE)` within the minibuffer, lists all possible input values to the interactive Emacs command.

The most important buffer commands are

<code>C-x b</code>	Switch buffer
<code>C-x k</code>	Kill current buffer

Alternatively, you can switch to or kill buffers using the **Buffer** menu.

Executing commands

Emacs commands are executed by typing `M-x <command-name>` (remember that `(SPACE)` completes partial command names). Important and frequently used commands have short-cuts for their execution: Key bindings or even menu entries. For example, a file can be loaded with `M-x load-file`, or `C-x C-f`, or with the **File->Open** menu.

How to exit

To end the Emacs (and, SINGULAR) session, type `C-x C-c` (two characters), or use the **File -> Exit** menu.

When Emacs hangs

If Emacs stops responding to your commands, you can stop it safely by typing `C-g`, or, if this fails, by typing `C-]`.

More help

Nearly all aspects of Emacs are very well documented: type `C-h` and then a character saying what kind of help you want. For example, typing `C-h i` enters the **Info** documentation browser.

Using the mouse

Emacs is fully integrated with the mouse. In particular, clicking the right mouse button brings up a pop-up menu which usually contains a few commonly used commands.

3.2.2 Running SINGULAR under Emacs

There are two ways to start the SINGULAR Emacs interface: Typing `ESingular` instead of `Singular` on the command shell launches a new Emacs process, initializes the interface and runs SINGULAR within Emacs. The other way is to start the interface in an already running Emacs, by typing `M-x singular` inside Emacs. This initializes the interface and runs SINGULAR within Emacs. Both ways are described in more detail below.

Note: To properly run the Emacs interface, several files are needed which usually reside in the `emacs` subdirectory of your SINGULAR distribution. This directory is called `singular-emacs-home-directory` in the following.

Starting the interface using ESingular

As mentioned above, `ESingular` is an "out-of-the-box" solution: You don't have to add special things to your `.emacs` startup file to initialize the interface; everything is done for you in a special file called `.emacs-singular` (which comes along with the SINGULAR distribution and resides in the `singular-emacs-home-directory`) which is automatically loaded on Emacs startup (and the loading of the `.emacs` file is automatically suppressed).

The customizable variables of the SINGULAR Emacs interface are set to defaults which give the novice user a very shell like feeling of the interface. Nevertheless, these default settings can be changed, see Section 3.2.4 [Customization of the Emacs interface], page 27. Besides other Emacs initializations, such as fontification or blinking parentheses, a new menu item called `Singular` is added to the main menu, providing menu items for starting SINGULAR. On XEmacs, a button starting SINGULAR is added to the main toolbar.

The SINGULAR interface is started automatically; once you see a buffer called `*singular*` and the SINGULAR prompt, you are ready to start your SINGULAR session.

`ESingular` inherits all `Singular` options. For a description of all these options, see Section 3.1.6 [Command line options], page 19. Additionally there are the following options which are special to `ESingular`:

command-line option / environment variable	functionality
<code>--emacs=EMACS</code> <code>ESINGULAR_EMACS</code>	Use <code>EMACS</code> as Emacs program to run the SINGULAR Emacs interface, where <code>EMACS</code> may e.g. be <code>emacs</code> or <code>xemacs</code> .
<code>--emacs-dir=DIR</code> <code>ESINGULAR_EMACS_DIR</code>	Set the <code>singular-emacs-home-directory</code> , which is the directory where <code>singular.el</code> can be found, to <code>DIR</code> .
<code>--emacs-load=FILE</code> <code>ESINGULAR_EMACS_LOAD</code>	Load <code>FILE</code> on Emacs start-up, instead of the default load file.
<code>--singular=PROG</code> <code>ESINGULAR_SINGULAR</code>	Start <code>PROG</code> as SINGULAR program within Emacs

Notice that values of these options can also be given by setting the above mentioned environment variables (where values given as command-line arguments take priority over values given by environment variables).

Starting the interface within a running Emacs

If you are a more experienced Emacs user and you already have your own local `.emacs` startup file, you might want to start the interface out of your running Emacs without using `ESingular`. For this, you should add the following lisp code to your `.emacs` file:

```
(setq load-path (cons "<singular-emacs-home-directory>" load-path))
(autoload 'singular "singular"
  "Start Singular using default values." t)
(autoload 'singular-other "singular"
  "Ask for arguments and start Singular." t)
```

Then typing `M-x singular` in a running Emacs session initializes the interface in a new buffer and launches a SINGULAR process. The SINGULAR prompt comes up and you are ready to start your SINGULAR session.

It is a good idea to take a look at the (well documented) file `.emacs-singular` in the `singular-emacs-home-directory`, which comes along with the distribution. In it you find some useful initializations of the SINGULAR interface as well as some lisp code, which, for example, adds a button to the XEmacs toolbar. Some of this code might be useful for your `.emacs` file, too. And if you are an Emacs wizard, it is of course a good idea to take a look at `singular.el` in the `singular-emacs-home-directory`.

CYGWIN and ESingular

X11 server install `xlaunch`, `emacs-X11`, `xterm` and all dependencies. Create with `xlaunch` a startup file for the X-server which also starts the client `xterm`. From that one can start ESingular.

fork problems The simplest way to overcome fork problem is to run `/usr/bin/rebase-trigger full`, then stop all Cygwin processes and services, and then run `setup-x86.exe`. The `_autorebase` postinstall script will then take care of the rebase. Occasionally it is necessary to reboot the computer before doing this.

Starting, interrupting and stopping SINGULAR

There are the following commands to start and stop SINGULAR:

- `singular-other` (or menu `Singular`, item `Start...`)

Starts a SINGULAR process and asks for the following four parameters in the minibuffer area:

1. The SINGULAR executable. This can either be a file name with complete path, e.g., `/local/bin/Singular`. Then exactly this executable is started. The path may contain the character `~` denoting your home directory. Or it can be the name of a command without path, e.g., `Singular`. Then the executable is searched for in your `$PATH` environment variable.
2. The default working directory. This is the path to an existing directory, e.g., `~/work`. The current directory is set to this directory before SINGULAR is started.
3. Command line options. You can set any SINGULAR command line option (see Section 3.1.6 [Command line options], page 19).
4. The buffer name. You can specify the name of the buffer the interface is running in.

- `singular` (or menu `Singular`, item `Start default`)

Starts SINGULAR with default settings for the executable, the working directory, command line switches, and the buffer name. You can customize this default settings, see Section 3.2.4 [Customization of the Emacs interface], page 27.

- `singular-exit-singular` (bound to `C-c $` or menu `Singular`, item `Exit`)

Kills the running SINGULAR process of the current buffer (but does not kill the buffer). Once you have killed a SINGULAR process you can start a new one in the same buffer with the command `singular` (or select the item `Start default` of the `Singular` menu).

- `singular-restart` (bound to `C-c C-r` or menu `Singular`, item `Restart`)
Kills the running SINGULAR process of the current buffer and starts a new process in the same buffer with exactly the same command line arguments as before.
- `singular-control-c` (bound to `C-c C-c` or menu `Singular`, item `Interrupt`)
Interrupt the SINGULAR process running in the current buffer. Asks whether to (a)abort the current SINGULAR command, (q)uit or (r)estart the current SINGULAR process, or (c)ontinue without doing anything (default).

Whenever a SINGULAR process is started within the Emacs interface, the contents of a special startup file (by default `~/ .emacs-singularrc`) is pasted as input to SINGULAR at the very end of the usual startup sequence (see Section 3.1.7 [Startup sequence], page 22). The name of the startup file can be changed, see Section 3.2.4 [Customization of the Emacs interface], page 27.

3.2.3 Demo mode

The Emacs interface can be used to run interactive SINGULAR demonstrations. A demonstration is started by loading a so-called SINGULAR demo file with the Emacs command `singular-demo-load`, bound to `C-c C-d`, or with the menu `Commands->Load Demo`.

A SINGULAR demo file should consist of SINGULAR commands separated by blank lines. When running a demo, the input up to the next blank line is echoed to the screen. Hitting `RETURN` executes the echoed commands and shows their output. Hitting `RETURN` again, echos the next commands to the screen, and so on, until all commands of the demo file are executed. While running a demo, you can execute other commands on the SINGULAR prompt: the next input from the demo file is then echoed again, if you hit `RETURN` on an empty input line.

A SINGULAR demo can prematurely be exited by either starting another demo, or by executing the Emacs command `singular-demo-exit` (menu: `Commands->Exit Demo`).

Some aspects of running SINGULAR demos can be customized. See Section 3.2.4 [Customization of the Emacs interface], page 27, for more info.

3.2.4 Customization of the Emacs interface

Emacs provides a convenient interface to customize the behavior of Emacs and the SINGULAR Emacs interface for your own needs. You enter the customize environment by either calling `M-x customize` (on XEmacs you afterwards have to enter `emacs` in the minibuffer area) or by selecting the menu item `Options->Customize->Emacs...` for XEmacs, and the menu item `Help->Customize->Toplevel Customization Group` for Emacs, resp. A brief introduction to the customization mode comes up with the customization buffer. All customizable parameters are hierarchically grouped and you can browse through all these groups and change the values of the parameters using the mouse. At the end you can save your settings to a file making your changes permanent.

To change the settings of the SINGULAR Emacs interface you can either select the item `Preferences` of the `Singular` menu, call `M-x customize-group` and give the argument `singular-interactive` in the minibuffer area, or browse from the top-level customization group through the path `External->Singular->Singular interactive`.

The SINGULAR interface customization buffer is divided into four groups:

- `Singular Faces`
Here you can specify various faces used if `font-lock-mode` is enabled (which, by default, is).

- Singular Sections And Foldings

Here you can specify special faces for SINGULAR input and output and change the text used as replacement for folded sections.

For doing this, you also might find handy the function `customize-face-at-point`, which lets you customize the face at the current position of point. This function is automatically defined if you run `ESingular`). Otherwise, you should add its definition (see below) to your personal `.emacs` file.

- Singular Interactive Miscellaneous

Here you can specify various things such as the behavior of the cursor keys, the name of the special SINGULAR startup file, the appearance of the help window, or the default values for the `singular` command.

- Singular Demo Mode

Here you can specify how chunks of the demo file are divided, or specify a default directory for demo files.

When you run `ESingular`, the settings of customized variables are saved in the file `$HOME/.emacs-singular-cust`. Otherwise, the settings are appended to your `.emacs` file. Among others, this means that the customized settings of `ESingular` are not automatically taken over by a "normal" Emacs, and vice versa.

3.2.5 Editing SINGULAR input files with Emacs

Since SINGULAR's programming language is similar to C, you should use the Emacs C/C++-mode to edit SINGULAR input files and SINGULAR libraries. Among others, this Emacs mode provides automatic indentation, line-breaking and keyword highlighting.

When running `ESingular`, the C/C++-mode is automatically turned on whenever a file with the suffix `.sing`, or `.lib` is loaded.

For Emacs sessions which were not started by `ESingular`, you should add the following to your `.emacs` file:

```
;; turn on c++-mode for files ending in ".sing" and ".lib"
(setq auto-mode-alist (cons '("\\.sing\\\\" . c++-mode) auto-mode-alist))
(setq auto-mode-alist (cons '("\\.lib\\\\" . c++-mode) auto-mode-alist))
;; turn-on fontification for c++-mode
(add-hook 'c++-mode-hook
          (function (lambda () (font-lock-mode 1))))
;; turn on aut-new line and hungry-delete
(add-hook 'c++-mode-hook
          (function (lambda () (c-toggle-auto-hungry-state 1))))
;; a handy function for customization
(defun customize-face-at-point ()
  "Customize face which point is at."
  (interactive)
  (let ((face (get-text-property (point) 'face)))
    (if face
        (customize-face face)
        (message "No face defined at point")))))
```

Notice that you can change the default settings for source-code highlighting (colors, fonts, etc.) by customizing the respective faces using the `Customize` feature of Emacs. For doing this, you might find handy the above given function `customize-face-at-point`, which lets you customize the face of the current position of point (this function is automatically defined if you run `ESingular`).

3.2.6 Top 20 Emacs commands

Here is a list of the 20 probably most useful commands when using the SINGULAR Emacs interface. Starting and stopping of SINGULAR:

- `singular` (menu `Singular->Start Default...`): starts SINGULAR using default arguments.
- `singular-other` (menu `Singular->Start`): starts SINGULAR asking for several arguments in the minibuffer area.
- `singular-exit` (key `C-c $` or menu `Singular->Exit`): kills the SINGULAR process running in the current buffer (but does not kill the buffer).
- `singular-restart` (key `C-c C-r` or menu `Singular->Restart`): kills the SINGULAR process running in the current buffer and starts a new SINGULAR process with exactly the same arguments as before.

Editing input and output:

- `singular-beginning-of-line` (key `C-a`): moves point to beginning of line, then skips past the SINGULAR prompt, if any.
- `singular-toggle-truncate-lines` (key `C-c C-t` or menu `Commands->Truncate lines`): toggles whether long lines should be truncated or not. If lines are not truncated, the commands `singular-scroll-left` and `singular-scroll-right` are useful to scroll left and right, resp.
- `singular-dynamic-complete` (key `TAB`): performs context specific completion. If point is inside a string, file name completion is done. If point is at the end of a help command (i.e., `help` or `?`), completion on SINGULAR help topics is done. If point is at the end of an example command (i.e., `example`), completion is done on SINGULAR examples. In all other cases, completion on SINGULAR commands is done.
- `singular-folding-toggle-fold-latest-output` (key `C-c C-o` or menu `Commands->Fold/Unfold Latest Output`): toggles folding of the latest output section. If your last SINGULAR command produced a huge output, simply type `C-c C-o` and it will be replaced by a single line.
- `singular-folding-toggle-fold-at-point` (key `C-c C-f` or menu `Commands->Fold/Unfold At Point`): toggles folding of the section the point currently is in.
- `singular-folding-fold-all-output` (menu `Commands->Fold All Output`): folds all SINGULAR output, replacing each output section by a single line.
- `singular-folding-unfold-all-output` (menu `Commands->Unfold All Output`): unfolds all SINGULAR output sections showing their true contents.

Loading of files and SINGULAR demo mode:

- `singular-load-library` (key `C-c C-l` or menu `Commands->Libraries->other...`): asks for a standard library name or a library file in the minibuffer (hit `TAB` for completion) and loads the library into SINGULAR. The submenu `Libraries` of the `Commands` menu also provides a separate menu item for each standard library.
- `singular-load-file` (key `C-c <` or menu `Commands->Load File...`): asks for a file name in the minibuffer (which is expanded using `expand-file-name` if given a prefix argument) and loads the file into SINGULAR.
- `singular-demo-load` (key `C-c C-d` or menu `Commands->Load Demo...`): asks for a file name of a SINGULAR demo file in the minibuffer area (hit `SPACE` for completion) and enters the SINGULAR demo mode showing the first chunk of the demo.
- `singular-demo-exit` (menu `Commands->Exit Demo`): exits from SINGULAR demo mode and cleans up everything that is left from the demo.

Help and Customization:

- `singular-help` (key `C-h C-s` or menu `Singular->Singular Help`): asks for a SINGULAR help topic in the minibuffer (hit `TAB` for completion) and shows the help text in a separate buffer.
- `singular-example` (key `C-c C-e` or menu `Singular->Singular Example`): asks for a SINGULAR command in the minibuffer (hit `TAB` for completion) and executes the example of this command in the current SINGULAR buffer.
- `customize-group` (menu `Singular->Preferences`): enters the customization group of the SINGULAR Emacs interface. (If called via `M-x customize-group` give argument `singular-interactive` in the minibuffer area.)

3.3 Rings and orderings

All non-trivial algorithms in SINGULAR require the prior definition of a ring. Such a ring can be

1. a polynomial ring over a field,
2. a polynomial ring over a ring
3. a localization of 1.
4. a quotient ring by an ideal of 1. or 2.,
5. a tensor product of 1. or 2.

Except for quotient rings, all of these rings are realized by choosing a coefficient field, ring variables, and an appropriate global or local monomial ordering on the ring variables. See Section 3.3.3 [Term orderings], page 34, Appendix C [Mathematical background], page 766.

The coefficient field of the rings may be

1. the field of rational numbers Q (`QQ`),
2. finite fields Z/p , p a prime ≤ 2147483647 ,
3. finite fields $GF(p^n)$ with p^n elements, p a prime, $p^n \leq 2^{16}$,
4. transcendental extension of Q or Z/p ,
5. simple algebraic extension of Q or Z/p ,
6. the field of real numbers represented by floating point numbers of a user defined precision,
7. the field of complex numbers represented by (pairs of) floating point numbers of a user defined precision,
8. the ring of integers (`ZZ`),
9. finite rings Z/m with $m \in Z$.

In case of coefficient rings, which are not fields, only the following functions are guaranteed to work:

- basic polynomial arithmetic, i.e. addition, multiplication, division, exponentiation
- `std`, i.e. computing standard bases (and related: `syz`, etc.)
- `interred`
- `reduce`

Throughout this manual, the current active ring in SINGULAR is called basering. The reserved name `basing` in SINGULAR is an alias for the current active ring. The basering can be set by declaring a new ring as described in the following subsections or by using the commands `setring` and `keepring`. See Section 5.2.11 [`keepring`], page 293, Section 5.1.139 [`setring`], page 255.

Objects of ring dependent types are local to a ring. To access them after a change of the basering they have to be mapped using `map` or by the functions `imap` or `fetch`. See Section 3.5.4 [Objects],

page 45, Section 5.1.38 [fetch], page 180, Section 5.1.59 [imap], page 195, Section 4.11 [map], page 103.

All changes of the basering in a procedure are local to this procedure unless a `keepring` command is used as the last statement of the procedure. See Section 3.7 [Procedures], page 50, Section 5.2.11 [keepring], page 293.

3.3.1 Examples of ring declarations

The exact syntax of a ring declaration is given in the next two subsections; this subsection lists some examples first. Note that the chosen ordering implies that a unit-elements of the ring will be among the elements with leading monomial 1. For more information, see Section B.2 [Monomial orderings], page 760.

Every floating point number in a ring consists of two parts, which may be chosen by the user. The leading part represents the number and the rest is for numerical stability. Two numbers with a difference only in the rest will be regarded equal.

- the ring $Z/32003[x, y, z]$ with degree reverse lexicographical ordering. The exact ring declaration may be omitted in the first example since this is the default ring:

```
ring r1;
ring r2 = 32003, (x,y,z), dp;
ring r3=(ZZ/32003)[x,y,z];
ring r4 = (ZZ/32003), (x,y,z), dp;
```

- similar examples with indexed variables. The ring variables of r1 are going to be $x(1)..x(10)$; in r2 they will be $x(1)(1), x(1)(2), \dots, x(1)(8), x(2)(1), \dots, x(5)(8)$:

```
ring r1 = 32003, (x(1..10)), dp;
ring r2 = 32003, (x(1..5)(1..8)), dp;
ring r3 = (ZZ/32003)[x(1..5)(1..8)];
ring r4 = (ZZ/32003), (x(1..5)(1..8)), dp;
```

- the ring $Q[a, b, c, d]$ with lexicographical ordering:

```
ring r1 = 0, (a,b,c,d), lp;
ring r2 = QQ, (a,b,c,d), lp;
```

- the ring $Z/7[x, y, z]$ with local degree reverse lexicographical ordering. The non-prime 10 is converted to the next lower prime in the second example:

```
ring r1 = 7, (x,y,z), ds;
ring r2 = 10, (x,y,z), ds;
ring r3 = (ZZ/7), (x,y,z), ds;
```

- the ring $Z/7[x_1, \dots, x_6]$ with lexicographical ordering for x_1, x_2, x_3 and degree reverse lexicographical ordering for x_4, x_5, x_6 :

```
ring r1 = 7, (x(1..6)), (lp(3), dp);
ring r2 = (ZZ/7), (x(1..6)), (lp(3), dp);
```

- the localization of $(Q[a, b, c])[x, y, z]$ at the maximal ideal (x, y, z) :

```
ring r1 = 0, (x,y,z,a,b,c), (ds(3), dp(3));
ring r2 = QQ, (x,y,z,a,b,c), (ds(3), dp(3));
```

- the ring $Q[x, y, z]$ with weighted reverse lexicographical ordering. The variables x , y , and z have the weights 2, 1, and 3, respectively, and vectors are first ordered by components (in descending order) and then by monomials:

```
ring r1 = 0, (x,y,z), (c, wp(2,1,3));
ring r2 = QQ, (x,y,z), (c, wp(2,1,3));
```

For ascending component order, the component ordering `C` has to be used.

- the ring $K[x, y, z]$, where $K = Z/7(a, b, c)$ denotes the transcendental extension of $Z/7$ by a , b and c with degree lexicographical ordering:

```
ring r = (7,a,b,c),(x,y,z),Dp;
```

- the ring $K[x, y, z]$, where $K = Z/7[a]$ denotes the algebraic extension of degree 2 of $Z/7$ by a . In other words, K is the finite field with 49 elements. In the first case, a denotes an algebraic element over $Z/7$ with minimal polynomial $\mu_a = a^2 + a + 3$, in the second case, a refers to some generator of the cyclic group of units of K :

```
ring r = (7,a),(x,y,z),dp; minpoly = a^2+a+3;
ring r = (7^2,a),(x,y,z),dp;
```

- the ring $R[x, y, z]$, where R denotes the field of real numbers represented by simple precision floating point numbers. This is a special case:

```
ring r = real,(x,y,z),dp;
```

- the ring $R[x, y, z]$, where R denotes the field of real numbers represented by floating point numbers of 50 valid decimal digits and the same number of digits for the rest:

```
ring r = (real,50),(x,y,z),dp;
```

- the ring $R[x, y, z]$, where R denotes the field of real numbers represented by floating point numbers of 10 valid decimal digits and with 50 digits for the rest:

```
ring r = (real,10,50),(x,y,z),dp;
```

- the ring $R(j)[x, y, z]$, where R denotes the field of real numbers represented by floating point numbers of 30 valid decimal digits and the same number for the rest. j denotes the imaginary unit.

```
ring r = (complex,30,j),(x,y,z),dp;
```

- the ring $R(i)[x, y, z]$, where R denotes the field of real numbers represented by floating point numbers of 6 valid decimal digits and the same number for the rest. i is the default for the imaginary unit.

```
ring r = complex,(x,y,z),dp;
```

- the quotient ring $Z/7[x, y, z]$ modulo the square of the maximal ideal (x, y, z) :

```
ring R = 7,(x,y,z), dp;
qring r = std(maxideal(2));
```

- the ring $Z[x, y, z]$:

```
ring R = integer,(x,y,z), dp;
```

- the ring $Z/6^3[x, y, z]$:

```
ring R = (integer, 6, 3),(x,y,z), dp;
```

- the ring $Z/100[x, y, z]$:

```
ring R = (integer, 100),(x,y,z), dp;
```

3.3.2 General syntax of a ring declaration

Rings

Syntax: `ring name = (coefficients), (names_of_ring_variables), (ordering);` or
`ring name = cring [names_of_ring_variables]`

Default: `(ZZ/32003)[x,y,z];`

Purpose: declares a ring and sets it as the current basering. The second form sets the ordering to `(dp,C)`. `cring` stands currently for `QQ` (the rationals), `ZZ` (the integers) or `(ZZ/m)` (the field (m prime and <2147483648) resp. ring of the integers modulo m).

The coefficients (for the first form) are given by one of the following:

1. a **cring** as given above
2. a non-negative `int_expression` less than 2147483648 (2^{31}).
The `int_expression` should either be 0, specifying the field of rational numbers \mathbb{Q} , or a prime number p , specifying the finite field with p elements. If it is not a prime number, `int_expression` is converted to the next lower prime number.
3. an `expression_list` of an `int_expression` and one or more names.
The `int_expression` specifies the characteristic of the coefficient field as described above. The names are used as parameters in transcendental or algebraic extensions of the coefficient field. Algebraic extensions are implemented for one parameter only. In this case, a minimal polynomial has to be defined by an assignment to `minpoly`. See Section 5.3.3 [`minpoly`], page 298.
4. an `expression_list` of an `int_expression` and a name.
The `int_expression` has to be a prime number p to the power of a positive integer n . This defines the Galois field $\text{GF}(p^n)$ with p^n elements, where p^n has to be less than or equal to 2^{15} . The given name refers to a primitive element of $\text{GF}(p^n)$ generating the multiplicative group. Due to a different internal representation, the arithmetic operations in these coefficient fields are faster than arithmetic operations in algebraic extensions as described above.
5. an `expression_list` of the name **real** and two optional `int_expressions` determining the precision in decimal digits and the size for the stabilizing rest. The default for the rest is the same size as for the representation. An exception is the name **real** without any integers. These numbers are implemented as machine floating point numbers of single precision. Note that computations over all these fields are not exact.
6. an `expression_list` of the name **complex**, two optional `int_expressions` and a name. This specifies the field of complex numbers represented by floating point numbers with a precision similar to **real**. An `expression_list` without `int_expressions` defines a precision and rest with length 6. The name of the imaginary unit is given by the last parameter. Note that computations over these fields are not exact.
7. an `expression_list` with the name **integer**. This specifies the ring of integers.
8. an `expression_list` with the name **integer** and one subsequent `int_expression`. This specifies the ring of integers modulo the given `int_expression`.
9. an `expression_list` with the name **integer** and two `int_expressions` **b** and **e**. This specifies the ring of integers modulo b^e . If $b = 2$ and $e < \text{int_bit_size}$ an optimized implementation is used.

'`names_of_ring_variables`' is a list of names or indexed names.

'`ordering`' is a list of block orderings where each block ordering is either

1. **lp**, **dp**, **Dp**, **ls**, **ds**, or **Ds** optionally followed by a size parameter in parentheses.
2. **wp**, **Wp**, **ws**, **Ws**, or **a** followed by a weight vector given as an `intvec_expression` in parentheses.
3. **M** followed by an `intmat_expression` in parentheses.
4. **c** or **C**.

For the definition of the orderings, see Section B.2 [Monomial orderings], page 760.

If one of `coefficients`, `names_of_ring_variables`, and `ordering` consists of only one entry, the parentheses around this entry may be omitted.

Quotient rings

Syntax: `qring name = ideal_expression ;`

Default: none

Purpose: declares a quotient ring as the basering modulo `ideal_expression`, and sets it as current basering.

`ideal_expression` has to be represented by a standard basis.

The most convenient way to map objects from a ring to its quotient ring and vice versa is to use the `fetch` function (see Section 5.1.38 [fetch], page 180).

SINGULAR computes in a quotient ring as long as possible with the given representative of a polynomial, say, `f`. I.e., it usually does not reduce `f` w.r.t. the quotient ideal. This is only done when necessary during standard bases computations or by an explicit reduction using the command `reduce(f, std(0))` (see Section 5.1.129 [reduce], page 246).

Operations based on standard bases (e.g. `std,groebner`, etc., `reduce`) and functions which require a standard basis (e.g. `dim,hilb`, etc.) operated with the residue classes; all others on the polynomial objects.

Example:

```
// definition and usage:
ring r=(ZZ/32003)[x,y];
poly f=x3+yx2+3y+4;
qring q=std(maxideal(2));
basing;
↳ // coefficients: ZZ/32003
↳ // number of vars : 2
↳ //      block  1 : ordering dp
↳ //      : names  x y
↳ //      block  2 : ordering C
↳ // quotient ring from ideal
↳ _[1]=y2
↳ _[2]=xy
↳ _[3]=x2
poly g=fetch(r, f);
g;
↳ x3+x2y+3y+4
reduce(g,std(0));
↳ 3y+4
// polynomial and residue class:
ring R=QQ[x,y];
qring Q=std(y);
poly p1=x;
poly p2=x+y;
// comparing polynomial objects:
p1==p2;
↳ 0
// comparing residue classes:
reduce(p1,std(0))==reduce(p2,std(0));
↳ 1
```

3.3.3 Term orderings

Any polynomial (resp. vector) in SINGULAR is ordered w.r.t. a term ordering (or, monomial ordering), which has to be specified together with the declaration of a ring. SINGULAR stores and displays a polynomial (resp. vector) w.r.t. this ordering, i.e., the greatest monomial (also called the

leading monomial) is the first one appearing in the output polynomial, and the smallest monomial is the last one.

Remark: The novice user should generally use the ordering `dp` for computations in the polynomial ring $K[x_1, \dots, x_n]$, resp. `ds` for computations in the localization $\text{Loc}_{(x)}K[x_1, \dots, x_n]$. For more details, see Appendix B [Polynomial data], page 759.

In a ring declaration, SINGULAR offers the following orderings (but see also Section B.2 [Monomial orderings], page 760):

1. Global orderings

- `lp` lexicographical ordering
- `rp` reverse lexicographical ordering, i.e. a lexicographical ordering from the right with $1 < x_1 < \dots < x_n$ (should not be used as it reverses the "natural" $x_1 > \dots > x_n$, reorder the variables instead)
- `dp` degree reverse lexicographical ordering
- `Dp` degree lexicographical ordering
- `wp(intvec_expression)`
 weighted reverse lexicographical ordering; the weight vector is expected to consist of positive integers only.
- `Wp(intvec_expression)`
 weighted lexicographical ordering; the weight vector is expected to consist of positive integers only.

Global orderings are well-orderings, i.e., $1 < x$ for each ring variable x . They are denoted by a `p` as the second character in their name.

2. Local orderings

- `ls` negative lexicographical ordering
- `rs` negative reverse lexicographical ordering, i.e. a lexicographical ordering from the right (should not be used as it reverses the "natural" $x_1 < \dots < x_n$, reorder the variables instead)
- `ds` negative degree reverse lexicographical ordering
- `Ds` negative degree lexicographical ordering
- `ws(intvec_expression)`
 (general) weighted reverse lexicographical ordering; the first element of the weight vector has to be non-zero.
- `Ws(intvec_expression)`
 (general) weighted lexicographical ordering; the first element of the weight vector has to be non-zero.

Local orderings are not well-orderings. They are denoted by an `s` as the second character in their name.

3. Matrix orderings

- `M(intmat_expression)`
 intmat_expression has to be an invertible square matrix

Using matrix orderings, SINGULAR can compute standard bases w.r.t. any monomial ordering which is compatible with the natural semi-group structure on the monomials. In practice, the

predefined global and local orderings together with the block orderings should be sufficient in most cases. These orderings are faster than their corresponding matrix orderings since evaluation of a matrix ordering is more time consuming.

4. Extra weight vector

`a(intvec_expression)`

an extra weight vector `a(intvec_expression)` may precede any monomial ordering

5. Product ordering

`(ordering [(int_expression)], ...)`

any of the above orderings and the extra weight vector may be combined to yield product or block orderings

The orderings `lp`, `dp`, `Dp`, `ls`, `ds`, `Ds` and `rp` may be followed by an `int_expression` in parentheses giving the size of the block. For the last block the size is calculated automatically. For weighted orderings, the size of the block is given by the size of the weight vector. The same holds analogously for matrix orderings.

6. Module orderings

`(ordering, ..., C)`

`(ordering, ..., c)`

sort polynomial vectors by the monomial ordering first, then by components

`(C, ordering, ...)`

`(c, ordering, ...)`

sort polynomial vectors by components first, then by the monomial ordering

Here a capital `C` sorts generators in ascending order, i.e., `gen(1) < gen(2) < ...`. A small `c` sorts in descending order, i.e., `gen(1) > gen(2) > ...`. It is not necessary to specify the module ordering explicitly since `(ordering, ..., C)` is the default.

In fact, `c` or `C` may be specified anywhere in a product ordering specification, not only at its beginning or end. All monomial block orderings preceding the component ordering have higher precedence, all monomial block orderings following after it have lower precedence.

For a mathematical description of these orderings, see Appendix B [Polynomial data], page 759.

3.3.4 Coefficient rings

SINGULAR supports coefficient ranges which are not fields, i.e. the integers Z and the finite rings Z/n for a number n . These coefficient rings were implemented in SINGULAR 3.0.5 and at the moment only limited functionality is available.

p-adic numbers

The p-adic integers Z_p are the projective limit of the finite rings Z/p^n for n to infinity. Therefore, computations in this ring can be approximated by computations in Z/p^n for large n .

3.4 Implemented algorithms

The basic algorithm in SINGULAR is a general standard basis algorithm for any monomial ordering which is compatible with the natural semi-group structure of the exponents. This includes well-orderings (Buchberger algorithm to compute a Groebner basis) and tangent cone orderings (Mora algorithm) as special cases.

Nonetheless, there are a lot of other important algorithms:

- Algorithms to compute the standard operations on ideals and modules: intersection, ideal quotient, elimination, etc.
- Different Syzygy algorithms and algorithms to compute free resolutions of modules.
- Combinatorial algorithms to compute dimensions, Hilbert series, multiplicities, etc.
- Algorithms for univariate and multivariate polynomial factorization, resultant and gcd computations.

Commands to compute standard bases

facstd	Section 5.1.34 [facstd], page 177 computes a list of Groebner bases via the Factorizing Groebner Basis Algorithm, i.e., their intersection has the same radical as the original ideal. It need not be a Groebner basis of the given ideal. The intersection of the zero-sets is the zero-set of the given ideal.
fglm	Section 5.1.39 [fglm], page 181 computes a Groebner basis provided that a reduced Groebner basis w.r.t. another ordering is given. Implements the so-called FGLM (Faugere, Gianni, Lazard, Mora) algorithm. The given ideal must be zero-dimensional.
groebner	[groebner], page 785 computes a standard resp. Groebner basis using a heuristically chosen method. This is the preferred method to compute a standard resp. Groebner bases.
mstd	Section 5.1.99 [mstd], page 223 computes a standard basis and a minimal set of generators.
std	Section 5.1.149 [std], page 266 computes a standard resp. Groebner basis.
stdfglm	[stdfglm], page 785 computes a Groebner basis in a ring with a “difficult” ordering (e.g., lexicographical) via std w.r.t. a “simple” ordering and fglm . The given ideal must be zero-dimensional.
stdhilb	[stdhilb], page 785 computes a Groebner basis in a ring with a “difficult” ordering (e.g., lexicographical) via std w.r.t. a “simple” ordering and a std computation guided by the Hilbert series.

Further processing of standard bases

The next commands require the input to be a standard basis.

degree	Section 5.1.20 [degree], page 169 computes the (Krull) dimension, codimension and the multiplicity. The result is only displayed on the screen.
dim	Section 5.1.25 [dim], page 171 computes the dimension of the ideal resp. module.
highcorner	Section 5.1.55 [highcorner], page 192 computes the smallest monomial not contained in the ideal resp. module. The ideal resp. module has to be finite dimensional as a vector space over the ground field.

hilb	Section 5.1.56 [hilb], page 193 computes the first, and resp. or, second Hilbert series of an ideal resp. module.
kbase	Section 5.1.69 [kbase], page 203 computes a vector space basis (consisting of monomials) of the quotient of a ring by an ideal resp. of a free module by a submodule. The ideal resp. module has to be finite dimensional as a vector space over the ground field and has to be represented by a standard basis w.r.t. the ring ordering.
mult	Section 5.1.100 [mult], page 224 computes the degree of the monomial ideal resp. module generated by the leading monomials of the input.
reduce	Section 5.1.129 [reduce], page 246 reduces a polynomial, vector, ideal or module to its normal form with respect to an ideal or module represented by a standard basis.
vdim	Section 5.1.166 [vdim], page 281 computes the vector space dimension of a ring (resp. free module) modulo an ideal (resp. module).

Commands to compute resolutions

res	[res], page 785 computes a free resolution of an ideal or module using a heuristically chosen method. This is the preferred method to compute free resolutions of ideals or modules.
fres	Section 5.1.48 [fres], page 186 improved version of Section 5.1.147 [sres], page 264, computes a free resolution of an ideal or module using Schreyer's method. The input has to be a standard basis.
lres	Section 5.1.83 [lres], page 212 computes a free resolution of an ideal or module with LaScala's method. The input needs to be homogeneous.
mres	Section 5.1.98 [mres], page 222 computes a minimal free resolution of an ideal or module with the Syzygy method.
sres	Section 5.1.147 [sres], page 264 computes a free resolution of an ideal or module with Schreyer's method. The input has to be a standard basis.
nres	Section 5.1.105 [nres], page 228 computes a free resolution of an ideal or module with the standard basis method.
syz	Section 5.1.154 [syz], page 275 computes the first Syzygy (i.e., the module of relations of the given generators).

Further processing of resolutions

betti	Section 5.1.4 [betti], page 157 computes the graded Betti numbers of a module from a free resolution.
minres	Section 5.1.93 [minres], page 220 minimizes a free resolution of an ideal or module.

regularity

Section 5.1.130 [regularity], page 247

computes the regularity of a homogeneous ideal resp. module from a given minimal free resolution.

Processing of polynomials**char_series**

Section 5.1.6 [char_series], page 159

computes characteristic sets of polynomial ideals.

extgcd

Section 5.1.33 [extgcd], page 176

computes the extended gcd of two polynomials.

This is implemented as extended Euclidean Algorithm, and applicable for univariate polynomials only.

factorize

Section 5.1.36 [factorize], page 178

computes factorization of univariate and multivariate polynomials into irreducible factors.

The most basic algorithm is univariate factorization in prime characteristic. The Cantor-Zassenhaus Algorithm is used in this case. For characteristic 0, a univariate Hensel-lifting is done to lift from prime characteristic to characteristic 0. For multivariate factorization in any characteristic, the problem is reduced to the univariate case first, then a multivariate Hensel-lifting is used to lift the univariate factorization.

Factorization of polynomials over algebraic extensions is provided by factoring the norm for univariate polynomials f (the gcd of f and the factors of the norm is a factorization of f) resp. by the extended Zassenhaus algorithm for multivariate polynomials.

gcd

Section 5.1.50 [gcd], page 188

computes greatest common divisors of univariate and multivariate polynomials.

In the univariate case NTL is used. For prime characteristic, a subresultant gcd is used. In characteristic 0, the EZGCD is used, except for a special case where a modular algorithm is used.

resultant

Section 5.1.134 [resultant], page 250

computes the resultant of two univariate polynomials using the subresultant algorithm.

Multivariate polynomials are considered as univariate polynomials in the main variable (which has to be specified by the user).

vandermonde

Section 5.1.162 [vandermonde], page 279

interpolates a polynomial from its values at several points

Matrix computations**bareiss**

Section 5.1.3 [bareiss], page 156

implements sparse Gauss-Bareiss method for elimination (matrix triangularization) in arbitrary integral domains.

det

Section 5.1.23 [det], page 170

computes the determinant of a square matrix.

For matrices with integer entries a modular algorithm is used. For other domains the Gauss-Bareiss method is used.

minor Section 5.1.92 [minor], page 218
computes all minors (=subdeterminants) of a given size for a matrix.

Numeric computations

laguerre Section 5.1.74 [laguerre], page 205
computes all (complex) roots of a univariate polynomial

uressolve
Section 5.1.161 [uressolve], page 278
finds all roots of a 0-dimensional ideal with multivariate resultants

Controlling computations

option Section 5.1.110 [option], page 230
allows setting of options for manipulating the behaviour of computations (such as reduction strategies) and for showing protocol information indicating the progress of a computation.

3.5 The SINGULAR language

SINGULAR interprets commands given interactively on the command line as well as given in the context of user-defined procedures. In fact, SINGULAR makes no distinction between these two cases. Thus, SINGULAR offers a powerful programming language as well as an easy-to-use command line interface without differences in syntax or semantics.

In the following, the basic language concepts such as commands, expressions, names, objects, etc., are discussed. See Section 3.7 [Procedures], page 50, and Section 3.8 [Libraries], page 54, for the concepts of procedures and libraries.

In many aspects, the SINGULAR language is similar to the C programming language. For a description of some of the subtle differences, see Section 6.3 [Major differences to the C programming language], page 304.

Elements of the language

The major building blocks of the SINGULAR language are expressions, commands, and control structures. The notion of expressions in the SINGULAR and the C programming language are identical, whereas the notion of commands and control structures only roughly corresponds to C statements.

- An “expression” is a sequence of operators, functions, and operands that specifies a computation. An expression always results in a value of a specific type. See Chapter 4 [Data types], page 72, and its subsections (e.g., Section 4.16.2 [poly expressions], page 118), for information on how to build expressions.
- A “command” is either a declaration, an assignment, a call to a function without return value, or a print command. For detailed information, see Section 3.5.1 [General command syntax], page 41.
- “Control structures” determine the execution sequence of commands. SINGULAR provides control structures for conditional execution (`if ... else`) and iteration (`for` and `while`). Commands may be grouped in pairs of `{ }` (curly brackets) to form blocks. See Section 5.2 [Control structures], page 285, for more information.

Other notational conventions

For user-defined functions, the notions of “procedure” and “function” are synonymous.

As already mentioned above, functions without return values are called commands. Furthermore, whenever convenient, the term “command” is used for a function, even if it does return a value.

3.5.1 General command syntax

In SINGULAR a command is either a declaration, an assignment, a call to a function without return value, or a print command. The general form of a command is described in the following subsections.

Declaration

1. `type name = expression ;`
declares a variable with the given name of the given type and assigns the expression as initial value to it. Expression is an expression of the specified type or one that can be converted to that type. See Section 3.5.5 [Type conversion and casting], page 46.
2. `alias type name`
Introduces name as an alternative, read-only name for another variable_name. Can only be used in procedure headings to avoid copying large data.
3. `type name_list = expression_list ;`
declares variables with the given names and assigns successively each expression of expression_list to the corresponding name of name_list. Both lists must be of the same length. Each expression in expression_list is an expression of the specified type or one that can be converted to that type. See Section 3.5.5 [Type conversion and casting], page 46.
4. `type name ;`
declares a variable with the given name of the given type and assigns the default value of the specific type to it.

See Section 3.5.3 [Names], page 44, for more information on declarations. See Chapter 4 [Data types], page 72, for a description of all data types known to SINGULAR.

```
ring r;                // the default ring
poly f,g = x^2+y^3,xy+z2; // the polynomials f=x^2+y^3 and g=x*y+z^2
ideal I = f,g;         // the ideal generated by f and g
matrix m[3][3];       // a 3 x 3 zero matrix
int i=2;              // the integer i=2
```

Assignment

4. `name = expression ;`
assigns expression to name.
5. `name_list = expression_list ;`
assigns successively each expression of expression_list to the corresponding name of name_list. Both lists must be of the same length. This is not a simultaneous assignment. Thus, `f, g = g, f;` does not swap the values of `f` and `g`, but rather assigns `g` to both `f` and `g`.

A type conversion of the type of expression to the type of name must be possible. See Section 3.5.5 [Type conversion and casting], page 46.

An assignment itself does not yield a value. Hence, compound assignments like `i = j = k;` are not allowed and result in an error.

```
f = x^2 + y^2 ;      // overrides the old value of f
I = jacob(f);
f,g = I[1],x^2+y^2 ; // overrides the old values of f and g
```

Function without return value

6. `function_name [(argument_list)] ;`
 calls function `function_name` with arguments `argument_list`.

The function may have output (not to be confused with a return value of type string). See Section 5.1 [Functions], page 154. Functions without a return value are specified there to have a return type 'none'.

Some of these functions have to be called without parentheses, e.g., `help`, `LIB`.

```
ring r;
ideal i=x2+y2,x;
i=std(i);
degree(i);          // degree has no return value but prints output
↳ // dimension (proj.) = 0
↳ // degree (proj.) = 2
```

Print command

7. `expression ;`
 prints the value of an expression, for example, of a variable.

Use the function `print` (or the procedure `show` from `inout.lib`) to get a pretty output of various data types, e.g., matrix or `intmat`. See Section 5.1.119 [`print`], page 238.

```
int i=2;
i;
↳ 2
intmat m[2][2]=1,7,10,0;
print(m);
↳      1      7
↳      10     0
```

3.5.2 Special characters

The following characters and operators have special meanings:

=	assignment
{, }	parentheses for block programming
(,)	in expressions, for indexed names and for argument lists
[,]	access operator for strings, integer vectors, ideals, matrices, polynomials, resolutions, and lists. Used to build vectors of polynomials. Example: <code>s[3]</code> , <code>m[1,3]</code> , <code>i[1..3]</code> , <code>[f,g+x,0,0,1]</code> .
+	addition operator
++	increment operator
-	subtraction operator
--	decrement operator

<code>*</code>	multiplication operator
<code>/</code>	division operator. See Section 6.4 [Miscellaneous oddities], page 308, for the difference between the division operators <code>/</code> and <code>div</code> .
<code>%</code>	modulo operator (<code>mod</code> is an alias to <code>%</code>): result is always non-negative
<code>^</code> or <code>**</code>	exponentiation operator
<code>==</code>	comparison operator equal
<code>!=</code> or <code><></code>	comparison operator not equal
<code>>=</code>	comparison operator larger than or equal to
<code>></code>	comparison operator larger
<code><=</code>	comparison operator smaller than or equal to
<code><</code>	comparison operator smaller. Also used for file input. See Section 5.1.41 [filecmd], page 182.
<code>!</code>	boolean operator not
<code>&&</code>	boolean operator and
<code> </code>	boolean operator or
<code>"</code>	delimiter for string constants
<code>'</code>	delimiter for name substitution
<code>?</code>	synonym for <code>help</code>
<code>//</code>	comment delimiter. Comment extends to the end of the line.
<code>/*</code>	comment delimiter. Starts a comment which ends with <code>*/</code> .
<code>*/</code>	comment delimiter. Ends a comment which starts with <code>/*</code> .
<code>;</code>	statement separator
<code>,</code>	separator for expression lists and function arguments
<code>\</code>	escape character for <code>"</code> and <code>\</code> within strings
<code>..</code>	interval specifier returning intvec. E.g., <code>1..3</code> which is equivalent to the intvec <code>1, 2, 3</code> .
<code>:</code>	repeated entry. E.g., <code>3:5</code> generates an intvec of length 5 with constant entries 3, i.e., <code>(3, 3, 3, 3, 3)</code> .
<code>::</code>	accessor for package members. E.g., <code>MyPackage::i</code> accesses variable <code>i</code> in package <code>MyPackage</code> .
<code>_</code>	value of expression displayed last
<code>~</code>	breakpoint in procedures
<code>#</code>	list of parameters in procedures without explicit parameter list
<code>\$</code>	terminates SINGULAR

3.5.3 Names

SINGULAR is a strongly typed language. This means that all names (= identifiers) have to be declared prior to their use. For the general syntax of a declaration, see the description of declaration commands (see Section 3.5.1 [General command syntax], page 41).

See Chapter 4 [Data types], page 72, for a description of SINGULAR's data types. See Section 5.1.159 [typeof], page 277, for a short overview of possible types. To get information on a name and the object named by it, the `type` command may be used (see Section 5.1.158 [type], page 277).

It is possible to redefine an already existing name if doing so does not change its type. A redefinition first sets the variable to the default value and then computes the expression. The difference between redefining and overriding a variable is shown in the following example:

```

int i=3;
i=i+1;          // overriding
i;
↳ 4
int i=i+1;      // redefinition
↳ // ** redefining i ( int i=i+1;    // redefinition) ./examples/Names.sin\
  g:4
  i;
↳ 1

```

User defined names should start with a letter and consist of letters and digits only. As an exception to this rule, the characters `@`, and `_` may be used as part of a name, too (`@` as the first letter is reserved for purposes of library routines). Capital and small letters are distinguished. Indexed names are built as a name followed by an `int-expression` in parentheses. A list of indexed names can be built as a name followed by an `intvec-expression` in parentheses. For multi-indices, append an `int-expression` in parentheses to an indexed name. An alternative multi-index construction is `name_prefix(index_1, index_2,...)` where the `name_prefix` must be an undefined name.

```

ring R;
int n=3;
ideal j(3);
ideal j(n);    // is equivalent to the above
↳ // ** redefining j(3) ( ideal j(n);    // is equivalent to the above) .\
  /examples/Names_1.sing:4
  ideal j(2)=x;
  j(2..3);
↳ j(2)[1]=x j(3)[1]=0
  ring r=0,(x(1..2)(1..3)(1..2)),dp;
  r;
↳ // coefficients: QQ
↳ // number of vars : 12
↳ //          block  1 : ordering dp
↳ //          : names  x(1)(1)(1) x(1)(1)(2) x(1)(2)(1) x(1)(2)(2)\
  ) x(1)(3)(1) x(1)(3)(2) x(2)(1)(1) x(2)(1)(2) x(2)(2)(1) x(2)(2)(2) x(2)\
  3)(1) x(2)(3)(2)
↳ //          block  2 : ordering C
  int i(1,2),i(2,3);
  i(2,3);
↳ 0

```

Names must not coincide with reserved names (keywords). Type `reservedName()`; to get a list of the reserved names. See Section 5.1.133 [reservedName], page 249. Names should not interfere with names of ring variables or, more generally, with monomials. See Section 6.5 [Identifier resolution],

page 310.

The command `listvar` provides a list of the names in use (see Section 5.1.82 [listvar], page 210).

The most recently printed expression is available under the special name `_`, e.g.,

```

ring r;
ideal i=x2+y3,y3+z4;
std(i);
↳ _[1]=y3+x2
↳ _[2]=z4-x2
ideal k=_;
k*k+x;
↳ _[1]=y6+2x2y3+x4
↳ _[2]=y3z4+x2z4-x2y3-x4
↳ _[3]=z8-2x2z4+x4
↳ _[4]=x
size(_[3]);
↳ 3

```

A string expression enclosed in ‘...’ (back ticks) evaluates to the value of the variable given by the string expression. This feature is referred to as name substitution.

```

int foo(1)=42;
string bar="foo";
‘bar+(1)’;
↳ 42

```

3.5.4 Objects

Every object in SINGULAR has a type and a value. In most cases it has also a name and in some cases an attribute list. The value of an object may be examined simply by printing it with a `print` command: `object;`. The type of an object may be determined by means of the `typeof` function, the attributes by means of the `attrib` function (Section 5.1.159 [typeof], page 277, Section 5.1.2 [attrib], page 154):

```

ring r=0,x,dp;
typeof(10);
↳ int
typeof(10000000000000000);
↳ bigint
typeof(r);
↳ ring
attrib(x);
↳ no attributes
attrib(std(ideal(x)));
↳ attr:isSB, type int

```

Each object of type `poly`, `ideal`, `vector`, `module`, `map`, `matrix`, `number`, or `resolution` belongs to a specific ring. This is also true for `list`, if at least one of the objects contained in the list belongs to a ring. These objects are local to the ring. Their names can be duplicated for other objects in other rings. Objects from one ring can be mapped to another ring using maps or the commands `fetch` or `imap`. See Section 4.11 [map], page 103, Section 5.1.38 [fetch], page 180, Section 5.1.59 [imap], page 195.

All other types do not belong to a ring and can be accessed within every ring and across rings. They can be declared even if there is no active basering.

3.5.5 Type conversion and casting

Type conversion

Assignments convert the type of the right-hand side to the type of the left-hand side of the assignment, if possible. Operators and functions which require certain types of operands can also implicitly convert the type of an expression. It is, for example, possible to multiply a polynomial by an integer because the integer is automatically converted to a polynomial. Type conversions do not act transitively. Possible conversions are:

1.	<code>intvec</code>	\mapsto <code>intmat</code>
2.	<code>poly</code>	\mapsto <code>ideal</code>
3.	<code>bigint</code>	\mapsto <code>ideal</code>
4.	<code>int</code>	\mapsto <code>ideal</code>
5.	<code>intmat</code>	\mapsto <code>matrix</code>
6.	<code>ideal</code>	\mapsto <code>matrix</code>
7.	<code>module</code>	\mapsto <code>matrix</code>
8.	<code>number</code>	\mapsto <code>matrix</code>
9.	<code>poly</code>	\mapsto <code>matrix</code>
10.	<code>vector</code>	\mapsto <code>matrix</code>
11.	<code>bigint</code>	\mapsto <code>matrix</code>
12.	<code>int</code>	\mapsto <code>matrix</code>
13.	<code>intvec</code>	\mapsto <code>matrix</code>
14.	<code>ideal</code>	\mapsto <code>module</code>
15.	<code>matrix</code>	\mapsto <code>module</code>
16.	<code>vector</code>	\mapsto <code>module</code>
17.	<code>bigint</code>	\mapsto <code>number</code>
18.	<code>int</code>	\mapsto <code>number</code>
19.	<code>number</code>	\mapsto <code>poly</code>
20.	<code>bigint</code>	\mapsto <code>poly</code>
21.	<code>int</code>	\mapsto <code>poly</code>
22.	<code>list</code>	\mapsto <code>resolution</code>
23.	<code>poly</code>	\mapsto <code>vector</code> (<code>p</code> \mapsto <code>p*gen(1)</code>)
24.	<code>bigint</code>	\mapsto <code>vector</code>
25.	<code>int</code>	\mapsto <code>vector</code> (<code>i</code> \mapsto <code>i*gen(1)</code>)
26.	<code>int</code>	\mapsto <code>bigint</code>
27.	<code>int</code>	\mapsto <code>intvec</code>
28.	<code>string</code>	\mapsto <code>link</code>
29.	<code>resolution</code>	\mapsto <code>list</code>

Type casting

An expression can be casted to another type by using a type cast expression:
`type (expression)`.

Possible type casts are:

<code>to</code>	<code>from</code>
<code>bigint</code>	expression <code>int</code> , <code>number</code> , <code>poly</code>
<code>ideal</code>	expression lists of <code>int</code> , <code>number</code> , <code>poly</code>
<code>ideal</code>	<code>int</code> , <code>matrix</code> , <code>module</code> , <code>number</code> , <code>poly</code> , <code>vector</code>
<code>int</code>	<code>number</code> , <code>poly</code>
<code>intvec</code>	expression lists of <code>int</code> , <code>intmat</code>

<code>intmat</code>	<code>intvec</code> (see Section 4.7.3 [intmat type cast], page 89)
<code>list</code>	expression lists of any type
<code>matrix</code>	<code>module</code> , <code>ideal</code> , <code>vector</code> , <code>matrix</code> . There are two forms to convert something to a matrix: if <code>matrix(expression)</code> is used then the size of the matrix is determined by the size of expression. But <code>matrix(expression , m , n)</code> may also be used - the result is a $m \times n$ matrix (see Section 4.12.3 [matrix type cast], page 107)
<code>module</code>	expression lists of <code>int</code> , <code>number</code> , <code>poly</code> , <code>vector</code>
<code>module</code>	<code>ideal</code> , <code>matrix</code> , <code>vector</code>
<code>number</code>	<code>poly</code>
<code>poly</code>	<code>int</code> , <code>number</code>
<code>ring</code>	<code>list</code> (the inverse of <code>ringlist</code>)
<code>string</code>	any type (see Section 4.21.3 [string type cast], page 128)

Example:

```

ring r=0,x,(c,dp);
number(3x);
↪ 0
number(poly(3));
↪ 3
ideal i=1,2,3,4,5,6;
print(matrix(i));
↪ 1,2,3,4,5,6
print(matrix(i,3,2));
↪ 1,2,
↪ 3,4,
↪ 5,6
vector v=[1,2];
print(matrix(v));
↪ 1,
↪ 2
module(matrix(i,3,2));
↪ _[1]=[1,3,5]
↪ _[2]=[2,4,6]
// generators are columns of a matrix

```

3.5.6 Flow control

A block is a sequence of commands surrounded by `{` and `}`.

```

{
  command;
  ...
}

```

Blocks are used whenever SINGULAR is used as a structured programming language. The `if` and `else` structures allow conditional execution of blocks (see Section 5.2.9 [if], page 291, Section 5.2.5 [else], page 287). `for` and `while` loops are available for a repeated execution of blocks (see Section 5.2.8 [for], page 290, Section 5.2.15 [while], page 296). In procedure definitions, the main part and the example section are blocks as well (see Section 4.17 [proc], page 121).

3.6 Input and output

SINGULAR's input and output (short, I/O) are realized using links. Links are the communication channels of SINGULAR, i.e., something SINGULAR can write to and read from. In this section, a short overview of the usage of links and of the different link types is given.

For loading of libraries, see Section 5.1.79 [LIB], page 208. For executing program scripts, see Section 5.1.41 [filecmd], page 182.

Monitoring

A special form of I/O is monitoring. When monitoring is enabled, SINGULAR makes a typescript of everything printed on your terminal to a file. This is useful to create a protocol of a SINGULAR session. The `monitor` command enables and disables this feature (see Section 5.1.95 [monitor], page 221).

How to use links

Recall that links are the communication channels of SINGULAR, i.e., something SINGULAR can write to and read from using the functions `write` and `read`. There are furthermore the functions `dump` and `getdump` which store resp. retrieve the content of an entire SINGULAR session to, resp. from, a link. The `dump` and `getdump` commands are not available for DBM links.

For more information, see Section 5.1.172 [write], page 284, Section 5.1.128 [read], page 245, Section 5.1.27 [dump], page 173, Section 5.1.52 [getdump], page 189.

Example:

```

ring r; poly p = x+y;
dump(":w test.sv"); // dump the session to the file test.sv
kill r;           // kill the basering
listvar();       // no output after killing the ring
getdump(":r test.sv");// read the dump from the file
listvar();
⇒ // r           [0] *ring
⇒ //      p     [0] poly

```

Specifying a link can be as easy as specifying a filename as a string. Links do not even need to be explicitly opened or closed before, resp. after, they are used. To explicitly open or close a link, the `open`, resp. `close`, commands may be used (see Section 5.1.109 [open], page 230, Section 5.1.10 [close], page 161).

Links have various properties which can be queried using the `status` function (see Section 5.1.148 [status], page 265).

Example:

```

link l = "ssi:fork";
l;
⇒ // type : ssi
⇒ // mode : fork
⇒ // name :
⇒ // open : no
⇒ // read : not open
⇒ // write: not open
open(l);
status(l, "open");
⇒ yes

```

```

    close(l);
    status(l, "open");
    ↪ no

```

ASCII links

Data that can be converted to a string can be written into files for storage or communication with other programs. The data are written in plain ASCII format. Reading from an ASCII link returns a string — conversion into other data is up to the user. This can be done, for example, using the command `execute` (see Section 5.1.32 [execute], page 176).

ASCII links should primarily be used for storing small amounts of data, especially if it might become necessary to manually inspect or manipulate the data.

See Section 4.9.4 [ASCII links], page 95, for more information.

Example:

```

    // (over)write file test.ascii, link is specified as string
    write(":w test.ascii", "int i =", 3, ";");
    // reading simply returns the string
    read("test.ascii");
    ↪ int i =
    ↪ 3
    ↪ ;
    ↪
    // but now test.ascii is "executed"
    execute(read("test.ascii"));
    i;
    ↪ 3

```

Ssi links

Data is communicated with other processes (e.g., SINGULAR processes) which may run on the same computer or on different ones. Data exchange is accomplished using TCP/IP links in the ssi format. Reading from an ssi link returns the written expressions (i.e., not a string, in general).

Ssi links should primarily be used for communicating with other programs or for parallel computations (see, for example, Section A.1.8 [Parallelization with ssi links], page 700).

See Section 4.9.5 [Ssi links], page 96, for more information.

Example:

```

    ring r;
    link l = "ssi:tcp localhost:"+system("Singular"); // declare a link explicitly
    open(l); // needs an open, launches another SINGULAR as a server
    write(l, x+y);
    kill r;
    def p = read(l);
    typeof(p); p;
    ↪ poly
    ↪ x+y
    close(l); // shuts down SINGULAR server

```

DBM links

Data is stored in and accessed from a data base. Writing is accomplished by a key and a value and associates the value with the key in the specified data base. Reading is accomplished w.r.t. a key,

the value associated to it is returned. Both the key and the value have to be specified as strings. Hence, DBM links may be used only for data which may be converted to or from strings.

DBM links should primarily be used when data needs to be accessed not in a sequential way (like with files) but in an associative way (like with data bases).

See Section 4.9.7 [DBM links], page 99, for more information.

Example:

```
ring r;
// associate "x+y" with "mykey"
write("DBM:w test.dbm", "mykey", string(x+y));
// get from data base what is stored under "mykey"
execute(read("DBM: test.dbm", "mykey"));
↳ x+y
```

3.7 Procedures

Procedures contain sequences of commands in the SINGULAR language. They are used to extend the set of commands by user defined commands. In a SINGULAR session, procedures are defined by either typing them on the command line or by loading them from a library file with the LIB or load command (see Section 3.8 [Libraries], page 54). A procedure is invoked like normal built-in commands, i.e., by typing its name followed by the list of arguments in parentheses. The invocation then executes the sequence of commands constituting the procedure. All procedures defined in a SINGULAR session can be displayed by entering `listvar(proc);`.

See also See Section 3.8.6 [Procedures in a library], page 57.

3.7.1 Procedure definition

Syntax:

```
[static] proc proc_name [(

```

Purpose:

- Defines a new function, the `proc proc_name`.
- The help string, the parameter list, and the example section are optional. They are, however, mandatory for the procedures listed in the header of a library. The help string is ignored and no example section is allowed if the procedure is defined interactively, i.e., if it is not loaded from a file by the LIB or load command (see Section 5.1.79 [LIB], page 208 and see Section 5.2.12 [load], page 294).
- Once loaded from a file into a SINGULAR session, the information provided in the help string will be displayed upon entering `help proc_name;`, while the example section will be executed upon entering `example proc_name;`. See Section 3.7.2 [Parameter list], page 52, Section 3.7.3 [Help string], page 53, and the example in Section 3.8.6 [Procedures in a library], page 57.
- In the body of a library, each procedure not meant to be accessible by users should be declared static. See Section 3.8.6 [Procedures in a library], page 57.

Example of an interactive procedure definition and its execution:

```

proc milnor_number (poly p)
{
  ideal i= std(jacob(p));
  int m_nr=vdim(i);
  if (m_nr<0)
  {
    "// not an isolated singularity";
  }
  return(m_nr);          // the value of m_nr is returned
}
ring r1=0,(x,y,z),ds;
poly p=x^2+y^2+z^5;
milnor_number(p);
↳ 4

```

Example of a procedure definition in a library:

First, we define the library (and store it as `sample.lib`):

```

// Example of a user accessible procedure
proc tab (int n)
"USAGE:   tab(n);  n integer
RETURNS:  string of n space tabs
EXAMPLE:  example tab; shows an example"
{ return(internal_tab(n)); }
example
{
  "EXAMPLE:"; echo=2;
  for(int n=0; n<=4; n=n+1)
  { tab(4-n)+"*"+tab(n)+" "+tab(n)+"*"; }
}

// Example of a static procedure
static proc internal_tab (int n)
{ return(" "[1,n]); }

```

Now, we load the library and execute its procedures:

```

LIB "sample.lib";          // load the library sample.lib
example tab;              // show an example
↳ // proc tab from lib sample.lib
↳ EXAMPLE:
↳   for(int n=0; n<=4; n=n+1)
↳   { tab(4-n)+"*"+tab(n)+" "+tab(n)+"*"; }
↳     ***
↳     * + *
↳     * + *
↳     * + *
↳     *   +   *
↳
↳     "*" + tab(3) + "*";          // use the procedure tab
↳ *   *
// the static procedure internal_tab is not accessible
" "+internal_tab(3) + "*";

```

```

↳    ? 'internal_tab(3)' is not defined
↳    ? error occurred in or before ./examples/Example_of_a_procedure_defini\
      tion_in_a_library:.sing line 5: '  "*" + internal_tab(3) + "*" ;'
      // show the help section for tab
      help tab;
↳ // ** Could not get 'IdxFile'.
↳ // ** Either set environment variable 'SINGULAR_IDX_FILE' to 'IdxFile',
↳ // ** or make sure that 'IdxFile' is at "%D/singular/singular.idx"
↳ // ** Displaying help in browser 'dummy'.
↳ // ** Use 'system("--browser", <browser>);' to change browser,
↳ // ** where <browser> can be: "dummy", "emacs".
↳    ? No functioning help browser available.
↳    ? error occurred in or before ./examples/Example_of_a_procedure_defini\
      tion_in_a_library:.sing line 7: '  help tab;'
```

3.7.2 Parameter list

Syntax: ()
 (parameter_definition)

Purpose:

- Defines the number, type and names of the arguments of a procedure.
- The parameter_list is optional.
- Adding list # as argument to a parameter list means to allow optional parameters. Furthermore, (list #) is the default for a parameter list (in case no list is explicitly given). Inside the procedure body, the arguments of list # are referenced by #[1], #[2], etc.
- If a procedure has optional parameters, the attribute default_arg gives the default values for the optional arguments. This provides in particular the possibility to also change the behaviour of all procedures nested inside the given procedure.

Example:

```

proc x0
{
    // can be called with
    ... // any number of arguments of any type: #[1], #[2], ...
        // number of arguments: size(#)
}

proc x1 ()
{
    ... // can only be called without arguments
}

proc x2 (ideal i, int j)
{
    ... // can only be called with 2 arguments,
        // which can be converted to ideal resp. int
}

proc x3 (i,j)
{
```

```

... // can only be called with 2 arguments
    // of any type
    // (i,j) is the same as (def i,def j)
}

proc x5 (i,list #)
{
... // can only be called with at least 1 argument
    // number of arguments: size(#)+1
}

attrib(x5,"default_arg",3);
x5(2); // is equivalent to
x5(2,3);

```

Note:

The parameter `list` may stretch across multiple lines.

A parameter may have any type (including the types `proc` and `ring`).

If a parameter is of type `ring`, then it can only be specified by name, but not with a type. For instance:

```

proc x6 (r)
{
... // this is correct, r may be of any type, even of type ring
}

proc x7 (ring r)
{
... // this is NOT CORRECT
}

```

3.7.3 Help string

Syntax: `string_constant`;

Purpose: Constitutes the help text of a procedure.

Format:

```

USAGE:    <proc_name>( <parameter list>);  <explanation of parameters>
ASSUME:   <description of assumptions made>
RETURN:   <description of what is returned>
SIDE EFFECTS: <description of global objects generated or manipulated,
but not returned>
REMARKS:  <information on theory and implemented algorithms,references>
NOTE:     <particularities, limitations, additional details>
KEYWORDS: <semicolon-separated phrases of index keys>
SEE ALSO: <comma-separated names of related procedures/cross references>
EXAMPLE:  example <proc_name>; shows an example

```

NOTE:

- ASSUME, SIDE EFFECTS, KEYWORDS, and SEE ALSO are optional. No help string is required for static procedures.
- EXAMPLE: refers to the example section of the procedure. In a SINGULAR session, the example will be carried out upon entering `example <proc_name>`; if the procedure is loaded from a file by the LIB or load command (see Section 5.1.79

[LIB], page 208 and see Section 5.2.12 [load], page 294). No example section is allowed if the procedure is defined interactively.

- See Section 3.8.10 [Typesetting of help and info strings], page 63 for help strings in the SINGULAR documentation.
- See the example in Section 3.8.6 [Procedures in a library], page 57 for an illustration.

3.7.4 Names in procedures

- All variables defined inside a procedure are local to the procedure and their names cannot interfere with names in other procedures. Without further action, they are automatically deleted after leaving the procedure.
- To keep local variables and their value after leaving the procedure, they have to be exported (i.e. made global) by a command like `export` or `exportto` (see Section 5.2.6 [export], page 287, see Section 5.2.7 [exportto], page 288, see Section 5.2.10 [importfrom], page 291; see Section 4.15 [package], page 117). To return the value of a local variable, use the `return` command (see Section 5.2.14 [return], page 295).

Example:

```

proc xxx
{
  int k=4;          //defines a local variable k
  int result=k+2;
  export(result);  //defines the global variable "result".
}
xxx();
listvar(all);
⇒ // result                [0] int 6

```

Note that the variable `result` became a global variable after the execution of `xxx`.

3.7.5 Procedure-specific commands

A few commands should only be used inside a procedure. They either make local objects global ones or return results to the level from where the procedure was called.

See Section 5.2.6 [export], page 287; Section 5.2.7 [exportto], page 288; Section 5.2.14 [return], page 295.

3.8 Libraries

- A library is a collection of SINGULAR procedures in a file.
- To load a library into a SINGULAR session, use the `LIB` or `load` command. Having loaded a library, its procedures can be used like any built-in SINGULAR function, and information on the library is obtained by entering `help libname.lib`;
- See Appendix D [SINGULAR libraries], page 785, for all libraries currently distributed with SINGULAR.
- When writing your own library, it is important to comply with the guidelines described in this section. Otherwise, due to potential parser errors, it may not be possible to load the library.
- Each library consists of a header and a body. The first line of a library must start with a double slash `//`.

- The library header consists of a version string, a category string, an info string, and LIB commands. The strings are mandatory. LIB commands are meant to load the additional libraries used by the library under consideration.
- The library body collects the procedures (declared static or not).
- No line of a library should consist of more than 60 characters.

3.8.1 Libraries in the SINGULAR Documentation

- The typesetting language in which the SINGULAR documentation is written is `texinfo`. The info string of a library included in the SINGULAR distribution will be parsed and automatically translated to the `texinfo` format. The same applies to the help string of each procedure listed in the PROCEDURE: section of the info string.
- Based on various tools, `info`, `dvi`, `p`, and `html` versions of the `texinfo` documentation are generated.
- For `texinfo` markup elements and other information facilitating optimal typesetting, see Section 3.8.10 [Typesetting of help and info strings], page 63.
- For the convenience of users checking directly the source code, the `texinfo` tools should be used economically. That is, the info and help texts should be well readable verbatim.
- The example of each procedure listed in the PROCEDURE: section of the info string is computed and its output is included in the documentation.

3.8.2 Version string

A version string is part of the header of a library.

Syntax: `version = string_constant;`

Purpose: Defines the version number of a library. It is displayed when the library is loaded.

Example: `version="version sample.lib 4.0.0.0 Dec_2013 ";`

Note: Syntax: `version<space><filename><space><version><space><date><space>`

3.8.3 Category string

A category string is part of the header of a library.

Syntax: `category = string_constant;`

Purpose: Defines the category of a library.

Example: `category="Algebraic geometry";`

Note: Reserved for sorting the libraries into categories.

3.8.4 Info string

Syntax: `info = string_constant;`

Purpose: Constitutes the help text of a library. Will be displayed in a SINGULAR session upon entering `help libname.lib; .` Will be part of the SINGULAR documentation if the library is distributed with SINGULAR. See Section 3.8.1 [Libraries in the SINGULAR Documentation], page 55.

Format:

```

info="
LIBRARY: <library_name> <one line description of the purpose>
AUTHOR: <name, and email address of author>
OVERVIEW: <concise, additional information on what is implemented>
REFERENCES: <references for further information>
KEYWORDS: <semicolon-separated phrases of index keys>
SEE ALSO: <comma-separated words of cross references>
PROCEDURES:
  <proc_name_1>();      <one line description of the purpose>
  .
  .
  <proc_name_N>();      <one line description of the purpose>
";

```

NOTE:

- In the documentation, the one line description of the purpose following LIBRARY: will be printed in its own line, starting with the prefix PURPOSE: .
- REFERENCES, KEYWORDS, and SEE ALSO are optional.
- Only non-static procedures should be listed in the PROCEDURES: section. A procedure parameter should be included between the brackets () only if the corresponding one line description of the purpose refers to it. See Section 3.8.6 [Procedures in a library], page 57.
- In the documentation, separate nodes (subsections in printed documents) are created precisely for those procedures of the library appearing in the PROCEDURES: section (that is, for some if not all non-static procedures of the library).

Example:

```

info="
LIBRARY: absfact.lib  Absolute factorization for characteristic 0
AUTHORS: Wolfram Decker,      decker at math.uni-sb.de
         Gregoire Lecerf,     lecerf at math.uvsq.fr
         Gerhard Pfister,     pfister at mathematik.uni-kl.de

OVERVIEW:
A library for computing the absolute factorization of multivariate
polynomials f with coefficients in a field K of characteristic zero.
Using Trager's idea, the implemented algorithm computes an absolutely
irreducible factor by factorizing over some finite extension field L
(which is chosen such that V(f) has a smooth point with coordinates in L).
Then a minimal extension field is determined making use of the
Rothstein-Trager partial fraction decomposition algorithm.

REFERENCES:
G. Cheze, G. Lecerf: Lifting and recombination techniques for absolute
factorization. Journal of Complexity, 23(3):380-420, 2007

KEYWORDS: factorization; absolute factorization.
SEE ALSO: factorize

PROCEDURES:
  absFactorize();      absolute factorization of poly
";

```

To see how this infostring appears in the documentation after typesetting, check Section D.4.1 [absfact_lib], page 809:

3.8.5 LIB commands

LIB commands are part of the header of a library.

Syntax: LIB "lib_1.lib";
 ...
 LIB "lib_r.lib";

Purpose: Loads libraries used by the library under consideration.

Example:

```
LIB "primdec.lib";
LIB "normal.lib";
```

Note: The keyword LIB must be followed by at least one space.

3.8.6 Procedures in a library

Here are hints and requirements on how procedures contained in a library should be implemented. For more on procedures, see Section 3.7 [Procedures], page 50.

1. Each procedure not meant to be accessible by users should be declared static.
2. The header of each procedure not declared static must comply with the guidelines described in Section 3.7.1 [Procedure definition], page 50 and Section 3.7.3 [Help string], page 53. In particular, it must have a help and example section, and assumptions made should be carefully explained. If the assumptions are checked by the procedure on run-time, errors may be reported using the Section 5.1.30 [ERROR], page 175 function.
3. Names of procedures should not be shorter than 4 characters and should not contain any special characters. In particular, the use of `_` in names of procedures is discouraged. If the name of the procedure is composed of more than one word, each new word should start with a capital letter, all other letters should be lower case (e.g. `linearMapKernel`).
4. No procedures should be defined within the body of another procedure.
5. A procedure may print out comments, for instance to explain results or to display intermediate computations. This is often helpful when calling the procedure directly, but it may also cause confusions in cases where the procedure is called by another procedure. The SINGULAR solution to this problem makes use of the function `dbprint` (see Section 5.1.17 [dbprint], page 167) and the reserved variables `printlevel` and `voice` (see Section 5.3.6 [printlevel], page 299 and see Section 5.3.11 [voice], page 303). Note that `printlevel` is a predefined, global variable whose value can be changed by the user, while `voice` is an internal variable, representing the nesting level of procedures. Accordingly, the value of Section 5.3.11 [voice], page 303 is 1 on the top level, 2 inside the first procedure, and so on. The default value of `printlevel` is 0, but `printlevel` can be set to any integer value by the user.

Example: If the procedure `Test` below is called directly from the top level, then ‘comment1’ is displayed, but not ‘comment2’. By default, nothing is displayed if `Test` is called from within any other procedure. However, if `printlevel` is set to a value `k` with `k>0`, then ‘comment1’ (resp. ‘comment2’) is displayed – provided `Test` is called from another procedure with nesting level at most `k` (resp. `k-1`).

The example part of a procedure behaves in this respect like the procedure on top level (the nesting level is 1, that is, the value of `voice` is 2). Therefore, due to the command

`printlevel=1;`, ‘comment1’ will be displayed when entering `example Test;`. However, since `printlevel` is a global variable, it should be reset to its old value at the end of the example part.

The predefined variable `echo` controls whether input lines are echoed or not. Its default is 0, but it can be reset by the user. Input is echoed if `echo>=voice`. At the beginning of the example part, `echo` is set to the value 2. In this way, the input lines of the example will be displayed when entering `example Test;`.

```

proc Test
"USAGE:  ...
      ...
EXAMPLE: example Test; shows an example
"
{  ...
  int p = printlevel - voice + 3;
  ...
  dbprint(p,"comment1");
  dbprint(p-1,"comment2");
  // dbprint prints only if p > 0
  ...
}
example
{ "EXAMPLE:"; echo = 2;
  int p = printlevel; //store old value of printlevel
  printlevel = 1;    //assign new value to printlevel
  ...
  Test();
  printlevel = p;    //reset printlevel to old value
}

```

Note: SINGULAR functions such as `pause` or `read` allow and require interactive user-input. They are, thus, in particular useful for debugging purposes. If such a command is used inside the procedure of a library to be distributed with SINGULAR, the example section of the procedure has to be written with some care – the procedure should only be called from within the example if the value of `printlevel` is 0. Otherwise, the automatic build process of SINGULAR will not run through since the examples are carried out during the build process. They are, thus, tested against changes in the code.

3.8.7 `template.lib`

First, we show the source-code of a template library:

```

////////////////////////////////////
version="version template.lib 4.1.2.0 Feb_2019 "; // $Id: 4d4a314bcbeaaaf113c4c4687bl
category="Miscellaneous";
// summary description of the library
info="
LIBRARY:  template.lib  A Template for a Singular Library
AUTHOR:   Olaf Bachmann, email: obachman@mathematik.uni-kl.de

SEE ALSO: standard_lib, Libraries,
          Typesetting of help and info strings

KEYWORDS: library, template.lib; template.lib; library, info string

```

```

PROCEDURES:
  mdouble(int)          return double of int argument
  mtriple(int)         return three times int argument
  msum([int,..,int])   sum of int arguments
";
/////////////////////////////////////////////////////////////////
proc mdouble(int i)
"USAGE:   mdouble(i); i int
RETURN:   int: i+i
NOTE:     Help string is in pure ASCII.
          This line starts on a new line since previous line is short.
          No new line here.
SEE ALSO: msum, mtriple, Typesetting of help and info strings
KEYWORDS: procedure, ASCII help
EXAMPLE:  example mdouble; shows an example"
{
  return (i + i);
}
example
{ "EXAMPLE:"; echo = 2;
  mdouble(0);
  mdouble(-1);
}
/////////////////////////////////////////////////////////////////
proc mtriple(int i)
"@c we do texinfo here
@table @asis
@item @strong{Usage:}
@code{mtriple(i)}; @code{i} int

@item @strong{Return:}
int: @math{i+i+i}
@item @strong{Note:}
Help is in pure Texinfo.
@*This help string is written in texinfo, which enables you to use,
among others, the @math command for mathematical typesetting
(for instance, to print @math{\alpha, \beta}).
@*Texinfo also gives more control over the layout, but is, admittedly,
more cumbersome to write.
@end table
@c use @c ref contstuct for references
@cindex procedure, texinfo help
@c ref
@strong{See also:}
@ref{mdouble}, @ref{msum}, @ref{Typesetting of help and info strings}
@c ref
"
{
  return (i + i + i);
}
example
{ "EXAMPLE:"; echo = 2;

```

```

    mtriple(0);
    mtriple(-1);
}
/////////////////////////////////////////////////////////////////
proc msum(list #)
"USAGE:  msum([i_1,..,i_n]); @code{i_1,..,i_n} def
RETURN:  Sum of int arguments
NOTE:    This help string is written in a mixture of ASCII and texinfo.
        @* Use @ref for references (e.g., @pxref{mtriple}).
        @* Use @code for typewriter font (e.g., @code{i_1}).
        @* Use @math for simple math mode typesetting (e.g., @math{i_1}).
        @* Warning: Parenthesis like } are not allowed inside @math and @code.
        @* Use @example for indented, preformatted text typesetting in
        typewriter font:
@example
    this --> that
@end example
        Use @format for preformatted text typesetting in normal font:
@format
    this --> that
@end format
        Use @texinfo for text in pure texinfo:
@texinfo
@expansion{}
@tex
${i_{1,1}}$
@end tex

@end texinfo
    Note that
        automatic linebreaking is still in affect (like in this line).
SEE ALSO:  mdouble, mtriple, Typesetting of help and info strings
KEYWORDS:  procedure, ASCII/Texinfo help
EXAMPLE:   example msum; shows an example"
{
    if (size(#) == 0) { return (0);}
    if (size(#) == 1) { return ([1]);}
    int i;
    def s = #[1];
    for (i=2; i<=size(#); i++)
    {
        s = s + #[i];
    }
    return (s);
}
example
{ "EXAMPLE:"; echo = 2;
    msum();
    msum(4);
    msum(1,2,3,4);
}

```

Second, we show how the library appears in the documentation after typesetting (with one subsection for each procedure):

Library: `template.lib`

Purpose: A Template for a Singular Library

Author: Olaf Bachmann, email: obachman@mathematik.uni-kl.de

Procedures: See also: Section 3.8 [Libraries], page 54; Section 3.8.10 [Typesetting of help and info strings], page 63; Section D.1 [standard.lib], page 785.

3.8.7.1 mdouble

Procedure from library `template.lib` (see Section 3.8.7 [template.lib], page 58).

Usage: `mdouble(i); i int`

Return: int: $i+i$

Note: Help string is in pure ASCII.
This line starts on a new line since previous line is short. No new line here.

Example:

```
LIB "template.lib";
mdouble(0);
↪ 0
mdouble(-1);
↪ -2
```

See also: Section 3.8.10 [Typesetting of help and info strings], page 63; Section 3.8.7.3 [msum], page 61; Section 3.8.7.2 [mtriple], page 61.

3.8.7.2 mtriple

Procedure from library `template.lib` (see Section 3.8.7 [template.lib], page 58).

Usage: `mtriple(i); i int`

Return: int: $i + i + i$

Note: Help is in pure Texinfo.
This help string is written in texinfo, which enables you to use, among others, the `@math` command for mathematical typesetting (for instance, to print α, β).
Texinfo also gives more control over the layout, but is, admittedly, more cumbersome to write.

See also:

Example:

```
LIB "template.lib";
mtriple(0);
↪ 0
mtriple(-1);
↪ -3
```

3.8.7.3 msum

Procedure from library `template.lib` (see Section 3.8.7 [template.lib], page 58).

Usage: `msum([i_1,..,i_n]); i_1,..,i_n def`

Return: Sum of int arguments

Note: This help string is written in a mixture of ASCII and texinfo.
 Use @ref for references (e.g., see Section 3.8.7.2 [mtriple], page 61).
 Use @code for typewriter font (e.g., `i_1`).
 Use @math for simple math mode typesetting (e.g., i_1).
 Warning: Parenthesis like } are not allowed inside @math and @code.
 Use @example for indented, preformatted text typesetting in typewriter font:

```
    this --> that
```

Use @format for preformatted text typesetting in normal font:

```
    this -> that
```

Use @texinfo for text in pure texinfo:

```
    ↦  $i_{1,1}$ 
```

Note that

automatic linebreaking is still in affect (like in this line).

Example:

```
LIB "template.lib";
msum();
↦ 0
msum(4);
↦ 4
msum(1,2,3,4);
↦ 10
```

See also: Section 3.8.10 [Typesetting of help and info strings], page 63; Section 3.8.7.1 [mdouble], page 61; Section 3.8.7.2 [mtriple], page 61.

3.8.8 Formal Checker

There is a formal library checker for SINGULAR which can be used online: see <https://www.singular.uni-kl.de/index.php/new-libraries/formal-library-checker.html>.

After uploading your library file, you will receive an output of hints, warnings, and errors which may help you to improve your library.

3.8.9 Documentation Tool

lib2doc is a utility to generate the stand-alone documentation for a SINGULAR library in various formats.

The lib2doc utility should be used by developers of SINGULAR libraries to check the generation of the documentation of their libraries.

lib2doc can be downloaded from

```
ftp://www.mathematik.uni-kl.de/pub/Math/Singular/misc/lib2doc.tar.gz
```

Important:

To use lib2doc, you need to have perl (version 5 or higher), texinfo (version 3.12 or higher) and Singular and libparse (version 1-3-4 or higher) installed on your system.

To generate the documentation for a library, follow these steps:

1. Unpack lib2doc.tar.gz

```
    gzip -dc lib2doc.tar.gz | tar -pxf -
```

and


```
cd lib2doc
```

2. Edit the beginning of the file `Makefile`, filling in the values for `SINGULAR` and `LIBPARSE`. Check also the values of `PERL` and `LATEX2HTML`.

3. Copy your library to the current directory:

```
cp <path-where-your-lib-is>/mylib.lib .
```

4. Now you can run the following commands:

```
make mylib.hlp
```

Generates the file `mylib.hlp` – the info file for the documentation of `mylib.lib`.
This file can be viewed using

```
info -f mylib.hlp
```

```
make mylib.dvi
```

Generates the file `mylib.dvi` – the dvi file for the documentation of `mylib.lib`.
This file can be viewed using

```
xdvi mylib.dvi
```

```
make mylib.ps
```

Generates the file `mylib.ps` – the PostScript file for the documentation of `mylib.lib`. This file can be viewed using (for example)

```
ghostview mylib.dvi
```

```
make mylib.html
```

Generates the file `mylib.html` – the HTML file for the documentation of `mylib.lib`. This file can be viewed using (for example)

```
firefox mylib.html
```

```
make clean
```

Deletes all generated files.

Note that you can safely ignore messages complaining about undefined references.

3.8.10 Typesetting of help and info strings

The info strings of the libraries which are included in the distribution of `SINGULAR` and the help strings of the corresponding procedures are parsed and automatically converted into the `texinfo` format (the typesetting language in which the documentation of `SINGULAR` is written).

The illustrative example given in Section 3.8.7 [template.lib], page 58 should provide sufficient information on how this works. For more details, check the following items:

- Users familiar with `texinfo` may write help and info strings directly in the `texinfo` format. The string should, then, start with the `@` sign. In this case, no parsing will be done.
- Help and info strings are typeset within a `@table @asis` environment (which is similar to the `latex description` environment).
- If a line starts with uppercase words up to a colon, then the text up to the colon is taken to be the description-string of an item, and the text following the colon is taken to be the content of the item.
- If the description-string of an item matches

SEE ALSO then the content of the item is assumed to consist of comma-separated words which are valid references to other `texinfo` nodes of the manual (e.g., all procedure and command names are also `texinfo` nodes).

KEYWORDS then the content of the item is assumed to be a semicolon-separated list of phrases which are taken as keys for the index of the manual (the name of a procedure/library is automatically added to the index keys).

- If the description-string of an item in the **info string of a library** matches

LIBRARY then the content of the item is assumed to be a one-line description of the library. If this one-line description consists of uppercase characters only, then it is typeset in lowercase characters (otherwise it is left as is).

PROCEDURES

then the content of the item is assumed to consist of lines of type

```
<proc_name>();    <one line description of the purpose>
```

Separate `texinfo` nodes (subsections in printed documents) are created precisely for those procedures of the library appearing here (that is, for some if not all non-static procedures of the library).

With respect to the content of an item, the following `texinfo` markup elements are recognized:

`@*` Enforces a line-break.

Example: `old line @* new line`
 \mapsto
 old line
 new line

`@ref{...}` For references to other parts of the SINGULAR manual, use one of the following `@ref{node}` constructs. Here, `node` must be the name of a section of the SINGULAR manual. In particular, it may be the name of a function, library or procedure in a library.

`@xref{node}`

for a reference to the node `node` at the beginning of a sentence.

`@ref{node}`

for a reference to the node `node` at the end of a sentence.

`@pxref{node}`

for a reference to the node `node` within parentheses.

Example: `@xref{Hurricanes}, for more info.`

\mapsto *Note Hurricanes::, for more info.

\mapsto See Section 3.1 [Hurricanes], page 24, for more info.

For more information, see `@ref{Hurricanes}`.

\mapsto For more information, see *Note Hurricanes::.

\mapsto For more information, see Section 3.1 [Hurricanes], page 24.

... storms cause flooding (`@pxref{Hurricanes}`) ...

\mapsto ... storms cause flooding (*Note Hurricanes::) ...

\mapsto ... storms cause flooding (see Section 3.1 [Hurricanes], page 24)

`@math{...}` Typeset short mathematical expressions in LaTeX math-mode syntax (short: does not cause expansion over multiple lines).

Example: `@math{\alpha}`

\mapsto

α

Note: The mathematical expressions inside `@math{...}` must not contain the characters `{,}`, and `@`.

`@code{..}` Typeset short strings in typewriter font (short: does not cause expansion over multiple lines).

Example: `@code{typewriter font}`
 \mapsto
 typewriter font

Note: The string inside `@code{..}` must not contain the characters `{,}`, and `@`.

Typeset pre-formatted text in typewriter font.

```
@example
...
@end example
```

Example:

```
before example
@example
in example
notice escape of special characters like @{,@},@@
@end example
after example

 $\mapsto$ 
before example
in example
notice escape of special characters like {,},@
after example
```

Note: Inside an `@example` environment, the characters `{,},@` have to be escaped by an `@` sign.

Typeset pre-formatted text in normal font.

```
@format
...
@end format
```

Example:

```
before format
@format
in format
notice escape of special characters like @{,@},@@
@end format
after format

 $\mapsto$ 
before format
in format
escape of special characters like {,},@
after format
```

Note: Inside an `@format` environment, the characters `{,},@` have to be escaped by an `@` sign.

Write text in pure `texinfo`.

```
@texinfo
...
@end texinfo
```

Example:

```
@texinfo
Among others, within a texinfo environment,
one can use the tex environment to typeset
more complex mathematical items like
@tex
i{1,1} $
@tex
@end texinfo
```

↦

Among others, within a texinfo environment, one can use the tex environment to typeset more complex mathematical items like $i_{1,1}$

Furthermore, a line-break is inserted before each line whose previous line is shorter than 60 characters and does not contain any of the above described recognized texinfo markup elements.

3.8.11 Loading a library

Libraries can be loaded with the `LIB` or the `load` command (see Section 5.1.79 [`LIB`], page 208 and see Section 5.2.12 [`load`], page 294).

Syntax: `LIB string_expression ;`
`load string_expression ;`

Type: none

Purpose: Reads a library from a file. If the given filename does not start with `.` or `/` and if the file cannot be located in the current directory, the `SearchPath` is checked for a directory containing a file with this name.

Note on SearchPath:

The `SearchPath` for a library is constructed at SINGULAR start-up time as follows:

1. the directories contained in the environment variable `SINGULARPATH` are appended.
2. the directories `$BinDir/LIB`, `$RootDir/LIB`, `$RootDir/./LIB`, `$DefaultDir/LIB`, `$DefaultDir/./LIB` are appended, where
 - `$BinDir` is the value of the environment variable `SINGULAR_BIN_DIR`, if set, or, if not set, the directory in which the SINGULAR program resides
 - `$RootDir` is the value of the environment variable `SINGULAR_ROOT_DIR`, if set, or, if not set, `$BinDir/./`.
 - `$DefaultDir` is the value of the environment variable `SINGULAR_DEFAULT_DIR`, if set, or `/usr/local`.
3. all directories which do not exist are removed from the `SearchPath`.

For setting environment variables, see Section 5.1.153 [`system`], page 270, or consult the manual of your shell.

The library `SearchPath` can be examined by starting up SINGULAR with the option `-v`, or by issuing the command `system("--version");`.

Note on standard.lib:

Unless SINGULAR is started with the `--no-stdlib` option, the library `standard.lib` is automatically loaded at start-up time.

Following a `LIB` or `load` command, only the names of the procedures in the library are loaded. The body of a particular procedure is only read upon the first call of the procedure. This minimizes memory consumption by unused procedures. Starting a SINGULAR session with the `-q` or `--quiet` option unsets the option `loadLib` and inhibits, thus, the monitoring of library loading (see option).

All libraries loaded in a SINGULAR session are displayed upon entering `listvar(package);` :

```

option(loadLib); // show loading of libraries;
                  // standard.lib is loaded

listvar(package);
↳ // Singmathic           [0] package Singmathic (C,singmathic.s\
  o)
↳ // Standard             [0] package Standard (S,standard.lib)
↳ // Top                  [0] package Top (T)
                          // the names of the procedures of inout.lib
LIB "inout.lib"; // are now known to Singular
↳ // ** loaded inout.lib (4.1.2.0, Feb_2019)
listvar(package);
↳ // Inout                [0] package Inout (S,inout.lib)
↳ // Singmathic           [0] package Singmathic (C,singmathic.s\
  o)
↳ // Standard             [0] package Standard (S,standard.lib)
↳ // Top                  [0] package Top (T)

```

See Section 3.1.6 [Command line options], page 19; Section 5.1.79 [LIB], page 208; Section 2.3.3 [Procedures and libraries], page 10; Appendix D [SINGULAR libraries], page 785; Section 4.17 [proc], page 121; Section D.1 [standard.lib], page 785; Section 4.21 [string], page 127; Section 5.1.153 [system], page 270.

3.9 Debugging tools

If SINGULAR does not come back to the prompt while calling a user defined procedure, probably a bracket or a " is missing. The easiest way to leave the procedure is to type some brackets or " and then `(RETURN)` .

3.9.1 ASSUME

Syntax: `ASSUME (int_constant , expression)`

Purpose: Tests the expression for correctness if the `int_constant` is smaller as a variable `assumeLevel`. If no such variable exist the `int` expression is compared against 0. It is possible to define an individual `assumeLevel` for each library and/or procedure. If the expression is evaluated and not true (i.e. does not evaluate to `int(0)`) an error is raised.

Note: `ASSUME` shall be used for documentation and debugging, production code of a library must never define `assumeLevel`.

Example:

```

ASSUME(0,2==2); // always tested
ASSUME(1,1==2); // not evaluated
int assumeLevel=2;
ASSUME(1,1==2);
↳ ? ASSUME failed: ASSUME(1,1==2);

```

```

↳ ? error occurred in or before ./examples/ASSUME.sing line 4: ' ASSU
    (1,1==2);'
    // setting a different assumeLevel for poly.lib:
    int Poly::assumeLevel=2;
↳ Poly of type 'ANY'. Trying load.
↳ ? 'Poly' no such package
↳ ? error occurred in or before ./examples/ASSUME.sing line 6: ' int l
    ly::assumeLevel=2;'
↳ ? wrong type declaration. type 'help int;'
```

3.9.2 Tracing of procedures

Setting the `TRACE` variable to 1 (resp. 3) results in reporting of all procedure entries and exits (resp. together with line numbers). If `TRACE` is set to 4, `Singular` displays each line before its interpretation and waits for the `RETURN` key being pressed. See Section 5.3.9 [TRACE var], page 301.

Example:

```

proc t1
{
  int i=2;
  while (i>0)
  { i=i-1; }
}
TRACE=3;
t1();
↳
↳ entering t1 (level 0)
↳ {1}{2}{3}{4}{5}{4}{5}{6}{7}{4}{5}{6}{7}{4}{6}{7}{8}
↳ leaving t1 (level 0)
```

3.9.3 Source code debugger

The source code debugger (`sdb`) is an experimental feature, its interface may change in future versions of `SINGULAR`.

To enable the use of the source code debugger `SINGULAR` has to be started with the option `-d` or `--sdb` (see Section 3.1.6 [Command line options], page 19).

sdb commands

Each `sdb` command consists of one character which may be followed by a parameter.

<code>b</code>	print backtrace of calling stack
<code>c</code>	continue
<code>e</code>	edit the current procedure and reload it (current call will be aborted) only available on UNIX systems
<code>h,?</code>	display help screen
<code>n</code>	execute current line, <code>sdb</code> break at next line
<code>p <identifier></code>	display type and value of the variable given by <code><identifier></code>
<code>Q</code>	quit this <code>SINGULAR</code> session

```
q <flags>  quit debugger, set debugger flags(0,1,2)
           0: continue, disable the debugger
           1: continue
           2: throw an error, return to toplevel
```

Syntactical errors in procedures

If SINGULAR was started using the command line option `-d` or `--sdb`, a syntactical error in a procedure will start the source code debugger instead of returning to the top level with an error message. The commands `q 1` and `q 2` are equivalent in this case.

SDB breakpoints in procedures

Up to seven SDB breakpoints can be set. To set a breakpoint at a procedure use `breakpoint`. (See Section 5.2.3 [breakpoint], page 286).

These breakpoints can be cleared with the command `d breakpoint_no` from within the debugger or with `breakpoint(proc_name , -1);`.

3.9.4 Break points

A break point can be put into a proc by inserting the command `~`. If Singular reaches a break point it asks for lines of commands (line-length must be less than 80 characters) from the user. It returns to normal execution if given an empty line. See Section 5.2.16 [`~`], page 297.

Example:

```
proc t
{
  int i=2;
  ~;
  return(i+1);
}
t();
↳ -- break point in t --
↳ -- 0: called from STDIN --
i;           // here local variables of the procedure can be accessed
↳ 2
↳ -- break point in t --

↳ 3
```

3.9.5 Printing of data

The procedure `dbprint` is useful for optional output of data: it takes 2 arguments and prints the second argument, if the first argument is positive; otherwise, it does nothing. See Section 5.1.17 [dbprint], page 167; Section 5.3.11 [voice], page 303.

3.9.6 libparse

`libparse` is a stand-alone program contained in the SINGULAR distribution (at the place where the SINGULAR executable program resides), which cannot be called inside SINGULAR. It is a debugging tool for libraries which performs exactly the same checks as the `load` command in SINGULAR, but generates more output during parsing. `libparse` is useful if an error occurs while loading the

library, but the whole block around the line specified seems to be correct. In these situations the real error might have occurred hundreds of lines earlier in the library.

Usage:

`libparse [options] singular-library`

Options:

`-d Debuglevel`

increases the amount of output during parsing, where Debuglevel is an integer between 0 and 4. Default is 0.

`-s` turns on reporting about violations of unenforced syntax rules

The following syntax checks are performed in any case:

- counting of pairs of brackets {,} , [,] and (,) (number of { has to match number of }, same for [,] and (,)).
- counting of " (number of " must be even).
- general library syntax (only LIB, static, proc (with parameters, help, body and example) and comments, i.e // and /* ... */ , are allowed).

Its output lists all procedures that have been parsed successfully:

```
$ libparse sample.lib
Checking library 'sample.lib'
  Library      function      line,start-eod  line,body-eob  line,example-eeo
Version:0.0.0;
g Sample          tab line      9,  149-165    13,  271-298    14,  300-402
l Sample          internal_tab line    24,  450-475    25,  476-496     0,    0-496
```

where the following abbreviations are used:

- g: global procedure (default)
- l: static procedure, i.e., local to the library.

each of the following is the position of the byte in the library.

- start: begin of 'proc'
- eod: end of parameters
- body: start of procedurebody '{'
- eob: end of procedurebody '}'
- example: position of 'example'
- eoe: end of example '}'

Hence in the above example, the first procedure of the library `sample.lib` is user-accessible and its name is `tab`. The procedure starts in line 9, at character 149. The head of the procedure ends at character 165, the body starts in line 13 at character 271 and ends at character 298. The example section extends from line 14 character 300 to character 402.

The following example shows the result of a missing close-bracket `}` in line 26 of the library `sample.lib`.

```
LIB "sample.lib";
↳ ? Library sample.lib: ERROR occurred: in line 26, 497.
↳ ? missing close bracket '}' at end of library in line 26.
↳ ? Cannot load library,... aborting.
↳ ? error occurred in STDIN line 1: 'LIB "sample.lib";'
```


3.9.7 option(warn)

If this option is set some constructs which **may** lead to bug will result in a warning. While there are legitimate uses for them and they are **not errors** is is worth thinking about it.

change of options during a procedure call: is this side effect intended?

use of **def**: avoids type checking, but useful if a procedure handles several types at once

ASSUME outside of procedures: while a failed **ASSUME** aborts the current procedures and return to the top level - what should it do at top level?

See Section 5.1.110 [option], page 230.

3.10 Dynamic loading

In addition to the concept of libraries, it is also possible to dynamically extend the functionality by loading functions written in C/C++ or some other higher programming language. A collection of such functions is called a dynamic module and can be loaded by the command **LIB** or **load**. It is basically handled in the same way as a library: upon loading, a new **package** is created which holds the contents of the dynamic module. General information about the loaded module can be displayed by the command **help package_name**. After loading the dynamic module, its functions can be used exactly like the built-in **SINGULAR** functions.

To have the full functionality of a built-in function, dynamic modules need to comply with certain requirements on their internal structure. As this would be beyond the scope of the **Singular** manual, a separate, more detailed guide on how to write and use dynamic modules is available.

4 Data types

This chapter explains all data types of SINGULAR in alphabetical order. For every type, there is a description of the declaration syntax as well as information about how to build expressions of certain types.

The term expression list in SINGULAR refers to any comma separated list of expressions.

For the general syntax of a declaration see Section 3.5.1 [General command syntax], page 41.

4.1 cring

Variables of type cring represent the ring of coefficients (see Section 4.14 [number], page 113)

4.1.1 cring declarations

Syntax: `cring name = cring-expression ;`

Purpose: defines a new coefficient ring resp. field to be used for a ring definition (see Section 4.19 [ring], page 124). Most objects of this type are predefined.

Default: none

Example:

```

ZZ;
↳ ZZ
ZZ/3;
↳ ZZ/3

```

4.1.2 cring expressions

A cring expression is:

1. an identifier of type cring:
 - QQ - the rational numbers
 - ZZ - the integers
2. a function returning cring
3. an expression involving crings and the arithmetic operations `/`.

Example:

```

ZZ/3;
↳ ZZ/3

```

See Section 4.19 [ring], page 124.

4.1.3 cring operations

`/` residue class ring

Example:

```

ZZ/101;
↳ ZZ/101

```

4.1.4 cring related functions

<code>crossprod</code>	cross product of several objects of type <code>cring</code> (see Section 5.1.15 [<code>crossprod</code>], page 166)
<code>Float</code>	several variants of Floating point (inexact) real and complex numbers (see Section 5.1.45 [<code>Float</code>], page 184).
<code>flintQ</code>	multivariate rational functions over \mathbb{Q} (via <code>flint</code> , requires $\geq 2.5.3$) (see Section 5.1.44 [<code>flintQ</code>], page 183).

See Section 5.1.45 [`Float`], page 184; Section 5.1.15 [`crossprod`], page 166; Section 5.1.44 [`flintQ`], page 183.

4.2 bigint

Variables of type `bigint` represent the arbitrary long integers. They can only be constructed from other types (`int`, `number`).

4.2.1 bigint declarations

Syntax: `bigint name = int_expression ;`

Purpose: defines a long integer variable

Default: 0

Example:

```

    bigint i = 42;
    ring r=0,x,dp;
    number n=2;
    bigint j = i + bigint(n)^50; j;
    ↪ 1125899906842666

```

4.2.2 bigint expressions

A `bigint` expression is:

1. an identifier of type `bigint`
2. a function returning `bigint`
3. an expression involving `bigints` and the arithmetic operations `+`, `-`, `*`, `div`, `% (mod)`, or `^`
4. a type cast to `bigint`.

Example:

```

// Note: 11*13*17*100*200*2000*503*1111*222222
// returns a machine integer:
11*13*17*100*200*2000*503*1111*222222;
↪ // ** int overflow(*), result may be wrong
↪ // ** int overflow(*), result may be wrong
↪ // ** int overflow(*), result may be wrong
↪ // ** int overflow(*), result may be wrong
↪ -1875651584
// using the type cast number for a greater allowed range
bigint(11)*13*17*100*200*2000*503*1111*222222;

```

↳ 1207574812868424000000

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.6 [int], page 82; Section 4.14 [number], page 113.

4.2.3 bigint operations

+	addition
-	negation or subtraction
*	multiplication
div	integer division (omitting the remainder ≥ 0)
mod, %	integer modulo (the remainder of the division div)
^, **	exponentiation (exponent must be non-negative)
<, >, <=, >=, ==, <>	comparators

Example:

```
bigint(5)*2, bigint(2)^100-10;
↳ 10 1267650600228229401496703205366
bigint(-5) div 2, bigint(-5) mod 2;
↳ -3 1
```

4.2.4 bigint related functions

gcd	greatest common divisor (see Section 5.1.50 [gcd], page 188)
memory	memory usage (see Section 5.1.89 [memory], page 216)

See Section 5.1.89 [memory], page 216.

4.3 bigintmat

Big integer matrices are matrices with big integer entries. No basering definition is required to use bigint matrices, for they do not belong to a ring. Bigintmat entries can have any size because of the use of bigint.

4.3.1 bigintmat declarations

Syntax: `bigintmat name = bigintmat_expression ;`
`bigintmat name [rows] [cols] = bigintmat_expression ;`
`bigintmat name [rows] [cols] = list_of_int_and_bigint_expressions ;`
rows and cols must be positive int expressions.

Purpose: defines a bigintmat variable.
Given a list of (big) integers, the matrix is filled up with the first row from the left to the right, then the second one and so on. If the (big-)int_list contains less than rows*cols elements, the remaining ones are set to zero; if it contains more elements, only the first rows*cols ones are considered.

Default: empty (1x0 matrix)

Example:

```

bigintmat bim[4][3]=2, 5, 224553233465, 232444, 434, 0, 0, 4544232222;
bim;
↳      2,          5,224553233465,
↳ 232444,          434,          0,
↳      0,4544232222,          0,
↳      0,          0,          0
bim[2, 1];
↳ 232444

```

4.3.2 bigintmat expressions

A bigintmat expression is:

1. an identifier of type bigintmat
2. a function returning bigintmat
3. a bigintmat operation involving (big-)ints and int operations (+, -, *)
4. an expression involving bigintmats and the operations (+, -, *)
5. a type cast to bigintmat (see Section 4.3.3 [bigintmat type cast], page 75)

Example:

```

bigintmat m1[2][2]=1, 2, 6, 3;
m1*3;
↳ 3,6,
↳ 18,9
intmat im[3][2] = intmat(m1*3);
bigintmat m2 = bigintmat(im); // cast intmat im to bigintmat
m2;
↳ 3,6,
↳ 18,9
m2*m1+m2;
↳ 42,30,
↳ 90,72
_+4;
↳ 46, 0,
↳ 0,76

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.3 [bigintmat], page 74.

4.3.3 bigintmat type cast

Syntax: bigintmat (expression)

Type: bigintmat

Purpose: Converts expression to a bigintmat, where expression must be of type intmat, or bigintmat. The size (resp. dimension) of the created bigintmat equals the size (resp. dimension) of the expression.

Example:

```

intmat im[2][1]=2, 3;
bigintmat(im);
↳ 2,

```

```

↳ 3
    bigintmat(_);
↳ 2,
↳ 3
    bigintmat(intmat(intvec(1,2,3,4), 2, 2)); //casts at first to intmat, th
↳ 1,2,
↳ 3,4

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.3 [bigintmat], page 74; Section 4.7.3 [intmat type cast], page 89.

4.3.4 bigintmat operations

- + addition with intmat, int, or bigint. In case of (big-)int, it is added to every entry of the matrix.
- negation or subtraction with intmat, int, or bigint. In case of (big-)int, it is subtracted from every entry of the matrix.
- * multiplication with intmat, int, or bigint; In case of (big-)int, every entry of the matrix is multiplied by the (big-)int
- <>, == comparators

bigintmat_expression [int, int]

is a bigintmat entry, where the first index indicates the row and the second the column

Example:

```

    bigintmat m[3][4] = 3,3,6,3,5,2,2,7,0,0,45,3;
    m;
↳ 3,3, 6,3,
↳ 5,2, 2,7,
↳ 0,0,45,3
    m[1,3]; // show entry at [row 1, col 3]
↳ 6
    m[1,3] = 10; // set entry at [row 1, col 3] to 10
    m;
↳ 3,3,10,3,
↳ 5,2, 2,7,
↳ 0,0,45,3
    size(m); // number of entries
↳ 12
    bigintmat n[2][3] = 2,6,0,4,0,5;
    n * m;
↳ 36,18, 32,48,
↳ 12,12,265,27
    typeof(_);
↳ bigintmat
    -m;
↳ -3,-3,-10,-3,
↳ -5,-2, -2,-7,
↳ 0, 0,-45,-3
    bigintmat o;
    o=n-10;

```

```

o;
↳ -8, 0,0,
↳ 0,-10,0
  m*2;          // double each entry of m
↳ 6,6,20, 6,
↳ 10,4, 4,14,
↳ 0,0,90, 6
  o-2*m;
↳      ? bigintmat/cmatrix not compatible
↳      ? error occurred in or before ./examples/bigintmat_operations.sing lin\
      e 15: ' o-2*m;'
```

4.4 def

Objects may be defined without a specific type: they inherit their type from the first assignment to them. E.g., `ideal i=x,y,z; def j=i^2;` defines the ideal i^2 with the name `j`.

Note: Unlike other assignments a ring as an untyped object is not a copy but another reference to the same (possibly unnamed) ring. This means that entries in one of these rings appear also in the other ones. The following defines a ring `s` which is just another reference (or name) for the basering `r`. The name `basing` is an alias for the current ring.

```

ring r=32003,(x,y,z),dp;
poly f = x;
def s=basing;
setring s;
nameof(basing);
↳ s
  listvar();
↳ // s          [0] *ring
↳ //      f          [0] poly
↳ // r          [0] ring(*)
  poly g = y;
  kill f;
  listvar(r);
↳ // r          [0] ring(*)
↳ // g          [0] poly
  ring t=32003,(u,w),dp;
  def rt=r+t;
  rt;
↳ // coefficients: ZZ/32003
↳ // number of vars : 5
↳ //      block  1 : ordering dp
↳ //                : names    x y z
↳ //      block  2 : ordering dp
↳ //                : names    u w
↳ //      block  3 : ordering C
```

This reference to a ring with `def` is useful if the basering is not local to the procedure (so it cannot be accessed by its name) but one needs a name for it (e.g., for a use with `setring` or `map`). `setring r;` does not work in this case, because `r` may not be local to the procedure.

4.4.1 def declarations

Syntax: `def name = expression ;`

Purpose: defines an object of the same type as the right-hand side.

Default: none

Note: This is useful if the right-hand side may be of variable type as a consequence of a computation (e.g., ideal or module or matrix). It may also be used in procedures to give the basering a name which is local to the procedure.

Example:

```
def i=2;
typeof(i);
↳ int
```

See Section 5.1.159 [typeof], page 277.

4.5 ideal

Ideals are represented as lists of polynomials which generate the ideal. Like polynomials they can only be defined or accessed with respect to a basering.

Note: `size` counts only the non-zero generators of an ideal whereas `ncols` counts all generators; see Section 5.1.142 [size], page 259, Section 5.1.103 [ncols], page 227.

4.5.1 ideal declarations

Syntax: `ideal name = list_of_poly_and_ideal_expressions ;`
`ideal name = ideal_expression ;`

Purpose: defines an ideal.

Default: 0

Example:

```
ring r=0,(x,y,z),dp;
poly s1 = x2;
poly s2 = y3;
poly s3 = z;
ideal i = s1, s2-s1, 0,s2*s3, s3^4;
i;
↳ i[1]=x2
↳ i[2]=y3-x2
↳ i[3]=0
↳ i[4]=y3z
↳ i[5]=z4
size(i);
↳ 4
ncols(i);
↳ 5
```

4.5.2 ideal expressions

An ideal expression is:

1. an identifier of type ideal
2. a function returning an ideal
3. a combination of ideal expressions by the arithmetic operations + or *

4. a power of an ideal expression (operator \wedge or $**$)
Note that the computation of the product $i*i$ involves all products of generators of i while i^2 involves only the different ones, and is therefore faster.
5. a type cast to ideal

Example:

```

ring r=0,(x,y,z),dp;
ideal m = maxideal(1);
m;
↳ m[1]=x
↳ m[2]=y
↳ m[3]=z
poly f = x2;
poly g = y3;
ideal i = x*y*z , f-g, g*(x-y) + f^4 ,0, 2x-z2y;
ideal M = i + maxideal(10);
timer =0;
i = M*M;
timer;
↳ 0
ncols(i);
↳ 505
timer =0;
i = M^2;
ncols(i);
↳ 505
timer;
↳ 0
i[ncols(i)];
↳ x20
vector v = [x,y-z,x2,y-x,x2yz2-y];
ideal j = ideal(v);

```

4.5.3 ideal operations

- + addition (concatenation of the generators and simplification)
- * multiplication (with ideal, poly, vector, module; simplification in case of multiplication with ideal)
- \wedge exponentiation (by a non-negative integer)

ideal_expression [intvec_expression]

are polynomial generators of the ideal, index 1 gives the first generator.

Note: For simplification of an ideal, see also Section 5.1.141 [simplify], page 258.

Example:

```

ring r=0,(x,y,z),dp;
ideal I = 0,x,0,1;
I;
↳ I[1]=0
↳ I[2]=x

```

```

↳ I[3]=0
↳ I[4]=1
  I + 0;    // simplification
↳ _[1]=1
  ideal J = I,0,x,x-z;;
  J;
↳ J[1]=0
↳ J[2]=x
↳ J[3]=0
↳ J[4]=1
↳ J[5]=0
↳ J[6]=x
↳ J[7]=x-z
  I * J;    // multiplication with simplification
↳ _[1]=1
  I*x;
↳ _[1]=0
↳ _[2]=x2
↳ _[3]=0
↳ _[4]=x
  vector V = [x,y,z];
  print(V*I);
↳ 0,x2,0,x,
↳ 0,xy,0,y,
↳ 0,xz,0,z
  ideal m = maxideal(1);
  m^2;
↳ _[1]=x2
↳ _[2]=xy
↳ _[3]=xz
↳ _[4]=y2
↳ _[5]=yz
↳ _[6]=z2
  ideal II = I[2..4];
  II;
↳ II[1]=x
↳ II[2]=0
↳ II[3]=1

```

4.5.4 ideal related functions

char_series

irreducible characteristic series (see Section 5.1.6 [char_series], page 159)

coeffs matrix of coefficients (see Section 5.1.12 [coeffs], page 163)

contract contraction by an ideal (see Section 5.1.13 [contract], page 165)

diff partial derivative (see Section 5.1.24 [diff], page 171)

degree multiplicity, dimension and codimension of the ideal of leading terms (see Section 5.1.20 [degree], page 169)

dim Krull dimension of basering modulo the ideal of leading terms (see Section 5.1.25 [dim], page 171)

<code>eliminate</code>	elimination of variables (see Section 5.1.28 [eliminate], page 174)
<code>facstd</code>	factorizing Groebner basis algorithm (see Section 5.1.34 [facstd], page 177)
<code>factorize</code>	ideal of factors of a polynomial (see Section 5.1.36 [factorize], page 178)
<code>fglm</code>	Groebner basis computation from a Groebner basis w.r.t. a different ordering (see Section 5.1.39 [fglm], page 181)
<code>finduni</code>	computation of univariate polynomials lying in a zero dimensional ideal (see Section 5.1.43 [finduni], page 183)
<code>fres</code>	free resolution of a standard basis (see Section 5.1.48 [fres], page 186)
<code>groebner</code>	Groebner basis computation (a wrapper around <code>std</code> , <code>stdhilb</code> , <code>stdfglm</code> ,...) (see [groebner], page 785)
<code>highcorner</code>	the smallest monomial not contained in the ideal. The ideal has to be zero-dimensional. (see Section 5.1.55 [highcorner], page 192)
<code>homog</code>	homogenization with respect to a variable (see Section 5.1.57 [homog], page 194)
<code>hilb</code>	Hilbert series of a standard basis (see Section 5.1.56 [hilb], page 193)
<code>indepSet</code>	sets of independent variables of an ideal (see Section 5.1.61 [indepSet], page 196)
<code>interred</code>	interreduction of an ideal (see Section 5.1.64 [interred], page 199)
<code>intersect</code>	ideal intersection (see Section 5.1.65 [intersect], page 199)
<code>jacob</code>	ideal of all partial derivatives resp. jacobian matrix (see Section 5.1.66 [jacob], page 200)
<code>jet</code>	Taylor series up to a given order (see Section 5.1.68 [jet], page 201)
<code>kbase</code>	vector space basis of basering modulo ideal of leading terms (see Section 5.1.69 [kbase], page 203)
<code>koszul</code>	Koszul matrix (see Section 5.1.73 [koszul], page 205)
<code>lead</code>	leading terms of a set of generators (see Section 5.1.75 [lead], page 206)
<code>lift</code>	lift-matrix (see Section 5.1.80 [lift], page 209)
<code>liftstd</code>	standard basis and transformation matrix computation (see Section 5.1.81 [liftstd], page 209)
<code>lres</code>	free resolution for homogeneous ideals (see Section 5.1.83 [lres], page 212)
<code>maxideal</code>	power of the maximal ideal at 0 (see Section 5.1.88 [maxideal], page 216)
<code>minbase</code>	minimal generating set of a homogeneous ideal, resp. module, or an ideal, resp. module, in a local ring (see Section 5.1.91 [minbase], page 218)
<code>minor</code>	set of minors of a matrix (see Section 5.1.92 [minor], page 218)
<code>modulo</code>	representation of $(h1 + h2)/h1 \cong h2/(h1 \cap h2)$ (see Section 5.1.94 [modulo], page 220)
<code>mres</code>	minimal free resolution of an ideal resp. module w.r.t. a minimal set of generators of the given ideal resp. module (see Section 5.1.98 [mres], page 222)
<code>mstd</code>	standard basis and minimal generating set of an ideal (see Section 5.1.99 [mstd], page 223)

mult	multiplicity, resp. degree, of the ideal of leading terms (see Section 5.1.100 [mult], page 224)
ncols	number of columns (see Section 5.1.103 [ncols], page 227)
nres	a free resolution of an ideal resp. module M which is minimized from the second free module on (see Section 5.1.105 [nres], page 228)
preimage	preimage under a ring map (see Section 5.1.116 [preimage], page 236)
qhweight	quasihomogeneous weights of an ideal (see Section 5.1.122 [qhweight], page 241)
quotient	ideal quotient (see Section 5.1.125 [quotient], page 243)
reduce	normalform with respect to a standard basis (see Section 5.1.129 [reduce], page 246)
res	free resolution of an ideal resp. module but not changing the given ideal resp. module (see [res], page 785)
simplify	simplification of a set of polynomials (see Section 5.1.141 [simplify], page 258)
size	number of non-zero generators (see Section 5.1.142 [size], page 259)
slimgb	Groebner basis computation with slim technique (see Section 5.1.143 [slimgb], page 260)
sortvec	permutation for sorting ideals resp. modules (see Section 5.1.144 [sortvec], page 261)
sres	free resolution of a standard basis (see Section 5.1.147 [sres], page 264)
std	standard basis computation (see Section 5.1.149 [std], page 266)
stdfglm	standard basis computation with fglm technique (see [stdfglm], page 785)
stdhilb	Hilbert driven standard basis computation (see [stdhilb], page 785)
subst	substitution of a ring variable (see Section 5.1.152 [subst], page 269)
syz	computation of the first syzygy module (see Section 5.1.154 [syz], page 275)
vdim	vector space dimension of basering modulo ideal of leading terms (see Section 5.1.166 [vdim], page 281)
weight	optimal weights (see Section 5.1.170 [weight], page 283)

4.6 int

Variables of type `int` represent the machine integers and are, therefore, limited in their range (e.g., the range is between -2147483647 and 2147483647 on 32-bit machines). They are mainly used to count things (dimension, rank, etc.), in loops (see Section 5.2.8 [for], page 290), and to represent boolean values (`FALSE` is represented by 0, every other value means `TRUE`, see Section 4.6.5 [boolean expressions], page 86).

Integers consist of a sequence of digits, possibly preceded by a sign. A space is considered as a separator, so it is not allowed between digits. A sequence of digits outside the allowed range is converted to the type `bigint`, see Section 4.2 [bigint], page 73.

4.6.1 int declarations

Syntax: `int name = int_expression ;`

Purpose: defines an integer variable.

Default: 0

Example:

```

    int i = 42;
    int j = i + 3; j;
    ↪ 45
    i = i * 3 - j; i;
    ↪ 81
    int k;    // assigning the default value 0 to k
    k;
    ↪ 0

```

4.6.2 int expressions

An int expression is:

1. a sequence of digits (if the number represented by this sequence is too large to fit into the range of integers it is automatically converted to the type number, if a basering is defined)
2. an identifier of type int
3. a function returning int
4. an expression involving ints and the arithmetic operations +, -, *, div (/), % (mod), or ^
5. a boolean expression
6. a type cast to int

Note: Variables of type int represent the compiler integers and are, therefore, limited in their range (see Section 6.1 [Limitations], page 304). If this range is too small the expression must be converted to the type number over a ring with characteristic 0.

Example:

```

12345678901; // too large
↪ 12345678901
typeof(_);
↪ bigint
ring r=0,x,dp;
12345678901;
↪ 12345678901
typeof(_);
↪ bigint
// Note: 11*13*17*100*200*2000*503*1111*222222
// returns a machine integer:
11*13*17*100*200*2000*503*1111*222222;
↪ // ** int overflow(*), result may be wrong
↪ // ** int overflow(*), result may be wrong
↪ // ** int overflow(*), result may be wrong
↪ // ** int overflow(*), result may be wrong
↪ -1875651584
// using the type cast number for a greater allowed range
number(11)*13*17*100*200*2000*503*1111*222222;
↪ 12075748128684240000000
ring rp=32003,x,dp;
12345678901;
↪ 12345678901
typeof(_);
↪ bigint

```

```

intmat m[2][2] = 1,2,3,4;
m;
↳ 1,2,
↳ 3,4
m[2,2];
↳ 4
typeof(_);
↳ int
det(m);
↳ -2
m[1,1] + m[2,1] == trace(m);
↳ 0
! 0;
↳ 1
1 and 2;
↳ 1
intvec v = 1,2,3;
def d =transpose(v)*v; // scalarproduct gives an 1x1 intvec
typeof(d);
↳ intvec
int i = d[1]; // access the first (the only) entry in the intvec
ring rr=31,(x,y,z),dp;
poly f = 1;
i = int(f); // cast to int
// Integers may be converted to constant polynomials by an assignment,
poly g=37;
// define the constant polynomial g equal to the image of
// the integer 37 in the actual coefficient field, here it equals 6
g;
↳ 6

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.14 [number], page 113.

4.6.3 int operations

++	changes its operand to its successor, is itself no int expression
--	changes its operand to its predecessor, is itself no int expression
+	addition
-	negation or subtraction
*	multiplication
div	integer division (omitting the remainder), rounding toward 0
%, mod	integer modulo (the remainder of the division)
~, **	exponentiation (exponent must be non-negative)
<, >, <=, >=, ==, <>	comparators

Note: An assignment `j=i++`; or `j=i--`; is not allowed, in particular it does not change the value of `j`, see Section 6.1 [Limitations], page 304.

Example:

```

    int i=1;
    int j;
    i++; i; i--; i;
↳ 2
↳ 1
    // ++ and -- do not return a value as in C, cannot assign
    j = i++;
↳ // ** right side is not a datum, assignment ignored
↳ // ** in line >> j = i++;<<
    // the value of j is unchanged
    j; i;
↳ 0
↳ 2
    i+2, 2-i, 5^2;
↳ 4 0 25
    5 div 2, 8%3;
↳ 2 2
    -5 div 2, -5 mod 2, -5 % 2;
↳ -2 -1 -1
    1<2, 2<=2;
↳ 1 1

```

4.6.4 int related functions

char	characteristic of the coefficient field of a ring (see Section 5.1.5 [char], page 159)
deg	degree of a polynomial resp. vector (see Section 5.1.19 [deg], page 168)
det	determinant (see Section 5.1.23 [det], page 170)
dim	Krull dimension of basering modulo ideal of leading terms, resp. dimension of module of leading terms (see Section 5.1.25 [dim], page 171)
extgcd	Bezout representation of gcd (see Section 5.1.33 [extgcd], page 176)
find	position of a substring in a string (see Section 5.1.42 [find], page 182)
gcd	greatest common divisor (see Section 5.1.50 [gcd], page 188)
koszul	Koszul matrix (see Section 5.1.73 [koszul], page 205)
memory	memory usage (see Section 5.1.89 [memory], page 216)
mult	multiplicity of an ideal, resp. module, of leading terms (see Section 5.1.100 [mult], page 224)
ncols	number of columns (see Section 5.1.103 [ncols], page 227)
npars	number of ring parameters (see Section 5.1.104 [npars], page 227)
nrows	number of rows of a matrix, resp. the rank of the free module where the vector or module lives (see Section 5.1.106 [nrows], page 228)
nvars	number of ring variables (see Section 5.1.108 [nvars], page 229)
ord	degree of the leading term of a polynomial resp. vector (see Section 5.1.111 [ord], page 234)
par	n-th parameter of the basering (see Section 5.1.113 [par], page 235)

pardeg	degree of a number considered as a polynomial in the ring parameters (see Section 5.1.114 [pardeg], page 235)
prime	the next lower prime (see Section 5.1.117 [prime], page 237)
random	a pseudo random integer between the given limits (see Section 5.1.126 [random], page 244)
regularity	regularity of a resolution (see Section 5.1.130 [regularity], page 247)
rvar	test, if the given expression or string is a ring variable (see Section 5.1.137 [rvar], page 253)
size	number of elements in an object (see Section 5.1.142 [size], page 259)
trace	trace of an integer matrix (see Section 5.1.156 [trace], page 276)
var	n-th ring variable of the basering (see Section 5.1.163 [var], page 279)
vdim	vector space dimension of basering modulo ideal of leading terms, resp. of freemodule modulo module of leading terms (see Section 5.1.166 [vdim], page 281)

4.6.5 boolean expressions

A boolean expression is an int expression used in a logical context:

An int expression $\langle \rangle 0$ evaluates to *TRUE* (represented by 1), 0 evaluates to *FALSE* (represented by 0).

The following is the list of available comparisons of objects of the same type.

Note: There are no comparisons for ideals and modules, resolutions and maps.

- integer comparisons:

```

i == j
i != j    // or    i <> j
i <= j
i >= j
i > j
i < j

```

- number comparisons:

```

m == n
m != n    // or    m <> n
m < n
m > n
m <= n
m >= n

```

For numbers from \mathbb{Z}/p or from field extensions not all operations are useful:

- 0 is always the smallest element,
- in \mathbb{Z}/p the representatives in the range $-(p-1)/2..(p-1)/2$ when $p > 2$ resp. 0 and 1 for $p = 2$ are used for comparisons,
- in field extensions the last two operations (\geq , \leq) yield always *TRUE* (1) and the $<$ and $>$ are equivalent to \neq .

- polynomial or vector comparisons:


```

f == g
f != g    // or    f <> g
f <= g    // comparing the leading term w.r.t. the monomial order
f < g
f >= g
f > g

```

4. intmat or matrix comparisons:

```

v == w
v != w    // or    v <> w

```

5. intvec or string comparisons:

```

f == g
f != g    // or    f <> g
f <= g    // comparing lexicographically
f >= g    // w.r.t. the order specified by ASCII
f > g
f < g

```

6. boolean expressions combined by boolean operations (**and**, **or**, **not**)

Note: All arguments of a logical expression are first evaluated and then the value of the logical expression is determined. For example, the logical expression $(a \ || \ b)$ is evaluated by first evaluating a *and* b , even though the value of b has no influence on the value of $(a \ || \ b)$, if a evaluates to true.

Note that this evaluation is different from the left-to-right, conditional evaluation of logical expressions (as found in most programming languages). For example, in these other languages, the value of $(1 \ || \ b)$ is determined without ever evaluating b .

See Section 6.3 [Major differences to the C programming language], page 304.

4.6.6 boolean operations

and logical **and**, may also be written as **&&**

or logical **or**, may also be written as **||**

not logical **not**, may also be written as **!**

The precedence of the boolean operations is:

1. parentheses
2. comparisons
3. not
4. and
5. or

Example:

```

(1>2) and 3;
↳ 0
1 > 2 and 3;
↳ 0
! 0 or 1;
↳ 1
!(0 or 1);
↳ 0

```

4.7 intmat

Integer matrices are matrices with integer entries. For the range of integers see Section 6.1 [Limitations], page 304. Integer matrices do not belong to a ring, they may be defined without a basering being defined. An intmat can be multiplied by and added to an int; in this case the int is converted into an intmat of the right size with the integer on the diagonal. The integer 1, for example, is converted into the unit matrix.

4.7.1 intmat declarations

Syntax: `intmat name = intmat_expression ;`
`intmat name [rows] [cols] = intmat_expression ;`
`intmat name [rows] [cols] = list_of_int_and_intvec_and_intmat_expressions ;`
 rows and cols must be positive int expressions.

Purpose: defines an intmat variable.
 Given a list of integers, the matrix is filled up with the first row from the left to the right, then the second row and so on. If the int_list contains less than rows*cols elements, the matrix is filled up with zeros; if it contains more elements, only the first rows*cols elements are used.

Default: 0 (1 x 1 matrix)

Example:

```
intmat im[3][5]=1,3,5,7,8,9,10,11,12,13;
im;
↳ 1,3,5,7,8,
↳ 9,10,11,12,13,
↳ 0,0,0,0,0
im[3,2];
↳ 0
intmat m[2][3] = im[1..2,3..5]; // defines a submatrix
m;
↳ 5,7,8,
↳ 11,12,13
```

4.7.2 intmat expressions

An intmat expression is:

1. an identifier of type intmat
2. a function returning intmat
3. an intmat operation involving ints and int operations (+, -, *, div, %)
4. an expression involving intmats and the operations (+, -, *)
5. a type cast to intmat (see Section 4.7.3 [intmat type cast], page 89)

Example:

```
intmat Idm[2][2];
Idm +1; // add the unit intmat
↳ 1,0,
↳ 0,1
intmat m1[3][2] = _,1,-2; // take entries from the last result
```

```

    m1;
    ↪ 1,0,
    ↪ 0,1,
    ↪ 1,-2
    intmat m2[2][3]=1,0,2,4,5,1;
    transpose(m2);
    ↪ 1,4,
    ↪ 0,5,
    ↪ 2,1
    intvec v1=1,2,4;
    intvec v2=5,7,8;
    m1=v1,v2;          // fill m1 with v1 and v2
    m1;
    ↪ 1,2,
    ↪ 4,5,
    ↪ 7,8
    trace(m1*m2);
    ↪ 56

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.14 [number], page 113.

4.7.3 intmat type cast

Syntax: `intmat (expression)`
 `intmat (expression, int_n, int_m)`

Type: `intmat`

Purpose: Converts expression to an `intmat`, where expression must be of type `intvec`, `intmat`, or `bigintmat`. If `int_n` and `int_m` are supplied, then they specify the dimension of the `intmat`. Otherwise, the size (resp. dimensions) of the `intmat` are determined by the size (resp. dimensions) of the expression. If expression is a `bigintmat` containing an entry larger than the limit of `int`, it is set to 0 in the returning `intmat`.

Example:

```

    intmat(intvec(1));
    ↪ 1
    intmat(intvec(1), 1, 2);
    ↪ 1,0
    intmat(intvec(1,2,3,4), 2, 2);
    ↪ 1,2,
    ↪ 3,4
    intmat(_, 2, 3);
    ↪ 1,2,3,
    ↪ 4,0,0
    intmat(_, 2, 1);
    ↪ 1,2
    bigintmat bim[2][3]=34, 64, 345553234, 35553, 6434, 6563335675;
    intmat(bim);
    ↪ 34,64,345553234,
    ↪ 35553,6434,-2026598917

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.7 [intmat], page 88; Section 4.12.3 [matrix type cast], page 107.

4.7.4 intmat operations

+	addition with intmat or int; the int is converted into a diagonal intmat
-	negation or subtraction with intmat or int; the int is converted into a diagonal intmat
*	multiplication with intmat, intvec, or int; the int is converted into a diagonal intmat
div,/	division of entries in the integers (omitting the remainder)
%, mod	entries modulo int (remainder of the division)
<>, ==	comparators

intmat_expression [intvec_expression, intvec_expression]

is an intmat entry, where the first index indicates the row and the second the column

Example:

```

intmat m[2][4] = 1,0,2,4,0,1,-1,0,3,2,1,-2;
m;
↳ 1,0,2,4,
↳ 0,1,-1,0
m[2,3];          // entry at row 2, col 3
↳ -1
size(m);         // number of entries
↳ 8
intvec v = 1,0,-1,2;
m * v;
↳ 7,1
typeof(_);
↳ intvec
intmat m1[4][3] = 0,1,2,3,v,1;
intmat m2 = m * m1;
m2;              // 2 x 3 intmat
↳ -2,5,4,
↳ 4,-1,-1
m2*10;          // multiply each entry of m with 10;
↳ -20,50,40,
↳ 40,-10,-10
-m2;
↳ 2,-5,-4,
↳ -4,1,1
m2 % 2;
↳ 0,1,0,
↳ 0,1,1
m2 div 2;
↳ -1,2,2,
↳ 2,-1,-1
m2[2,1];        // entry at row 2, col 1
↳ 4
m1[2..3,2..3];  // submatrix
↳ 1 0 2 1
m2[nrows(m2),ncols(m2)]; // the last entry of intmat m2
↳ -1

```

4.7.5 intmat related functions

bet	Betti numbers of a free resolution (see Section 5.1.4 [bet], page 157)
det	determinant (see Section 5.1.23 [det], page 170)
ncols	number of cols (see Section 5.1.103 [ncols], page 227)
nrows	number of rows (see Section 5.1.106 [nrows], page 228)
random	pseudo random intmat (see Section 5.1.126 [random], page 244)
size	total number of entries (see Section 5.1.142 [size], page 259)
transpose	transpose of an intmat (see Section 5.1.157 [transpose], page 276)
trace	trace of an intmat (see Section 5.1.156 [trace], page 276)

4.8 intvec

Variables of type `intvec` are lists of integers. For the range of integers see Section 6.1 [Limitations], page 304. They may be used for simulating sets of integers (and other sets if the `intvec` is used as an index set for other objects). Addition and subtraction of an `intvec` with an `int` or an `intvec` is done element-wise.

4.8.1 intvec declarations

Syntax: `intvec name = intvec_expression ;`
`intvec name = list_of_int_and_intvec_expressions ;`

Purpose: defines an `intvec` variable.
 An `intvec` consists of an ordered list of integers.

Default: 0

Example:

```

intvec iv=1,3,5,7,8;
iv;
↳ 1,3,5,7,8
iv[4];
↳ 7
iv[3..size (iv)];
↳ 5 7 8

```

4.8.2 intvec expressions

An `intvec` expression is:

1. a range: `int expression .. int expression`
2. a repeated entry: `int expression : positive int expression`
`(a:b` generates an `intvec` of length `b>0` with identical entries `a`)
3. a function returning `intvec`
4. an expression involving `intvec` operations with `int` (+, -, *, /, %)
5. an expression of `intvecs` involving `intvec` operations (+, -)
6. an expression involving an `intvec` operation with `intmat` (*)

7. a type cast to intvec

Example:

```

intvec v=-1,2;
intvec w=v,v;          // concatenation
w;
↳ -1,2,-1,2
w=2:3;                // repetition
w;
↳ 2,2,2
int k = 3;
v = 7:k;
v;
↳ 7,7,7
v=-1,2;
w=-2..2,v,1;
w;
↳ -2,-1,0,1,2,-1,2,1
intmat m[3][2] = 0,1,2,-2,3,1;
m*v;
↳ 2,-6,-1
typeof(_);
↳ intvec
v = intvec(m);
v;
↳ 0,1,2,-2,3,1
ring r;
poly f = x2z + 2xy-z;
f;
↳ x2z+2xy-z
v = leadexp(f);
v;
↳ 2,0,1

```

4.8.3 intvec operations

- + addition with intvec or int (component-wise)
- negation or subtraction with intvec or int (component-wise)
- * multiplication with int (component-wise)
- /, div division by int (component-wise)
- %, mod modulo (component-wise)
- <>, ==, <=, >=, >, < comparison (done lexicographically, different length will be filled with 0 at the right)

intvec_expression [int_expression]

is an element of the intvec; the first element has index one.

Example:

```

    intvec iv = 1,3,5,7,8;
    iv+1;           // add 1 to each entry
↳ 2,4,6,8,9
    iv*2;
↳ 2,6,10,14,16
    iv;
↳ 1,3,5,7,8
    iv-10;
↳ -9,-7,-5,-3,-2
    iv=iv,0;
    iv;
↳ 1,3,5,7,8,0
    iv div 2;
↳ 0,1,2,3,4,0
    iv+iv;         // component-wise addition
↳ 2,6,10,14,16,0
    iv[size(iv)-1]; // last-1 entry
↳ 8
    intvec iw=2,3,4,0;
    iv==iw;       // lexicographic comparison
↳ 0
    iv < iw;
↳ 1
    iv != iw;
↳ 1
    iv[2];
↳ 3
    iw = 4,1,2;
    iv[iw];
↳ 7 1 3

```

4.8.4 intvec related functions

hilb	Hilbert series as intvec (see Section 5.1.56 [hilb], page 193)
indepSet	sets of independent variables of an ideal (see Section 5.1.61 [indepSet], page 196)
leadexp	the exponent vector of the leading monomial (see Section 5.1.77 [leadexp], page 207)
monomial	the power product corresponding to the exponent vector (see Section 5.1.96 [monomial], page 221)
nrows	number of rows (see Section 5.1.106 [nrows], page 228)
qhweight	quasihomogeneous weights (see Section 5.1.122 [qhweight], page 241)
size	length of the intvec (see Section 5.1.142 [size], page 259)
sortvec	permutation for sorting ideals/modules (see Section 5.1.144 [sortvec], page 261)
transpose	transpose of an intvec, returns an intmat (see Section 5.1.157 [transpose], page 276)
weight	weights for the weighted ecart method (see Section 5.1.170 [weight], page 283)

4.9 link

Links are the communication channels of SINGULAR, i.e., something SINGULAR can write to and/or read from. Currently, SINGULAR supports four different link types:

- ASCII links (see Section 4.9.4 [ASCII links], page 95)
- ssi links (see Section 4.9.5 [Ssi links], page 96)
- pipe links (see Section 4.9.6 [Pipe links], page 99)
- DBM links (see Section 4.9.7 [DBM links], page 99)

4.9.1 link declarations

Syntax: `link name = string_expression ;`

Purpose: defines a new communication link.

Default: none

Example:

```

link l=":w example.txt";
int i=22;           // cf. ASCII links for explanation
string s="An int follows:";
write(l,s,i);
l;
↳ // type : ASCII
↳ // mode : w
↳ // name : example.txt
↳ // open : yes
↳ // read : not ready
↳ // write: ready
  close(l);        //
  read(l);
↳ An int follows:
↳ 22
↳
  close(l);

```

4.9.2 link expressions

A link expression is:

1. an identifier of type link
2. a string describing the link

A link is described by a string which consists of two parts: a property string followed by a name string. The property string describes the type of the link (`ASCII`, `ssi` or `DBM`) and the mode of the link (e.g., open for read, write or append). The name string describes the filename of the link, resp. a network connection for ssi links.

For a detailed format description of the link describing string see:

- for ASCII links: Section 4.9.4 [ASCII links], page 95
- ssi links (see Section 4.9.5 [Ssi links], page 96)
- pipe links (see Section 4.9.6 [Pipe links], page 99)
- for DBM links: Section 4.9.7 [DBM links], page 99

4.9.3 link related functions

close	closes a link (see Section 5.1.10 [close], page 161)
dump	generates a dump of all variables and their values (see Section 5.1.27 [dump], page 173)
getdump	reads a dump (see Section 5.1.52 [getdump], page 189)
open	opens a link (see Section 5.1.109 [open], page 230)
read	reads from a link (see Section 5.1.128 [read], page 245)
status	gets the status of a link (see Section 5.1.148 [status], page 265)
write	writes to a link (see Section 5.1.172 [write], page 284)
kill	closes and kills a link (see Section 5.1.71 [kill], page 204)
waitall	wait till all links of a list of links become ready (only ssi:tcp links) (see Section 5.1.167 [waitall], page 281)
waitfirst	wait till at least one link of a list of links become ready (only ssi:tcp links) (see Section 5.1.168 [waitfirst], page 282)

4.9.4 ASCII links

Via ASCII links data that can be converted to a string can be written into files for storage or communication with other programs. The data is written in plain ASCII format. The output format of polynomials is done w.r.t. the value of the global variable `short` (see Section 5.3.7 [short], page 300). Reading from an ASCII link returns a string — conversion into other data is up to the user. This can be done, for example, using the command `execute` (see Section 5.1.32 [execute], page 176).

The ASCII link describing string has to be one of the following:

1. "ASCII: " + filename
the mode (read or append) is set by the first `read` or `write` command.
2. "ASCII:r " + filename
opens the file for reading.
3. "ASCII:w " + filename
opens the file for overwriting.
4. "ASCII:a " + filename
opens the file for appending.

There are the following default values:

- the type `ASCII` may be omitted since ASCII links are the default links.
- if non of `r`, `w`, or `a` is specified, the mode of the link is set by the first `read` or `write` command on the link. If the first command is `write`, the mode is set to `a` (append mode).
- if the filename is omitted, `read` reads from `stdin` and `write` writes to `stdout`.

Using these default rules, the string `":r temp"` describes a link which is equivalent to the link `"ASCII:r temp"`: an ASCII link to the file `temp` which is opened for reading. The string `"temp"` describes an ASCII link to the file `temp`, where the mode is set by the first `read` or `write` command. See also the example below.

Note that the filename may contain a path. On Microsoft Windows (resp. MS-DOS) platforms, names of a drive can precede the filename, but must be started with a `//` (as in `//c/temp/ex`. An

ASCII link can be used either for reading or for writing, but not for both at the same time. A `close` command must be used before a change of I/O direction. Types without a conversion to `string` cannot be written.

Example:

```

ring r=32003,(x,y,z),dp;
link l=":w example.txt"; // type is ASCII, mode is overwrite
l;
⇒ // type : ASCII
⇒ // mode : w
⇒ // name : example.txt
⇒ // open : no
⇒ // read : not ready
⇒ // write: not ready
  status(l, "open", "yes"); // link is not yet opened
⇒ 0
  ideal i=x2,y2,z2;
  write (l,1,";"2,";"ideal i="i,"");
  status(l, "open", "yes"); // now link is open
⇒ 1
  status(l, "mode"); // for writing
⇒ w
  close(l); // link is closed
  write("example.txt","int j=5;");// data is appended to file
  read("example.txt"); // data is returned as string
⇒ 1
⇒ ;
⇒ 2
⇒ ;
⇒ ideal i=
⇒ x2,y2,z2;
⇒ int j=5;
⇒
  execute(read(l)); // read string is executed
⇒ 1
⇒ 2
⇒ // ** redefining i (ideal i=) ./examples/ASCII_links.sing:14
  close(l); // link is closed

```

4.9.5 Ssi links

Ssi (simple singular interface) links give the possibility to store and communicate data betweenm Singular processes: Read and write access is very fast compared to ASCII links. Ssi links can be established using files or using TCP sockets. For ring-dependent data, a ring description is written together with the data. Reading from an Ssi link returns an expression (not a string) which was evaluated after the read operation. If the expression read from an Ssi link is not from the same ring as the current ring, then a `read` changes the current ring.

Currently under development - not everything is implemtented.

4.9.5.1 Ssi file links

Ssi file links provide the possibility to store data in a file using the ssi format. For storing large amounts of data, ssi file links should be used instead of ASCII links. Unlike ASCII links, data read from ssi file links is returned as expressions one at a time.

The ssi file link describing string has to be one of the following:

1. "ssi:r " + filename
opens the file for reading.
2. "ssi:w " + filename
opens the file for overwriting.
3. "ssi:a " + filename
opens the file for appending.

Note that the filename may contain a path. An ssi file link can be used either for reading or for writing, but not for both at the same time. A `close` command must be used before a change of I/O direction.

Example:

```

ring r;
link l="ssi:w example.ssi"; // type=ssi, mode=overwrite
l;
⇒ // type : ssi
⇒ // mode : w
⇒ // name : example.ssi
⇒ // open : no
⇒ // read : not open
⇒ // write: not open
ideal i=x2,y2,z2;
write (l,1, i, "hello world");// write three expressions
write(l,4); // append one more expression
close(l); // link is closed
// open the file for reading now
read(l); // only first expression is read
⇒ 1
kill r; // no basering active now
def i = read(l); // second expression
// notice that current ring was set, the name was assigned
// automatically
listvar(ring);
⇒ // ssiRing0 [0] *ring
⇒ // ZZ [0] cring
⇒ // QQ [0] cring
def s = read(l); // third expression
listvar();
⇒ // s [0] string hello world
⇒ // ssiRing0 [0] *ring
⇒ // i [0] ideal, 3 generator(s)
⇒ // l [0] link
close(l); // link is closed

```

4.9.5.2 Ssi tcp links

Ssi tcp links give the possibility to exchange data between two processes which may run on the same or on different computers. Ssi tcp links can be opened in four different modes:

tcp SINGULAR acts as a server.

connect SINGULAR acts as a client.

tcp <host>:<program>

SINGULAR acts as a client, launching an application as server. This requires `ssh/ssh` to be installed on the computers (and preferably an automatic login via `.ssh/authorized_keys`).

fork SINGULAR acts as a client, forking another SINGULAR as server.

The Ssi tcp link describing string has to be

- tcp mode:

1. `"ssi:tcp"`

SINGULAR becomes a server and waits at the first free port (>1024) for a connect call.

- connect mode:

2. `"ssi:connect " + host:port`

SINGULAR becomes a client and connects to a server waiting at the host and port.

- launch mode:

4. `"ssi:tcp" + host:application`

SINGULAR becomes a client and starts (launches) the application using `ssh` on a (possibly) different host which then acts as a server.

- fork mode:

8. `"ssi:fork"`

SINGULAR becomes a client and forks another SINGULAR on the same host which acts as a server.

To open an ssi tcp link in launch mode, the application to launch must either be given with an absolute pathname, or must be in a directory contained in the search path. The launched application acts as a server, whereas the SINGULAR that actually opened the link acts as a client. The client "listens" at the some free port until the server application does a connect call.

If the ssi tcp link is opened in fork mode a child of the current SINGULAR is forked. All variables and their values are inherited by the child. The child acts as a server whereas the SINGULAR that actually opened the link acts as a client.

To arrange the evaluation of an expression by a server, the expression must be quoted using the command `quote` (see Section 5.1.124 [quote], page 242), so that a local evaluation is prevented. Otherwise, the expression is evaluated first, and the result of the evaluation is written, instead of the expression which is to be evaluated.

If SINGULAR is in server mode, the value of the variable `link_11` is the ssi link connecting to the client and SINGULAR is in an infinite read-eval-write loop until the connection is closed from the client side (by closing its connecting link). Reading and writing is done to the link `link_11`: After an expression is read, it is evaluated and the result of the evaluation is written back. That is, for each expression which was written to the server, there is exactly one expression written back. This might be an "empty" expression, if the evaluation on the server side does not return a value.

Ssi tcp links should explicitly be opened before being used. Ssi tcp links are bidirectional, i.e. can be used for both, writing and reading. Reading from an ssi tcp link blocks until data was written to that link. The `status` command can be used to check whether there is data to read.

Example:

```

    int i=7;
    link l = "ssi:fork";      // fork link declaration
    open(l); l;
    ↪ // type : ssi
    ↪ // mode : fork
    ↪ // name :
    ↪ // open : yes
    ↪ // read : not ready
    ↪ // write: ready

    write(l,quote(i)); // Child inherited vars and their values
    read(l);
    ↪ 7
    close(l);             // shut down forked child

```

4.9.6 Pipe links

Pipe links provide access to stdin and stdout of any program. Pipe links are bidirectional. **Syntax:** `"|: " + string_for_system`

The `string_for_system` will be passed to `system` after connecting the input and output to the corresponding stdout and stdin.

Example:

```

    link l="|: date";
    open(l); l;
    ↪ // type : pipe
    ↪ // mode :
    ↪ // name : date
    ↪ // open : yes
    ↪ // read : not ready
    ↪ // write: ready
    read(l);
    ↪ Mo 14. Dez 12:59:51 2020
    l;
    ↪ // type : pipe
    ↪ // mode :
    ↪ // name : date
    ↪ // open : yes
    ↪ // read : not ready
    ↪ // write: ready
    close(l);

```

4.9.7 DBM links

DBM links provide access to data stored in a data base. Each entry in the data base consists of a (`key_string`, `value_string`) pair. Such a pair can be inserted with the command `write(link, key_string, value_string)`. By calling `write(link, key_string)`, the entry with key `key_string` is

deleted from the data base. The value of an entry is returned by the command `read(link, key_string)`. With only one argument, `read(link)` returns the next key in the data base. Using this feature a data base can be scanned in order to access all entries of the data base.

If a data base with name `name` is opened for writing for the first time, two files (`name.pag` and `name.dir`), which contain the data base, are automatically created.

The DBM link describing string has to be one of the following:

1. "DBM: " + name
opens the data base for reading (default mode).
2. "DBM:r " + name
opens the data base for reading.
3. "DBM:rw " + name
opens the data base for reading and writing.

Note that `name` must be given without the suffix `.pag` or `.dir`. The name may contain an (absolute) path.

Example:

```

link l="DBM:rw example";
write(l,"1","abc");
write(l,"3","XYZ");
write(l,"2","ABC");
l;
⇒ // type : DBM
⇒ // mode : rw
⇒ // name : example
⇒ // open : yes
⇒ // read : ready
⇒ // write: ready
close(l);
// read all keys (till empty string):
read(l);
⇒ 1
read(l);
⇒ 3
read(l);
⇒ 2
read(l);
⇒
// read data corresponding to key "1"
read(l,"1");
⇒ abc
// read all data:
read(l,read(l));
⇒ abc
read(l,read(l));
⇒ XYZ
read(l,read(l));
⇒ ABC
// close
close(l);

```

4.10 list

Lists are arrays whose elements can be of different types (including ring). If one element belongs to a ring the whole list belongs to that ring. This applies also to the special list `#`. The expression `list()` is the empty list.

Note that a list stores the objects itself and not the names. Hence, if `L` is a list, `L[1]` for example has no name. A name, say `R`, can be created for `L[1]` by `def R=L[1];`. To store also the name of an object, say `r`, it can be added to the list with `nameof(r);`. Rings may be objects of a list.

Note: Unlike other assignments a ring as an element of a list is not a copy but another reference to the same ring.

4.10.1 list declarations

Syntax: `list name = expression_list;`
`list name = list_expression;`

Purpose: defines a list (of objects of possibly different types).

Default: empty list

Example:

```

list l=1,"str";
l[1];
↳ 1
l[2];
↳ str
ring r;
listvar(r);
↳ // r                               [0] *ring
ideal i = x^2, y^2 + z^3;
l[3] = i;
l;
↳ [1]:
↳ 1
↳ [2]:
↳ str
↳ [3]:
↳ _[1]=x2
↳ _[2]=z3+y2
listvar(r); // the list l belongs now to the ring r
↳ // r                               [0] *ring
↳ // l                               [0] list, size: 3
↳ // i                               [0] ideal, 2 generator(s)

```

4.10.2 list expressions

A list expression is:

1. the empty list `list()`
2. an identifier of type list
3. a function returning list
4. list expressions combined by the arithmetic operation `+`
5. a type cast to list

See Section 3.5.5 [Type conversion and casting], page 46.

Example:

```

list l = "hello",1;
l;
↳ [1]:
↳ hello
↳ [2]:
↳ 1
l = list();
l;
↳ empty list
ring r =0,x,dp;
factorize((x+1)^2);
↳ [1]:
↳ _[1]=1
↳ _[2]=x+1
↳ [2]:
↳ 1,2
list(1,2,3);
↳ [1]:
↳ 1
↳ [2]:
↳ 2
↳ [3]:
↳ 3

```

4.10.3 list operations

+ concatenation

delete deletes one element from list, returns new list

insert inserts or appends a new element to list, returns a new list

list_expression [int_expression]
 is a list entry; the index 1 gives the first element.

Example:

```

list l1 = 1,"hello",list(-1,1);
list l2 = list(1,5,7);
l1 + l2; // a new list
↳ [1]:
↳ 1
↳ [2]:
↳ hello
↳ [3]:
↳ [1]:
↳ -1
↳ [2]:
↳ 1
↳ [4]:
↳ 1

```



```

↳ [5]:
↳ 5
↳ [6]:
↳ 7
12 = delete(12, 2); // delete 2nd entry
12;
↳ [1]:
↳ 1
↳ [2]:
↳ 7

```

4.10.4 list related functions

bareiss returns a list of a matrix (lower triangular) and of an intvec (permutations of columns, see Section 5.1.3 [bareiss], page 156)

betti Betti numbers of a resolution (see Section 5.1.4 [betti], page 157)

delete deletion of an element from a list (see Section 5.1.21 [delete], page 169)

facstd factorizing Groebner basis algorithm (see Section 5.1.34 [facstd], page 177)

factorize
list of factors of a polynomial (see Section 5.1.36 [factorize], page 178)

insert insertion of a new element into a list (see Section 5.1.62 [insert], page 197)

minres minimization of a free resolution (see Section 5.1.93 [minres], page 220)

names list of all user-defined variable names (see Section 5.1.102 [names], page 225)

size number of entries (see Section 5.1.142 [size], page 259)

conversion from resolution
(see Section 4.18 [resolution], page 123)

4.11 map

Maps are ring maps from a preimage ring into the basering.

Note:

- The target of a map is **ALWAYS** the actual basering
- The preimage ring has to be stored "by its name", that means, maps can only be used in such contexts, where the name of the preimage ring can be resolved (this has to be considered in subprocedures). See also Section 6.5 [Identifier resolution], page 310, Section 3.7.4 [Names in procedures], page 54.

Maps between rings with different coefficient fields are possible and listed below.

Canonically realized are

- $Q \rightarrow Q(a, \dots)$ (Q : the rational numbers)
- $Q \rightarrow R$ (R : the real numbers)
- $Q \rightarrow C$ (C : the complex numbers)
- $Z/p \rightarrow (Z/p)(a, \dots)$ (Z : the integers)
- $Z/p \rightarrow GF(p^n)$ (GF : the Galois field)
- $Z/p \rightarrow R$

- $R \rightarrow C$

Possible are furthermore

- $Z/p \rightarrow Q$, $[i]_p \mapsto i \in [-p/2, p/2] \subseteq Z$
- $Z/p \rightarrow Z/p'$, $[i]_p \mapsto i \in [-p/2, p/2] \subseteq Z$, $i \mapsto [i]_{p'} \in Z/p'$
- $C \rightarrow R$, by taking the real part

Finally, in SINGULAR we allow the mapping from rings with coefficient field Q to rings whose ground fields have finite characteristic:

- $Q \rightarrow Z/p$
- $Q \rightarrow (Z/p)(a, \dots)$

In these cases the denominator and the numerator of a number are mapped separately by the usual map from Z to Z/p , and the image of the number is built again afterwards by division. It is thus not allowed to map numbers whose denominator is divisible by the characteristic of the target ground field, or objects containing such numbers. We, therefore, strongly recommend using such maps only to map objects with integer coefficients.

4.11.1 map declarations

Syntax: `map name = preimage_ring_name , ideal_expression ;`
`map name = preimage_ring_name , list_of_poly_and_ideal_expressions ;`
`map name = map_expression ;`

Purpose: defines a ring map from `preimage_ring` to `basing`.
 Maps the variables of the preimage ring to the generators of the ideal. If the ideal contains less elements than variables in the `preimage_ring` the remaining variables are mapped to 0, if the ideal contains more elements these are ignored. The image ring is always the current `basing`. For the mapping of coefficients from different fields see Section 4.11 [map], page 103.

Default: none

Note: There are standard mappings for maps which are close to the identity map: `fetch` and `imap`.

The name of a map serves as the function which maps objects from the `preimage_ring` into the `basing`. These objects must be defined by names (no evaluation in the preimage ring is possible).

Example:

```
ring r1=32003,(x,y,z),dp;
ideal i=x,y,z;
ring r2=32003,(a,b),dp;
map f=r1,a,b,a+b;
// maps from r1 to r2,
// x -> a
// y -> b
// z -> a+b
f(i);
↪ _[1]=a
↪ _[2]=b
↪ _[3]=a+b
// operations like f(i[1]) or f(i*i) are not allowed
ideal i=f(i);
```

```

// objects in different rings may have the same name
map g = r2,a2,b2;
map phi = g(f);
// composition of map f and g
// maps from r1 to r2,
// x -> a2
// y -> b2
// z -> a2+b2
phi(i);
↳ _[1]=a2
↳ _[2]=b2
↳ _[3]=a2+b2

```

See Section 5.1.38 [fetch], page 180; Section 4.5.2 [ideal expressions], page 78; Section 5.1.59 [imap], page 195; Section 4.11 [map], page 103; Section 4.19 [ring], page 124.

4.11.2 map expressions

A map expression is:

1. an identifier of type map
2. a function returning map
3. map expressions combined by composition using parentheses (,)

4.11.3 map operations

() composition of maps. If, for example, f and g are maps, then $f(g)$ is a map expression giving the composition $f \circ g$ of f and g , provided the target ring of g is the basering of f .

map_expression [int_expressions]
 is a map entry (the image of the corresponding variable)

Example:

```

ring r=0,(x,y),dp;
map f=r,y,x; // the map f permutes the variables
f;
↳ f[1]=y
↳ f[2]=x
poly p=x+2y3;
f(p);
↳ 2x3+y
map g=f(f); // the map g defined as f^2 is the identity
g;
↳ g[1]=x
↳ g[2]=y
g(p) == p;
↳ 1

```

4.11.4 map related functions

fetch the identity map between rings (see Section 5.1.38 [fetch], page 180)

imap a convenient map procedure for inclusions and projections of rings (see Section 5.1.59 [imap], page 195)

preimage preimage under a ring map (see Section 5.1.116 [preimage], page 236)

subst substitution of a ring variable (see Section 5.1.152 [subst], page 269)

See also the libraries Section D.4.2 [algebra_lib], page 809 and Section D.2.12 [ring_lib], page 803, which contain more functions, related to maps.

4.12 matrix

Objects of type matrix are matrices with polynomial entries. Like polynomials they can only be defined or accessed with respect to a basering. In order to compute with matrices having integer or rational entries, define a ring with characteristic 0 and at least one variable.

A matrix can be multiplied by and added to a poly; in this case the polynomial is converted into a matrix of the right size with the polynomial on the diagonal.

If A is a matrix then the assignment `module M=A;` or `module M=module(A);` creates a module generated by the columns of A. Note that the trailing zero columns of A may be deleted by module operations with M.

4.12.1 matrix declarations

Syntax: `matrix name[rows][cols] = list_of_poly_expressions ;`
`matrix name = matrix_expression ;`

Purpose: defines a matrix (of polynomials).

The given poly_list fills up the matrix beginning with the first row from the left to the right, then the second row and so on. If the poly_list contains less than rows*cols elements, the matrix is filled up with zeros; if it contains more elements, then only the first rows*cols elements are used. If the right-hand side is a matrix expression the matrix on the left-hand side gets the same size as the right-hand side, otherwise the size is determined by the left-hand side. If the size is omitted a 1x1 matrix is created.

Default: 0 (1 x 1 matrix)

Example:

```
int ro = 3;
ring r = 32003, (x,y,z).dp;
poly f=xyz;
poly g=z*f;
ideal i=f,g,g^2;
matrix m[ro][3] = x3y4, 0, i, f ; // a 3 x 3 matrix
m;
↪ m[1,1]=x3y4
↪ m[1,2]=0
↪ m[1,3]=xyz
↪ m[2,1]=xyz2
↪ m[2,2]=x2y2z4
↪ m[2,3]=xyz
↪ m[3,1]=0
↪ m[3,2]=0
↪ m[3,3]=0
print(m);
```

```

↳ x3y4,0,      xyz,
↳ xyz2,x2y2z4,xyz,
↳ 0,  0,      0
  matrix A; // the 1 x 1 zero matrix
  matrix B[2][2] = m[1..2, 2..3]; //defines a submatrix
  print(B);
↳ 0,      xyz,
↳ x2y2z4,xyz
  matrix C=m; // defines C as a 3 x 3 matrix equal to m
  print(C);
↳ x3y4,0,      xyz,
↳ xyz2,x2y2z4,xyz,
↳ 0,  0,      0

```

4.12.2 matrix expressions

A matrix expression is:

1. an identifier of type matrix
2. a function returning matrix
3. matrix expressions combined by the arithmetic operations +, - or *
4. a type cast to matrix (see Section 4.12.3 [matrix type cast], page 107)

Example:

```

ring r=0,(x,y),dp;
poly f= x3y2 + 2x2y2 +2;
matrix H = jacob(jacob(f)); // the Hessian of f
matrix mc = coef(f,y);
print(mc);
↳ y2,  1,
↳ x3+2x2,2
module MD = [x+y,1,x],[x+y,0,y];
matrix M = MD;
print(M);
↳ x+y,x+y,
↳ 1,  0,
↳ x,  y

```

4.12.3 matrix type cast

Syntax: matrix (expression)
matrix (expression, int_n, int_m)

Type: matrix

Purpose: Converts expression to a matrix, where expression must be of type int, intmat, intvec, number, poly, ideal, vector, module, or matrix. If int_n and int_m are supplied, then they specify the dimension of the matrix. Otherwise, the size (resp. dimensions) of the matrix is determined by the size (resp. dimensions) of the expression.

Example:

```

ring r=32003,(x,y,z),dp;
matrix(x);

```

```

↳ _[1,1]=x
   matrix(x, 1, 2);
↳ _[1,1]=x
↳ _[1,2]=0
   matrix(intmat(intvec(1,2,3,4), 2, 2));
↳ _[1,1]=1
↳ _[1,2]=2
↳ _[2,1]=3
↳ _[2,2]=4
   matrix(_, 2, 3);
↳ _[1,1]=1
↳ _[1,2]=2
↳ _[1,3]=0
↳ _[2,1]=3
↳ _[2,2]=4
↳ _[2,3]=0
   matrix(_, 2, 1);
↳ _[1,1]=1
↳ _[2,1]=3

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.7.3 [intmat type cast], page 89; Section 4.12 [matrix], page 106.

4.12.4 matrix operations

- + addition with matrix or poly; the polynomial is converted into a diagonal matrix
- negation or subtraction with matrix or poly (the first operand is expected to be a matrix); the polynomial is converted into a diagonal matrix
- * multiplication with matrix or poly; the polynomial is converted into a diagonal matrix
- / division by poly
- ==, <>, != comparators

matrix-expression [int-expression, int-expression]

is a matrix entry, where the first index indicates the row and the second the column

Example:

```

ring r=32003,x,dp;
matrix A[3][3] = 1,3,2,5,0,3,2,4,5; // define a matrix
print(A); // nice printing of small matrices
↳ 1,3,2,
↳ 5,0,3,
↳ 2,4,5
   A[2,3]; // matrix entry
↳ 3
   A[2,3] = A[2,3] + 1; // change entry
   A[2,1..3] = 1,2,3; // change 2nd row
print(A);
↳ 1,3,2,
↳ 1,2,3,
↳ 2,4,5
   matrix E[3][3]; E = E + 1; // the unit matrix

```

```

    matrix B =x*E - A;
    print(B);
    ↪ x-1,-3, -2,
    ↪ -1, x-2,-3,
    ↪ -2, -4, x-5
    // the same (but x-A does not work):
    B = -A+x;
    print(B);
    ↪ x-1,-3, -2,
    ↪ -1, x-2,-3,
    ↪ -2, -4, x-5
    det(B);          // the characteristic polynomial of A
    ↪ x3-8x2-2x-1
    A*A*A - 8 * A*A - 2*A == E; // Cayley-Hamilton
    ↪ 1
    vector v =[x,-1,x2];
    A*v; // multiplication of matrix and vector
    ↪ _[1,1]=2x2+x-3
    ↪ _[2,1]=3x2+x-2
    ↪ _[3,1]=5x2+2x-4
    matrix m[2][2]=1,2,3;
    print(m-transpose(m));
    ↪ 0,-1,
    ↪ 1,0

```

4.12.5 matrix related functions

<code>bareiss</code>	Gauss-Bareiss algorithm (see Section 5.1.3 [bareiss], page 156)
<code>coef</code>	matrix of coefficients and monomials (see Section 5.1.11 [coef], page 162)
<code>coeffs</code>	matrix of coefficients (see Section 5.1.12 [coeffs], page 163)
<code>det</code>	determinant (see Section 5.1.23 [det], page 170)
<code>diff</code>	partial derivative (see Section 5.1.24 [diff], page 171)
<code>jacob</code>	Jacobi matrix (see Section 5.1.66 [jacob], page 200)
<code>koszul</code>	Koszul matrix (see Section 5.1.73 [koszul], page 205)
<code>lift</code>	lift-matrix (see Section 5.1.80 [lift], page 209)
<code>liftstd</code>	standard basis and transformation matrix computation (see Section 5.1.81 [liftstd], page 209)
<code>minor</code>	set of minors of a matrix (see Section 5.1.92 [minor], page 218)
<code>ncols</code>	number of columns (see Section 5.1.103 [ncols], page 227)
<code>nrows</code>	number of rows (see Section 5.1.106 [nrows], page 228)
<code>print</code>	nice print format (see Section 5.1.119 [print], page 238)
<code>size</code>	number of matrix entries (see Section 5.1.142 [size], page 259)
<code>subst</code>	substitute a ring variable (see Section 5.1.152 [subst], page 269)
<code>trace</code>	trace of a matrix (see Section 5.1.156 [trace], page 276)

transpose

transposed matrix (see Section 5.1.157 [transpose], page 276)

wedge

wedge product (see Section 5.1.169 [wedge], page 282)

See also the library Section D.3.1 [matrix_lib], page 806, which contains more matrix-related functions.

4.13 module

Modules are submodules of a free module over the basering with basis `gen(1)`, `gen(2)`, `...`. They are represented by lists of vectors which generate the submodule. Like vectors they can only be defined or accessed with respect to a basering.

If R is the basering, and M is a submodule of R^n

generated by vectors v_1, \dots, v_k , then v_1, \dots, v_k

may be considered as the generators of relations of R^n/M between the canonical generators `gen(1), ..., gen(n)`. Hence any finitely generated R -module can be represented in SINGULAR by its module of relations. The assignments `module M=v1,...,vk; matrix A=M;` create the presentation matrix of size $n \times k$ for R^n/M , i.e., the columns of A are the vectors v_1, \dots, v_k which generate M (cf. Section B.1 [Representation of mathematical objects], page 759).

4.13.1 module declarations

Syntax: `module name = list_of_vector_expressions ;`
`module name = module_expression ;`

Purpose: defines a module.

Default: `[0]`

Example:

```
ring r=0,(x,y,z),(c,dp);
vector s1 = [x2,y3,z];
vector s2 = [xy,1,0];
vector s3 = [0,x2-y2,z];
poly f = xyz;
module m = s1, s2-s1,f*(s3-s1);
m;
↳ m[1]=[x2,y3,z]
↳ m[2]=[-x2+xy,-y3+1,-z]
↳ m[3]=[-x3yz,-xy4z+x3yz-xy3z]
// show m in matrix format (columns generate m)
print(m);
↳ x2,-x2+xy,-x3yz,
↳ y3,-y3+1,-xy4z+x3yz-xy3z,
↳ z,-z, 0
```

4.13.2 module expressions

A module expression is:

1. an identifier of type module
2. a function returning module
3. module expressions combined by the arithmetic operation +

4. multiplication of a module expression with an ideal or a poly expression: *
5. a type cast to module

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.5 [ideal], page 78; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

4.13.3 module operations

- + addition (concatenation of the generators and simplification)
- * multiplication with ideal or poly (but not ‘module’ * ‘module!’)

`module_expression [int_expression , int_expression]`

is a module entry, where the first index indicates the row and the second the column

`module_expressions [int_expression]`

is a vector, where the index indicates the column (generator)

Example:

```

ring r=0,(x,y,z),dp;
module m=[x,y],[0,0,z];
print(m*(x+y));
↳ x2+xy,0,
↳ xy+y2,0,
↳ 0,    xz+yz
// this is not distributive:
print(m*x+m*y);
↳ x2,0, xy,0,
↳ xy,0, y2,0,
↳ 0, xz,0, yz

```

4.13.4 module related functions

- `coeffs` matrix of coefficients (see Section 5.1.12 [coeffs], page 163)
- `degree` multiplicity, dimension and codimension of the module of leading terms (see Section 5.1.20 [degree], page 169)
- `diff` partial derivative (see Section 5.1.24 [diff], page 171)
- `dim` Krull dimension of free module over the basering modulo the module of leading terms (see Section 5.1.25 [dim], page 171)
- `eliminate` elimination of variables (see Section 5.1.28 [eliminate], page 174)
- `freemodule` the free module of given rank (see Section 5.1.47 [freemodule], page 186)
- `fres` free resolution of a standard basis (see Section 5.1.48 [fres], page 186)
- `groebner` Groebner basis computation (a wrapper around `std`, `stdhilb`, `stdfglm`,...) (see [groebner], page 785)
- `hilb` Hilbert function of a standard basis (see Section 5.1.56 [hilb], page 193)
- `homog` homogenization with respect to a variable (see Section 5.1.57 [homog], page 194)

interred	interreduction of a module (see Section 5.1.64 [interred], page 199)
intersect	module intersection (see Section 5.1.65 [intersect], page 199)
jet	Taylor series up to a given order (see Section 5.1.68 [jet], page 201)
kbase	vector space basis of free module over the basering modulo the module of leading terms (see Section 5.1.69 [kbase], page 203)
lead	initial module (see Section 5.1.75 [lead], page 206)
lift	lift-matrix (see Section 5.1.80 [lift], page 209)
liftstd	standard basis and transformation matrix computation (see Section 5.1.81 [liftstd], page 209)
lres	free resolution (see Section 5.1.83 [lres], page 212)
minbase	minimal generating set of a homogeneous ideal, resp. module, or an ideal, resp. module, over a local ring
modulo	represents $(h_1 + h_2)/h_1 = h_2/(h_1 \cap h_2)$ (see Section 5.1.94 [modulo], page 220)
mres	minimal free resolution of an ideal resp. module w.r.t. a minimal set of generators of the given module (see Section 5.1.98 [mres], page 222)
mult	multiplicity, resp. degree, of the module of leading terms (see Section 5.1.100 [mult], page 224)
nres	computation of a free resolution of an ideal resp. module M which is minimized from the second free module on (see Section 5.1.105 [nres], page 228)
ncols	number of columns (see Section 5.1.103 [ncols], page 227)
nrows	number of rows (see Section 5.1.106 [nrows], page 228)
print	nice print format (see Section 5.1.119 [print], page 238)
prune	minimization of the embedding into a free module (see Section 5.1.121 [prune], page 241)
qhweight	quasihomogeneous weights of an ideal, resp. module (see Section 5.1.122 [qhweight], page 241)
quotient	module quotient (see Section 5.1.125 [quotient], page 243)
reduce	normalform with respect to a standard basis (see Section 5.1.129 [reduce], page 246)
res	free resolution of an ideal, resp. module, but not changing the given ideal, resp. module (see [res], page 785)
simplify	simplification of a set of vectors (see Section 5.1.141 [simplify], page 258)
size	number of non-zero generators (see Section 5.1.142 [size], page 259)
sortvec	permutation for sorting ideals/modules (see Section 5.1.144 [sortvec], page 261)
sres	free resolution of a standard basis (see Section 5.1.147 [sres], page 264)
std	standard basis computation (see Section 5.1.149 [std], page 266, Section 5.1.81 [liftstd], page 209)
subst	substitution of a ring variable (see Section 5.1.152 [subst], page 269)
syz	computation of the first syzygy module (see Section 5.1.154 [syz], page 275)

vdim vector space dimension of free module over the basering modulo module of leading terms (see Section 5.1.166 [vdim], page 281)

weight "optimal" weights (see Section 5.1.170 [weight], page 283)

4.14 number

Numbers are elements from the coefficient ring (or ground ring). They can only be defined or accessed with respect to a basering which determines the coefficient field. See Section 4.19.2 [ring declarations], page 124 for declarations of coefficient fields.

Warning: Beware of the special meaning of the letter *e* (immediately following a sequence of digits) if the field is real (or complex), Section 6.4 [Miscellaneous oddities], page 308.

4.14.1 number declarations

Syntax: number name = number_expression ;

Purpose: defines a number.

Default: 0

Note: Numbers may only be declared w.r.t. the coefficient field of the current basering, i.e., a ring has to be defined prior to any number declaration. See Section 3.3 [Rings and orderings], page 30 for a list of the available coefficient fields.

Example:

```

// finite field Z/p, p<= 32003
ring r = 32003,(x,y,z),dp;
number n = 4/6;
n;
↳ -10667
// finite field GF(p^n), p^n <= 32767
// z is a primitive root of the minimal polynomial
ring rg= (7^2,z),x,dp;
number n = 4/9+z;
n;
↳ z38
// the rational numbers
ring r0 = 0,x,dp;
number n = 4/6;
n;
↳ 2/3
// algebraic extensions of Z/p or Q
ring ra=(0,a),x,dp;
minpoly=a^2+1;
number n=a3+a2+2a-1;
n;
↳ (a-2)
a^2;
↳ -1
// transcendental extensions of Z/p or Q
ring rt=(0,a),x,dp;
number n=a3+a2+2a-1;
n;

```



```

    2 /2;    // the notation of / for div might change in the future
    ↪ // ** int division with '/': use 'div' instead in line >> 2 /2;    // the\
        notation of / for div might change in the future<<
    ↪ 1
    ring r0=0,x,dp;
    2/3, 4/8, 2/2 ; // are numbers
    ↪ 2/3 1/2 1

    poly f = 2x2 +1;
    leadcoef(f);
    ↪ 2
    typeof(_);
    ↪ number
    ring rr =real,x,dp;
    1.7e-2; 1.7e+2; // are valid (but 1.7e2 not), if the field is 'real'
    ↪ (1.700e-02)
    ↪ (1.700e+02)
    ring rp = (31,t),x,dp;
    2/3, 4/8, 2/2 ; // are numbers
    ↪ 11 -15 1
    poly g = (3t2 +1)*x2 +1;
    leadcoef(g);
    ↪ (3t2+1)
    typeof(_);
    ↪ number
    par(1);
    ↪ (t)
    typeof(_);
    ↪ number

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.19 [ring], page 124.

4.14.3 number operations

+	addition
-	negation or subtraction
*	multiplication
/	division
%, mod	modulo
^, **	power, exponentiation (by an integer)
<=, >=, ==, <>	comparison
mod	integer modulo (the remainder of the division div), always non-negative

Note: Quotient and exponentiation is only recognized as a number expression if it is already a number, see Section 6.4 [Miscellaneous oddities], page 308.

For the behavior of comparison operators in rings with ground field different from real or the rational numbers, see Section 4.6.5 [boolean expressions], page 86.

Example:

```

    ring r=0,x,dp;
    number n = 1/2 +1/3;
    n;
    ↪ 5/6
    n/2;
    ↪ 5/12
    1/2/3;
    ↪ 1/6
    1/2 * 1/3;
    ↪ 1/6
    n = 2;
    n^-2;
    ↪ 1/4
    // the following oddities appear here
    2/(2+3);
    ↪ // ** int division with '/': use 'div' instead in line >> 2/(2+3);<<
    ↪ 0
    number(2)/(2+3);
    ↪ 2/5
    2^-2; // for int's exponent must be non-negative
    ↪ ? exponent must be non-negative
    ↪ ? error occurred in or before ./examples/number_operations.sing line 1\
    2: ' 2^-2; // for int's exponent must be non-negative'
    number(2)^-2;
    ↪ 1/4
    3/4>=2/5;
    ↪ 1
    2/6==1/3;
    ↪ 1

```

4.14.4 number related functions

cleardenom

cancellation of denominators of numbers in polyomial and divide it by its content (see Section 5.1.9 [cleardenom], page 161)

impart imaginary part of a complex number, 0 otherwise (see Section 5.1.60 [impart], page 196, Section 5.1.131 [repart], page 248)

numerator, denominator

the numerator/denominator of a rational number (see Section 5.1.107 [numerator], page 229, Section 5.1.22 [denominator], page 170)

leadcoef coefficient of the leading term (see Section 5.1.76 [leadcoef], page 207)

par n-th parameter of the basering (see Section 5.1.113 [par], page 235)

pardeg degree of a number in ring parameters (see Section 5.1.114 [pardeg], page 235)

parstr string form of ring parameters (see Section 5.1.115 [parstr], page 236)

repart real part of a complex number (see Section 5.1.60 [impart], page 196, Section 5.1.131 [repart], page 248)

4.15 package

The data type package is used to group identifiers into collections. It is mainly used as an internal means to avoid collisions of names of identifiers in libraries with variable names defined by the user. The most important package is the toplevel package, called `Top`. It contains all user defined identifiers as well as all user accessible library procedures. Identifiers which are local to a library are contained in a package whose name is obtained from the name of the library, where the first letter is converted to uppercase, the remaining ones to lowercase. Another reserved package name is `Current` which denotes the current package name in use. See also Section 3.8 [Libraries], page 54.

4.15.1 package declarations

Syntax: package name ;

Purpose: defines a package (Only relevant in very special situations).

Example:

```

package Test;
int i=3; exportto(Test,i);
Test::i+2;
↳ 5
i;
↳ ? 'i' is undefined
↳ ? error occurred in or before ./examples/package_declarations.sing 1
e 4: ' i;'
listvar();
listvar(Test);
↳ // Test [0] package Test (N)
↳ // ::i [0] int 3
package dummy = Test;
kill Test;
listvar(dummy);
↳ // dummy [0] package dummy (N)
↳ // ::i [0] int 3

```

4.15.2 package related functions

`exportto` transfer an identifier to the specified package (see Section 5.2.7 [exportto], page 288)

`importfrom`

generate a copy of an identifier from the specified package in the current package (see Section 5.2.10 [importfrom], page 291)

`listvar` list variables currently defined in a given package (see Section 5.1.82 [listvar], page 210)

`load` load a library or dynamic module (see Section 5.2.12 [load], page 294)

`LIB` load a library or dynamic module (see Section 5.1.79 [LIB], page 208)

4.16 poly

Polynomials are the basic data for all main algorithms in SINGULAR. They consist of finitely many terms (coefficient*monomial) which are combined by the usual polynomial operations (see Section 4.16.2 [poly expressions], page 118). Polynomials can only be defined or accessed with respect to a basering which determines the coefficient type, the names of the indeterminates and the monomial ordering.

```
ring r=32003,(x,y,z),dp;
poly f=x3+y5+z2;
```

4.16.1 poly declarations

Syntax: poly name = poly_expression ;

Purpose: defines a polynomial.

Default: 0

Example:

```
ring r = 32003,(x,y,z),dp;
poly s1 = x3y2+151x5y+186xy6+169y9;
poly s2 = 1*x^2*y^2*z^2+3z8;
poly s3 = 5/4x4y2+4/5*x*y^5+2x2y2z3+y7+11x10;
int a,b,c,t=37,5,4,1;
poly f=3*x^a+x*y^(b+c)+t*x^a*y^b*z^c;
f;
↳ x37y5z4+3x37+xy9
short = 0;
f;
↳ x^37*y^5*z^4+3*x^37+x*y^9
```

Section 5.3.7 [short], page 300

4.16.2 poly expressions

A polynomial expression is (optional parts in square brackets):

1. a monomial (there are NO spaces allowed inside a monomial)

[coefficient] ring_variable [exponent] [ring_variable [exponent] ...].

Monomials which contain an indexed ring variable must be built from ring_variable and coefficient with the operations * and ^

2. an identifier of type poly
3. a function returning poly
4. polynomial expressions combined by the arithmetic operations +, -, *, /, or ^
5. an int expression (see Section 3.5.5 [Type conversion and casting], page 46)
6. a type cast to poly

Example:

```
ring S=0,(x,y,z,a(1)),dp;
2x, x3, 2x2y3, xyz, 2xy2; // are monomials
2*x, x^3, 2*x^2*y^3, x*y*z, 2*x*y^2; // are poly expressions
2*a(1); // is a valid polynomial expression (a(1) is a name of a variable),
// but not 2a(1) (is a syntax error)
2*x^3; // is a valid polynomial expression equal to 2x3 (a valid monomial)
// but not equal to 2x^3 which will be interpreted as (2x)^3
// since 2x is a monomial
ring r=0,(x,y),dp;
poly f = 10x2y3 +2x2y2-2xy+y -x+2;
lead(f);
```



```

↳ 10x2y3
  leadmonom(f);
↳ x2y3
  simplify(f,1);      // normalize leading coefficient
↳ x2y3+1/5x2y2-1/5xy-1/10x+1/10y+1/5
  poly g = 1/2x2 + 1/3y;
  cleardenom(g);
↳ 3x2+2y
  int i = 102;
  poly(i);
↳ 102
  typeof(_);
↳ poly

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.19 [ring], page 124.

4.16.3 poly operations

+	addition
-	negation or subtraction
*	multiplication
/, div	division by a polynomial, ignoring the remainder (only implemented for polynomials over QQ, ZZ/p and field extensions of them) (See also Section 5.1.125 [quotient], page 243, Section 5.1.26 [division], page 172, Section 5.1.129 [reduce], page 246)
%, mod	the remainder from the division by a polynomial (only implemented for polynomials over QQ, ZZ/p and field extensions of them) (See also Section 5.1.125 [quotient], page 243, Section 5.1.26 [division], page 172, Section 5.1.129 [reduce], page 246)
~, **	power by a positive integer
<, <=, >, >=, ==, <>	comparators (considering leading monomials w.r.t. monomial ordering)
poly_expression [intvec_expression]	the sum of monomials at the indicated places w.r.t. the monomial ordering

Example:

```

ring R=0, (x,y), dp;
poly f = x3y2 + 2x2y2 + xy - x + y + 1;
f;
↳ x3y2+2x2y2+xy-x+y+1
  f + x5 + 2;
↳ x5+x3y2+2x2y2+xy-x+y+3
  f * x2;
↳ x5y2+2x4y2+x3y-x3+x2y+x2
  (x+y)/x;
↳ 1
  f/3x2;
↳ 1/3xy2+2/3y2
  x5 > f;

```

```

↳ 1
  x<=y;
↳ 0
  x>y;
↳ 1
  ring r=0,(x,y),ds;
  poly f = fetch(R,f);
  f;
↳ 1-x+y+xy+2x2y2+x3y2
  x5 > f;
↳ 0
  f[2..4];
↳ -x+y+xy
  size(f);
↳ 6
  f[size(f)+1]; f[-1]; // monomials out of range are 0
↳ 0
↳ 0
  intvec v = 6,1,3;
  f[v]; // the polynom built from the 1st, 3rd and 6th monomial of f
↳ 1+y+x3y2

```

4.16.4 poly related functions

<code>cleardenom</code>	cancellation of denominators of numbers in polynomial and divide it by its content (see Section 5.1.9 [cleardenom], page 161; [content], page 798)
<code>coef</code>	matrix of coefficients and monomials (see Section 5.1.11 [coef], page 162)
<code>coeffs</code>	matrix of coefficients (see Section 5.1.12 [coeffs], page 163)
<code>deg</code>	degree (see Section 5.1.19 [deg], page 168)
<code>diff</code>	partial derivative (see Section 5.1.24 [diff], page 171)
<code>extgcd</code>	Bezout representation of gcd (see Section 5.1.33 [extgcd], page 176)
<code>factorize</code>	factorization of polynomial (see Section 5.1.36 [factorize], page 178)
<code>finduni</code>	univariate polynomials in a zero-dimensional ideal (see Section 5.1.43 [finduni], page 183)
<code>gcd</code>	greatest common divisor (see Section 5.1.50 [gcd], page 188)
<code>homog</code>	homogenization (see Section 5.1.57 [homog], page 194)
<code>jacob</code>	ideal, resp. matrix, of all partial derivatives (see Section 5.1.66 [jacob], page 200)
<code>lead</code>	leading term (see Section 5.1.75 [lead], page 206)
<code>leadcoef</code>	coefficient of the leading term (see Section 5.1.76 [leadcoef], page 207)
<code>leadexp</code>	the exponent vector of the leading monomial (see Section 5.1.77 [leadexp], page 207)
<code>leadmonom</code>	leading monomial (see Section 5.1.78 [leadmonom], page 208)
<code>jet</code>	monomials of degree at most k (see Section 5.1.68 [jet], page 201)

<code>ord</code>	degree of the leading monomial (see Section 5.1.111 [<code>ord</code>], page 234)
<code>qhweight</code>	quasihomogeneous weights (see Section 5.1.122 [<code>qhweight</code>], page 241)
<code>reduce</code>	normal form with respect to a standard base (see Section 5.1.129 [<code>reduce</code>], page 246)
<code>rvar</code>	test for ring variable (see Section 5.1.137 [<code>rvar</code>], page 253)
<code>simplify</code>	normalization of a polynomial (see Section 5.1.141 [<code>simplify</code>], page 258)
<code>size</code>	number of monomials (see Section 5.1.142 [<code>size</code>], page 259)
<code>subst</code>	substitution of a ring variable (see Section 5.1.152 [<code>subst</code>], page 269)
<code>trace</code>	trace of a matrix (see Section 5.1.156 [<code>trace</code>], page 276)
<code>var</code>	the indicated variable of the ring (see Section 5.1.163 [<code>var</code>], page 279)
<code>varstr</code>	variable(s) in string form (see Section 5.1.165 [<code>varstr</code>], page 280)

4.17 proc

Procedures are sequences of SINGULAR commands in a special format. They are used to extend the set of SINGULAR commands with user defined commands. Once a procedure is defined it can be used as any other SINGULAR command. Procedures may be defined by either typing them on the command line or by loading them from a file. For a detailed description on the concept of procedures in SINGULAR see Section 3.7 [Procedures], page 50. A file containing procedure definitions which comply with certain syntax rules is called a library. Such a file is loaded using the command `LIB`. For more information on libraries see Section 3.8 [Libraries], page 54.

4.17.1 proc declaration

Syntax: `[static] proc proc_name [(<parameter_list>)
<help_string>
{
 <procedure_body>
}]
[example
{
 <sequence_of_commands>
}]`

Purpose: Defines a new function, the `proc proc_name`. Once loaded in a SINGULAR session, the information provided in the help string will be displayed upon entering `help proc_name;`, while the `example` section will be executed upon entering `example proc_name;`. See Section 3.7.2 [Parameter list], page 52, Section 3.7.3 [Help string], page 53, and the example in Section 3.8.6 [Procedures in a library], page 57.

The help string, the parameter list, and the example section are optional. They are, however, mandatory for the procedures listed in the header of a library. The help string is ignored and no example section is allowed if the procedure is defined interactively, i.e., if it is not loaded from a file by the `LIB` or `load` command (see Section 5.1.79 [`LIB`], page 208 and see Section 5.2.12 [`load`], page 294).

In the body of a library, each procedure not meant to be accessible by users should be declared static. See Section 3.8.6 [Procedures in a library], page 57.

Example:

```

proc milnor_number (poly p)
{
  ideal i= std(jacob(p));
  int m_nr=vdim(i);
  if (m_nr<0)
  {
    "// not an isolated singularity";
  }
  return(m_nr);          // the value of m_nr is returned
}
ring r1=0,(x,y,z),ds;
poly p=x^2+y^2+z^5;
milnor_number(p);
↳ 4

```

See Section 5.1.79 [LIB], page 208; Section 3.8 [Libraries], page 54; Section 5.2.1 [apply], page 285.

4.17.2 proc expression

Syntax: variable_name -> { expression(s) }

Purpose: Defines a new function, within apply or for assigning.

Example:

```

apply(1..3,x->{x**2});
↳ 1 4 9

```

See Section 5.2.1 [apply], page 285; Section 4.17 [proc], page 121.

4.17.3 procs with different argument types

Syntax: branchTo (string_expression , ... proc_name)

Purpose: branch to the given procedure if the argument types matches the types given as strings (which may be empty - matching the empty argument list). The main procedure (p in the example) must be defined without an argument list, and branchTo statement must be the first statement within the procedure body.

Example:

```

proc p1(int i) { "int:",i; }
proc p21(string s) { "string:",s; }
proc p22(string s1, string s2) { "two strings:",s1,s2; }
proc p()
{ branchTo("int",p1);
  branchTo("string","string",p22);
  branchTo("string",p21);
  ERROR("not defined for these argument types");
}
p(1);
↳ int: 1
p("hu");
↳ string: hu
p("ha","ha");
↳ two strings: ha ha
p(1,"hu");

```

```

↳ ? not defined for these argument types
↳ ? leaving ::p (0)

```

See Section 4.17 [proc], page 121.

4.18 resolution

The type resolution is intended as an intermediate representation which internally retains additional information obtained during computation of resolutions. It furthermore enables the use of partial results to compute, for example, Betti numbers or minimal resolutions. Like ideals and modules, a resolution can only be defined w.r.t. a basering (see Section C.3 [Syzygies and resolutions], page 767).

Note: To access the elements of a resolution, it has to be assigned to a list. This assignment also completes computations and may therefore take time, (resp. an access directly with the brackets [,] causes implicitly a cast to a list).

4.18.1 resolution declarations

Syntax: `resolution name = resolution_expression ;`

Purpose: defines a resolution.

Default: none

Example:

```

ring R;
ideal i=z2,x;
resolution re=res(i,0);
re;
↳ 1      2      1
↳ R <--  R <--  R
↳
↳ 0      1      2
↳
  betti(re);
↳ 1,1,0,
↳ 0,1,1
  list l = re;
  l;
↳ [1]:
↳   _[1]=x
↳   _[2]=z2
↳ [2]:
↳   _[1]=-z2*gen(1)+x*gen(2)
↳ [3]:
↳   _[1]=0

```

4.18.2 resolution expressions

A resolution expression is:

1. an identifier of type resolution
2. a function returning a resolution
3. a type cast to resolution from a list of ideals, resp. modules..

See Section 3.5.5 [Type conversion and casting], page 46.

4.18.3 resolution related functions

betti	Betti numbers of a resolution (see Section 5.1.4 [betti], page 157)
fres	free resolution of a standard basis (see Section 5.1.48 [fres], page 186)
lres	free resolution (see Section 5.1.83 [lres], page 212)
minres	minimize a free resolution (see Section 5.1.93 [minres], page 220)
mres	minimal free resolution of an ideal, resp. module and a minimal set of generators of the given ideal, resp. module (see Section 5.1.98 [mres], page 222)
res	free resolution of an ideal, resp. module, but not changing the given ideal, resp. module (see [res], page 785)
sres	free resolution of a standard basis (see Section 5.1.147 [sres], page 264)

4.19 ring

Rings are used to describe properties of polynomials, ideals etc. Almost all computations in SINGULAR require a basering. For a detailed description of the concept of rings see Section 3.3 [Rings and orderings], page 30.

4.19.1 qring

SINGULAR offers the opportunity to calculate in quotient rings (factor rings), i.e., rings modulo an ideal. The ideal has to be given as a standard basis. For a detailed description of the concept of rings and quotient rings see Section 3.3 [Rings and orderings], page 30. Beside the construction, an object describing a quotient ring is of type `ring`.

See Section 4.19.5 [qring declaration], page 126.

4.19.2 ring declarations

Syntax: `ring name = (coefficients), (names_of_ring_variables), (ordering);` or
`ring name = cring [names_of_ring_variables]`

Default: `(ZZ/32003)[x,y,z]`

Purpose: declares a ring and sets it as the actual basering. The second form sets the ordering to `(dp,C)`.

For the second form: `cring` stands currently for `QQ` (the rationals), `ZZ` (the integers) or `(ZZ/m)` (the field (m prime and <2147483648) resp. ring of the integers modulo m).

The coefficients for the first form are given by one of the following:

1. a `cring` as given above
2. a non-negative `int_expression` less or equal 2147483647.
3. an `expression_list` of an `int_expression` and one or more names.
4. the name `real`
5. an `expression_list` of the name `real` and an `int_expression`.
6. an `expression_list` of the name `complex`, an optional `int_expression` and a name.
7. an `expression_list` of the name `integer`.
8. an `expression_list` of the name `integer` and following `int_expressions`.

9. an `expression_list` of the name `integer` and two `int_expressions`.

For the definition of the 'coefficients', see Section 3.3 [Rings and orderings], page 30.

'names_of_ring_variables' must be a list of names or (multi-)indexed names.

'ordering' is a list of block orderings where each block ordering is either

1. `lp`, `dp`, `Dp`, `rp`, `ls`, `ds`, `Ds`, or `rs` optionally followed by a size parameter in parentheses.
2. `wp`, `Wp`, `ws`, `Ws`, or `a` followed by a weight vector given as an `intvec_expression` in parentheses.
3. `M` followed by an `intmat_expression` in parentheses.
4. `c` or `C`.

For the definition of the orderings, see Section 3.3.3 [Term orderings], page 34, Section B.2 [Monomial orderings], page 760.

If one of `coefficients`, `names_of_ring_variables`, and `ordering` consists of only one entry, the parentheses around this entry may be omitted.

See also Section 3.3.1 [Examples of ring declarations], page 31; Section 4.19 [ring], page 124; Section 5.1.135 [ringlist], page 250.

4.19.3 ring related functions

<code>charstr</code>	description of the coefficient field of a ring (see Section 5.1.7 [charstr], page 160)
<code>keepring</code>	move ring to next upper level (see Section 5.2.11 [keepring], page 293)
<code>npars</code>	number of ring parameters (see Section 5.1.104 [npars], page 227)
<code>nvars</code>	number of ring variables (see Section 5.1.108 [nvars], page 229)
<code>ordstr</code>	monomial ordering of a ring (see Section 5.1.112 [ordstr], page 235)
<code>parstr</code>	names of all ring parameters or the name of the n-th ring parameter (see Section 5.1.115 [parstr], page 236)
<code>qring</code>	quotient ring (see Section 4.19.1 [qring], page 124)
<code>ringlist</code>	decomposition of a ring into a list of its components (see Section 5.1.135 [ringlist], page 250)
<code>setring</code>	setting of a new basering (see Section 5.1.139 [setring], page 255)
<code>varstr</code>	names of all ring variables or the name of the n-th ring variable (see Section 5.1.165 [varstr], page 280)

4.19.4 ring operations

`+` construct a new ring $k[X, Y]$ from $k_1[X]$ and $k_2[Y]$. (The sets of variables must be distinct).

`==,<>` compare two rings

Note: Concerning the ground fields k_1 and k_2 take the following guide lines into consideration:

- Neither k_1 nor k_2 may be R or C .
- If the characteristic of k_1 and k_2 differs, then one of them must be Q .
- At most one of k_1 and k_2 may have parameters.
- If one of k_1 and k_2 is an algebraic extension of Z/p it may not be defined by a `charstr` of type (p^n, a) .

Example:

```

ring R1=0,(x,y),dp;
ring R2=32003,(a,b),dp;
def R=R1+R2;
R;
↳ // coefficients: ZZ/32003
↳ // number of vars : 4
↳ //      block 1 : ordering dp
↳ //      : names x y
↳ //      block 2 : ordering dp
↳ //      : names a b
↳ //      block 3 : ordering C

```

Section D.2.12 [ring_lib], page 803

4.19.5 qring declaration

Syntax: qring name = ideal_expression ;

Default: none

Purpose: declares a quotient ring as the basering modulo ideal_expression and sets it as current basering.

Operations based on standard bases (e.g. `std,groebner`, etc., `reduce`) and functions which require a standard basis (e.g. `dim,hilb`, etc.) operated with the residue classes; all others on the polynomial objects.

Example:

```

ring r=0,(x,y,z),dp;
ideal i=xy;
qring q=std(i);
basing;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ //      block 1 : ordering dp
↳ //      : names x y z
↳ //      block 2 : ordering C
↳ // quotient ring from ideal
↳ _[1]=xy
// simplification is not immediate:
(x+y)^2;
↳ x2+2xy+y2
reduce(_,std(0));
↳ x2+y2
// polynomial and residue class:
ring R=0,(x,y),dp;
qring Q=std(y);
poly p1=x;
poly p2=x+y;
// comparing polynomial objects:
p1==p2;
↳ 0
// comparing residue classes:
reduce(p1,std(0))==reduce(p2,std(0));
↳ 1

```


4.20 smatrix

An experimental type:

Objects of type `smatrix` are (sparse) matrices with polynomial entries. Like polynomials they can only be defined or accessed with respect to a basering.

Objects of type `smatrix` can be converted to and from `matrix` and `module`. Operations are `+`, `-`, `*`, `==`, `<>`. Functions are `ncols`, `nrows`, `std`, `transpose`, `tensor`. Additional `flatten(m)` and `system("unflatten",m,col)`.

Resizing can be done via `smatrix(m,r,c)` where `m` is of type `module` or `smatrix`.

Access to single entries: `m[i,j]`

See `[flatten]`, page 806; Section 4.12 `[matrix]`, page 106; Section 4.13 `[module]`, page 110; Section 5.1.103 `[ncols]`, page 227; Section 5.1.106 `[nrows]`, page 228; Section 4.19 `[ring]`, page 124; Section 5.1.149 `[std]`, page 266; Section 5.1.155 `[tensor]`, page 275; Section 5.1.157 `[transpose]`, page 276.

4.21 string

Variables of type `string` are used for output (almost every type can be "converted" to `string`) and for creating new commands at runtime see Section 5.1.32 `[execute]`, page 176. They are also return values of certain interpreter related functions (see Section 5.1 `[Functions]`, page 154). String constants consist of a sequence of ANY characters (including newline!) between a starting `"` and a closing `"`. There is also a string constant `newline`, which is the newline character. The `+` sign "adds" strings, `""` is the empty string (hence strings form a semigroup). Strings may be used to comment the output of a computation or to give it a nice format. Strings may also be used for intermediate conversion of one type into another.

```

string s="Hi";
string s1="a string with new line at the end"+newline;
string s2="another string with new line at the end
";
s;s1;s2;
↳ Hi
↳ a string with new line at the end
↳
↳ another string with new line at the end
↳
ring r; ideal i=std(ideal(x,y^3));
"dimension of i =",dim(i)," , multiplicity of i =",mult(i);
↳ dimension of i = 1 , multiplicity of i = 3
"dimension of i = "+string(dim(i))+", multiplicity of i = "+string(mult(i));
↳ dimension of i = 1, multiplicity of i = 3
"a"+"b","c";
↳ ab c

```

A comma between two strings makes an expression list out of them (such a list is printed with a separating blank in between), while a `+` concatenates strings.

4.21.1 string declarations

Syntax: `string name = string_expression ;`
`string name = list_of_string_expressions ;`

Purpose: defines a string variable.

Default: "" (the empty string)

Example:

```
string s1="Now I know";
string s2="how to encode a \" in a string...";
string s=s1+" "+s2; // concatenation of 3 strings
s;
↳ Now I know how to encode a " in a string...
s1,s2; // 2 strings, separated by a blank in the output:
↳ Now I know how to encode a " in a string...
```

4.21.2 string expressions

A string expression is:

1. a sequence of characters between two unescaped quotes (")
2. an identifier of type string
3. a function returning string
4. a substring (using the bracket operator)
5. a type cast to string (see Section 4.21.3 [string type cast], page 128)
6. string expressions combined by the operation +.

Example:

```
// string_expression[start, length] : a substring
// (possibly filled up with blanks)
// the substring of s starting at position 2
// with a length of 4
string s="123456";
s[2,4];
↳ 2345
"abcd"[2,2];
↳ bc
// string_expression[position] : a character from a string
s[3];
↳ 3
// string_expression[position..position] :
// a substring starting at the first position up to the second
// given position
s[2..4];
↳ 2 3 4
// a function returning a string
typeof(s);
↳ string
```

See Section 3.5.5 [Type conversion and casting], page 46

4.21.3 string type cast

Syntax: string (expression [, expression_2, ... expression_n])

Type: string

Purpose: Converts each expression to a string, where expression can be of any type. The concatenated string of all converted expressions is returned.

The elements of `intvec`, `intmat`, `ideal`, `module`, `matrix`, and `list`, are separated by a comma. No newlines are inserted.

Not defined elements of a list are omitted.

For `link`, the name of the link is used.

For `map`, the ideal defining the mapping is converted.

Note: When applied to a list, elements of type `intvec`, `intmat`, `ideal`, `module`, `matrix`, and `list` become indistinguishable.

Example:

```

    string("1+1=", 2);
    ↪ 1+1=2
    string(intvec(1,2,3,4));
    ↪ 1,2,3,4
    string(intmat(intvec(1,2,3,4), 2, 2));
    ↪ 1,2,3,4
    ring r;
    string(r);
    ↪ (ZZ/32003),(x,y,z),(dp(3),C)
    string(ideal(x,y));
    ↪ x,y
    qring R = std(ideal(x,y));
    string(R);
    ↪ (ZZ/32003),(x,y,z),(dp(3),C)
    map phi = r, ideal(x,z);
    string(phi);
    ↪ x,z
    list l;
    string(l);
    ↪
    l[3] = 1;
    string(l); // notice that l[1],l[2] are omitted
    ↪ 1
    l[2] = 1;
    l;
    ↪ [2]:
    ↪   [3]:
    ↪     1
    ↪ [3]:
    ↪   1
    string(l); // notice that lists of list is flattened
    ↪ 1,1
    l[1] = intvec(1,2,3);
    l;
    ↪ [1]:
    ↪   1,2,3
    ↪ [2]:
    ↪   [3]:
    ↪     1
    ↪ [3]:

```

```

      ↪      1
      string(1); // notice that intvec elements are not distinguishable
      ↪ 1,2,3,1,1

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 5.1.119 [print], page 238; Section 4.21 [string], page 127.

4.21.4 string operations

`+` concatenation

`<=`, `>=`, `==`, `<>`

comparison (lexicographical with respect to the ASCII encoding)

`string_expression [int_expression]`

is a character of the string; the index 1 gives the first character.

`string_expression [int_expression, int_expression]`

is a substring, where the first argument is the start index and the second is the length of the substring, filled up with blanks if the length exceeds the total size of the string

`string_expression [intvec_expression]`

is a expression list of characters from the string

Example:

```

      string s="abcde";
      s[2];
      ↪ b
      s[3,2];
      ↪ cd
      ">>" + s[1,10] + "<<";
      ↪ >>abcde <<
      s[2]="BC"; s;
      ↪ aBcde
      intvec v=1,3,5;
      s=s[v]; s;
      ↪ ace
      s="654321"; s=s[3..5]; s;
      ↪ 432

```

4.21.5 string related functions

<code>charstr</code>	description of the coefficient field of a ring (see Section 5.1.7 [charstr], page 160)
<code>execute</code>	executing string as command (see Section 5.1.32 [execute], page 176)
<code>find</code>	position of a substring in a string (see Section 5.1.42 [find], page 182)
<code>names</code>	list of strings of all user-defined variable names (see Section 5.1.102 [names], page 225)
<code>nameof</code>	name of an object (see Section 5.1.101 [nameof], page 224)
<code>option</code>	lists all defined options (see Section 5.1.110 [option], page 230)
<code>ordstr</code>	monomial ordering of a ring (see Section 5.1.112 [ordstr], page 235)
<code>parstr</code>	names of all ring parameters or the name of the n-th ring parameter (see Section 5.1.115 [parstr], page 236)

read	read a file (see Section 5.1.128 [read], page 245)
size	length of a string (see Section 5.1.142 [size], page 259)
sprintf	string formatting (see [sprintf], page 785)
typeof	type of an object (see Section 5.1.159 [typeof], page 277)
varstr	names of all ring variables or the name of the n-th ring variable (see Section 5.1.165 [varstr], page 280)

4.22 vector

Vectors are elements of a free module over the basering with basis `gen(1)`, `gen(2)`, `...`. Like polynomials they can only be defined or accessed with respect to the basering. Each vector belongs to a free module of rank equal to the biggest index of a generator with non-zero coefficient. Since generators with zero coefficients need not be written any vector may be considered also as an element of a free module of higher rank. (E.g., if `f` and `g` are polynomials then `f*gen(1)+g*gen(3)+gen(4)` may also be written as `[f,0,g,1]` or as `[f,0,g,1,0]`.) Note that the elements of a vector have to be surrounded by square brackets (`[,]`) (cf. Section B.1 [Representation of mathematical objects], page 759).

4.22.1 vector declarations

Syntax: `vector name = vector_expression ;`

Purpose: defines a vector of polynomials (an element of a free module).

Default: `[0]`

Example:

```
ring r=0,(x,y,z),(c,dp);
poly s1 = x2;
poly s2 = y3;
poly s3 = z;
vector v = [s1, s2-s1, s3-s1]+ s1*gen(5);
// v is a vector in the free module of rank 5
v;
↦ [x2,y3-x2,-x2+z,0,x2]
```

4.22.2 vector expressions

A vector expression is:

1. an identifier of type vector
2. a function returning vector
3. a polynomial expression (via the canonical embedding $p \mapsto p*\text{gen}(1)$)
4. vector expressions combined by the arithmetic operations `+` or `-`
5. a polynomial expression and a vector expression combined by the arithmetic operation `*`
6. a type cast to vector using the brackets `[,]`

Example:

```

// ordering gives priority to components:
ring rr=0,(x,y,z),(c,dp);
vector v=[x2+y3,2,0,x*y]+gen(6)*x6;
v;
↳ [y3+x2,2,0,xy,0,x6]
vector w=[z3-x,3y];
v-w;
↳ [y3-z3+x2+x,-3y+2,0,xy,0,x6]
v*(z+x);
↳ [xy3+y3z+x3+x2z,2x+2z,0,x2y+xyz,0,x7+x6z]
// ordering gives priority to monomials:
// this results in a different output
ring r=0,(x,y,z),(dp,c);
imap(rr,v);
↳ x6*gen(6)+y3*gen(1)+x2*gen(1)+xy*gen(4)+2*gen(2)

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 4.19 [ring], page 124.

4.22.3 vector operations

+ addition
 - negation or subtraction
 / division by a monomial, not divisible terms yield 0
 <, <=, >, >=, ==, <>
 comparators (considering leading terms w.r.t. monomial ordering)
 vector_expression [int_expressions]
 is a vector entry; the index 1 gives the first entry.

Example:

```

ring R=0,(x,y),(c,dp);
[x,y]-[1,x];
↳ [x-1,-x+y]
[1,2,x,4][3];
↳ x

```

4.22.4 vector related functions

cleardenom quotient of a vector by its content (see Section 5.1.9 [cleardenom], page 161)
coeffs matrix of coefficients (see Section 5.1.12 [coeffs], page 163)
deg degree (see Section 5.1.19 [deg], page 168)
diff partial derivative (see Section 5.1.24 [diff], page 171)
gen i-th generator (see Section 5.1.51 [gen], page 188)
homog homogenization (see Section 5.1.57 [homog], page 194)
jet k-jet: monomials of degree at most k (see Section 5.1.68 [jet], page 201)
lead leading term (see Section 5.1.75 [lead], page 206)

<code>leadcoef</code>	leading coefficient (see Section 5.1.76 [<code>leadcoef</code>], page 207)
<code>leadexp</code>	the exponent vector of the leading monomial (see Section 5.1.77 [<code>leadexp</code>], page 207)
<code>leadmonom</code>	leading monomial (see Section 5.1.78 [<code>leadmonom</code>], page 208)
<code>nrows</code>	number of rows (see Section 5.1.106 [<code>nrows</code>], page 228)
<code>ord</code>	degree of the leading monomial (see Section 5.1.111 [<code>ord</code>], page 234)
<code>reduce</code>	normal form with respect to a standard base (see Section 5.1.129 [<code>reduce</code>], page 246)
<code>simplify</code>	normalize a vector (see Section 5.1.141 [<code>simplify</code>], page 258)
<code>size</code>	number of monomials (see Section 5.1.142 [<code>size</code>], page 259)
<code>subst</code>	substitute a ring variable (see Section 5.1.152 [<code>subst</code>], page 269)

4.23 User defined types

User defined types are (non-empty) lists with a fixed size whose element can be accessed by names (and not indices). These elements have a predefined type (which can also be a user defined type). If these elements depend on a ring they can only be accessed if their base ring is the current base ring. In contrast to usual lists the elements of a user defined type may belong to different rings.

4.23.1 Definition of a user defined type

Syntax: `newstruct(name , string-expression);`
`newstruct(name , name , string-expression);`

Purpose: defines a new type with elements given by the last argument (`string-expression`). The name of the new type is the first argument (of type `string`) and must be longer than one character.

The second name (of type `string`) is an already defined type which should be extended by the new type.

The last argument (of type `string`) must be an comma separated list of a type followed by a name. If there are duplicate member names, the last one wins.

(User defined) member names are restricted to alphanumeric characters and must start with a letter.

Operations:

the only operations of user defined types are:

assignment (between objects of the same or extended type)

`typeof`

`string` and printing

operator `.` to access the elements

Example:

```
newstruct("nt","int a,poly b,string c");
nt A;
nt B;
A.a=3;
A.c=string(A.a);
B=A;
```

```

newstruct("t2","nt","string c");
t2 C; C.c="t2-c";
A=C;
typeof(A);
↳ t2
A;
↳ c=t2-c
↳ c=
↳ b=??
↳ a=0
// a motivating example -----
newstruct("IDEAL","ideal I,proc prettyprint");
newstruct("HOMOGENEOUS_IDEAL","IDEAL","intvec weights,proc prettyprint")
proc IDEAL_pretty_print(IDEAL I)
{
  "ideal generated by";
  I.I;
}
proc H_IDEAL_pretty_print(HOMOGENEOUS_IDEAL I)
{
  "homogeneous ideal generated by";
  I.I;
  "with weights";
  I.weights;
}
proc p_print(IDEAL I) { I.prettyprint(I); }
ring r;
IDEAL I;
I.I=ideal(x+y2,z);
I.prettyprint=IDEAL_pretty_print;
HOMOGENEOUS_IDEAL H;
H.I=ideal(x,y,z);
H.prettyprint=H_IDEAL_pretty_print;
H.weights=intvec(1,1,1);
p_print(I);
↳ ideal generated by
↳ _[1]=y2+x
↳ _[2]=z
  p_print(H);
↳ homogeneous ideal generated by
↳ _[1]=x
↳ _[2]=y
↳ _[3]=z
↳ with weights
↳ 1,1,1

```

4.23.2 Declaration of objects of a user defined type

Example:

```

newstruct("nt","int a,poly b,string c");
nt A;
// as long as there is no value assigned to A.b, no ring is needed

```



```
nt B=A;
```

4.23.3 Access to elements of a user defined type

Access to elements of a user defined type via `.: <object>.<element_name>`. The `<element_names>` are from the definition of the type. Additionally, all (potentially) ring dependent elements have an additional entry `r_<element_name>` for the corresponding ring.

Example:

```
newstruct("nt","int a,poly b,string c");
nt A;
3+A.a;
↳ 3
A.c="example string";
ring r;
A.b=poly(1); // assignment: expression must be of the given type
A;
↳ c=example string
↳ b=1
↳ a=0
A.r_b;
↳ // coefficients: ZZ/32003
↳ // number of vars : 3
↳ //      block 1 : ordering dp
↳ //      : names x y z
↳ //      block 2 : ordering C
```

4.23.4 Commands for user defined types

User defined types are normal data types (which do not belong to a ring, even if they have ring dependent parts), so they can be passed as argument to procedures, and received as result from procedures.

In order to apply kernel commands to these types (like `string`, `+`), provide a usual procedure (say `proc p..`) for that task and install it via `system("install", user.type , kernel_command , p, number_of_args)`; . The `user.type` and `kernel_command` have to be given as strings. For `kernel_command` having a variable number of arguments (internal `CMD_M`) use 4 independent of the number of really supplied arguments.

List of available kernel commands and the required `number_of_args`, some accept several variants and appear therefor at several places:

inplace binary operands: `+, -, *, /, div, %, &, |, [,` `number_of_args:2`

unary functions: `attrib, bareiss, betti, char, char_series, charstr, cleardenom, close, convhull, defined, deg, degree, denominator, det, dim, dump, ERROR, envelope, execute, facstd, factorize, finduni, gen, getdump, hilb, impart, indepSet, interred, jacob, janet, kbase, killattrib, lead, leadcoef, leadexp, leadmonom, load, ludecomp, maxideal, memory, minbase, minres, monitor, monomial, mult, mstd, nameof, ncols, npars, nrows, numerator, nvars, open, opposite, ord, ordstr, par, pardeg, parstr, preimage, prime, primefactors, prune, qhweight, rank, read, regularity, repart, ringlist, rvar, sba, size, slimgb, sortvec, sqrfree, syz, trace, transpose, twostd, typeof, univariate, var, variables, varstr, vdim, waitfirst, waitall, weight`

functions with 2 arguments: `attrib, betti, bracket, chinrem, coeffs, contract, deg, delete, diff, dim, extgcd, eliminate, exportto, facstd, factorize, farey, fetch,`

fglm, fglmquot, find, fres, frwalk, gcd, hilb, homog, hres, imap, importfrom, indepSet, insert, interpolation, janet, kbase, kernel, killattrib, koszul, lift, liftstd, load, lres, modulo, mpresmat, mres, newstruct, nc_algebra, nres, oppose, parstr, primefactors, quotient, random, rank, read, sba, simplify, sqrfree, sres, varstr, waitfirst, waitall, wedge

functions with 3 arguments: attrib, bareiss, coeffs, eliminate, find, fres, frwalk, hilb, homog, insert, koszul, laguerre, lift, liftstd, newstruct, preimage, random, resultant, sba, vandermonde

functions with variable number of arguments arguments (number_of_args:4): breakpoint, coef, dbprint, division, factmodd, intersect, jet, luinverse, lusolve, minor, names, option, qrds, reduce, reservedName, simplex, status, std, subst, system, test, uressolve, write

Example:

```

newstruct("nt","int a,poly b,string c");
nt A;
A;
↳ c=
↳ b=??
↳ a=0
ring r;
// a pretty print routine for nt:
proc pretty_print(nt A)
{
    "nt with string c:"+A.c+" and poly:"+string(A.b);
}
system("install","nt","print",pretty_print,1); // default printing uses print
A;
↳ nt with string c: and poly:0
↳
// a custom add for nt:
proc nt_add(nt A,nt B)
{
    nt C;
    C.a=A.a+B.a; C.b=A.b+B.b; C.c=A.c+B.c;
    return(C);
}
system("install","nt","+","nt_add,2);
A.b=x;
nt B; B.c="B"; B.b=y;
A+B;
↳ nt with string c:B and poly:x+y
↳

```

4.23.5 Assignments for user defined types

By default, only objects of the same (user defined) type can be assigned, there is no automatic type conversion as for the kernel data types.

But the operator = can be overridden in order to write custom constructors (the custom constructor does not apply to assignments of the same type): via `system("install", user_type, "=", p, 1);`. The user_type has to be given as a string.

Example:

```

newstruct("wrapping","poly p");
proc wrap(poly p)
{
  wrapping w; w.p = p;
  return (w);
}
system("install", "wrapping", "=", wrap, 1);
ring r = 0,x,dp;
wrapping w = x+1;
w;
↳ p=x+1
w = int(1); // via conversion int->poly
w;
↳ p=1
w=number(2); // via conversion number->poly
w;
↳ p=2

```

The user defined procedure for = provides also generic type conversions: `hh A=hh(b)`; is equivalent to `hh tmp=b; hh A=tmp; kill tmp;`.

4.24 cone

In order to use convex objects in Singular, Singular uses `gfanlib`, a C++ library for convex geometry by Anders N. Jensen. Please check the readme file for installation instructions. The library Section D.13.2 [`gfan.lib`], page 901 provides this C++ library and an interface to it. Some functions require `polymake`: Section D.13.4 [`polymake.lib`], page 906 provides an interface to it.

In the finite dimensional real vector space \mathbb{R}^n , a convex rational polyhedral cone, in short “cone”, is the convex set generated by finitely many half-lines, which in turn are generated by rational, and hence integer, points. It may or may not contain whole subspace of \mathbb{R}^n (e.g. entire lines). The biggest subspace contained in a cone is called “lineality space”. Modulo its lineality space, each cone is generated by a distinct minimal set of half lines, which are referred to as “rays”. Alternatively, a cone can be represented as a set of points satisfying a system homogeneous rational, and hence integer, linear inequalities and equations. These two characterizations of cones are the two main ways of defining cones in Singular (see [`coneViaPoints`], page 901, see [`coneViaInequalities`], page 901).

```

LIB"gfan.lib";
cone c; // ambient dim 0, no equations,
// no inequalities
cone c = 17; // ambient dim 17, no equations,
// no inequalities

```

See Section D.13.2 [`gfan.lib`], page 901.

4.25 fan

In order to use convex objects in Singular, Singular uses `gfanlib`, a C++ library for convex geometry by Anders N. Jensen. Please check the readme file for installation instructions. The library Section D.13.2 [`gfan.lib`], page 901 provides this C++ library and an interface to it. Some functions require `polymake`: Section D.13.4 [`polymake.lib`], page 906 provides an interface to it.

A polyhedral fan is a collection of cones closed under taking intersections and faces.

```

fan f;                                // ambient dim 0, no cones, lineality
                                        // space = ambient space
fan f = emptyFan(int n);              // ambient dim = n, no cones,
                                        // symmetry subgroup = <e>
fan f = n;                            // synonymous convenience method for
                                        // 'fan f = emptyFan(n)'
fan f = emptyFan(
    optional intmat G);               // symmetry group generated by rows of
                                        // G; ambient dim = number of columns
                                        // of G
    // Here a row r = r_1, ...r_n represents the permutation
    // which takes i to r_i. SINGULAR will check for the validity
    // of the provided rows, i.e. whether {r_1,...,r_n} = {1,...,n}
fan f = fullFan(int n);               // ambient dim = n, the fan consists of
                                        // one cone which equals the ambient
                                        // space, symmetry subgroup = <e>
fan f = fullFan(
    optional intmat G);               // symmetry group generated by rows of
                                        // G; ambient dim = number of columns
                                        // of G; the fan consists of one cone
                                        // which equals the ambient space

```

See also Section D.13.2 [gfan.lib], page 901.

4.26 polytope

In order to use convex objects in Singular, Singular uses gfanlib, a C++ library for convex geometry by Anders N. Jensen. Please check the readme file for installation instructions. The library Section D.13.2 [gfan.lib], page 901 provides this C++ library and an interface to it. Some functions require polymake: Section D.13.4 [polymake.lib], page 906 provides an interface to it.

In \mathbb{R}^n , a rational convex polytope (in short “polytope”) is a set of points satisfying a system rational, and hence integer, linear inequalities and equations. It is internally realised as a cone in $\mathbb{R}^{(n+1)}$ intersected with the hyperplane, in which the first coordinate equals 1.

See Section D.13.2 [gfan.lib], page 901.

4.27 pyobject

The pyobject is a black box data type in SINGULAR for handling objects from the programming language python. It needs the python support of SINGULAR to be installed.

Together with some basic operations and funtions, pyobject instances access python functionality from within SINGULAR and store the results for re-use:

Note that this feature is automatically loaded on demand when initializing an object of type pyobject. For accessing pyobject-related functions before using any python object, please type LIB("pyobject.so"); at the SINGULAR prompt.

```

pyobject pystr = "Hello";
pyobject pyint = 2;
string singstr = string(pystr + " World!");
singstr;
↳ 'Hello World!'
pystr + pyint; // Error: not possible
↳ ? pyobject error occurred

```

```

↳      ? cannot concatenate 'str' and 'int' objects
↳      ? error occurred in or before ./examples/pyobject.sing line 5: 'pystr \
      + pyint; // Error: not possible'
pystr * pyint; // But this is allowed,
↳ 'HelloHello'
pystr * 3;      // as well as this;
↳ 'HelloHelloHello'

python_run("def newfunc(*args): return list(args)"); // syncs contexts!
newfunc(1, 2, 3);      // newfunc also knowd to SINGULAR
↳ [1, 2, 3]

def pylst = python_eval("[3, 7, 1]");
proc(attrib(pylst, "sort"))(); // Access python member routines as attributes
pylst.sort();      // <- equivalent short-notation
pylst."sort"();   // <- alternative short-notation
pylst;
↳ [1, 3, 7]

python_import("os");      // Gets stuff from python module 'os'
name;      // The identifier of the operating system
↳ 'posix'

```

4.27.1 pyobject declarations

Syntax: `pyobject name = pyobject_expression ;`

Purpose: defines a python object.

Default: None

Example:

```

      pyobject empty;
      empty;
↳ None

      pyobject pystr = "Hello World!";
      pyobject pyone = 17;
      pyobject pylst = list(pystr, pyone);
      pylst;
↳ ['Hello World!', 17]

```

4.27.2 pyobject expressions

A pyobject expression is (optional parts in square brackets):

1. an identifier of type pyobject
2. a function returning pyobject
3. pyobject expressions combined by the arithmetic operations `+`, `-`, `*`, `/`, or `^`, and the member-of operators `.` and `::`
4. an list expression with elements made of pyobject expressions (see Section 3.5.5 [Type conversion and casting], page 46)
5. an string expression (see Section 3.5.5 [Type conversion and casting], page 46)

6. an int expression (see Section 3.5.5 [Type conversion and casting], page 46)

Example:

```

pyobject pystr = "python string ";
pystr;
↳ 'python string '
pyobject pyint = 2;
pyint;
↳ 2
pyobject pylst = list(pystr, pyint);
pylst;
↳ ['python string ', 2]
pyint + pyint;
↳ 4
pyint * pyint;
↳ 4
pystr + pystr;
↳ 'python string python string '
pystr * pyint;
↳ 'python string python string '
python_eval("17 + 4");
↳ 21
typeof(_);
↳ pyobject

```

4.27.3 pyobject operations

+	addition
-	negation or subtraction
*	multiplication
/	division
~, **	power by a positive integer
<, <=, >, >=, ==, <>	comparators (considering leading monomials w.r.t. monomial ordering)

pyobject_expression [int_expression]
get the item from the pyobject by index

pyobject_expression (pyobject_expression_sequence)
call the pyobject with a sequence of python arguments (the latter may be empty)

pyobject_expression . (string_expression | name),
pyobject_expression :: (string_expression | name) get attribute (class member) of a python object

Example:

```

pyobject two = 2;
pyobject three = 3;

two + three;

```

```
↳ 5
two - three;
↳ -1
two * three;
↳ 6
two / three;
↳ 0
two ^ three;
↳ 8
two ** three;
↳ 8

three < two;
↳ 0
two < three;
↳ 1
three <= two;
↳ 0
two <= three;
↳ 1
two == three;
↳ 0
two == two;
↳ 1
three > two;
↳ 1
two > three;
↳ 0
three >= two;
↳ 1
two >= three;
↳ 0
two != two;
↳ 0
two != three;
↳ 1

pyobject pystr = "Hello";
pystr + " World!";
↳ 'Hello World!'
pystr * 3;
↳ 'HelloHelloHello'
pystr[1];
↳ 'e'

python_run("def newfunc(*args): return list(args)");
newfunc();
↳ []
newfunc(two, three);
↳ [2, 3]

newfunc.__class__;
↳ <type 'function'>
```

```

newfunc::"__class__";
↳ <type 'function'>
newfunc.func_name;
↳ 'newfunc'
newfunc::func_name;
↳ 'newfunc'

```

4.27.4 pyobject related functions

attrib list, get and set attributes (class members) of a pyobject (see Section 5.1.2 [attrib], page 154)

Example:

```

pyobject pystr = "Kublai Khan";

// Additional functionality through attrib
attrib(pystr, "__doc__");
↳ "str(object='') -> string\n\nReturn a nice string representation of the
  bject.\nIf the argument is a string, the return value is the same objec
  "
proc(attrib(pystr, "count"))("K");
↳ 2

pystr.__doc__; // <- Short notations
↳ "str(object='') -> string\n\nReturn a nice string representation of the
  bject.\nIf the argument is a string, the return value is the same objec
  "
pystr.count("a"); // Even shorter (if attribute's name is valid)
↳ 2

python_run("def func(): return 17");
attrib(func);
↳ ['__call__', '__class__', '__closure__', '__code__', '__defaults__', '__
  elattr__', '__dict__', '__doc__', '__format__', '__get__', '__getattrib
  e__', '__globals__', '__hash__', '__init__', '__module__', '__name__',
  _new__', '__reduce__', '__reduce_ex__', '__repr__', '__setattr__', '__s
  eof__', '__str__', '__subclasshook__', 'func_closure', 'func_code', 'fu
  _defaults', 'func_dict', 'func_doc', 'func_globals', 'func_name']
attrib(func, "func_name");
↳ 'func'
attrib(func, "func_name", "byAnyOtherName");
attrib(func, "func_name");
↳ 'byAnyOtherName'

```

killattrib

deletes an attribute from a pyobject (see Section 5.1.72 [killattrib], page 204)

Example:

```

LIB("pyobject.so");
python_run("def new_pyobj(): pass");
attrib(new_pyobj, "new_attr", "something");

```



```

attrib(new_pyobj, "new_attr");
↳ 'something'
attrib(new_pyobj);
↳ ['__call__', '__class__', '__closure__', '__code__', '__defaults__', '__delattr__', '__dict__', '__doc__', '__format__', '__get__', '__getattr__', '__getitem__', '__globals__', '__hash__', '__init__', '__module__', '__name__', '__new__', '__reduce__', '__reduce_ex__', '__repr__', '__setattr__', '__sizeof__', '__str__', '__subclasshook__', 'func_closure', 'func_code', 'func_defaults', 'func_dict', 'func_doc', 'func_globals', 'func_name', 'new_attr']

killattrib(new_pyobj, "new_attr");
attrib(new_pyobj);
↳ ['__call__', '__class__', '__closure__', '__code__', '__defaults__', '__delattr__', '__dict__', '__doc__', '__format__', '__get__', '__getattr__', '__getitem__', '__globals__', '__hash__', '__init__', '__module__', '__name__', '__new__', '__reduce__', '__reduce_ex__', '__repr__', '__setattr__', '__sizeof__', '__str__', '__subclasshook__', 'func_closure', 'func_code', 'func_defaults', 'func_dict', 'func_doc', 'func_globals', 'func_name']

```

python_run

execute string-given `python` commands and import new symbols from `python` to SINGULAR's context (see Section 4.27.7 [python_run], page 144).

python_eval

evaluate a string-given `python` expression and return the result to SINGULAR (see Section 4.27.5 [python_eval], page 143).

python_import

import `python` module into SINGULAR's context (see Section 4.27.6 [python_import], page 143)

4.27.5 python_eval

Syntax: `python_eval (string-expression)`

Type: `pyobject`

Purpose: Evaluates a `python` expression (given as a string) and returns the result as `pyobject`.

Example:

```

LIB("pyobject.so");
python_eval("17 + 4");
↳ 21
typeof(_);
↳ pyobject
list l1 = python_eval("range(10)");

```

4.27.6 python_import

Syntax: `python_import (string-expression)`

Type: `pyobject`

Purpose: Imports `python` module (given as a string) in the SINGULAR context.

Example:

```

LIB("pyobject.so");
python_import("os");
name;                               // e. g. 'posix'
↳ 'posix'
sep;                                 // pathname separator
↳ '/'
linesep;                             // end of line marker
↳ '\n'

```

4.27.7 python_run

Syntax: python_run (string_expression)

Type: none

Purpose: Executes python commands (given as a string) in python context and syncs the contexts afterwards.

Example:

```

LIB("pyobject.so");
python_run("def newfunc(*args): return list(args)");
newfunc(1, 2, 3);                    // newfunc also known to SINGULAR now
↳ [1, 2, 3]

python_run("import os");
os;
↳ <module 'os' from '/usr/lib64/python2.7/os.pyc'>
attrib(os, "name");
↳ 'posix'

```

4.28 reference and shared (experimental)

The black box data types `reference` and `shared` in SINGULAR allow for concurrently accessing SINGULAR object data. Copying such object will only add an additional handle which allows you to define multiple identifiers for the same object instance.

Both experimental features are hidden by default, please activate them by typing `system("reference");` or `system("shared");`, respectively, at the SINGULAR prompt.

You must initialize a `reference` using a named identifier or a subexpression of the latter. The resulting object can be stored to gain read and write access from sophisticated data structures.

```

system("reference"); system("shared");
int i = 17;
reference ref = i;

ref;
↳ 17
↳
ref = 19;
ref;
↳ 19
↳
i; // original handle changed!
↳ 19

```

```

kill ref;
i;           // 'i' stays alive
↳ 19

reference uninitialized;
uninitialized; // not initialized
↳ <unassigned reference or shared memory>
// error: not a named identifier:
uninitialized = 17;
↳ ? Can only take reference from identifier
↳ ? error occurred in or before ./examples/reference_and_shared__experim\
    ental_.sing line 16: 'uninitialized = 17;

// but subexpressions of named identifiers will do
list ll = list(3,4,5);
reference ref = ll[2];
ref;
↳ 4
↳
ref = 12;
ref;
↳ 12
↳
ll;
↳ [1]:
↳ 3
↳ [2]:
↳ 12
↳ [3]:
↳ 5

```

In contrast, the type `shared` can be used to avoid the initial identifier definition. Each copy has equal rights for manipulating the data.

```

system("reference"); system("shared");
shared ll= list(2,3);

ll[1];
↳ 2
↳
ll[1]= 17;
ll;
↳ [1]:
↳ 17
↳ [2]:
↳ 3
↳

```

In most cases the value look-up is done automatically, but sometimes you have to disambiguate the input.

```

system("reference"); system("shared");
int i = 0;
reference ref = i;
shared sh = 12;

```

```

ref + sh;    // automated 'dereferencing'
↳ 12
ref + 4;
↳ 4
4 + sh;
↳ 16

list ll = list(ref, ref, ref, ref, ref, ref, ref);
string(ll);
↳ 0,0,0,0,0,0,0
ref = 1;
string(ll); // all one now
↳ 1,1,1,1,1,1,1

ll[3] = 0;
string(ll); // only third element changed
↳ 1,1,0,1,1,1,1

reference(ll[1]) = 9;
string(ll); // all others changed
↳ 9,9,0,9,9,9,9

def(ll[1]) = 11; // alternative (generic) syntax
string(ll);
↳ 11,11,0,11,11,11,11

```

The previous example had shown that `reference` and `shared` objects can store highly structured data without duplicating data all over again. As an additional feature, you can use `reference` objects for implementing procedures having side-effects.

```

system("reference"); system("shared");
list changeme;
changeme;
↳ empty list

proc setfirst(reference ll, def arg) { ll[1] = arg; }

setfirst(changeme, 17);
changeme;
↳ [1]:
↳ 17

```

If you do not need write-access to `proc` parameters, your code will usually perform better using the `alias` statement in the parameter list, see Section 4.17 [proc], page 121.

4.28.1 reference declarations

Syntax: `reference name = identifier ;`

Purpose: defines a `reference` object.

Default: None

Example:

```

system("reference"); system("shared");
reference empty;

```

```

    empty;
    ↪ <unassigned reference or shared memory>

    string str = "Hello World!";
    reference ref = str;
    ref;
    ↪ Hello World!
    ↪
    ref = 17;    // cannot change type of 'i'
    ↪ ? 'string'(str) = 'int' is not supported
    ↪ ? expected 'string' = 'string'
    ↪ ? error occurred in or before ./examples/reference_declarations.sing
        ine 8: ' ref = 17;    // cannot change type of 'i''
    list ll= list(4, 5, 6);
    reference lref = ll[2];
    lref;
    ↪ 5
    ↪
    lref = str; // change list element
    ll;
    ↪ [1]:
    ↪ 4
    ↪ [2]:
    ↪ Hello World!
    ↪ [3]:
    ↪ 6

```

4.28.2 reference expressions

A reference expression:

1. any identifier
2. any subexpression of an identifier
3. an object of type `reference` (result will reference the original identifier, too)

Example:

```

system("reference"); system("shared");
int i = 17;
reference ref = i; // new reference
ref;
↪ 17
↪
reference second = ref;
second;
↪ 17
↪
second = 9;      // also tied to 'i'
i;
↪ 9
typeof(ref);
↪ reference

list ll = list(1, 2, 3);

```

```

reference lref = ll[1];
lref;
↳ 1
↳
lref = 12;
ll;
↳ [1]:
↳ 12
↳ [2]:
↳ 2
↳ [3]:
↳ 3

```

4.28.3 shared declarations

Syntax: shared name = expression ;

Purpose: defines a shared object.

Default: None

Example:

```

system("reference"); system("shared");
shared empty;
empty;
↳ ' _ '
↳

shared str = "Hello World!";
str;
↳ Hello World!
↳

shared ll= list(4, 5, 6);
ll;
↳ [1]:
↳ 4
↳ [2]:
↳ 5
↳ [3]:
↳ 6
↳

ll[2] = str; // change list element
ll;
↳ [1]:
↳ 4
↳ [2]:
↳ Hello World!
↳ [3]:
↳ 6
↳

```

4.28.4 shared expressions

shared expression:

1. any expression
2. an object of type `shared` (result will reference the same data)

Example:

```

system("reference"); system("shared");
  shared sh = 17; // new shared
  shared second = sh;
  second;
↳ 17
↳
  second = 9;      // also tied to 'sh'
  sh;
↳ 9
↳
  typeof(sh);
↳ shared

  shared ll = list(1, 2, 3);
  shared lref = ll[1];
  lref;
↳ 1
↳
  lref = 12;
  ll;
↳ [1]:
↳ 12
↳ [2]:
↳ 2
↳ [3]:
↳ 3
↳

```

4.28.5 reference and shared operations

All operations of the underlying objects are forwarded by `reference` and `shared` objects. This kind of dereferencing is done automatically in most cases:

Example:

```

system("reference"); system("shared");
int i = 2;
reference two = i;
shared three = 3;

two * three;
↳ 6
two ^ three;
↳ 8
two ** three;
↳ 8

two + two;

```

```

↳ 4
two - two;
↳ 0

ring r = 0, (x,y,z), dp;
poly p = x + y + z;
reference ref = p;
shared zvar =z;
subst(ref, x,1, y,2, zvar,3);
↳ 6

```

In some cases **references** have to be dereferenced explicitly. For instance, this is the case for n-ary function calls not starting with a **reference** or **shared** object. You can use the **link** operator or a type cast to work around this. In contrast, some constructs like left-hand subexpressions prematurely evaluate. You can avoid this by using the **def** operator or by explicitly type casting to **reference**.

```

system("reference"); system("shared");
ring r = 0, (x,y,z), dp;
poly p = x + y + z;
shared xsh = x;
subst(p, xsh,1, y,2, z,3);          // fails
↳ ? subst('poly','shared','int') failed
↳ ? expected subst('poly','poly','poly')
↳ ? expected subst('matrix','poly','int')
↳ ? error occurred in or before ./examples/reference_and_shared_operatio\
    ns_1.sing line 5: 'subst(p, xsh,1, y,2, z,3);          // fails'
subst(p, poly(xsh),1, y,2, z,3); // good
↳ 6
subst(p, link(xsh),1, y,2, z,3); // fine
↳ 6

list ll = list(xsh, xsh, xsh);
ll[1] = y;          // replaced only first entry
ll;
↳ [1]:
↳ y
↳ [2]:
↳ x
↳
↳ [3]:
↳ x
↳
shared(ll[2]) = z;  // replaces the others
ll;
↳ [1]:
↳ y
↳ [2]:
↳ z
↳
↳ [3]:
↳ z
↳
def(ll[2]) = x;    // generic alternative

```



```

ll;
↳ [1]:
↳   y
↳ [2]:
↳   x
↳
↳ [3]:
↳   x
↳

```

In particular, explicit dereferencing is useful to distinguish between typecasting and nested constructions.

```

system("reference"); system("shared");
shared shl = list(1);
shl;
↳ [1]:
↳   1
↳
list(shl); // wraps 'shl' by a list
↳ [1]:
↳   [1]:
↳     1
↳
link(shl); // extract the list in 'shl'
↳ [1]:
↳   1

```

4.28.6 reference and shared related functions

def explicitly type casts to `reference` or `shared`, respectively. (Note: For the `def` declaration, see Section 4.4 [def], page 77.)

Example:

```

system("reference"); system("shared");
int i = 1;
reference ref = i;
shared sh = 17;
list ll = list(ref, ref, ref, sh, sh);
ll[1] = 2; // replace only one entry
ll;
↳ [1]:
↳   2
↳ [2]:
↳   1
↳
↳ [3]:
↳   1
↳
↳ [4]:
↳   17
↳
↳ [5]:
↳   17

```

```

↳
def(l1[2]) = 3;      // change the others
l1;
↳ [1]:
↳ 2
↳ [2]:
↳ 3
↳
↳ [3]:
↳ 3
↳
↳ [4]:
↳ 17
↳
↳ [5]:
↳ 17
↳
def(l1[4]) = 19;    // same here
l1;
↳ [1]:
↳ 2
↳ [2]:
↳ 3
↳
↳ [3]:
↳ 3
↳
↳ [4]:
↳ 19
↳
↳ [5]:
↳ 19
↳

```

`link` explicitly dereference a reference or shared object. (Note: For the `link` declaration, see Section 4.9 [link], page 94.)

Example:

```

system("reference"); system("shared");
ring r = 0, (x,y,z), dp;
poly p = x + y + z;
def x_=x;
reference xref=x_;
xref;
↳ x
↳
subst(p, xref,1, y,2, z,3);          // fails
↳ ? subst('poly','reference','int') failed
↳ ? expected subst('poly','poly','poly')
↳ ? expected subst('matrix','poly','int')
↳ ? error occurred in or before ./examples/reference_and_shared_relate
functions_1.sing line 7: 'subst(p, xref,1, y,2, z,3);          // fails'
subst(p, link(xref),1, y,2, z,3); // fine

```

↳ 6

system The **reference** and **shared** objects overload the **system** command to gain extended features, see `system(ref, "help")` for more details. (Note: For the general **system** command, see Section 5.1.153 [system], page 270.)

Example:

```
system("reference"); system("shared");
shared sh;
system(sh, "help");
↳ system(<ref>, ...): extended functionality for reference/shared data <ref>
>
↳ system(<ref>, count)           - number of references pointing to <ref>
↳ system(<ref>, enumerate)       - unique number for identifying <ref>
↳ system(<ref>, undefined)       - checks whether <ref> had been assigned
↳ system(<ref>, "help")          - prints this information message
↳ system(<ref>, "typeof")        - actual type referenced by <ref>
↳ system(<ref1>, same, <ref2>)  - tests for identic reference objects
```

5 Functions and system variables

5.1 Functions

This section gives a complete reference of all functions, commands and special variables of the SINGULAR kernel (i.e., all built-in commands). See Section D.1 [standard.lib], page 785, for those functions from the `standard.lib` (this library is automatically loaded at start-up time) which extend the functionality of the kernel and are written in the SINGULAR programming language.

The general syntax of a function is

```
[target =] function_name (<arguments>);
```

If no target is specified, the result is printed. In some cases (e.g., `export`, `keepring`, `setring`, `type`) the brackets are optional. For the commands `kill`, `help`, `break`, `quit`, `exit` and `LIB` no brackets are allowed.

5.1.1 align

Syntax: `align (vector_expression, int_expression)`
`align (module_expression, int_expression)`

Type: type of the first argument

Purpose: maps module generators `gen(i)` to `gen(i+s)` for all `i`.

Example:

```
ring r=0,(x,y,z),(c,dp);
align([1,2,3],3);
↪ [0,0,0,1,2,3]
align([0,0,1,2,3],-1);
↪ [0,1,2,3]
align(freemodule(2),1);
↪ _[1]=[0,1]
↪ _[2]=[0,0,1]
```

5.1.2 attrib

Syntax: `attrib (name)`

Type: none

Purpose: displays the attribute list of the object called `name`.

Example:

```
ring r=0,(x,y,z),dp;
ideal I=std(maxideal(2));
attrib(I);
↪ attr:isSB, type int
```

Syntax: `attrib (name , string_expression)`

Type: any

Purpose: returns the value of the attribute `string_expression` of the variable `name`. If the attribute is not defined for this variable, `attrib` returns the empty string.

Example:

```

ring r=0,(x,y,z),dp;
ideal I=std(maxideal(2));
attrib(I,"isSB");
↳ 1
// maxideal(2) is a standard basis,
// SINGULAR does know it for maxideal:
attrib(maxideal(2), "isSB");
↳ 1

```

Syntax: `attrib (name, string_expression, expression)`

Type: none

Purpose: sets the attribute string_expression of the variable name to the value expression.

Example:

```

ring r=0,(x,y,z),dp;
ideal I=maxideal(2); // the attribute "isSB" is not set
vdim(I);
↳ 4
attrib(I,"isSB",0); // the standard basis attribute is reset here
vdim(I);
↳ // ** I is no standard basis
↳ 4

```

Remark: An attribute may be described by any string_expression. Some of these are used by the kernel of SINGULAR and referred to as reserved attributes. Non-reserved attributes may be used, however, in procedures and can considerably speed up computations.

Reserved attributes:

(cf.class, global, isSB, isHomog, rank, ring_cf, rowShift are used by the kernel, the other are used by libraries)

`cf_class` (for ring)

the internal type of the coefficients (see `n_coeffType`)

`global` (for ring)

1, if the ordering is global

`isSB` (for ideal, module)

the standard basis property is set by all commands computing a standard basis like `groebner`, `std`, `stdhilb` etc.; used by `lift`, `dim`, `degree`, `mult`, `hilb`, `vdim`, `kbase`

`isHomog` (for ideal, module)

the weight vector of module generators for homogeneous or quasihomogeneous ideals/modules, used by `betti`, `degree`, `highcorner`, `hilbert`, `homog`, `prune`, `sba`, `slimgb`, `std`, `syz`, `kbase`, `modulo`, `mres`, `nres`, `stdhilb`.

`isCI` complete intersection property

`isCM` Cohen-Macaulay property

`maxExp` (for ring/list from ringlist)

limit for each exponent (32767 by default)

`rank` (for module)

set/get the rank of a module (see Section 5.1.106 [nrows], page 228)

`ring_cf` (for `ring`)
 the coefficients of the polynomial ring are considered to be a ring
`withSB` value of type `ideal`, resp. `module`, is `std`
`withHilb` value of type `intvec` is `hilb(.,1)` (see Section 5.1.56 [hilb], page 193)
`withRes` value of type `list` is a free resolution
`withDim` value of type `int` is the dimension (see Section 5.1.25 [dim], page 171)
`withMult` value of type `int` is the multiplicity (see Section 5.1.100 [mult], page 224)

See Section 5.1.72 [killattrib], page 204.

5.1.3 bareiss

qcindex Gauss

Syntax: `bareiss (module_expression)`
 `bareiss (matrix_expression)`
 `bareiss (module_expression, int_expression, int_expression)`
 `bareiss (matrix_expression, int_expression, int_expression)`

Type: list of `module` and `intvec`

Purpose: applies the sparse Gauss-Bareiss algorithm (see Section C.9 [References], page 783, Lee and Saunders) to a `module` (or with type conversion to a `matrix`) with an 'optimal' pivot strategy. The vectors of the `module` are the columns of the `matrix`, hence elimination takes place w.r.t. rows.

With only one parameter a complete elimination is done. Result is a list: the first entry is a `module` with a minimal independent set of vectors (as a `matrix` lower triangular), the second entry an `intvec` with the permutation of the rows w.r.t. the original `matrix`, that is, a `k` at position `l` indicates that row `k` was carried over to the row `l`.

The further parameters control the algorithm. `bareiss(M,i,j)` does not attempt to diagonalize the last `i` rows in the elimination procedure and stops computing when the remaining number of vectors (columns) to reduce is at most `j`.

Example:

```

ring r=0,(x,y,z),(c,dp);
module mm;
// ** generation of the module mm **
int d=7;
int b=2;
int db=d-b;
int i;
for(i=d;i>0;i--){ mm[i]=3*x*gen(i); }
for(i=db;i;i--){ mm[i]=mm[i]+7*y*gen(i+b); }
for(i=d;i>db;i--){ mm[i]=mm[i]+7*y*gen(i-db); }
for(i=d;i>b;i--){ mm[i]=mm[i]+11*z*gen(i-b); }
for(i=b;i;i--){ mm[i]=mm[i]+11*z*gen(i+db); }
// ** the generating matrix of mm **
print(mm);
↳ 3x, 0, 11z,0, 0, 7y, 0,
↳ 0, 3x, 0, 11z,0, 0, 7y,
↳ 7y, 0, 3x, 0, 11z,0, 0,
↳ 0, 7y, 0, 3x, 0, 11z,0,

```

```

↳ 0, 0, 7y, 0, 3x, 0, 11z,
↳ 11z,0, 0, 7y, 0, 3x, 0,
↳ 0, 11z,0, 0, 7y, 0, 3x
  // complete elimination
  list ss=bareiss(mm);
  print(ss[1]);
↳ 7y, 0, 0, 0, 0, 0, 0,
↳ 3x, -33xz, 0, 0, 0, 0, 0,
↳ 11z,-121z2,1331z3,0, 0, 0, 0,
↳ 0, 0, 0, 9317yz3,0, 0, 0,
↳ 0, 21xy, _[5,3],14641z4,-43923xz4,0, 0,
↳ 0, 0, 0, 0, 65219y2z3,_[6,6],0,
↳ 0, 49y2, _[7,3],3993xz3,_[7,5], _[7,6],_[7,7]
  ss[2];
↳ 2,7,5,1,4,3,6
  // elimination up to 3 vectors
  ss=bareiss(mm,0,3);
  print(ss[1]);
↳ 7y, 0, 0, 0, 0, 0, 0,
↳ 3x, -33xz, 0, 0, 0, 0, 0,
↳ 11z,-121z2,1331z3,0, 0, 0, 0,
↳ 0, 0, 0, 9317yz3,0, 0, 0,
↳ 0, 0, 0, 0, 27951xyz3,102487yz4,65219y2z3,
↳ 0, 21xy, _[6,3],14641z4,_[6,5], _[6,6], -43923xz4,
↳ 0, 49y2, _[7,3],3993xz3,_[7,5], _[7,6], _[7,7]
  ss[2];
↳ 2,7,5,1,3,4,6
  // elimination without the last 3 rows
  ss=bareiss(mm,3,0);
  print(ss[1]);
↳ 7y, 0, 0, 0, 0, 0, 0,
↳ 0, 77yz,0, 0, 0, 0, 0,
↳ 0, 0, 231xyz, 0, 0, 0, 0,
↳ 0, 0, 0, 1617xy2z,0, 0, 0,
↳ 11z,21xy,-1331z3,14641z4, _[5,5],_[5,6],_[5,7],
↳ 0, 0, 539y2z, _[6,4], _[6,5],_[6,6],-3773y3z,
↳ 3x, 49y2,-363xz2,3993xz3, _[7,5],_[7,6],_[7,7]
  ss[2];
↳ 2,3,4,1

```

See Section 5.1.23 [det], page 170; Section 4.12 [matrix], page 106.

5.1.4 betti

Syntax: betti (list_expression)
 betti (resolution_expression)
 betti (list_expression , int_expression)
 betti (resolution_expression , int_expression)

Type: intmat

Purpose: with 1 argument: computes the graded Betti numbers of a minimal resolution of R^n/M , if R denotes the basering, M is a homogeneous submodule of R^n and the argument represents a resolution of R^n/M .

The entry d of the `intmat` at place (i,j) is the minimal number of generators in degree $i+j$ of the j -th syzygy module (= module of relations) of R^n/M , i.e. the 0th (resp. 1st) syzygy module of R^n/M is R^n (resp. M). The argument is considered to be the result of a `res/fres/sres/mres/nres/lres` command. This implies that a zero is only allowed (and counted) as a generator in the first module.

For the computation `betti` uses only the initial monomials. This could lead to confusing results for a non-homogeneous input.

If the optional second argument is non-zero, the Betti numbers will be minimized. `betti` sets the attribute `rowShift`.

Example:

```

ring r=32003,(a,b,c,d),dp;
ideal j=bc-ad,b3-a2c,c3-bd2,ac2-b2d;
list T=mres(j,0); // 0 forces a full resolution
// a minimal set of generators for j:
print(T[1]);
↳ bc-ad,
↳ c3-bd2,
↳ ac2-b2d,
↳ b3-a2c
// second syzygy module of r/j which is the first
// syzygy module of j (minimal generating set):
print(T[2]);
↳ bd,c2,ac,b2,
↳ -a,-b,0, 0,
↳ c, d, -b,-a,
↳ 0, 0, -d,-c
// the second syzygy module (minimal generating set):
print(T[3]);
↳ -b,
↳ a,
↳ -c,
↳ d
print(T[4]);
↳ 0
betti(T);
↳ 1,0,0,0,
↳ 0,1,0,0,
↳ 0,3,4,1
// most useful for reading off the graded Betti numbers:
print(betti(T),"betti");
↳          0    1    2    3
↳ -----
↳    0:    1    -    -    -
↳    1:    -    1    -    -
↳    2:    -    3    4    1
↳ -----
↳ total:    1    4    4    1
↳

```

Hence,

- the 0th syzygy module of r/j (which is r) has 1 generator in degree 0 (which is 1),

- the 1st syzygy module $T[1]$ (which is j) has 4 generators (one in degree 2 and three in degree 3),
- the 2nd syzygy module $T[2]$ has 4 generators (all in degree 4),
- the 3rd syzygy module $T[3]$ has 1 generator in degree 5,

where the generators are the columns of the displayed matrix and degrees are assigned such that the corresponding maps have degree 0:

$$0 \leftarrow r/j \leftarrow r(1) \xleftarrow{T[1]} r(2) \oplus r^3(3) \xleftarrow{T[2]} r^4(4) \xleftarrow{T[3]} r(5) \leftarrow 0 \quad .$$

See Section C.3 [Syzygies and resolutions], page 767; Section 5.1.48 [fres], page 186; Section 5.1.58 [hres], page 195; Section 5.1.83 [lres], page 212; Section 5.1.98 [mres], page 222; Section 5.1.119 [print], page 238; [res], page 785; Section 4.18 [resolution], page 123; Section 5.1.147 [sres], page 264.

5.1.5 char

Syntax: `char (ring_name)`

Type: `int`

Purpose: returns the characteristic of the coefficient field of a ring.

Example:

```

ring r=32003,(x,y),dp;
char(r);
↳ 32003
ring s=0,(x,y),dp;
char(s);
↳ 0
ring ra=(7,a),(x,y),dp;
minpoly=a^3+a+1;
char(ra);
↳ 7
ring rp=(49,a),(x,y),dp;
char(rp);
↳ 7
ring rr=real,x,dp;
char(rr);
↳ 0

```

See Section 5.1.7 [charstr], page 160; Section 4.19 [ring], page 124.

5.1.6 char_series

Syntax: `char_series (ideal_expression)`

Type: `matrix`

Purpose: the rows of the matrix represent the irreducible characteristic series of the ideal with respect to the current ordering of variables.

One application is the decomposition of the zero-set.

Example:

```

ring r=32003,(x,y,z),dp;
print(char_series(ideal(xyz,xz,y)));
↳ y,z,
↳ x,y

```

See Section C.4 [Characteristic sets], page 768.

5.1.7 charstr

Syntax: charstr (ring_name)

Type: string

Purpose: returns the description of the coefficient field of a ring. (Tests for certain types of coefficients should use the routines from `ring.lib` as the string representation may change.)

Example:

```

ring r=32003,(x,y),dp;
charstr(r);
↳ ZZ/32003
ring s=0,(x,y),dp;
charstr(s);
↳ QQ
ring ra=(7,a),(x,y),dp;
minpoly=a^3+a+1;
charstr(ra);
↳ 7,a
ring rp=(49,a),(x,y),dp;
charstr(rp);
↳ 49,a
ring rr=real,x,dp;
charstr(rr);
↳ Float()

```

See Section 5.1.5 [char], page 159; Section 5.1.112 [ordstr], page 235; Section 4.19 [ring], page 124; Section D.2.12 [ring.lib], page 803; Section 5.1.165 [varstr], page 280.

5.1.8 chinrem

Syntax: chinrem (list, intvec)
chinrem (list, list)
chinrem (intvec, intvec)

Type: the same type as the elements of the first argument
If the elements of the first argument are lists again, chinrem is applied recursively.

Purpose: applies chinese remainder theorem to the first argument w.r.t. the moduli given in the second. The elements in the first list must be of same type which can be `bigint/int`, `poly`, `ideal`, `module` or `matrix`. The moduli, if given by a list, must be of type `bigint` or `int`.

If data depending on a ring are involved, the coefficient field must be \mathbb{Q} .

Example:

```

chinrem(intvec(2,-3),intvec(7,11));
↳ 30
chinrem(list(2,-3),list(7,11));
↳ 30
ring r=0,(x,y),dp;
ideal i1=5x+2y,x2+3y2+xy;
ideal i2=2x-3y,2x2+4y2+5xy;
chinrem(list(i1,i2),intvec(7,11));

```

```

↳ _[1]=-9x+30y
↳ _[2]=-20x2-6xy-18y2
  chinrem(list(i1,i2),list(bigint(7),bigint(11)));
↳ _[1]=-9x+30y
↳ _[2]=-20x2-6xy-18y2
  chinrem(list(list(i1,i2),list(i1,i2)),list(bigint(7),bigint(11)));
↳ [1]:
↳   _[1]=-9x+30y
↳   _[2]=-20x2-6xy-18y2
↳ [2]:
↳   _[1]=-9x+30y
↳   _[2]=-20x2-6xy-18y2

```

See Section D.4.16 [modstd_lib], page 818.

5.1.9 cleardenom

Syntax: cleardenom (poly_expression)
cleardenom (vector_expression)

Type: same as the input type

Purpose: multiplies a polynomial, resp. vector, by a suitable constant to cancel all denominators from its coefficients and then divide it by its content.

Example:

```

ring r=0,(x,y,z),dp;
poly f=(3x+6y)^5;
f/5;
↳ 243/5x5+486x4y+1944x3y2+3888x2y3+3888xy4+7776/5y5
  cleardenom(f/5);
↳ x5+10x4y+40x3y2+80x2y3+80xy4+32y5
  vector w= [4x2+20,6x+2,0,8]; // application to a vector
  print(cleardenom(w));
↳ [2x2+10,3x+1,0,4]

```

See [content], page 798.

5.1.10 close

Syntax: close (link_expression)

Type: none

Purpose: closes a link.

Example:

```

link l="ssi:tcp localhost:"+system("Singular");
open(l); // start SINGULAR "server" on localhost in batchmode
close(l); // shut down SINGULAR server

```

See Section 4.9 [link], page 94; Section 5.1.109 [open], page 230.

5.1.11 coef

Syntax: `coef (poly_expression, product_of_ringvars)`
`coef (ideal_expression, product_of_ringvars)`

Type: matrix

Syntax: `coef (vector_expression, product_of_ringvars, matrix_name, matrix_name)`

Type: none

Purpose: determines the monomials in f divisible by a ring variable of m (where f is the first argument and m the second argument) and the coefficients of these monomials as polynomials in the remaining variables. First case: returns a $2 \times n$ matrix M , n being the number of the determined monomials. The first row consists of these monomials, the second row of the corresponding coefficients of the monomials in f . Thus, $f = M[1,1] \cdot M[2,1] + \dots + M[1,n] \cdot M[2,n]$.

Second case: apply to all generators of the ideal and combine the results into one matrix.

Third case: the second matrix (i.e., the 4th argument) contains the monomials, the first matrix (i.e., the 3rd argument) the corresponding coefficients of the monomials in the vector.

Note: `coef` considers only monomials which really occur in f (i.e., which are not 0), while `coeffs` (see Section 5.1.12 [`coeffs`], page 163) returns the coefficient 0 at the appropriate place if a monomial is not present.

Example:

```

ring r=32003,(x,y,z),dp;
poly f=x5+5x4y+10x2y3+y5;
matrix m=coef(f,y);
print(m);
↳ y5,y3, y, 1,
↳ 1, 10x2,5x4,x5
f=x20+xyz+xy+x2y+z3;
print(coef(f,xy));
↳ x20,x2y,xy, 1,
↳ 1, 1, z+1,z3
print(coef(maxideal(3),yz));
↳ y3,y2z,yz2,z3,y2,yz,z2,y, z, 1,
↳ 0, 0, 0, 1, 0, 0, 0, 0, 0, 0,
↳ 0, 0, 1, 0, 0, 0, 0, 0, 0, 0,
↳ 0, 1, 0, 0, 0, 0, 0, 0, 0, 0,
↳ 1, 0, 0, 0, 0, 0, 0, 0, 0, 0,
↳ 0, 0, 0, 0, 0, 0, x, 0, 0, 0,
↳ 0, 0, 0, 0, 0, x, 0, 0, 0, 0,
↳ 0, 0, 0, 0, x, 0, 0, 0, 0, 0,
↳ 0, 0, 0, 0, 0, 0, 0, 0, x2,0,
↳ 0, 0, 0, 0, 0, 0, 0, x2,0, 0,
↳ 0, 0, 0, 0, 0, 0, 0, 0, 0, x3
vector v=[f,zy+77+xy];
print(v);
↳ [x20+x2y+xyz+z3+xy,xy+yz+77]
matrix mc; matrix mm;
coef(v,y,mc,mm);

```

```

    print(mc);
    ↪ x2+xz+x,x20+z3,
    ↪ x+z,      77
    print(mm);
    ↪ y,1,
    ↪ y,1

```

See Section 5.1.12 [coeffs], page 163.

5.1.12 coeffs

Syntax: coeffs (poly_expression , ring_variable)
 coeffs (ideal_expression , ring_variable)
 coeffs (vector_expression , ring_variable)
 coeffs (module_expression , ring_variable)
 coeffs (poly_expression , ring_variable , matrix_name)
 coeffs (ideal_expression , ring_variable , matrix_name)
 coeffs (vector_expression , ring_variable , matrix_name)
 coeffs (module_expression , ring_variable , matrix_name)

Type: matrix

Purpose: develops each polynomial of the first argument J as a univariate polynomial in the given ring_variable z, and returns the coefficients as a matrix M.

With e denoting the maximal z-degree occurring in the polynomials of J, and $d:=e+1$, $M = (m_{ij})$ satisfies the following conditions:

- (i) If J is a single polynomial f, then M is a $(d \times 1)$ -matrix and $m_{i+1,j}, 0 \leq i \leq e$, is the coefficient of z^i in f.
- (ii) If J is an ideal with generators f_1, f_2, \dots, f_k then M is a $(d \times k)$ -matrix and $m_{i+1,j}, 0 \leq i \leq e, 1 \leq j \leq k$, is the coefficient of z^i in f_j .
- (iii) If J is a k-dimensional vector with entries f_1, f_2, \dots, f_k then M is a $(dk \times 1)$ -matrix and $m_{(j-1)d+i+1,1}, 0 \leq i \leq e, 1 \leq j \leq k$, is the coefficient of z^i in f_j .
- (iV) If J is a module generated by s vectors v_1, v_2, \dots, v_s of dimension k then M is a $(dk \times s)$ -matrix and $m_{(j-1)d+i+1,r}, 0 \leq i \leq e, 1 \leq j \leq k, 1 \leq r \leq s$, is the coefficient of z^i in the j-th entry of v_r .

The optional third argument T can be used to return the matrix of powers of z such that $\text{matrix}(J) = T * M$ holds in each of the previous four cases.

Note: coeffs returns the coefficient 0 at the appropriate matrix entry if a monomial is not present, while coef considers only monomials which actually occur in the given expression.

Example:

```

    ring r;
    poly f = (x+y)^3;
    poly g = xyz+z10y4;
    ideal i = f, g;
    matrix M = coeffs(i, y);
    print(M);
    ↪ x3, 0,
    ↪ 3x2,xz,
    ↪ 3x, 0,

```

```

↳ 1, 0,
↳ 0, z10
vector v = [f, g];
M = coeffs(v, y);
print(M);
↳ x3,
↳ 3x2,
↳ 3x,
↳ 1,
↳ 0,
↳ 0,
↳ xz,
↳ 0,
↳ 0,
↳ z10

```

Syntax: `coeffs (ideal_expression, ideal_expression)`
`coeffs (module_expression, module_expression)`
`coeffs (ideal_expression, ideal_expression, product_of_ringvars)`
`coeffs (module_expression, module_expression, product_of_ringvars)`

Type: matrix

Purpose: expresses each polynomial of the first argument M as a sum $\sum_{i=1}^k m_i \cdot a_i \cdot x^{e_i}$, where the m_i come from a specified set of monomials, the a_i are from the underlying coefficient ring (or field), and the x^{e_i} are powers of a specified ring variable x.

The second parameter K provides the set of monomials which should be sufficient to generate all entries of M.

Both M and K can be thought of as the matrices obtained by `matrix(M)` and `matrix(K)`, respectively. (If M and K are given by ideals, then this matrix has just one row.)

The optional parameter `product_of_ringvars` determines the variable x: It is expected to be either the product of all ring variables (then x is 1, and each polynomial will be expressed as $\sum_{i=1}^k m_i \cdot a_i$, or `product_of_ringvars` is the product of all ring variables except one variable (which then determines x). If `product_of_ringvars` is omitted then `x = 1` as default.

If K contains all monomials that are necessary to express the entries of M, then the returned matrix A satisfies $K \cdot A = M$. Otherwise only a subset of entries of $K \cdot A$ and M will coincide. In this case, the valid entries start at `M[1,1]` and run from left to right, top to bottom.

Note: Note that in general not all entries of $K \cdot A$ and M will coincide, depending on the set of monomials provided by K.

Example:

```

ring r=32003,(x,y,z),dp;
module M = [y3+x2z, xy], [-xy, y2+x2z];
print(M);
↳ y3+x2z,-xy,
↳ xy, x2z+y2
module K = [x2, xy], [y3, xy], [xy, x];
print(K);
↳ x2,y3,xy,
↳ xy,xy,x
matrix A = coeffs(M, K, xy); // leaving z as variable of interest

```

```

    print(A); // attention: only the first row of M is reproduced by K*A
    ↦ z,0,
    ↦ 1,0,
    ↦ 0,-1

```

See Section 5.1.11 [coef], page 162; Section 5.1.69 [kbase], page 203.

5.1.13 contract

Syntax: `contract (ideal-expression, ideal-expression)`

Type: matrix

Purpose: contracts each of the n elements of the second ideal J by each of the m elements of the first ideal I , producing an $m \times n$ matrix. Contraction is defined on monomials by:

$$\text{contract}(x^A, x^B) := \begin{cases} x^{(B-A)}, & \text{if } B \geq A \text{ componentwise} \\ 0, & \text{otherwise.} \end{cases}$$

where A and B are the multiexponents of the ring variables represented by x . `contract` is extended bilinearly to all polynomials.

Example:

```

    ring r=0,(a,b,c,d),dp;
    ideal I=a2,a2+bc,abc;
    ideal J=a2-bc,abcd;
    print(contract(I,J));
    ↦ 1,0,
    ↦ 0,ad,
    ↦ 0,d

```

See Section 5.1.24 [diff], page 171.

5.1.14 create_ring

Procedure from library `standard.lib` (see Section D.1 [standard.lib], page 785).

Usage: `create_ring(l1, l2, l3[, l4, "no_minpoly"]);`
 $l1$ int or list, $l2$ list or string, $l3$ list or string, $l4$ ideal

Return: `ring(list(l1, l2, l3, l4))`

Note: $l1, l2, l3, l4$ are assumed to be the four entries of `ringlist(R)` where R is the ring to be returned.

Optional arguments: If $l4$ is not given, it is assumed to be `ideal(0)`. If "no_minpoly" is given, then the minimal polynomial in $l1$, if present, is set to 0.

Shortcuts: Strings such as "0", "(32003)" or "(0,a,b,c)" can be given as $l1$. Indexed parameters as in "(0,a(1..3))" are not supported. Strings such as "(x,y,z)" can be given as $l2$. Indexed variables as in "(x(1..3),y,z)" are not supported. Strings representing orderings such as "dp" or "(lp(3), ds(2))" can be given as $l3$, except matrix orderings given by "M([intmat_expression])".

Example:

```

ring R = (0,a), x, lp;
ringlist(R);
↳ [1]:
↳   [1]:
↳     0
↳   [2]:
↳     [1]:
↳       a
↳   [3]:
↳     [1]:
↳       [1]:
↳         lp
↳     [2]:
↳       1
↳   [4]:
↳     _[1]=0
↳ [2]:
↳   [1]:
↳     x
↳ [3]:
↳   [1]:
↳     [1]:
↳       lp
↳     [2]:
↳       1
↳   [2]:
↳     [1]:
↳       C
↳     [2]:
↳       0
↳ [4]:
↳   _[1]=0
minpoly = a^2+1;
cring Q = ideal(x^3-2);
ring S = create_ring(ringlist(Q)[1], "(x,y,t)", "dp", "no_minpoly");
basing;
↳ // coefficients: QQ(a)
↳ // number of vars : 3
↳ //           block 1 : ordering dp
↳ //           : names  x y t
↳ //           block 2 : ordering C

```

5.1.15 crossprod

Syntax: crossprod (cring-expression, ...)

Type: cring

Purpose: cross product of several objects of type cring

Example:

```

crossprod(ZZ/32003,Float());
↳ ZZ/32003 x Float()

```

See Section 4.1 [cring], page 72.

5.1.16 `datetime`

Procedure from library `standard.lib` (see Section D.1 [`standard.lib`], page 785).

Syntax: `datetime ()`

Return: string

Purpose: return the current date and time as a string

Example:

```
datetime();
↳ Mo 14. Dez 12:59:22 2020
```

5.1.17 `dbprint`

Syntax: `dbprint (int_expression, expression_list)`

Type: none

Purpose: applies the print command to each expression in the `expression_list` if `int_expression` is positive. `dbprint` may also be used in procedures in order to print results subject to certain conditions.

Syntax: `dbprint (expression)`

Type: none

Purpose: The print command is applied to the expression if `printlevel` ≥ `voice`.

Note: See Section 3.8 [Libraries], page 54, for an example how this is used for displaying comments while procedures are executed.

Example:

```
int debug=0;
intvec i=1,2,3;
dbprint(debug,i);
debug=1;
dbprint(debug,i);
↳ 1,
↳ 2,
↳ 3
voice;
↳ 1
printlevel;
↳ 0
dbprint(i);
```

See Section 3.9 [Debugging tools], page 67; Section 5.1.119 [`print`], page 238; Section 5.3.6 [`printlevel`], page 299; Section 5.3.11 [`voice`], page 303.

5.1.18 `defined`

Syntax: `defined (name)`

Type: int

Purpose: returns a value $\neq 0$ (TRUE) if there is a user-defined object with this name, and 0 (FALSE) otherwise.

A non-zero return value is the level where the object is defined (level 1 denotes the top level, level 2 the level of a first procedure, level 3 the level of a procedure called by a first procedure, etc.). For ring variables and other constants, -1 is returned.

Note: A local object m may be identified by `if (defined(m)==voice)`.

Example:

```

ring r=(0,t),(x,y),dp;
matrix m[5][6]=x,y,1,2,0,x+y;
defined(mm);
↳ 0
defined(r) and defined(m);
↳ 1
defined(m)==voice; // m is defined in the current level
↳ 1
defined(x);
↳ -1
defined(z);
↳ 0
defined("z");
↳ -1
defined(t);
↳ -1
defined(42);
↳ -1

```

See Section 5.1.137 [rvar], page 253; Section 5.3.11 [voice], page 303.

5.1.19 deg

Syntax: `deg (poly_expression)`
`deg (vector_expression)`
`deg (poly_expression , intvec_expression)`
`deg (vector_expression , intvec_expression)`

Type: int

Purpose: returns the maximal (weighted) degree of the terms of a polynomial or a vector; `deg(0)` is -1.

The optional second argument gives the weight vector, otherwise weight 1 is used for lex orderings and block ordering, the default weights of the base ring are used for orderings consisting of one block.

Example:

```

ring r=0,(x,y,z),lp;
deg(0);
↳ -1
deg(x3+y4+xyz3);
↳ 5
ring rr=7,(x,y),wp(2,3);
poly f=x2+y3;
deg(f);
↳ 9

```

```

ring R=7,(x,y),ws(2,3);
poly f=x2+y3;
deg(f);
↳ 9
vector v=[x2,y];
deg(v);
↳ 4

```

See Section 5.1.68 [jet], page 201; Section 5.1.111 [ord], page 234; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.20 degree

Syntax: degree (ideal_expression)
 degree (module_expression)

Type: string

Purpose: computes the (Krull) dimension and the multiplicity of the ideal, resp. module, generated by the leading monomials of the input and prints it. This is equal to the dimension and multiplicity of the ideal, resp. module, if the input is a standard basis with respect to a degree ordering.

Example:

```

ring r3=32003,(x,y,z),ds;
int a,b,c,t=11,10,3,1;
poly f=xa+yb+z(3*c)+x(c+2)*y(c-1)+x(c-1)*y(c-1)*z3
+x(c-2)*yc*(y2+t*x)2;
ideal i=jacob(f);
ideal i0=std(i);
degree(i0);
↳ // dimension (local) = 0
↳ // multiplicity = 314

```

See Section 5.1.25 [dim], page 171; Section 4.5 [ideal], page 78; Section 5.1.100 [mult], page 224; Section 5.1.149 [std], page 266; Section 5.1.166 [vdim], page 281.

5.1.21 delete

Syntax: delete (list_expression, int_expression)
 delete (intvec_expression, int_expression)
 delete (ideal_expression, int_expression)
 delete (module_expression, int_expression)

Type: type of the first argument

Purpose: deletes the element with the given index from a list/intvec/ideal/module (the input is not changed).

Example:

```

list l="a","b","c";
list l1=delete(l,2);l1;
↳ [1]:
↳ a
↳ [2]:
↳ c

```

```

1;
↪ [1]:
↪ a
↪ [2]:
↪ b
↪ [3]:
↪ c
delete(1..5,2);
↪ 1,3,4,5
ring r=0,(x,y,z),dp;
delete(maxideal(1),1);
↪ _[1]=y
↪ _[2]=z

```

See Section 4.5 [ideal], page 78; Section 5.1.62 [insert], page 197; Section 4.8 [intvec], page 91; Section 4.10 [list], page 101; Section 4.13 [module], page 110.

5.1.22 denominator

Syntax: denominator (number_expression)

Type: number

Purpose: returns the denominator of a number.

Example:

```

ring r = 0, x, dp;
number n = 3/2;
denominator(n);
↪ 2

```

See Section 5.1.9 [cleardenom], page 161; [content], page 798; Section 5.1.107 [numerator], page 229.

5.1.23 det

Syntax: det (intmat_expression)
det (matrix_expression)
det (smatrix_expression)
det (matrix_expression , string_expression)
det (smatrix_expression , string_expression)

Type: int, resp. poly

Purpose: returns the determinant of a square matrix. The applied algorithms depend on type of input or the optional second argument.

The optional second argument specifies the algorithm to use. Possible values are "Bareiss", "SBareiss", "Mu" and "Factory".

Example:

```

ring r=7,(x,y),wp(2,3);
matrix m[3][3]=1,2,3,4,5,6,7,8,x;
det(m);
↪ -3x-1

```

See Section 4.7 [intmat], page 88; Section 4.12 [matrix], page 106; Section 5.1.92 [minor], page 218.

5.1.24 diff

Syntax: diff (poly_expression , ring_variable)
diff (vector_expression , ring_variable)
diff (ideal_expression , ring_variable)
diff (module_expression , ring_variable)
diff (matrix_expression , ring_variable)

Type: the same as the type of the first argument

Syntax: diff (ideal_expression , ideal_expression)

Type: matrix

Syntax: diff (number_expression , ring_parameter)

Type: number

Purpose: computes the partial derivative of a polynomial object by a ring variable (first forms) respectively differentiates each polynomial (1..n) of the second ideal by the differential operator corresponding to each polynomial (1..m) in the first ideal, producing an m x n matrix.
respectively if the coefficient ring is a transcendental field extension, differentiates a number (that is, a rational function) by a transcendental variable (ring parameter).

Example:

```

ring r=0,(x,y,z),dp;
poly f=2x3y+3z5;
diff(f,x);
↳ 6x2y
vector v=[f,y2+z];
diff(v,z);
↳ 15z4*gen(1)+gen(2)
ideal j=x2-yz,xyz;
ideal i=x2,x2+yz,xyz;
// corresponds to differential operators
// d2/dx2, d2/dx2+d2/dydz, d3/dxdydz:
print(diff(i,j));
↳ 2,0,
↳ 1,x,
↳ 0,1
// differentiation of rational functions:
ring R=(0,t),(x),dp;
number f = t^2/(1-t)^2;
diff(f,t);
↳ (-2t)/(t3-3t2+3t-1)

```

See Section 5.1.13 [contract], page 165; Section 4.5 [ideal], page 78; Section 5.1.66 [jacob], page 200; Section 4.12 [matrix], page 106; Section 4.13 [module], page 110; Section 4.16 [poly], page 117; Section 5.1.163 [var], page 279; Section 4.22 [vector], page 131.

5.1.25 dim

Syntax: dim (ideal_expression)
dim (module_expression)
dim (resolution_expression)

```
dim ( ideal_expression , ideal_expression )
dim ( module_expression , ideal_expression )
```

Type: int

Purpose: computes the dimension of the ideal, resp. module, generated by the leading monomials of the given generators of the ideal, resp. module. This is also the dimension of the ideal if it is represented by a standard basis.

`dim(I, J)` is the dimension of I/J .

`dim(res)` computes the cohomological dimension of `res[1]`.

Note: The dimension of an ideal I means the Krull dimension of the basering modulo I . The dimension of a module is the dimension of its annihilator ideal. In the case of ideal (1), -1 is returned.

Example:

```
ring r=32003,(x,y,z),dp;
ideal I=x2-y,x3;
dim(std(I));
↪ 1
dim(std(ideal(1)));
↪ -1
```

See Section 5.1.20 [degree], page 169; Section 4.5 [ideal], page 78; Section 5.1.100 [mult], page 224; Section 5.1.149 [std], page 266; Section 5.1.166 [vdim], page 281.

5.1.26 division

Syntax: `division (ideal_expression, ideal_expression)`
`division (module_expression, module_expression)`
`division (ideal_expression, ideal_expression, int_expression)`
`division (module_expression, module_expression, int_expression)`
`division (ideal_expression, ideal_expression, int_expression, intvec_expression)`
`division (module_expression, module_expression, int_expression, intvec_expression)`

Type: list

Purpose: `division` computes a division with remainder. For two ideals resp. modules M (first argument) and N (second argument), it returns a list T, R, U where T is a matrix, R is an ideal resp. a module, and U is a diagonal matrix of units such that $\text{matrix}(M) * U = \text{matrix}(N) * T + \text{matrix}(R)$ is a standard representation for the normal form R of M with respect to a standard basis of N . `division` uses different algorithms depending on whether N is represented by a standard basis. For a polynomial basering, the matrix U is the identity matrix. A matrix T as above is also computed by `lift`.

For additional arguments n (third argument) and w (fourth argument), `division` returns a list T, R as above such that $\text{matrix}(M) = \text{matrix}(N) * T + \text{matrix}(R)$ is a standard representation for the normal form R of M with respect to N up to weighted degree n with respect to the weight vector w . The weighted degree of T and R respect to w is at most n . If the weight vector w is not given, `division` uses the standard weight vector $w = 1, \dots, 1$.

Example:

```
ring R=0,(x,y),ds;
poly f=x5+x2y2+y5;
```

```

division(f,jacob(f)); // automatic conversion: poly -> ideal
↳ [1]:
↳   _[1,1]=1/5x
↳   _[2,1]=3/10y
↳ [2]:
↳   _[1]=-1/2y5
↳ [3]:
↳   _[1,1]=1
division(f^2,jacob(f));
↳ [1]:
↳   _[1,1]=1/20x6-9/80xy5-5/16x7y+5/8x2y6
↳   _[2,1]=1/8x2y3+1/5x5y+1/20y6-3/4x3y4-5/4x6y2-5/16xy7
↳ [2]:
↳   _[1]=0
↳ [3]:
↳   _[1,1]=1/4-25/16xy
division(ideal(f^2),jacob(f),10);
↳ // ** _ is no standard basis
↳ [1]:
↳   _[1,1]=-75/8y9
↳   _[2,1]=1/2x2y3+x5y-1/4y6-3/2x3y4+15/4xy7+375/16x2y8
↳ [2]:
↳   _[1]=x10+9/4y10

```

See Section 4.5 [ideal], page 78; Section 5.1.80 [lift], page 209; Section 4.13 [module], page 110; Section 4.16.3 [poly operations], page 119; Section 5.1.129 [reduce], page 246.

5.1.27 dump

Syntax: dump (link.expression)

Type: none

Purpose: dumps (i.e., writes in a "message" or "block") the state of the SINGULAR session (i.e., all defined variables and their values) to the specified link (which must be either an ASCII or ssi link) such that a `getdump` can retrieve it later on.

Example:

```

ring r;
// write the whole session to the file dump.ascii
// in ASCII format
dump(":w dump.ascii");
kill r; // kill the basering
// reread the session from the file
// redefining everything which was not explicitly killed before
getdump("dump.ascii");
r;
↳ // coefficients: ZZ/32003
↳ // number of vars : 3
↳ //      block 1 : ordering dp
↳ //      : names x y z
↳ //      block 2 : ordering C

```

Restrictions:

For ASCII links, integer matrices contained in lists are dumped as integer list elements (and not as integer matrices), and lists of lists are dumped as one flattened list. Furthermore, links themselves are not dumped.

See Section 5.1.52 [getdump], page 189; Section 4.9 [link], page 94; Section 5.1.172 [write], page 284.

5.1.28 eliminate

Syntax: `eliminate (ideal-expression, product_of_ring_variables)`
`eliminate (module-expression, product_of_ring_variables)`
`eliminate (ideal-expression, intvec-expression)`
`eliminate (module-expression, intvec-expression)`
`eliminate (ideal-expression, product_of_ring_variables, intvec_hilb)`
`eliminate (module-expression, product_of_ring_variables, intvec_hilb)`

Type: the same as the type of the first argument

Purpose: eliminates variables occurring as factors/entries of the second argument from an ideal (resp. a submodule of a free module), by intersecting it (resp. each component of the submodule) with the subring not containing these variables.
`eliminate` does not need a special ordering nor a standard basis as input.

Note: Since elimination is expensive, for homogeneous input it might be useful first to compute the Hilbert function of the ideal (first argument) with a fast ordering (e.g., `dp`). Then make use of it to speed up the computation: a Hilbert-driven elimination uses the `intvec` provided as the third argument.
 If the ideal (resp. module) is not homogeneous with weights 1, this `intvec` will be silently ignored.

Example:

```

ring r=32003,(x,y,z),dp;
ideal i=x2,xy,y5;
eliminate(i,x);
↳ _[1]=y5
ring R=0,(x,y,t,s,z),dp;
ideal i=x-t,y-t2,z-t3,s-x+y3;
eliminate(i,ts);
↳ _[1]=y2-xz
↳ _[2]=xy-z
↳ _[3]=x2-y
ideal j=x2,xy,y2;
intvec v=hilb(std(j),1);
eliminate(j,y,v);
↳ _[1]=x2

```

See Section 5.1.56 [hilb], page 193; Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 5.1.149 [std], page 266.

5.1.29 eval

Syntax: `eval (expression)`

Type: none

Purpose: evaluates (quoted) expressions. Within a quoted expression, the quote can be "undone" by an `eval` (i.e., each `eval` "undoes" the effect of exactly one quote). Used only when receiving a quoted expression from an ssi link, with `quote` and `write` to prevent local evaluations when writing to an ssi link.

Example:

```

link l="ssi:w example.ssi";
ring r=0,(x,y,z),ds;
ideal i=maxideal(3);
ideal j=x7+x3,x2,z;
// compute i+j before writing, but not std
// this writes 'std(ideal(x3,...,z))'
write (l, quote(std(eval(i+j))));
option(prot);
close(l);
// now read it in again and evaluate
// read(l) forces to compute 'std(ideal(x3,...,z))'
read(l);
↳ _[1]=z
↳ _[2]=x2
↳ _[3]=xy2
↳ _[4]=y3
close(l);

```

See Section 4.9.5 [Ssi links], page 96; Section 5.1.124 [quote], page 242; Section 5.1.172 [write], page 284.

5.1.30 ERROR

Syntax: `ERROR (string_expression)`

Type: none

Purpose: Immediately interrupts the current computation, returns to the top-level, and displays the argument `string_expression` as error message.

Note: This should be used as an emergency, resp. failure, exit within procedures.

Example:

```

int i=1;
proc myError() {ERROR("Need to leave now");i=2;}
myError();
↳ ? Need to leave now
↳ ? leaving ::myError (0)
i;
↳ 1

```

5.1.31 example

Syntax: `example topic ;`

Purpose: computes an example for `topic`. Examples are available for all SINGULAR kernel and library functions. Where available (e.g., within Emacs), use <TAB> completion for a list of all available example topics.

Example:

```
example prime;
example intvec_declarations;
```

Section 5.1.54 [help], page 191

5.1.32 execute

Syntax: `execute (string_expression)`

Type: none

Purpose: executes a string containing a sequence of SINGULAR commands.

Note: The command `return` cannot appear in the string. `execute` should be avoided in procedures whenever possible, since it may give rise to name conflicts. Moreover, such procedures cannot be precompiled (a feature which SINGULAR will provide in the future).

Example:

```
ring r=32003,(x,y,z),dp;
ideal i=x+y,z3+22y;
write(":w save_i",i);
ring r0=0,(x,y,z),Dp;
string s="ideal k="+read("save_i")+";";
s;
↳ ideal k=x+y,z3+22y;
   execute(s); // define the ideal k
   k;
↳ k[1]=x+y
↳ k[2]=z3+22y
```

5.1.33 extgcd

Syntax: `extgcd (int_expression, int_expression)`
`extgcd (bigint_expression, bigint_expression)`
`extgcd (poly_expression, poly_expression)`

Type: list of 3 objects of the same type as the type of the arguments

Purpose: computes extended gcd: the first element is the greatest common divisor of the two arguments, the second and third are factors such that if `list L=extgcd(a,b)`; then `L[1]=a*L[2]+b*L[3]`.

Note: Polynomials must be univariate (in the same variable) to apply `extgcd`.

Example:

```
extgcd(24,10);
↳ [1]:
↳ 2
↳ [2]:
↳ -2
↳ [3]:
↳ 5
ring r=0,(x,y),lp;
extgcd(x4-x6,(x2+x5)*(x2+x3));
↳ [1]:
```

```

↳      x5+x4
↳ [2]:
↳      1/2x2+1/2x+1/2
↳ [3]:
↳      1/2

```

See Section 5.1.50 [gcd], page 188; Section 4.6 [int], page 82.

5.1.34 facstd

Syntax: `facstd (ideal_expression)`
`facstd (ideal_expression, ideal_expression)`

Type: list of ideals

Purpose: returns a list of ideals computed by the factorizing Groebner basis algorithm. The intersection of these ideals has the same zero-set as the input, i.e., the radical of the intersection coincides with the radical of the input ideal. In many (but not all!) cases this is already a decomposition of the radical of the ideal. (Note however that in general, no inclusion between the input and output ideals holds.) The second, optional argument gives a list of polynomials which define non-zero constraints: those ideals which contain one of the constraint polynomials are omitted from the output list. Thus the zero set of the intersection of the output ideals is contained in the zero set V of the first input ideal and contains the complement in V of the zero set of the second input ideal.

Note: Not implemented for baserings over real ground fields and galois fields (that is, only implemented for ground fields for which Section 5.1.36 [factorize], page 178 is implemented).

Example:

```

ring r=32003, (x,y,z), (c,dp);
ideal I=xyz,x2z;
facstd(I);
↳ [1]:
↳      _[1]=z
↳ [2]:
↳      _[1]=x
facstd(I,x);
↳ [1]:
↳      _[1]=z

```

See Section 4.5 [ideal], page 78; Section 4.19 [ring], page 124; Section 5.1.149 [std], page 266.

5.1.35 factmodd

Syntax: `factmodd (poly_expression, int_expression`
`[, poly_expression, poly_expression]`
`[, int_expression, int_expression]`
`)`

Type: list of polys

Purpose: Computes a factorization of a polynomial $h(x, y)$ in $K[[x]][y]$ up to a certain degree in x , whenever a factorization of $h(0, y)$ is provided or can be computed.

The algorithm is based on Hensel's lemma: Let $h(x, y)$ denote a monic polynomial in y of degree $m + n$ with coefficients in $K[[x]]$. Suppose there are two monic factors $f_0(y)$ (of degree n) and $g_0(y)$ of degree (m) such that $h(0, y) = f_0(y) * g_0(y)$ and $\langle f_0, g_0 \rangle = K[y]$.

Fix an integer $d \geq 0$. Then there are monic polynomials in y with coefficients in $K[[x]]$, namely $f(x, y)$ of degree n and $g(x, y)$ of degree m such that $h(x, y) = f(x, y) * g(x, y)$ modulo $\langle x^{(d+1)} \rangle$ (*).

The function's six arguments are $h, d, f_0, g_0, xIndex$, and $yIndex$, where $xIndex$ and $yIndex$ denote indices of ring variables that are to play the roles of x and y as above. h must be provided as an element of $K[x, y]$ since all terms of h with x -degree larger than d can be ignored due to (*).

If f_0 and g_0 are not given, the algorithm computes the factorization of $h(0, y)$ and is expected to find exactly two distinct factors (which may appear with multiplicities larger than 1) and uses these as f_0 and g_0 .

If $xIndex$ and $yIndex$ are missing they will be expected to be 1 and 2, respectively.

Note: The function expects the ground ring to contain at least two variables.

Example:

```
ring r = 0, (x,y), dp;
poly f0 = y240; poly g0 = y102+1;
poly h = y342+14x260+7x140y110+2x120y130+y240;
int d = 260;
list L = factmodd(h, d, f0, g0); L;
⇒ [1]:
⇒ -14x260y204-4x240y224-14x260y102-7x140y212-2x120y232+14x260+7x140y110
  2x120y130+y240
⇒ [2]:
⇒ 42x260y66+8x240y86+7x140y74+2x120y94+y102+1
  // check result: next output should be zero
  reduce(h - L[1] * L[2], std(x^(d+1)));
⇒ 0
```

See Section 5.1.36 [factorize], page 178.

5.1.36 factorize

Syntax: `factorize (poly_expression)`
`factorize (poly_expression, 0)`
`factorize (poly_expression, 2)`

Type: list of ideal and intvec

Syntax: `factorize (poly_expression, 1)`

Type: ideal

Purpose: computes the irreducible factors (as an ideal) of the polynomial together with or without the multiplicities (as an intvec) depending on the second argument:

0: returns factors and multiplicities, first factor is a constant.

May also be written with only one argument.

1: returns non-constant factors (no multiplicities).

2: returns non-constant factors and multiplicities.

Note: Not implemented for the coefficient fields real, finite fields of type (p^n, a) and \mathbb{Z}/m .

Example:

```

ring r=32003,(x,y,z),dp;
factorize(9*(x-1)^2*(y+z));
↳ [1]:
↳  _[1]=9
↳  _[2]=y+z
↳  _[3]=x-1
↳ [2]:
↳  1,1,2
factorize(9*(x-1)^2*(y+z),1);
↳ _[1]=y+z
↳ _[2]=x-1
factorize(9*(x-1)^2*(y+z),2);
↳ [1]:
↳  _[1]=y+z
↳  _[2]=x-1
↳ [2]:
↳  1,2
ring rQ=0,x,dp;
poly f = x2+1;           // irreducible in Q[x]
factorize(f);
↳ [1]:
↳  _[1]=1
↳  _[2]=x2+1
↳ [2]:
↳  1,1
ring rQi = (0,i),x,dp;
minpoly = i2+1;
poly f = x2+1;           // splits into linear factors in Q(i)[x]
factorize(f);
↳ [1]:
↳  _[1]=1
↳  _[2]=x+(-i)
↳  _[3]=x+(i)
↳ [2]:
↳  1,1,1

```

See [absFactorize], page 809; Section 4.16 [poly], page 117.

5.1.37 farey

Syntax: farey (bigint_expression , bigint_expression)
 farey (ideal_expression , bigint_expression)
 farey (module_expression , bigint_expression)
 farey (matrix_expression , bigint_expression)
 farey (list_expression , bigint_expression)

Type: type of the first argument (unless it is list)

Purpose: lift the first argument modulo the second to the rationals.

The (coefficients of the) result a/b is the best approximation under the condition $|a|, |b| \leq \sqrt{(N-1)}/2$ `farey(list(a,b,..),B)` is equivalent to `list(farey(a,B),farey(b,B),...)`.

Note: The current coefficient field must be the rationals.

Example:

```

ring r=0,x,dp;
farey(2,32003);
↪ 2

```

See Section 5.1.8 [chinrem], page 160.

5.1.38 fetch

Syntax: `fetch (ring_name, name)`
`fetch (ring_name, name, intvec_expression)`
`fetch (ring_name, name, intvec_expression, intvec_expression)`

Type: number, poly, vector, ideal, module, matrix or list (the same type as the second argument)

Purpose: maps objects between rings. `fetch` is the identity map between rings and q rings, in the first case the *i*-th variable of the source ring is mapped to the *i*-th variable of the basering. If the basering has less variables than the source ring these variables are mapped to zero. In the 2nd and 3rd form the intvec describes the permutation of the variables: an *i* at position *j* maps the variable `var(j)` of the source to the variable `var(i)` of the destination. Negative numbers (and the fourth argument) describe mapping of parameters.

A zero means that that variable/parameter is mapped to 0.

The coefficient fields must be compatible. (See Section 4.11 [map], page 103 for a description of possible mappings between different ground fields).

`fetch` offers a convenient way to change variable names or orderings, or to map objects from a ring to a quotient ring of that ring or vice versa.

Note: Compared with `imap`, `fetch` uses the position of the ring variables, not their names.

Example:

```

ring r=0,(x,y,z),dp;
ideal i=maxideal(2);
ideal j=std(i);
poly f=x+y2+z3;
vector v=[f,1];
qring q=j;
poly f=fetch(r,f);
f;
↪ z3+y2+x
vector v=fetch(r,v);
v;
↪ z3*gen(1)+y2*gen(1)+x*gen(1)+gen(2)
ideal i=fetch(r,i);
i;
↪ i[1]=z2
↪ i[2]=yz
↪ i[3]=y2
↪ i[4]=xz
↪ i[5]=xy
↪ i[6]=x2
ring rr=0,(a,b,c),lp;
poly f=fetch(q,f);

```

```

    f;
    ↪ a+b2+c3
    vector v=fetch(r,v);
    v;
    ↪ a*gen(1)+b2*gen(1)+c3*gen(1)+gen(2)
    ideal k=fetch(q,i);
    k;
    ↪ k[1]=c2
    ↪ k[2]=bc
    ↪ k[3]=b2
    ↪ k[4]=ac
    ↪ k[5]=ab
    ↪ k[6]=a2
    fetch(q,i,1..nvars(q)); // equivalent to fetch(q,i)
    ↪ _[1]=c2
    ↪ _[2]=bc
    ↪ _[3]=b2
    ↪ _[4]=ac
    ↪ _[5]=ab
    ↪ _[6]=a2

```

See Section 5.1.59 [imap], page 195; Section 4.11 [map], page 103; Section 4.19.1 [qring], page 124; Section 4.19 [ring], page 124.

5.1.39 fglm

Syntax: fglm (ring_name, ideal_name)

Type: ideal

Purpose: computes for the given ideal in the given ring a reduced Groebner basis in the current ring, by applying the so-called FGLM (Faugere, Gianni, Lazard, Mora) algorithm. The main application is to compute a lexicographical Groebner basis from a reduced Groebner basis with respect to a degree ordering. This can be much faster than computing a lexicographical Groebner basis directly.

Assume: The ideal must be zero-dimensional and given as a reduced Groebner basis in the given ring. The monomial ordering must be global.

Note: The only permissible differences between the given ring and the current ring are the monomial ordering and a permutation of the variables, resp. parameters.

Example:

```

ring r=0,(x,y,z),dp;
ideal i=y3+x2, x2y+x2, x3-x2, z4-x2-y;
option(redSB); // force the computation of a reduced SB
i=std(i);
vdim(i);
↪ 28
ring s=0,(z,x,y),lp;
ideal j=fglm(r,i);
j;
↪ j[1]=y4+y3
↪ j[2]=xy3-y3
↪ j[3]=x2+y3
↪ j[4]=z4+y3-y

```

See Section 5.1.40 [fglmquot], page 182; Section 5.1.110 [option], page 230; Section 4.19.1 [qring], page 124; Section 4.19 [ring], page 124; Section 5.1.149 [std], page 266; [stdfglm], page 785; Section 5.1.166 [vdim], page 281.

5.1.40 fglmquot

Syntax: `fglmquot (ideal_expression, poly_expression)`

Type: ideal

Purpose: computes a reduced Groebner basis of the ideal quotient $I:p$ of a zero-dimensional ideal I and a polynomial p using FGLM-techniques.

Assume: The ideal must be zero-dimensional and given as a reduced Groebner basis in the given ring. The polynomial must be reduced with respect to the ideal.

Example:

```
ring r=0,(x,y,z),lp;
ideal i=y3+x2,x2y+x2,x3-x2,z4-x2-y;
option(redSB); // force the computation of a reduced SB
i=std(i);
poly p=reduce(x+yz2+z10,i);
ideal j=fglmquot(i,p);
j;
↳ j[1]=z12
↳ j[2]=yz4-z8
↳ j[3]=y2+y-z8-z4
↳ j[4]=x+y-z10-z6-z4
```

See Section 5.1.39 [fglm], page 181; Section 5.1.110 [option], page 230; Section 5.1.125 [quotient], page 243; Section 4.19 [ring], page 124; Section 5.1.149 [std], page 266; Section 5.1.166 [vdim], page 281.

5.1.41 files, input from

Syntax: `< "filename"`

Type: none

Purpose: Read and execute the content of the file filename. Shorthand for `execute(read(filename))`.

Example:

```
< "example"; //read in the file example and execute it
```

See Section 5.1.32 [execute], page 176; Section 5.1.128 [read], page 245.

5.1.42 find

Syntax: `find (string_expression, substring_expression)`
`find (string_expression, substring_expression, int_expression)`

Type: int

Purpose: returns the first position of the substring in the string or 0 (if not found), starts the search at the position given in the 3rd argument.

Example:

```

    find("Aac","a");
    ↪ 2
    find("abab","a"+"b");
    ↪ 1
    find("abab","a"+"b",2);
    ↪ 3
    find("abab","ab",3);
    ↪ 3
    find("0123","abcd");
    ↪ 0

```

See Section 4.21 [string], page 127.

5.1.43 finduni

Syntax: finduni (ideal-expression)

Type: ideal

Purpose: returns an ideal which is contained in the ideal-expression, such that the i-th generator is a univariate polynomial in the i-th ring variable.
The polynomials have minimal degree w.r.t. this property.

Assume: The ideal must be zero-dimensional and given as a reduced Groebner basis in the current ring.

Example:

```

ring r=0,(x,y,z), dp;
ideal i=y3+x2,x2y+x2,z4-x2-y;
option(redSB); // force computation of reduced basis
i=std(i);
ideal k=finduni(i);
print(k);
↪ x4-x2,
↪ y4+y3,
↪ z12

```

See Section 5.1.110 [option], page 230; Section 4.19 [ring], page 124; Section 5.1.149 [std], page 266; Section 5.1.166 [vdim], page 281.

5.1.44 flintQ

Syntax: flintQ (list_of_names)

Type: cring

Purpose: returns a coefficient ring of multivariate rational functions over \mathbb{Q} to be used in ring definitions. Require flint $\geq 2.5.3$.

Example:

```

LIB "flint.so";
ring R1=flintQ(a,b),(x,y),dp;
R1;
↪ // coefficients: flintQQ(a,b)
↪ // number of vars : 2

```

```

    ↪ //      block  1 : ordering dp
    ↪ //                : names  x y
    ↪ //      block  2 : ordering C

```

See Section 4.1 [cring], page 72; Section 4.19 [ring], page 124.

5.1.45 Float

Syntax: Float ()
 Float (int_expression)
 Float (int_expression , int_expression)

Type: cring

Purpose: returns a coefficient ring of floating point (inexact) real number to be used in ring definitions.

Example:

```

    ring R1=Float(),(x,y),dp;
    R1;
    ↪ // coefficients: Float()
    ↪ // number of vars : 2
    ↪ //      block  1 : ordering dp
    ↪ //                : names  x y
    ↪ //      block  2 : ordering C
    ring R2=Float(10,20),(a,b),dp;
    R2;
    ↪ // coefficients: Float(10,20)
    ↪ // number of vars : 2
    ↪ //      block  1 : ordering dp
    ↪ //                : names  a b
    ↪ //      block  2 : ordering C

```

See Section 4.1 [cring], page 72; Section 4.19 [ring], page 124.

5.1.46 fprintf

Procedure from library `standard.lib` (see Section D.1 [standard.lib], page 785).

Syntax: fprintf (link_expression, string_expression [, any_expressions])

Return: none

Purpose: fprintf(l,fmt,...); performs output formatting. The second argument is a format control string. Additional arguments may be required, depending on the content of the control string. A series of output characters is generated as directed by the control string; these characters are written to the link l. The control string `fmt` is simply text to be copied, except that the string may contain conversion specifications.

Type `help print`; for a listing of valid conversion specifications. As an addition to the conversions of `print`, the `%n` and `%2` conversion specification does not consume an additional argument, but simply generates a newline character.

Note: If one of the additional arguments is a list, then it should be enclosed once more into a `list()` command, since passing a list as an argument flattens the list by one level.

Example:

```

    ring r=0,(x,y,z),dp;
    module m=[1,y],[0,x+z];
    intmat M=betti(mres(m,0));
    list l=r,m,M;
    link li=""; // link to stdout
    fprintf(li,"s:%s,l:%l",1,2);
    ↪ s:1,l:int(2)
    fprintf(li,"s:%s",l);
    ↪ s:(QQ),(x,y,z),(dp(3),C)
    fprintf(li,"s:%s",list(l));
    ↪ s:(QQ),(x,y,z),(dp(3),C),y*gen(2)+gen(1),x*gen(2)+z*gen(2),1,1
    fprintf(li,"2l:%2l",list(l));
    ↪ 2l:list("(QQ),(x,y,z),(dp(3),C)",
    ↪ module(y*gen(2)+gen(1),
    ↪ x*gen(2)+z*gen(2)),
    ↪ intmat(intvec(1,1),1,2))
    ↪
    fprintf(li,"%p",list(l));
    ↪ [1]:
    ↪ // coefficients: QQ
    ↪ // number of vars : 3
    ↪ //          block 1 : ordering dp
    ↪ //          : names    x y z
    ↪ //          block 2 : ordering C
    ↪ [2]:
    ↪   _[1]=y*gen(2)+gen(1)
    ↪   _[2]=x*gen(2)+z*gen(2)
    ↪ [3]:
    ↪   1,1
    fprintf(li,"%;",list(l));
    ↪ [1]:
    ↪ // coefficients: QQ
    ↪ // number of vars : 3
    ↪ //          block 1 : ordering dp
    ↪ //          : names    x y z
    ↪ //          block 2 : ordering C
    ↪ [2]:
    ↪   _[1]=y*gen(2)+gen(1)
    ↪   _[2]=x*gen(2)+z*gen(2)
    ↪ [3]:
    ↪   1,1
    ↪
    fprintf(li,"%b",M);
    ↪           0      1
    ↪ -----
    ↪    0:      1      1
    ↪ -----
    ↪ total:    1      1
    ↪

```

See also: Section 5.1.119 [print], page 238; [printf], page 785; [sprintf], page 785; Section 4.21 [string], page 127.

5.1.47 freemodule

Syntax: `freemodule (int_expression)`

Type: module

Purpose: creates the free module of rank n generated by `gen(1), \dots, gen(n)`.

Example:

```

ring r=32003,(x,y),(c,dp);
freemodule(3);
↳ _[1]=[1]
↳ _[2]=[0,1]
↳ _[3]=[0,0,1]
matrix m=freemodule(3); // generates the 3x3 unit matrix
print(m);
↳ 1,0,0,
↳ 0,1,0,
↳ 0,0,1

```

See Section 5.1.51 [gen], page 188; Section 4.13 [module], page 110.

5.1.48 fres

Syntax: `fres (ideal_expression/module_expression , int_expression , [string_expression])`

Type: resolution

Purpose: computes a (not necessarily minimal) free resolution of the input ideal/module, using Schreyer's algorithm, see reference.

If the second argument is $n > 0$, then the resolution is computed up to step n . If it is 0, `fres` computes the whole resolution.

The optional third argument can be set to

- "complete" (default) to compute the whole syzygy module in each step,
- "frame" to compute only the so-called frame,
- "extended frame" to compute only the first two terms of each generator w.r.t. the induced monomial ordering, or
- "single module" to return only the frame of each module except the last one and to return the last module in its entirety. This option can be used to reduce the amount of memory needed for the computation.

Note: The input ideal/module must be a standard basis.

Reference:

B. Erocal, O. Motsak, F.-O. Schreyer, A. Steenpass: Refined Algorithms to Compute Syzygies. J. Symb. Comput. 74 (2016), 308-327. <http://arxiv.org/abs/1502.01654>

Example:

```

ring r = 0, (w,x,y,z), dp;
ideal I = w2-xz, wx-yz, x2-wy, xy-z2, y2-wz;
attrib(I, "isSB", 1);
resolution s = fres(I, 0);
s;
↳ 1      5      6      2
↳ r <--  r <--  r <--  r

```

```

↳
↳ 0      1      2      3
↳ resolution not minimized yet
↳
↳   print(betti(s, 0), "betti");
↳           0      1      2      3
↳ -----
↳    0:      1      -      -      -
↳    1:      -      5      5      1
↳    2:      -      -      1      1
↳ -----
↳ total:      1      5      6      2
↳
↳   list l = s;
↳   print(l[1]);
↳ w2-xz,
↳ wx-yz,
↳ x2-wy,
↳ xy-z2,
↳ y2-wz
↳   print(l[2]);
↳ -x,y, 0, -z,0, -y2+wz,
↳ w, -x,-y,0, z, z2,
↳ -z,w, 0, -y,0, 0,
↳ 0, 0, w, x, -y,-yz,
↳ 0, 0, -z,-w,x, w2
↳   print(l[3]);
↳ 0, -y2+wz,
↳ y, z2,
↳ -x,-wy,
↳ w, yz,
↳ -z,-w2,
↳ 1, x

```

See Section A.3.4 [Free resolution], page 712; Section 5.1.93 [minres], page 220; [res], page 785; Section 5.1.147 [sres], page 264; Section 5.1.154 [syz], page 275.

5.1.49 frwalk

Syntax: `frwalk (ring_name, ideal_name)`
`frwalk (ring_name, ideal_name , int-expression)`

Type: ideal

Purpose: computes for the ideal `ideal_name` in the ring `ring_name` a Groebner basis in the current ring, by applying the fractal walk algorithm.

The main application is to compute a lexicographical Groebner basis from a reduced Groebner basis with respect to a degree ordering. This can be much faster than computing a lexicographical Groebner basis directly.

Note: When calling `frwalk`, the only permissible difference between the ring `ring_name` and the active base ring is the monomial ordering.

Example:

```

ring r=0,(x,y,z),dp;
ideal i=y3+x2, x2y+x2, x3-x2, z4-x2-y;
i=std(i);
ring s=0,(x,y,z),lp;
ideal j=frwalk(r,i);
j;
↳ j[1]=z12
↳ j[2]=yz4-z8
↳ j[3]=y2+y-z8-z4
↳ j[4]=xy-xz4-y+z4
↳ j[5]=x2+y-z4

```

See Section 5.1.39 [fglm], page 181; [groebner], page 785; Section 4.19.1 [qring], page 124; Section 4.19 [ring], page 124; Section 5.1.149 [std], page 266.

5.1.50 gcd

Syntax: gcd (int_expression, int_expression)
gcd (bigint_expression, bigint_expression)
gcd (number_expression, number_expression)
gcd (poly_expression, poly_expression)

Type: the same as the type of the arguments

Purpose: computes the greatest common divisor.

Note: Not implemented for the coefficient fields real and finite fields of type (p^n, a) .
The gcd of two numbers is their gcd as integer numbers or polynomials, otherwise it is not defined.

Example:

```

gcd(2,3);
↳ 1
gcd(bigint(2)^20,bigint(3)^23); // also applicable for bigints
↳ 1
typeof(_);
↳ bigint
ring r=0,(x,y,z),lp;
gcd(3x2*(x+y),9x*(y2-x2));
↳ x2+xy
gcd(number(6472674604870),number(878646537247372));
↳ 2

```

See Section 4.2 [bigint], page 73; Section 5.1.33 [extgcd], page 176; Section 4.6 [int], page 82; Section 4.14 [number], page 113.

5.1.51 gen

Syntax: gen (int_expression)

Type: vector

Purpose: returns the i-th free generator of a free module.

Example:

```

ring r=32003,(x,y,z),(c,dp);
gen(3);
↳ [0,0,1]
vector v=gen(5);
poly f=xyz;
v=v+f*gen(4); v;
↳ [0,0,0,xyz,1]
ring rr=32003,(x,y,z),dp;
fetch(r,v);
↳ xyz*gen(4)+gen(5)

```

See Section 5.1.47 [freemodule], page 186; Section 4.6 [int], page 82; Section 4.22 [vector], page 131.

5.1.52 getdump

Syntax: `getdump (link_expression)`

Type: none

Purpose: reads the content of the entire file, resp. link, and restores all variables from it. For ASCII links, `getdump` is equivalent to an `execute(read(link))` command. For ssi links, `getdump` should only be used on data which were previously `dump`'ed.

Example:

```

int i=3;
dump(":w example.txt");
kill i;
option(noredefine);
getdump("example.txt");
i;
↳ 3

```

Restrictions:

`getdump` is not supported for DBM links, or for a link connecting to `stdin` (standard input).

See Section 5.1.27 [dump], page 173; Section 4.9 [link], page 94; Section 5.1.128 [read], page 245.

5.1.53 groebner

Procedure from library `standard.lib` (see Section D.1 [standard_lib], page 785).

Syntax: `groebner (ideal_expression)`
`groebner (module_expression)`
`groebner (ideal_expression, list of string_expressions)`
`groebner (ideal_expression, list of string_expressions and int_expression)`

Type: type of the first argument

Purpose: computes a standard basis of the first argument `I` (ideal or module) by a heuristically chosen method (default) or by a method specified by further arguments of type string. Possible methods are:

- the direct methods `"std"` or `"slimgb"` without conversion,
- conversion methods `"hilb"` or `"fglm"` where a Groebner basis is first computed with an "easy" ordering and then converted to the ordering of the basering by the Hilbert driven Groebner basis computation or by linear algebra. The actual computation of

the Groebner basis can be specified by "std" or by "slimgb" (not for all orderings implemented).

A further string "par2var" converts parameters to an extra block of variables before a Groebner basis computation (and afterwards back). `option(prot)` informs about the chosen method.

Hint: Since there exists no uniform best method for computing standard bases, and since the difference in performance of a method on different examples can be huge, it is recommended to test, for hard examples, first various methods on a simplified example (e.g. use characteristic 32003 instead of 0 or substitute a subset of parameters/variables by integers, etc.).

Example:

```

intvec opt = option(get);
option(prot);
ring r = 0, (a,b,c,d), dp;
ideal i = a+b+c+d, ab+ad+bc+cd, abc+abd+acd+bcd, abcd-1;
groebner(i);
↳ std in (QQ), (a,b,c,d), (dp(4), C)
↳ [65535:2] 1(3)s2(2)s3s4-s5ss6-s7--
↳ product criterion:8 chain criterion:5
↳ _[1]=a+b+c+d
↳ _[2]=b2+2bd+d2
↳ _[3]=bc2+c2d-bd2-d3
↳ _[4]=bcd2+c2d2-bd3+cd3-d4-1
↳ _[5]=bd4+d5-b-d
↳ _[6]=c3d2+c2d3-c-d
↳ _[7]=c2d4+bc-bd+cd-2d2
ring s = 0, (a,b,c,d), lp;
ideal i = imap(r,i);
groebner(i, "hilb");
↳ compute hilbert series with std in ring (QQ), (a,b,c,d,@), (dp(5), C)
↳ weights used for hilbert series: 1,1,1,1,1
↳ [1048575:2] 1(3)s2(2)s3s4-s5ss6-s7--
↳ product criterion:8 chain criterion:5
↳ std with hilb in (QQ), (a,b,c,d,@), (lp(4), dp(1), C)
↳ [1048575:2] 1(6)s2(5)s3(4)s4-s5ssh6(3)shhhhh8shh
↳ product criterion:9 chain criterion:8
↳ hilbert series criterion:9
↳ dehomogenization
↳ simplification
↳ imap to ring (QQ), (a,b,c,d), (lp(4), C)
↳ _[1]=c2d6-c2d2-d4+1
↳ _[2]=c3d2+c2d3-c-d
↳ _[3]=bd4-b+d5-d
↳ _[4]=bc-bd5+c2d4+cd-d6-d2
↳ _[5]=b2+2bd+d2
↳ _[6]=a+b+c+d
ring R = (0,a), (b,c,d), lp;
minpoly = a2+1;
ideal i = a+b+c+d, ab+ad+bc+cd, abc+abd+acd+bcd, d2-c2b2;
groebner(i, "par2var", "slimgb");

```



```

⇒ //add minpoly to input
⇒ compute hilbert series with slimgb in ring (QQ),(b,c,d,a,@),(dp(5),C)
⇒ weights used for hilbert series: 1,1,1,1,1
⇒ slimgb in ring (QQ),(b,c,d,a,@),(dp(5),C)
⇒ CC2M[2,2](2)C3M[1,1](2)4M[2,e1](2)C5M[2,e2](3)C6M[1,1](0)
⇒ NF:8 product criterion:15, ext_product criterion:3
⇒ std with hilb in (QQ),(b,c,d,a,@),(lp(3),dp(1),dp(1),C)
⇒ [1048575:2]1(7)s2(6)s(5)s3(4)s4-s5ssh6(3)shhhh
⇒ product criterion:15 chain criterion:5
⇒ hilbert series criterion:7
⇒ dehomogenization
⇒ simplification
⇒ imap to ring (QQ),(b,c,d,a),(lp(3),dp(1),C)
⇒ //simplification
⇒ (S:4)rtrtrtr
⇒ //imap to original ring
⇒ _[1]=d2
⇒ _[2]=c+(a)
⇒ _[3]=b+c+d+(a)
groebner(i,"fglm"); //computes a reduced standard basis
⇒ std in (0,a),(b,c,d),(dp(3),C)
⇒ [65535:2]1(3)s2(2)s3s4-s5ss6-s7
⇒ (S:2)--
⇒ product criterion:9 chain criterion:1
⇒ ..+++--
⇒ vdim= 2
⇒ ..++++-
⇒ _[1]=d2
⇒ _[2]=c+(a)
⇒ _[3]=b+d
option(set,opt);

```

See also: Section D.4.8 [ffmodstd_lib], page 813; Section D.4.16 [modstd_lib], page 818; Section D.4.20 [nfmodstd_lib], page 822; Section 5.1.143 [slimgb], page 260; Section 5.1.149 [std], page 266; [stdfglm], page 785; [stdhilb], page 785.

5.1.54 help

Syntax: help;
help topic ;

Type: none

Purpose: displays online help information for topic using the currently set help browser. If no topic is given, the title page of the manual is displayed.

Note:

- ? may be used instead of help.
- topic can be an index entry of the SINGULAR manual or the name of a (loaded) procedure which has a help section.
- topic may contain wildcard characters (i.e., * characters).
- If a (possibly "wildcarded") topic cannot be found (or uniquely matched) a warning is displayed and no help information is provided.

- If `topic` is the name of a (loaded) procedure whose help section has changed w.r.t. the help available in the manual then, instead of displaying the respective help section of the manual in the help browser, the "newer" help section of the procedure is simply printed to the terminal.
- The browser in which the help information is displayed can be either set with the command-line option `--browser=<browser>` (see Section 3.1.6 [Command line options], page 19), or with the command `system("--browser", "<browser>")`. Use the command `system("browsers")`; for a list of all available browsers. See Section 3.1.3 [The online help system], page 15, for more details about help browsers.

Example:

```

help;          // display title page of manual
help ring;    // display help for 'ring'
?ring;       // equivalent to 'help ringe;'
⇒ // ** No help for topic 'ringe' (not even for '*ringe*')
⇒ // ** Try '?' for general help
⇒ // ** or '?Index;' for all available help topics
?ring*;
⇒ // ** No unique help for 'ring*'
⇒ // ** Try one of
⇒ ?Rings and orderings; ?Rings and standard bases; ?ring;
⇒ ?ring declarations; ?ring operations; ?ring related functions;
⇒ ?ring.lib; ?ring_lib; ?ringtensor; ?ringweights;
help Rings and orderings;
help standard.lib; // displays help for library 'standard.lib'

```

See Section 3.1.6 [Command line options], page 19; Section 3.8 [Libraries], page 54; Section 3.7.1 [Procedure definition], page 50; Section 3.1.3 [The online help system], page 15; Section 5.1.153 [system], page 270.

5.1.55 highcorner

Syntax: `highcorner (ideal_expression)`
`highcorner (module_expression)`

Type: poly, resp. vector

Purpose: returns the smallest monomial not contained in the ideal, resp. module, generated by the initial terms of the given generators. If the generators are a standard basis, this is also the smallest monomial not contained in the ideal, resp. module. If the ideal, resp. module, is not zero-dimensional, 0 is returned. The command works also in global orderings, but is not very useful there.

Note: Let the ideal I be given by a standard basis. Then `highcorner(I)` returns 0 if and only if $\dim(I) > 0$ or $\dim(I) = -1$. Otherwise it returns the smallest monomial m not in I which has the following properties (with x_i the variables of the basering):

- if $x_i > 1$ then x_i does not divide m (hence, $m=1$ if the ordering is global)
- given any set of generators f_1, \dots, f_k of I , let f'_i be obtained from f_i by deleting the terms divisible by $x_i \cdot m$ for all i with $x_i < 1$. Then f'_1, \dots, f'_k generate I .

Example:

```

ring r=0, (x,y), ds;
ideal i=x3,x2y,y3;
highcorner(std(i));

```

```

↳ xy2
highcorner(std(ideal(1)));
↳ 0

```

See Section 5.1.25 [dim], page 171; Section 5.1.149 [std], page 266; Section 5.1.166 [vdim], page 281.

5.1.56 hilb

Syntax: hilb (ideal_expression)
 hilb (module_expression)
 hilb (ideal_expression, int_expression)
 hilb (module_expression, int_expression)
 hilb (ideal_expression, int_expression , intvec_expression)
 hilb (module_expression, int_expression , intvec_expression)

Type: none (if called with one argument)
 intvec (if called with two or three arguments)

Purpose: computes the (weighted) Hilbert series of the base ring R modulo the ideal, resp. R^k modulo the module, defined by the leading terms of the generators of the given ideal, resp. module.

If `hilb` is called with one argument, then the first and second Hilbert series together with some additional information are displayed.

If `hilb` is called with two arguments, then the n -th Hilbert series is returned as an intvec, where $n = 1, 2$ is the second argument.

If a weight vector w is given as 3rd argument, then the Hilbert series is computed w.r.t. these weights w (by default all weights are set to 1).

Caution: The last entry of the returned intvec is not part of the actual Hilbert series, but is used in the Hilbert driven standard basis computation (see [stdhilb], page 785). (It is the minimum weight of the module generators or 0).

Syntax: hilb (intvec_expression)

Type: intvec

Purpose: computes the second Hilbert series from the first, i.e. if intvec $v = \text{hilb}(I, 1)$; then `hilb(v)` yields the same result as `hilb(I, 2)`.

Note: If the input is homogeneous w.r.t. the weights and a standard basis, the result is the (weighted) Hilbert series of the original ideal, resp. module.

Example:

```

ring R=32003,(x,y,z),dp;
ideal i=x2,y2,z2;
ideal s=std(i);
hilb(s);
↳ //      1 t^0
↳ //      -3 t^2
↳ //      3 t^4
↳ //      -1 t^6
↳
↳ //      1 t^0
↳ //      3 t^1
↳ //      3 t^2

```

```

↳ //          1 t^3
↳ // dimension (affine) = 0
↳ // degree (affine) = 8
  hilb(s,1);
↳ 1,0,-3,0,3,0,-1,0
  hilb(s,2);
↳ 1,3,3,1,0
  intvec w=2,2,2;
  hilb(s,1,w);
↳ 1,0,0,0,-3,0,0,0,3,0,0,0,-1,0

```

See Section C.2 [Hilbert function], page 766; Section 4.5 [ideal], page 78; Section 4.8 [intvec], page 91; Section 4.13 [module], page 110; Section 5.1.149 [std], page 266; [stdhilb], page 785.

5.1.57 homog

Syntax: `homog (ideal_expression)`
`homog (module_expression)`

Type: `int`

Purpose: tests for homogeneity: returns 1 for homogeneous input, 0 otherwise.

Note: If the current ring has a weighted monomial ordering, `homog` tests for weighted homogeneity w.r.t. the given weights.

Syntax:
`homog (polynomial_expression, ring_variable)`
`homog (vector_expression, ring_variable)`
`homog (ideal_expression, ring_variable)`
`homog (module_expression, ring_variable)`

Type: same as first argument

Purpose: homogenizes polynomials, vectors, ideals, or modules by multiplying each monomial with a suitable power of the given ring variable.

Note: If the current ring has a weighted monomial ordering, `homog` computes the weighted homogenization w.r.t. the given weights.
The homogenizing variable must have weight 1.

Example:

```

ring r=32003,(x,y,z),ds;
poly s1=x3y2+x5y+3y9;
poly s2=x2y2z2+3z8;
poly s3=5x4y2+4xy5+2x2y2z3+y7+11x10;
ideal i=s1,s2,s3;
homog(s2,z);
↳ x2y2z4+3z8
homog(i,z);
↳ _[1]=3y9+x5yz3+x3y2z4
↳ _[2]=x2y2z4+3z8
↳ _[3]=11x10+y7z3+5x4y2z4+4xy5z4+2x2y2z6
homog(i);
↳ 0
homog(homog(i,z));
↳ 1

```

See Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.58 hres

Syntax: `hres (ideal_expression, int_expression)`

Type: resolution

Purpose: computes a free resolution of an ideal using the Hilbert-driven algorithm. More precisely, let R be the basering and I be the given ideal. Then `hres` computes a minimal free resolution of R/I

$$\dots \longrightarrow F_2 \xrightarrow{A_2} F_1 \xrightarrow{A_1} R \longrightarrow R/I \longrightarrow 0.$$

If the `int_expression` k is not zero then the computation stops after k steps and returns a list of modules $M_i = \text{module}(A_i)$, $i=1..k$.

`list L=hres(I,0)`; returns a list L of n modules (where n is the number of variables of the basering) such that $L[i] = M_i$ in the above notation.

Note: The `ideal_expression` has to be homogeneous. Accessing single elements of a resolution may require some partial computations to be finished. Therefore, it may take some time.

Example:

```

ring r=0,(x,y,z),dp;
ideal I=xz,yz,x3-y3;
def L=hres(I,0);
print(betti(L),"betti");
↪          0    1    2
↪ -----
↪    0:    1    -    -
↪    1:    -    2    1
↪    2:    -    1    1
↪ -----
↪ total:    1    3    2
↪
↪ L[2]; // the first syzygy module of r/I
↪ _[1]=-x*gen(1)+y*gen(2)
↪ _[2]=-x2*gen(2)+y2*gen(1)+z*gen(3)

```

See Section 5.1.4 [betti], page 157; Section 5.1.48 [fres], page 186; Section 4.5 [ideal], page 78; Section 4.6 [int], page 82; Section 5.1.83 [lres], page 212; Section 5.1.93 [minres], page 220; Section 4.13 [module], page 110; Section 5.1.98 [mres], page 222; [res], page 785; Section 5.1.147 [sres], page 264.

5.1.59 imap

Syntax: `imap (ring_name, name)`

Type: number, poly, vector, ideal, module, matrix or list (the same type as the second argument)

Purpose: identity map on common subrings. `imap` is the map between rings and qrings with compatible ground fields which is the identity on variables and parameters of the same name and 0 otherwise. (See Section 4.11 [map], page 103 for a description of possible

mappings between different ground fields). Useful for mapping from a homogenized ring to the original ring or for mappings from/to rings with/without parameters. Compared with `fetch`, `imap` uses the names of variables and parameters. Unlike `map` and `fetch` `imap` can map parameters to variables.

Mapping rational functions which are not polynomials to polynomials is undefined (i.e. the result depends on the version).

Example:

```

ring r=0,(x,y,z,a,b,c),dp;
ideal i=xy2z3a4b5+1,homog(xy2z3a4b5+1,c); i;
↪ i[1]=xy2z3a4b5+1
↪ i[2]=xy2z3a4b5+c15
ring r1=0,(a,b,x,y,z),lp;
ideal j=imap(r,i); j;
↪ j[1]=a4b5xy2z3+1
↪ j[2]=a4b5xy2z3
ring r2=(0,a,b),(x,y,z),ls;
ideal j=imap(r,i); j;
↪ j[1]=1+(a4b5)*xy2z3
↪ j[2]=(a4b5)*xy2z3

```

See Section 5.1.38 [fetch], page 180; Section 5.1.57 [homog], page 194; Section 4.11 [map], page 103; Section 4.19.1 [qring], page 124; Section 4.19 [ring], page 124.

5.1.60 `impart`

Syntax: `impart (number_expression)`

Type: number

Purpose: returns the imaginary part of a number in a complex ground field, returns 0 otherwise.

Example:

```

ring r=(complex,i),x,dp;
impart(1+2*i);
↪ 2

```

See Section 5.1.131 [repart], page 248.

5.1.61 `indepSet`

Syntax: `indepSet (ideal_expression)`

Type: intvec

Purpose: computes a maximal set U of independent variables (in the sense defined in the note below) of the ideal given by a standard basis. If v is the result then $v[i]$ is 1 if and only if the i -th variable of the ring, $x(i)$, is an independent variable. Hence, the set U consisting of all variables $x(i)$ with $v[i]=1$ is a maximal independent set.

Note: U is a set of independent variables for I if and only if $I \cap K[U] = (0)$, i.e., eliminating the remaining variables gives (0) . U is maximal if $\dim(I)=\#U$.

Syntax: `indepSet (ideal_expression, int_expression)`

Type: list

Purpose: computes a list of all maximal independent sets of the leading ideal (if the flag is 0), resp. of all those sets of independent variables of the leading ideal which cannot be enlarged.

Example:

```

ring r=32003,(x,y,u,v,w),dp;
ideal I=xyw,yvw,uyw,xv;
attrib(I,"isSB",1);
indepSet(I);
↳ 1,1,1,0,0
eliminate(I,vw);
↳ _[1]=0
indepSet(I,0);
↳ [1]:
↳ 1,1,1,0,0
↳ [2]:
↳ 0,1,1,1,0
↳ [3]:
↳ 1,0,1,0,1
↳ [4]:
↳ 0,0,1,1,1
indepSet(I,1);
↳ [1]:
↳ 1,1,1,0,0
↳ [2]:
↳ 0,1,1,1,0
↳ [3]:
↳ 1,0,1,0,1
↳ [4]:
↳ 0,0,1,1,1
↳ [5]:
↳ 0,1,0,0,1
eliminate(I,xuv);
↳ _[1]=0

```

See Section 4.5 [ideal], page 78; Section 5.1.149 [std], page 266.

5.1.62 insert

Syntax: `insert (list_expression, expression)`
`insert (list_expression, expression, int_expression)`

Type: list

Purpose: inserts a new element (expression) into a list at the beginning, or (if called with 3 arguments) after the given position (the input is not changed).

Example:

```

list L=1,2;
insert(L,4,2);
↳ [1]:
↳ 1
↳ [2]:
↳ 2
↳ [3]:

```

```

↳      4
      insert(L,4);
↳ [1]:
↳      4
↳ [2]:
↳      1
↳ [3]:
↳      2

```

See Section 5.1.21 [delete], page 169; Section 4.10 [list], page 101.

5.1.63 interpolation

Syntax: interpolation (list, intvec)

Type: ideal

Purpose: interpolation(l,v) computes the reduced Groebner basis of the intersection of ideals $l[1]^v[1], \dots, l[N]^v[N]$ by applying linear algebra methods.

Assume: Every ideal from the list l must be a maximal ideal of a point and should have the following form: variable_1-coordinate_1, ..., variable_n-coordinate_n, where n is the number of variables in the ring.

The ring should be a polynomial ring over \mathbb{Z}_p or \mathbb{Q} with global ordering.

Example:

```

ring r=0,(x,y),dp;
ideal p_1=x,y;
ideal p_2=x+1,y+1;
ideal p_3=x+2,y-1;
ideal p_4=x-1,y+2;
ideal p_5=x-1,y-3;
ideal p_6=x,y+3;
ideal p_7=x+2,y;
list l=p_1,p_2,p_3,p_4,p_5,p_6,p_7;
intvec v=2,1,1,1,1,1,1;
ideal j=interpolation(l,v);
// generator of degree 3 gives the equation of the unique
// singular cubic passing
// through p_1,...,p_7 with singularity at p_1
j;
↳ j[1]=-4x3-4x2y-2xy2+y3-8x2-4xy+3y2
↳ j[2]=-y4+8x2y+6xy2-2y3+10xy+3y2
↳ j[3]=-xy3+2x2y+xy2+4xy
↳ j[4]=-2x2y2-2x2y-2xy2+y3-4xy+3y2
// computes values of generators of j at p_4, results should be 0
subst(j,x,1,y,-2);
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0
↳ _[4]=0
// computes values of derivatives d/dx of generators at (0,0)
subst(diff(j,x),x,0,y,0);
↳ _[1]=0
↳ _[2]=0

```



```

↳ _[3]=0
↳ _[4]=0

```

See Section 5.1.24 [diff], page 171; Section 5.1.39 [fglm], page 181; Section 5.1.65 [intersect], page 199; Section 5.1.149 [std], page 266; Section 5.1.152 [subst], page 269.

5.1.64 interred

Syntax: `interred (ideal_expression)`
`interred (module_expression)`

Type: the same as the input type

Purpose: interreduces a set of polynomials/vectors.

Input: f_1, \dots, f_n

Output: g_1, \dots, g_s with $s \leq n$ and the properties

- $(f_1, \dots, f_n) = (g_1, \dots, g_s)$,
- $L(g_i) \neq L(g_j)$ for all $i \neq j$,
- in the case of a global ordering (polynomial ring) and `option(redSB);`:
 $L(g_i)$ does not divide m for all monomials m of $\{g_1, \dots, g_{i-1}, g_{i+1}, \dots, g_s\}$,
- in the case of a local ordering (localization of polynomial ring) and `option(redSB);`:
if $L(g_i) | L(g_j)$ for any $i \neq j$, then $ecart(g_i) > ecart(g_j)$.

Here, $L(g)$ denotes the leading term of g and $ecart(g) := deg(g) - deg(L(g))$.

Example:

```

ring r=0,(x,y,z),dp;
ideal i=zx+y3,z+y3,z+xy;
interred(i);
↳ _[1]=xz-z
↳ _[2]=xy+z
↳ _[3]=y3+xz
ring R=0,(x,y,z),ds;
ideal i=zx+y3,z+y3,z+xy;
interred(i);
↳ _[1]=z+xy
↳ _[2]=xy-y3
↳ _[3]=x2y-y3

```

See Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 5.1.149 [std], page 266.

5.1.65 intersect

Syntax: `intersect (expression_list of ideal_expression)`
`intersect (expression_list of module_expression)`

Type: ideal, resp. module

Purpose: computes the intersection of ideals, resp. modules.

Note: If the option `prot` is enabled then the result the used method (elimination/syzygies) is displayed.

An optional last argument specifies the Groebner base algorithm to use. Possible values are "std" and "slimgb".

Example:

```

ring R=0,(x,y),dp;
ideal i=x;
ideal j=y;
intersect(i,j);
↳ _[1]=xy
ring r=181,(x,y,z),(c,ls);
ideal id1=maxideal(3);
ideal id2=x2+xyz,y2-z3y,z3+y5xz;
ideal id3=intersect(id1,id2,ideal(x,y));
ideal id4=intersect(id1,id2,"slimgb");
id3;
↳ id3[1]=yz3+xy6z
↳ id3[2]=yz4-y2z
↳ id3[3]=y2z3-y3
↳ id3[4]=xz3+x2y5z
↳ id3[5]=xyz2+x2z
↳ id3[6]=xy2+x2z2
↳ id3[7]=xy2z+x2y
↳ id3[8]=x2yz+x3
id4;
↳ id4[1]=xyz2+x2z
↳ id4[2]=xy2z+x2y
↳ id4[3]=x2yz+x3
↳ id4[4]=-yz4+y2z
↳ id4[5]=-y2z3+y3
↳ id4[6]=-xyz3+xy2
↳ id4[7]=z3+xy5z

```

See Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 5.1.110 [option], page 230.

5.1.66 jacob

Syntax: jacob (poly_expression)
 jacob (ideal_expression)
 jacob (module_expression)

Type: ideal, if the input is a polynomial
 matrix, if the input is an ideal
 module, if the input is a module

Purpose: computes the Jacobi ideal, resp. Jacobi matrix, generated by all partial derivatives of the input.

Note: In a ring with n variables, jacob of a module or an ideal (considered as matrix with a single a row) or a polynomial (considered as a matrix with a single entry) is the matrix consisting of horizontally concatenated blocks (in this order): diff(MT,var(1)), ... , diff(MT,var(n)), where MT is the transposed input argument considered as a matrix.

Example:

```

ring R;
poly f = x2yz + xy3z + xyz5;
ideal i = jacob(f); i;
↳ i[1]=yz5+y3z+2xyz

```

```

↳ i[2]=xz5+3xy2z+x2z
↳ i[3]=5xyz4+xy3+x2y
  matrix m = jacob(i);
  print(m);
↳ 2yz,          z5+3y2z+2xz, 5yz4+y3+2xy,
↳ z5+3y2z+2xz,6xyz,          5xz4+3xy2+x2,
↳ 5yz4+y3+2xy,5xz4+3xy2+x2,20xyz3
  print(jacob(m));
↳ 0, 2z,          2y,          2z,          6yz,5z4+3y2+2x,2y,          5z4+3y2+2x,
  20yz3,
↳ 2z,6yz,          5z4+3y2+2x,6yz,          6xz,6xy,          5z4+3y2+2x,6xy,
  20xz3,
↳ 2y,5z4+3y2+2x,20yz3,          5z4+3y2+2x,6xy,20xz3,          20yz3,          20xz3,
  60xyz2

```

See Section 5.1.24 [diff], page 171; Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 5.1.108 [nvars], page 229.

5.1.67 Janet

Syntax: Janet (ideal_expression)
 Janet (ideal_expression , int_expression)

Type: ideal

Purpose: computes the Janet basis of the given ideal, resp. the standard basis if 1 is given as the second argument.

Remark: It works only with global orderings.

Example:

```

ring r=0,(x,y,z),dp;
ideal i=x*y*z-1,x+y+z,x*y+x*z+y*z; // cyclic 3
Janet(i);
↳ Length of Janet basis: 4
↳ _[1]=x+y+z
↳ _[2]=y2+yz+z2
↳ _[3]=z3-1
↳ _[4]=yz3-y

```

See [groebner], page 785; Section 4.5 [ideal], page 78; Section 5.1.149 [std], page 266.

5.1.68 jet

Syntax: jet (poly_expression , int_expression)
 jet (vector_expression , int_expression)
 jet (ideal_expression , int_expression)
 jet (module_expression , int_expression)
 jet (poly_expression , int_expression , intvec_expression)
 jet (vector_expression , int_expression , intvec_expression)
 jet (ideal_expression , int_expression , intvec_expression)
 jet (module_expression , int_expression , intvec_expression)
 jet (poly_expression , poly_expression , int_expression)
 jet (vector_expression , poly_expression , int_expression)
 jet (ideal_expression , matrix_expression , int_expression)
 jet (module_expression , matrix_expression , int_expression)

Type: the same as the type of the first argument

Purpose: deletes from the first argument all terms of degree bigger than the second argument. If a third argument w of type `intvec` is given, the degree is replaced by the weighted degree defined by w . If a second argument u of type `poly` or `matrix` is given, the first argument p is replaced by p/u . In this case, the coefficient must be from a field.

Example:

```

ring r=32003,(x,y,z),(c,dp);
jet(1+x+x2+x3+x4,3);
↳ x3+x2+x+1
poly f=1+x+x2+xz+y2+x3+y3+x2y2+z4;
jet(f,3);
↳ x3+y3+x2+y2+xz+x+1
intvec iv=2,1,1;
jet(f,3,iv);
↳ y3+y2+xz+x+1
// the part of f with (total) degree >3:
f-jet(f,3);
↳ x2y2+z4
// the homogeneous part of f of degree 2:
jet(f,2)-jet(f,1);
↳ x2+y2+xz
// the part of maximal degree:
jet(f,deg(f))-jet(f,deg(f)-1);
↳ x2y2+z4
// the absolute term of f:
jet(f,0);
↳ 1
// now for other types:
ideal i=f,x,f*f;
jet(i,2);
↳ _[1]=x2+y2+xz+x+1
↳ _[2]=x
↳ _[3]=3x2+2y2+2xz+2x+1
vector v=[f,1,x];
jet(v,1);
↳ [x+1,1,x]
jet(v,0);
↳ [1,1]
v=[f,1,0];
module m=v,v,[1,x2,z3,0,1];
jet(m,2);
↳ _[1]=[x2+y2+xz+x+1,1]
↳ _[2]=[x2+y2+xz+x+1,1]
↳ _[3]=[1,x2,0,0,1]
ring rs=0,x,ds;
// 1/(1+x) till degree 5
jet(1,1+x,5);
↳ 1-x+x2-x3+x4-x5

```

See Section 5.1.19 [deg], page 168; Section 4.5 [ideal], page 78; Section 4.6 [int], page 82; Section 4.8 [intvec], page 91; Section 4.13 [module], page 110; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.69 kbase

Syntax: `kbase (ideal_expression)`
`kbase (module_expression)`
`kbase (ideal_expression, int_expression)`
`kbase (module_expression, int_expression)`

Type: the same as the input type of the first argument

Purpose: With one argument: computes a vector space basis (consisting of monomials) of the quotient ring by the ideal, resp. of a free module by the module, in case it is finite dimensional and if the input is a standard basis with respect to the ring ordering. Note that, if the input is not a standard basis, the leading terms of the input are used and the result may have no meaning.
 With two arguments: computes the part of a vector space basis of the respective quotient with degree of the monomials equal to the second argument. Here, the quotient does not need to be finite dimensional. If an attribute `isHomog` (of type `intvec`) is present, it is used as module weight.

Example:

```
ring r=32003,(x,y,z),ds;
ideal i=x2,y2,z;
kbase(std(i));
↳ _[1]=xy
↳ _[2]=y
↳ _[3]=x
↳ _[4]=1
i=x2,y3,xyz; // quotient not finite dimensional
kbase(std(i),2);
↳ _[1]=z2
↳ _[2]=yz
↳ _[3]=xz
↳ _[4]=y2
↳ _[5]=xy
```

See Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 5.1.166 [vdim], page 281.

5.1.70 kernel

Syntax: `kernel (ring_name, map_name)`
`preimage (ring_name, ideal_expression)`

Type: ideal

Purpose: returns the kernel of a given map.
 The second argument has to be a map from the basering to the given ring (or an ideal defining such a map).

Example:

```
ring r1=32003,(x,y,z,w),lp;
ring r=32003,(x,y,z),dp;
```

```

ideal i=x,y,z;
map f=r1,i;
setring r1;
// the kernel of f
kernel(r,f);
↳ _[1]=w

```

See [alg_kernel], page 810; [hom_kernel], page 816; Section 4.5 [ideal], page 78; Section 4.11 [map], page 103; Section 5.1.94 [modulo], page 220; Section 5.1.116 [preimage], page 236; Section 4.19 [ring], page 124.

5.1.71 kill

Syntax: kill name
kill list_of_names

Type: none

Purpose: deletes objects.

Example:

```

int i=3;
ring r=0,x,dp;
poly p;
listvar();
↳ // r [0] *ring
↳ // p [0] poly
↳ // i [0] int 3
kill i,r;
// the variable 'i' does not exist any more
i;
↳ ? 'i' is undefined
↳ ? error occurred in or before ./examples/kill.sing line 7: ' i;'
listvar();

```

See Section 5.1.18 [defined], page 167; Section D.2.3 [general_lib], page 790.

5.1.72 killattrib

Syntax: killattrib (name)
killattrib (name, string_expression)

Type: none

Purpose: deletes all attributes respective the attribute given as the second argument.

Example:

```

ring r=32003,(x,y),lp;
ideal i=maxideal(1);
attrib(i,"isSB",1);
attrib(i);
↳ attr:isSB, type int
killattrib(i,"isSB");
attrib(i);
↳ no attributes
attrib(i,"isSB",1);

```

```

killattrib(i);
attrib(i);
↳ no attributes

```

See Section 5.1.2 [attrib], page 154; Section 5.1.110 [option], page 230.

5.1.73 koszul

Syntax: `koszul (int_expression , int_expression)`
`koszul (int_expression , ideal_expression)`
`koszul (int_expression , int_expression , ideal_expression)`

Type: matrix

Purpose: `koszul(d,n)` computes a matrix of the Koszul relations of degree d of the first n ring variables.

`koszul(d,id)` computes a matrix of the Koszul relations of degree d of the generators of the ideal `id`.

`koszul(d,n,id)` computes a matrix of the Koszul relations of degree d of the first n generators of the ideal `id`.

Note: `koszul(1,id)`, `koszul(2,id)`, ... form a complex, that is, the product of the matrices `koszul(i,id)` and `koszul(i+1,id)` equals zero.

Example:

```

ring r=32003,(x,y,z),dp;
print(koszul(2,3));
↳ -y,-z,0,
↳ x, 0, -z,
↳ 0, x, y
ideal I=xz2+yz2+z3,xyz+y2z+yz2,xy2+y3+y2z;
print(koszul(1,I));
↳ xz2+yz2+z3,xyz+y2z+yz2,xy2+y3+y2z
print(koszul(2,I));
↳ -xyz-y2z-yz2,-xy2-y3-y2z,0,
↳ xz2+yz2+z3, 0, -xy2-y3-y2z,
↳ 0, xz2+yz2+z3, xyz+y2z+yz2
print(koszul(2,I)*koszul(3,I));
↳ 0,
↳ 0,
↳ 0

```

See Section 4.6 [int], page 82; Section 4.12 [matrix], page 106.

5.1.74 laguerre

Syntax: `laguerre (poly_expression , int_expression , int_expression)`

Type: list

Purpose: In characteristic 0:
computes all complex roots of a univariate polynomial using Laguerre's algorithm. The second argument defines the precision of the fractional part if the ground field is the field of rational numbers, otherwise it will be ignored. The third argument (can be 0, 1 or 2) gives the number of extra runs for Laguerre's algorithm (with corrupted roots),

leading to better results.

In characteristic p:

computes all roots of a univariate polynomial using factorization

Note: If the ground field is the field of complex numbers, the elements of the list are of type number, otherwise of type string.

Example:

```
ring rs1=0,(x,y),lp;
poly f=15x5+x3+x2-10;
laguerre(f,10,2);
↳ [1]:
↳ 0.8924637479
↳ [2]:
↳ (-0.7392783383+I*0.5355190078)
↳ [3]:
↳ (-0.7392783383-I*0.5355190078)
↳ [4]:
↳ (0.2930464644-I*0.9003002396)
↳ [5]:
↳ (0.2930464644+I*0.9003002396)
```

5.1.75 lead

Syntax: lead (poly_expression)
 lead (vector_expression)
 lead (ideal_expression)
 lead (module_expression)

Type: the same as the input type

Purpose: returns the leading (or initial) term(s) of a polynomial, a vector, resp. of the generators of an ideal or module with respect to the monomial ordering.

Note: IN may be used instead of lead.

Example:

```
ring r=32003,(x,y,z),(c,ds);
poly f=2x2+3y+4z3;
vector v=[2x10,f];
ideal i=f,z;
module m=v,[0,0,2+x];
lead(f);
↳ 3y
lead(v);
↳ [2x10]
lead(i);
↳ _[1]=3y
↳ _[2]=z
lead(m);
↳ _[1]=[2x10]
↳ _[2]=[0,0,2]
lead(0);
↳ 0
```


See Section 4.5 [ideal], page 78; Section 5.1.76 [leadcoef], page 207; Section 5.1.77 [leadexp], page 207; Section 5.1.78 [leadmonom], page 208; Section 4.13 [module], page 110; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.76 leadcoef

Syntax: leadcoef (poly_expression)
 leadcoef (vector_expression)

Type: number

Purpose: returns the leading (or initial) coefficient of a polynomial or a vector with respect to the monomial ordering.

Example:

```

ring r=32003,(x,y,z),(c,ds);
poly f=x2+y+z3;
vector v=[2*x^10,f];
leadcoef(f);
↳ 1
leadcoef(v);
↳ 2
leadcoef(0);
↳ 0

```

See Section 5.1.75 [lead], page 206; Section 5.1.77 [leadexp], page 207; Section 5.1.78 [leadmonom], page 208; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.77 leadexp

Syntax: leadexp (poly_expression)
 leadexp (vector_expression)

Type: intvec

Purpose: returns the exponent vector of the leading monomial of a polynomial or a vector. In the case of a vector the last component is the index in the vector. (The inverse to monomial.)

Example:

```

ring r=32003,(x,y,z),(c,ds);
poly f=x2+y+z3;
vector v=[2*x^10,f];
leadexp(f);
↳ 0,1,0
leadexp(v);
↳ 10,0,0,1
leadexp(0);
↳ 0,0,0

```

See Section 4.8 [intvec], page 91; Section 5.1.75 [lead], page 206; Section 5.1.76 [leadcoef], page 207; Section 5.1.78 [leadmonom], page 208; Section 5.1.96 [monomial], page 221; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.78 leadmonom

Syntax: `leadmonom (poly_expression)`
 `leadmonom (vector_expression)`

Type: the same as the input type

Purpose: returns the leading monomial of a polynomial or a vector as a polynomial or vector whose coefficient is one.

Example:

```

ring r=32003,(x,y,z),(c,ds);
poly f=2x2+3y+4z3;
vector v=[0,2x10,f];
leadmonom(f);
↳ y
leadmonom(v);
↳ [0,x10]
leadmonom(0);
↳ 0

```

See Section 4.8 [intvec], page 91; Section 5.1.75 [lead], page 206; Section 5.1.76 [leadcoef], page 207; Section 5.1.77 [leadexp], page 207; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.79 LIB

Syntax: `LIB string_expression;`

Type: none

Purpose: reads a library of procedures from a file. In contrast to the command `load`, the procedures from the library are added to the package `Top` as well as the package corresponding to the library. If the given filename does not start with `.` or `/` and cannot be located in the current directory, each directory contained in the library `SearchPath` is searched for file of this name. See Section 3.8.11 [Loading a library], page 66, for more info on `SearchPath`.

Note on standard.lib:

Unless SINGULAR is started with the `--no-stdlib` option, the library `standard.lib` is automatically loaded at start-up time.

Example:

```

option(loadLib); // show loading of libraries

// the names of the procedures of inout.lib
LIB "inout.lib"; // are now known to Singular
↳ // ** loaded inout.lib (4.1.2.0, Feb_2019)

```

See Section 3.1.6 [Command line options], page 19; Section 2.3.3 [Procedures and libraries], page 10; Appendix D [SINGULAR libraries], page 785; Section 5.2.12 [load], page 294; Section 4.15 [package], page 117; Section 4.17 [proc], page 121; Section D.1 [standard.lib], page 785; Section 4.21 [string], page 127; Section 5.1.153 [system], page 270.

5.1.80 lift

Syntax: lift (ideal_expression, subideal_expression)
 lift (module_expression, submodule_expression)
 lift (ideal_expression, subideal_expression, matrix_name)
 lift (module_expression, submodule_expression, matrix_name)
 lift (ideal_expression, subideal_expression, matrix_name, string_expression)
 lift (module_expression, submodule_expression, matrix_name, string_expression)

Type: matrix

Purpose: computes the transformation matrix which expresses the generators of a submodule in terms of the generators of a module. Depending on which algorithm is used, modules are represented by a standard basis, or not.
 More precisely, if m is the module (or ideal), sm the submodule (or ideal), and T the transformation matrix returned by lift, then $\text{matrix}(sm)*U = \text{matrix}(m)*T$ and $\text{module}(sm*U) = \text{module}(\text{matrix}(m)*T)$ (resp. $\text{ideal}(sm) = \text{ideal}(\text{matrix}(m)*T)$), where U is a diagonal matrix of units.
 U is always the identity if the basering is a polynomial ring (not power series ring). U is stored in the optional third argument.

Note: Gives a warning if sm is not a submodule.
 An optional 4th argument specifies the Groebner base algorithm to use. Possible values are "std" and "slimgb".

Example:

```
ring r=32003,(x,y,z),(dp,C);
ideal m=3x2+yz,7y6+2x2y+5xz;
poly f=y7+x3+xyz+z2;
ideal i=jacob(f);
matrix T=lift(i,m);
matrix(m)-matrix(i)*T;
↪ _[1,1]=0
↪ _[1,2]=0
```

See Section 5.1.26 [division], page 172; Section 4.5 [ideal], page 78; Section 5.1.81 [liftstd], page 209; Section 4.13 [module], page 110; Section 5.1.149 [std], page 266; Section 5.1.154 [syz], page 275.

5.1.81 liftstd

Syntax: liftstd (ideal_expression, matrix_name)
 liftstd (module_expression, matrix_name)
 liftstd (ideal_expression, matrix_name, module_name)
 liftstd (module_expression, matrix_name, module_name)
 liftstd (ideal_expression, matrix_name, module_name, string_expression)
 liftstd (module_expression, matrix_name, module_name, string_expression)

Type: ideal or module

Purpose: returns a standard basis of an ideal or module and the transformation matrix from the given ideal, resp. module, to the standard basis.
 That is, if m is the ideal or module, sm the standard basis returned by liftstd, and T the transformation matrix then $\text{matrix}(sm)=\text{matrix}(m)*T$ and $sm=\text{ideal}(\text{matrix}(m)*T)$, resp. $sm=\text{module}(\text{matrix}(m)*T)$.
 In an optional third argument the syzygy module will be returned. if

working in a quotient ring, then `matrix(sm)=reduce(matrix(m)*T,0)` and `sm=reduce(ideal(matrix(m)*T),0)`.

An optional 4th argument specifies the Groebner base algorithm to use. Possible values are "std" and "slimgb".

Example:

```

ring R=0,(x,y,z),dp;
poly f=x3+y7+z2+xyz;
ideal i=jacob(f);
matrix T;
ideal sm=liftstd(i,T);
sm;
↳ sm[1]=xy+2z
↳ sm[2]=3x2+yz
↳ sm[3]=yz2+3048192z3
↳ sm[4]=3024xz2-yz2
↳ sm[5]=y2z-6xz
↳ sm[6]=3097158156288z4+2016z3
↳ sm[7]=7y6+xz
print(T);
↳ 0,1,T[1,3], T[1,4],y, T[1,6],0,
↳ 0,0,-3x+3024z,3x, 0, T[2,6],1,
↳ 1,0,T[3,3], T[3,4],-3x,T[3,6],0
matrix(sm)-matrix(i)*T;
↳ _[1,1]=0
↳ _[1,2]=0
↳ _[1,3]=0
↳ _[1,4]=0
↳ _[1,5]=0
↳ _[1,6]=0
↳ _[1,7]=0
module s;
sm=liftstd(i,T,s);
print(s);
↳ -xy-2z,0, s[1,3],s[1,4],s[1,5],s[1,6],
↳ 0, -xy-2z,s[2,3],s[2,4],s[2,5],s[2,6],
↳ 3x2+yz,7y6+xz,s[3,3],s[3,4],s[3,5],s[3,6]

```

See Section 5.1.26 [division], page 172; Section 4.5 [ideal], page 78; Section 5.1.80 [lift], page 209; Section 4.12 [matrix], page 106; Section 5.1.110 [option], page 230; Section 4.19 [ring], page 124; Section 5.1.149 [std], page 266; Section 5.1.154 [syz], page 275.

5.1.82 listvar

Syntax: `listvar ([package])`
`listvar ([package,] type)`
`listvar ([package,] ring_name)`
`listvar ([package,] name)`
`listvar ([package,] all)`

Type: none

Purpose: lists all (user-)defined names:

- `listvar()`: all currently visible names except procedures in the current namespace,
- `listvar(type)`: all currently visible names of the given type,
- `listvar(ring_name)`: all names which belong to the given ring,
- `listvar(name)`: the object with the given name,
- `listvar(all)`: all names except procedures in the current and Top namespace.

The current basering is marked with a *. The nesting level of variables in procedures is shown in square brackets.

package can be `Current`, `Top` or any other identifier of type package.

Example:

```

proc t1 { }
proc t2 { }
ring s;
poly ss;
ring r;
poly f=x+y+z;
int i=7;
ideal I=f,x,y;
listvar();
↳ // i                [0] int 7
↳ // r                [0] *ring
↳ //      I          [0] ideal, 3 generator(s)
↳ //      f          [0] poly
↳ // s                [0] ring
  listvar(r);
↳ // r                [0] *ring
↳ // I                [0] ideal, 3 generator(s)
↳ // f                [0] poly
  listvar(t1);
↳ // t1                [0] proc
  listvar(proc);
↳ // t2                [0] proc
↳ // t1                [0] proc
↳ // mathicgb_prOrder [0] proc from singmathic.so (C)
↳ // mathicgb         [0] proc from singmathic.so (C)
↳ // create_ring      [0] proc from standard.lib
↳ // min               [0] proc from standard.lib
↳ // max               [0] proc from standard.lib
↳ // datetime         [0] proc from standard.lib
↳ // weightKB         [0] proc from standard.lib
↳ // fprintf           [0] proc from standard.lib
↳ // printf            [0] proc from standard.lib
↳ // sprintf           [0] proc from standard.lib
↳ // quotient4        [0] proc from standard.lib
↳ // quotient5        [0] proc from standard.lib
↳ // quotient3        [0] proc from standard.lib
↳ // quotient2        [0] proc from standard.lib
↳ // quotient1        [0] proc from standard.lib
↳ // quot              [0] proc from standard.lib
↳ // res              [0] proc from standard.lib

```

```

↳ // groebner [0] proc from standard.lib
↳ // qslimb [0] proc from standard.lib
↳ // hilbRing [0] proc from standard.lib
↳ // par2varRing [0] proc from standard.lib
↳ // quotientList [0] proc from standard.lib
↳ // stdhilb [0] proc from standard.lib
↳ // stdfglm [0] proc from standard.lib
↳ // Float [0] proc from kernel (C)
↳ // crossprod [0] proc from kernel (C)
LIB "polylib.lib";
listvar(Poly);
↳ ? Poly is undefined
↳ ? error occurred in or before ./examples/listvar.sing line 14: ' li
var(Poly);'

```

See Section 3.5.3 [Names], page 44; Section 3.7.4 [Names in procedures], page 54; Section 5.1.18 [defined], page 167; Section 5.1.102 [names], page 225; Section 4.15 [package], page 117; Section 5.1.158 [type], page 277.

5.1.83 lres

Syntax: `lres (ideal_expression, int_expression)`

Type: resolution

Purpose: computes a free resolution of an ideal using LaScala's algorithm.

More precisely, let R be the basering and I be the given ideal. Then `lres` computes a minimal free resolution of R/I

$$\dots \longrightarrow F_2 \xrightarrow{A_2} F_1 \xrightarrow{A_1} R \longrightarrow R/I \longrightarrow 0.$$

If the `int_expression` k is not zero then the computation stops after k steps and returns a list of modules $M_i = \text{module}(A_i)$, $i=1..k$.

`list L=lres(I,0)`; returns a list L of n modules (where n is the number of variables of the basering) such that $L[i] = M_i$ in the above notation.

Note: The `ideal_expression` has to be homogeneous.

Accessing single elements of a resolution may require that some partial computations have to be finished and may therefore take some time.

Example:

```

ring r=0,(x,y,z),dp;
ideal I=xz,yz,x3-y3;
def L=lres(I,0);
print(betti(L),"betti");
↳          0    1    2
↳ -----
↳    0:    1    -    -
↳    1:    -    2    1
↳    2:    -    1    1
↳ -----
↳ total:    1    3    2
↳
↳ L[2]; // the first syzygy module of r/I
↳ _[1]=-x*gen(1)+y*gen(2)
↳ _[2]=-x2*gen(2)+y2*gen(1)+z*gen(3)

```

See Section 5.1.4 [betti], page 157; Section 5.1.48 [fres], page 186; Section 5.1.58 [hres], page 195; Section 4.5 [ideal], page 78; Section 4.6 [int], page 82; Section 5.1.93 [minres], page 220; Section 4.13 [module], page 110; Section 5.1.98 [mres], page 222; [res], page 785; Section 5.1.147 [sres], page 264.

5.1.84 ludecomp

qcindex Gauss

Syntax: `ludecomp (matrix_expression)`

Type: list

Purpose: Computes the LU-decomposition of an $(m \times n)$ matrix.

The matrix, A say, must consist of numbers, only. This means that when the basering represents some $K[x_1, x_2, \dots, x_r]$, then all entries of A must come from the ground field K .

The LU-decomposition of A is a triple of matrices P , L , and U such that

- $P * A = L * U$,

- P is an $(m \times m)$ permutation matrix, i.e., its rows/columns form the standard basis of K^m ,

- L is an $(m \times m)$ matrix in lower triangular form with all diagonal entries equal to 1, and

- U is an $(m \times n)$ matrix in upper row echelon form.

From these conditions, it easily follows that also $A = P * L * U$ holds, since P is self-inverse.

`list L=ludecomp(A)`; fills a list L with the three above entries P , L , and U .

Example:

```

ring r=0,(x),dp;
matrix A[3][4]=1,2,3,4,1,1,1,1,2,2,1,1;
list plu = ludecomp(A);
print(plu[3]); // the matrix U of the decomposition
↳ 1,2, 3, 4,
↳ 0,-1,-2,-3,
↳ 0,0, -1,-1
print(plu[1]*A-plu[2]*plu[3]); // should be the zero matrix
↳ 0,0,0,0,
↳ 0,0,0,0,
↳ 0,0,0,0

```

See Section 5.1.85 [luinverse], page 213; Section 5.1.86 [lusolve], page 214.

5.1.85 luinverse

qcindex Gauss

Syntax: `luinverse (matrix_expression)`

Type: matrix

Syntax: `luinverse (matrix_expression, matrix_expression, matrix_expression)`

Type: matrix

Purpose: Computes the inverse of a matrix A , if A is invertible.

The matrix A must be given either directly, or by its LU-decomposition. In the latter case, three matrices P , L , and U are expected, in this order, which satisfy

- $P * A = L * U$,
- P is an $(m \times m)$ permutation matrix, i.e., its rows/columns form the standard basis of K^m ,
- L is an $(m \times m)$ matrix in lower triangular form with all diagonal entries equal to 1, and
- U is an $(m \times m)$ matrix in upper row echelon form.

Then, the inverse of A exists if and only if U is invertible, and one has $A^{-1} = U^{-1} \cdot L^{-1} \cdot P$, since P is self-inverse.

In the case of A being given directly, `luinverse` first computes its LU-decomposition, and then proceeds as in the case when P , L , and U are provided.

`list L=luinverse(A)`; fills the list L with either one entry $= 0$ (signaling that A is not invertible), or with the two entries $1, A^{-1}$. Thus, in either case the user may first check the condition `L[1]==1` to find out whether A is invertible.

Note: The method will give a warning for any non-quadratic matrix A .

Example:

```
ring r=0,(x),dp;
matrix A[3][3]=1,2,3,1,1,1,2,2,1;
list L = luinverse(A);
if (L[1] == 1)
{
  print(L[2]);
  "----- next should be the (3 x 3)-unit matrix:";
  print(A*L[2]);
}
↳ -1,4, -1,
↳ 1, -5,2,
↳ 0, 2, -1
↳ ----- next should be the (3 x 3)-unit matrix:
↳ 1,0,0,
↳ 0,1,0,
↳ 0,0,1
```

See Section 5.1.84 [ludecomp], page 213; Section 5.1.86 [lusolve], page 214.

5.1.86 lusolve

qcindex Gauss

Syntax: `lusolve (matrix_expression, matrix_expression, matrix_expression, matrix_expression)`

Type: `matrix`

Purpose: Computes all solutions of a linear equation system $A*x = b$, if solvable
The $(m \times n)$ matrix A must be given by its LU-decomposition, that is, by three matrices P , L , and U , in this order, which satisfy

- $P * A = L * U$,
- P is an $(m \times m)$ permutation matrix, i.e., its rows/columns form the standard basis of K^m ,
- L is an $(m \times m)$ matrix in lower triangular form with all diagonal entries equal to 1, and
- U is an $(m \times n)$ matrix in upper row echelon form.

The fourth argument, b , is expected to be an $(m \times 1)$ matrix.

`list Q=lusolve(P,L,U,b)`; fills the list `Q` with either one entry $= 0$ (signaling that $A*x=b$ has no solution), or with the three entries $1, x, H$, where x is any $(n \times 1)$ solution of the given linear system, and H is a matrix the columns of which span the solution space of the homogeneous linear system. (I.e., `ncols(H)` is the dimension of the solution space.)

If there is exactly one solution, then H is the 1×1 matrix with entry zero.

Note: The method will give a warning if the matrices violate the above conditions regarding row and column numbers, or if the number of rows of the vector `b` does not equal `m`. The method expects matrices with entries coming from the ground field of the given polynomial ring, only.

Example:

```

ring r=0,(x),dp;
matrix A[4][4]=1,1,1,0,1,2,3,1,1,3,5,2,1,4,7,3;
matrix b[4][1]=2,5,8,11;
list L=ludecomp(A);
list Q=lusolve(L[1],L[2],L[3],b);
if (Q[1] == 1)
{
  "one solution:";
  print(Q[2]);
  "check whether result is correct (iff next is zero vector):";
  print(A*Q[2]-b);
  if ((nrows(Q[3])==1) and (ncols(Q[3])==1) and (Q[3][1,1]==0))
  { "printed solution is the only solution to given linear system" }
  else
  {
    "homogeneous solution space is spanned by columns of:";
    print(Q[3]);
  }
}
↪ one solution:
↪ -1,
↪ 3,
↪ 0,
↪ 0
↪ check whether result is correct (iff next is zero vector):
↪ 0,
↪ 0,
↪ 0,
↪ 0
↪ homogeneous solution space is spanned by columns of:
↪ -1,-1,
↪ 1, 2,
↪ 0, -1,
↪ -1,0

```

See Section 5.1.84 [`ludecomp`], page 213; Section 5.1.85 [`luinverse`], page 213.

5.1.87 `max`

Procedure from library `standard.lib` (see Section D.1 [`standard.lib`], page 785).

Syntax: `max (i_1, ..., i_k)`

Type: same as type of i_1, \dots, i_k resp.

Purpose: returns the maximum for any arguments of a type for which ' $>$ ' is defined

Example:

```
// biggest int
max(2,3);
↳ 3
max(1,4,3);
↳ 4
// lexicographically biggest intvec
max(intvec(1,2),intvec(0,1),intvec(1,1));
↳ 1,2
// polynomial with biggest leading monomial
ring r = 0,x,dp;
max(x+1,x2+x);
↳ x2+x
```

See also: [min], page 786.

5.1.88 maxideal

Syntax: maxideal (int_expression)

Type: ideal

Purpose: returns the power given by int_expression of the maximal ideal generated by all ring variables (maxideal(i)=1 for $i \leq 0$).

Example:

```
ring r=32003,(x,y,z),dp;
maxideal(2);
↳ _[1]=z2
↳ _[2]=yz
↳ _[3]=y2
↳ _[4]=xz
↳ _[5]=xy
↳ _[6]=x2
```

See Section 4.5 [ideal], page 78; Section 4.19 [ring], page 124.

5.1.89 memory

Syntax: memory (int_expression)

Type: bigint

Purpose: returns statistics concerning the memory management:

- memory(0) is the number of active (used) bytes,
- memory(1) is the number of bytes allocated from the operating system,
- memory(2) is the maximal number of bytes ever allocated from the operating system during the current SINGULAR session.

Note: To monitor the memory usage during ongoing computations the option mem should be set (using the command option(mem);, see also Section 5.1.110 [option], page 230).

Example:

```

ring r=0,(x(1..500)),dp;
poly p=(x(1)+x(500))^50;
proc ReportMemoryUsage()
{ "Memory currently used by SINGULAR      :",memory(0),"Byte (",
  int(memory(0) div 1024), "KByte)" +newline+
  "Memory currently allocated from system:",memory(1), "Byte (",
  int(memory(1) div 1024), "KByte)";
  "Maximal memory allocated from system  :",memory(2), "Byte (",
  int(memory(2) div 1024), "KByte)";
}
ReportMemoryUsage();
↳ Memory currently used by SINGULAR      : 154464 Byte ( 150 KByte)
↳ Memory currently allocated from system: 2236416 Byte ( 2184 KByte)
↳ Maximal memory allocated from system  : 2236416 Byte ( 2184 KByte)
kill p;
ReportMemoryUsage(); // less memory used: p killed
↳ Memory currently used by SINGULAR      : 83896 Byte ( 82 KByte)
↳ Memory currently allocated from system: 2236416 Byte ( 2184 KByte)
↳ Maximal memory allocated from system  : 2236416 Byte ( 2184 KByte)
kill r;
ReportMemoryUsage(); // even less memory: r killed
↳ Memory currently used by SINGULAR      : 71656 Byte ( 70 KByte)
↳ Memory currently allocated from system: 2236416 Byte ( 2184 KByte)
↳ Maximal memory allocated from system  : 2236416 Byte ( 2184 KByte)

```

See Section 5.1.110 [option], page 230; Section 5.1.153 [system], page 270.

5.1.90 min

Procedure from library `standard.lib` (see Section D.1 [standard.lib], page 785).

Syntax: `max (i_1, ..., i_k)`

Type: same as type of `i_1, ..., i_k` resp.

Purpose: returns the maximum for any arguments of a type for which `'>'` is defined

Example:

```

// biggest int
max(2,3);
↳ 3
max(1,4,3);
↳ 4
// lexicographically biggest intvec
max(intvec(1,2),intvec(0,1),intvec(1,1));
↳ 1,2
// polynomial with biggest leading monomial
ring r = 0,x,dp;
max(x+1,x2+x);
↳ x2+x

```

See also: [min], page 786.

5.1.91 minbase

Syntax: `minbase (ideal_expression)`
`minbase (module_expression)`

Type: the same as the type of the argument

Purpose: returns a minimal set of generators of an ideal, resp. module, if the input is either homogeneous or if the ordering is local.

Note: this command is not available over coefficient rings.

Example:

```

ring r=181,(x,y,z),(c,1s);
ideal id2=x2+xyz,y2-z3y,z3+y5xz;
ideal id4=maxideal(3)+id2;
size(id4);
↳ 13
minbase(id4);
↳ _[1]=x2
↳ _[2]=xyz+x2
↳ _[3]=xz2
↳ _[4]=y2
↳ _[5]=yz2
↳ _[6]=z3

```

See Section 5.1.99 [mstd], page 223.

5.1.92 minor

Syntax: `minor (matrix_expression M, int_expression mSize,`
`[ideal_expression I],`
`[int_expression k],`
`[string_expression algorithm],`
`[int_expression cachedP],`
`[int_expression cachedM])`

Type: ideal

Purpose: returns the specified set of (mSize x mSize)-minors (= subdeterminants) of the given matrix M. These minors form the list of generators of the returned ideal.

If the optional ideal I is given, it is assumed to capture a standard basis. In this case, all computations will be performed modulo I.

If k is not given, all minors will be computed. Otherwise, if $k > 0$, the first k non-zero minors will be computed; for $k < 0$, the first $|k|$ minors will be computed regardless whether they are zero or not. Here, "first k minors" is with respect to a fixed ordering among all minors. (To understand the ordering, run the below example, type `minor(m,2,i,18)`; and inspect the ordering among the returned 18 minors. Note that this ordering is only enforced when some $k \neq 0$ is provided. Otherwise, no ordering among the returned minors can be guaranteed. This is due to the fact that in this case, `minor` may call a specially tuned implementation of Bareiss's algorithm.)

If no algorithm is given, a heuristic will pick the best-suited algorithm among Bareiss's algorithm (which is only applicable over integral domains), Laplace's algorithm, and Laplace's algorithm combined with caching of subdeterminantes. In the heuristic setting, `cacheP` and `cacheM` must also be absent.

If the argument `algorithm` is present it must be one of `B/bareiss`, `L/laplace`, and `C/cache`. For, `B/bareiss` and `L/laplace` the optional arguments `cacheP` and `cacheM` must again be absent, whereas for `C/cache`, they may be provided: `cachedP` determines the maximum number of cached subdeterminantes (=polynomials), and `cachedM` the total number of cached monomials (counted over all cached polynomials). If, for algorithm = `C/cache` `cachedP` and `cachedM` are not provided by the user, the values 200 and 100000, respectively, will be used as defaults.

Note: If `mSize` is larger than the given matrix, `minor` returns 0, if `mSize` is smaller than 1, `minor` returns 1.

Example:

```

ring r=0,(a,b,c,d,e,f,g,h,s,t,u,v),ds;
matrix m[3][4]=a,b,c,d,e,f,g,h,s,t,u,v;
print(m);
↳ a,b,c,d,
↳ e,f,g,h,
↳ s,t,u,v
// let's compute all non-zero minors;
// here we do not guarantee any ordering:
minor(m,2);
↳ _[1]=-hu+gv
↳ _[2]=-ht+fv
↳ _[3]=-hs+ev
↳ _[4]=-du+cv
↳ _[5]=-dt+bv
↳ _[6]=-ds+av
↳ _[7]=gt-fu
↳ _[8]=gs-eu
↳ _[9]=ct-bu
↳ _[10]=cs-au
↳ _[11]=-fs+et
↳ _[12]=-bs+at
↳ _[13]=-dg+ch
↳ _[14]=-df+bh
↳ _[15]=-de+ah
↳ _[16]=cf-bg
↳ _[17]=ce-ag
↳ _[18]=-be+af
ideal i=a,c; i=std(i);
// here come the first 4 non-zero minors mod I;
// this time, a fixed ordering is guaranteed:
minor(m,2,i,4);
↳ _[1]=-be
↳ _[2]=bg
↳ _[3]=-de
↳ _[4]=-df+bh
// and here the first 4 minors mod I (possibly zero)
// using Laplace's algorithm,
// again, the fixed ordering is guaranteed:
minor(m,2,i,-4,"Laplace");
↳ _[1]=-be
↳ _[2]=0
↳ _[3]=bg

```

```
↳ _[4]=--de
```

See Section 5.1.23 [det], page 170.

5.1.93 minres

Syntax: minres (list_expression)

Type: list

Syntax: minres (resolution_expression)

Type: resolution

Purpose: minimizes a free resolution of an ideal or module given by the list_expression, resp. resolution_expression.

Example:

```
ring r1=32003,(x,y),dp;
ideal i=x5+xy4,x3+x2y+xy2+y3;
resolution rs=lres(i,0);
rs;
↳ 1      2      1
↳ r1 <--  r1 <--  r1
↳
↳ 0      1      2
↳
list(rs);
↳ [1]:
↳ _[1]=x3+x2y+xy2+y3
↳ _[2]=xy4
↳ [2]:
↳ _[1]=xy4*gen(1)-x3*gen(2)-x2y*gen(2)-xy2*gen(2)-y3*gen(2)
minres(rs);
↳ 1      2      1
↳ r1 <--  r1 <--  r1
↳
↳ 0      1      2
↳
list(rs);
↳ [1]:
↳ _[1]=x3+x2y+xy2+y3
↳ _[2]=xy4
↳ [2]:
↳ _[1]=xy4*gen(1)-x3*gen(2)-x2y*gen(2)-xy2*gen(2)-y3*gen(2)
```

See Section 5.1.48 [fres], page 186; Section 5.1.98 [mres], page 222; [res], page 785; Section 5.1.147 [sres], page 264.

5.1.94 modulo

Syntax: modulo (ideal_expression, ideal_expression)
modulo (module_expression, module_expression)

Type: module

Purpose: `modulo(h1,h2)` represents $h_1/(h_1 \cap h_2) \cong (h_1 + h_2)/h_2$ where h_1 and h_2 are considered as submodules of the same free module R^l ($l=1$ for ideals). Let H_1 , resp. H_2 , be the matrices of size $l \times k$, resp. $l \times m$, having the generators of h_1 , resp. h_2 , as columns. Then $h_1/(h_1 \cap h_2) \cong R^k/\ker(\overline{H_1})$ where $\overline{H_1} : R^k \rightarrow R^l/Im(H_2) = R^l/h_2$ is the induced map.

`modulo(h1,h2)` returns generators of the kernel of this induced map.

Note: If for at least one of `h1` or `h2` the attribute `"isHomog"` is set, `modulo(h1,h2)` also sets the attribute `"isHomog"` (if possible, that is, if the weights are compatible).

Example:

```
ring r;
ideal h1=x,y,z;
ideal h2=x;
module m=modulo(h1,h2);
print(m);
↪ 1,0, 0,0,
↪ 0,-z,x,0,
↪ 0,y, 0,x
```

See [hom_kernel], page 816; Section 5.1.154 [syz], page 275.

5.1.95 monitor

Syntax: `monitor (link_expression)`
`monitor (link_expression, string_expression)`

Type: none

Purpose: controls the recording of all user input and/or program output into a file. The second argument describes what to log: `"i"` means input, `"o"` means output, `"io"` for both. The default for the second argument is `"i"`.

Each `monitor` command closes a previous monitor file and opens the file given by the first string expression.

`monitor ("")` turns off recording.

Example:

```
monitor("doe.tmp","io"); // log input and output to doe.tmp
ring r;
poly f=x+y+z;
int i=7;
ideal I=f,x,y;
monitor(""); // stop logging:
// doe.tmp contains now all input and output from the example above
```

See Section 4.9.2 [link expressions], page 94.

5.1.96 monomial

Syntax: `monomial (intvec_expression)`

Type: poly resp. vector

Purpose: converts an integer vector to a power product (the inverse to `leadexp`). Returns a `vector` iff the length of the argument is number of variables +1.

Example:

```

ring r=0,(x,y,z),dp;
monomial(intvec(2,3));
↪ x2y3
monomial(intvec(2,3,0,1));
↪ x2y3*gen(1)
leadexp(monomial(intvec(2,3,0,1)));
↪ 2,3,0,1

```

See Section 4.8 [intvec], page 91; Section 5.1.77 [leadexp], page 207.

5.1.97 mpresmat

Syntax: mpresmat (ideal_expression, int_expression)

Type: module

Purpose: computes the multipolynomial resultant matrix of the input system. Uses the sparse resultant matrix method of Gelfand, Kapranov and Zelevinsky (second parameter = 0) or the resultant matrix method of Macaulay (second parameter = 1).

Note: When using the resultant matrix method of Macaulay the input system must be homogeneous. The number of elements in the input system must be the number of variables in the basering plus one.

Example:

```

ring rsq=(0,s,t,u),(x,y),lp;
ideal i=s+tx+uy,x2+y2-10,x2+xy+2y2-16;
module m=mpresmat(i,0);
print(m);
↪ -16,0, -10,0, (s),0, 0, 0, 0, 0,
↪ 0, -16,0, -10,(u),(s),0, 0, 0, 0,
↪ 2, 0, 1, 0, 0, (u),0, 0, 0, 0,
↪ 0, 2, 0, 1, 0, 0, 0, 0, 0, 0,
↪ 0, 0, 0, 0, (t),0, -10,(s),0, -16,
↪ 1, 0, 0, 0, 0, (t),0, (u),(s),0,
↪ 0, 1, 0, 0, 0, 0, 1, 0, (u),2,
↪ 1, 0, 1, 0, 0, 0, 0, (t),0, 0,
↪ 0, 1, 0, 1, 0, 0, 0, 0, (t),1,
↪ 0, 0, 0, 0, 0, 0, 1, 0, 0, 1

```

See Section 5.1.161 [uressolve], page 278.

5.1.98 mres

Syntax: mres (ideal_expression, int_expression)
mres (module_expression, int_expression)

Type: resolution

Purpose: computes a minimal free resolution of an ideal or module M with the standard basis method. More precisely, let $A = \text{matrix}(M)$, then mres computes a free resolution of $\text{coker}(A) = F_0/M$

$$\dots \longrightarrow F_2 \xrightarrow{A_2} F_1 \xrightarrow{A_1} F_0 \longrightarrow F_0/M \longrightarrow 0,$$

where the columns of the matrix A_1 are a minimal set of generators of M if the basering is local or if M is homogeneous. If the int expression k is not zero, then the computation

stops after k steps and returns a list of modules $M_i = \text{module}(A_i)$, $i=1\dots k$.
 $\text{mres}(M,0)$ returns a resolution consisting of at most $n+2$ modules, where n is the number of variables of the basering. Let `list L=mres(M,0)`; then $L[1]$ consists of a minimal set of generators of the input, $L[2]$ consists of a minimal set of generators for the first syzygy module of $L[1]$, etc., until $L[p+1]$, such that $L[i] \neq 0$ for $i \leq p$, but $L[p+1]$, the first syzygy module of $L[p]$, is 0 (if the basering is not a qring).

Note: Accessing single elements of a resolution may require some partial computations to be finished and may therefore take some time.

Example:

```
ring r=31991,(t,x,y,z,w),ls;
ideal M=t2x2+tx2y+x2yz,t2y2+ty2z+y2zw,
      t2z2+tz2w+xz2w,t2w2+txw2+xyw2;
resolution L=mres(M,0);
L;
↳ 1      4      15      18      7      1
↳ r <--  r <--  r <--  r <--  r <--  r
↳
↳ 0      1      2      3      4      5
↳
// projective dimension of M is 5
```

See Section 5.1.48 [fres], page 186; Section 5.1.58 [hres], page 195; Section 4.5 [ideal], page 78; Section 5.1.83 [lres], page 212; Section 4.13 [module], page 110; [res], page 785; Section 5.1.147 [sres], page 264.

5.1.99 mstd

Syntax: `mstd (ideal_expression)`
`mstd (module_expression)`

Type: list

Purpose: returns a list whose first entry is a standard basis for the ideal, resp. module, whose second entry is a generating set for the ideal, resp. module. If the coefficient ring is a field and either the ideal/module is homogeneous or the ordering is local, this second entry is a minimal generating set.

Example:

```
ring r=0,(x,y,z,t),dp;
poly f=x3+y4+z6+xyz;
ideal j=jacob(f),f;
j=homog(j,t);j;
↳ j[1]=3x2+yz
↳ j[2]=4y3+xzt
↳ j[3]=6z5+xyt3
↳ j[4]=0
↳ j[5]=z6+y4t2+x3t3+xyzt3
mstd(j);
↳ [1]:
↳ _[1]=3x2+yz
↳ _[2]=4y3+xzt
↳ _[3]=6z5+xyt3
↳ _[4]=xyzt3
```

```

↳   _[5]=y2z2t3
↳   _[6]=yz3t4
↳   _[7]=xz3t4
↳   _[8]=yz2t7
↳   _[9]=xz2t7
↳   _[10]=y2zt7
↳   _[11]=xy2t7
↳ [2]:
↳   _[1]=3x2+yz
↳   _[2]=4y3+xzt
↳   _[3]=6z5+xyt3
↳   _[4]=xyzt3

```

See Section 4.5 [ideal], page 78; Section 5.1.91 [minbase], page 218; Section 4.13 [module], page 110; Section 5.1.149 [std], page 266.

5.1.100 mult

Syntax: `mult (ideal_expression)`
`mult (module_expression)`

Type: `int`

Purpose: computes the degree of the monomial ideal, resp. module, generated by the leading monomials of the input.

If the input is a standard basis of a homogeneous ideal then it returns the degree of this ideal.

If the input is a standard basis of an ideal in a (local) ring with respect to a local degree ordering then it returns the multiplicity of the ideal (in the sense of Samuel, with respect to the maximal ideal).

Example:

```

ring r=32003,(x,y),ds;
poly f=(x3+y5)^2+x2y7;
ideal i=std(jacob(f));
mult(i);
↳ 46
mult(std(f));
↳ 6

```

See Section 5.1.20 [degree], page 169; Section 5.1.25 [dim], page 171; Section 4.5 [ideal], page 78; Section 5.1.149 [std], page 266; Section 5.1.166 [vdim], page 281.

5.1.101 nameof

Syntax: `nameof (expression)`

Type: `string`

Purpose: returns the name of an expression as string.

Example:

```

int i=9;
string s=nameof(i);
s;
↳ i

```

```

    nameof(s);
    ↪ s
    nameof(i+1); //returns the empty string:
    ↪
    nameof(basering);
    ↪ basering
    basering;
    ↪    ? 'basering' is undefined
    ↪    ? error occurred in or before ./examples/nameof.sing line 7: ' base
    ng;'
    ring r;
    nameof(basering);
    ↪ r

```

See Section 5.1.102 [names], page 225; Section 5.1.133 [reservedName], page 249; Section 5.1.159 [typeof], page 277.

5.1.102 names

Syntax: names ()
 names (ring_name)
 names (package_name)
 names (level)

Type: list of strings

Purpose: returns the names of all user-defined variables which are ring independent (this includes the names of procedures) or, in the second case, which belong to the given ring. The third case restricts the variables to the given level.

package_name can be Current, Top or any other identifier of type package.

Example:

```

    int i=9;
    ring r;
    poly f;
    package p;
    int j; exportto(p,j);
    poly g;
    setring r;
    list l=names();
    l[1..3];
    ↪ l p r
    names(r);
    ↪ [1]:
    ↪    g
    ↪ [2]:
    ↪    f
    names(p);
    ↪ [1]:
    ↪    j
    names(0);
    ↪ [1]:
    ↪    l
    ↪ [2]:

```

```
↳ p
↳ [3]:
↳ r
↳ [4]:
↳ i
↳ [5]:
↳ mathicgb_prOrder
↳ [6]:
↳ mathicgb
↳ [7]:
↳ Singmathic
↳ [8]:
↳ create_ring
↳ [9]:
↳ min
↳ [10]:
↳ max
↳ [11]:
↳ datetime
↳ [12]:
↳ weightKB
↳ [13]:
↳ fprintf
↳ [14]:
↳ printf
↳ [15]:
↳ sprintf
↳ [16]:
↳ quotient4
↳ [17]:
↳ quotient5
↳ [18]:
↳ quotient3
↳ [19]:
↳ quotient2
↳ [20]:
↳ quotient1
↳ [21]:
↳ quot
↳ [22]:
↳ res
↳ [23]:
↳ groebner
↳ [24]:
↳ qslingb
↳ [25]:
↳ hilbRing
↳ [26]:
↳ par2varRing
↳ [27]:
↳ quotientList
↳ [28]:
↳ stdhilb
```

```

↳ [29]:
↳   stdfglm
↳ [30]:
↳   Standard
↳ [31]:
↳   Float
↳ [32]:
↳   crossprod
↳ [33]:
↳   ZZ
↳ [34]:
↳   QQ
↳ [35]:
↳   Top

```

See Section 5.1.101 [nameof], page 224; Section 5.1.133 [reservedName], page 249.

5.1.103 ncols

Syntax: ncols (matrix_expression)
 ncols (smatrix_expression)
 ncols (intmat_expression)
 ncols (ideal_expression)

Type: int

Purpose: returns the number of columns of a matrix, an intmat, or the number of given generators of the ideal, including zeros.

Note: size(ideal) counts the number of generators which are different from zero. (Use nrows to get the number of rows of a given matrix or intmat.)

Example:

```

      ring r;
      matrix m[5][6];
      ncols(m);
↳ 6
      ideal i=x,0,y;
      ncols(i);
↳ 3
      size(i);
↳ 2

```

See Section 4.12 [matrix], page 106; Section 5.1.106 [nrows], page 228; Section 5.1.142 [size], page 259; Section 4.20 [smatrix], page 127.

5.1.104 npars

Syntax: npars (ring_name)

Type: int

Purpose: returns the number of parameters of a ring.

Example:

```

ring r=(23,t,v),(x,a(1..7)),lp;
// the parameters are t,v
npars(r);
↳ 2

```

See Section 5.1.113 [par], page 235; Section 5.1.115 [parstr], page 236; Section 4.19 [ring], page 124.

5.1.105 nres

Syntax: nres (ideal_expression, int_expression)
nres (module_expression, int_expression)

Type: resolution

Purpose: computes a free resolution of an ideal or module M which is minimized from the second module on (by the standard basis method).

More precisely, let $A_1 = \text{matrix}(M)$, then **nres** computes a free resolution of $\text{coker}(A_1) = F_0/M$

$$\dots \longrightarrow F_2 \xrightarrow{A_2} F_1 \xrightarrow{A_1} F_0 \longrightarrow F_0/M \longrightarrow 0,$$

where the columns of the matrix A_1 are the given set of generators of M. If the int expression k is not zero then the computation stops after k steps and returns a list of modules $M_i = \text{module}(A_i)$, $i = 1, \dots, k$.

nres(M,0) returns a list of n modules where n is the number of variables of the basering. Let list L=**nres**(M,0); then L[1]=M is identical to the input, L[2] is a minimal set of generators for the first syzygy module of L[1], etc. (L[i] = M_i in the notations from above).

Example:

```

ring r=31991,(t,x,y,z,w),ls;
ideal M=t2x2+tx2y+x2yz,t2y2+ty2z+y2zw,
        t2z2+tz2w+xz2w,t2w2+txw2+xyw2;
resolution L=nres(M,0);
L;
↳ 1      4      15      18      7      1
↳ r <--  r <--  r <--  r <--  r <--  r
↳
↳ 0      1      2      3      4      5
↳ resolution not minimized yet
↳

```

See Section 5.1.48 [fres], page 186; Section 5.1.58 [hres], page 195; Section 4.5 [ideal], page 78; Section 5.1.83 [lres], page 212; Section 4.13 [module], page 110; Section 5.1.98 [mres], page 222; [res], page 785; Section 4.18 [resolution], page 123; Section 5.1.147 [sres], page 264.

5.1.106 nrows

Syntax: nrows (matrix_expression)
nrows (smatrix_expression)
nrows (intmat_expression)
nrows (intvec_expression)
nrows (module_expression)
nrows (vector_expression)

Type: int

Purpose: returns the number of rows of a matrix, an `intmat` or an `intvec`, resp. the minimal rank of a free module in which the given module or vector lives (the index of the last non-zero component).

Note: Use `ncols` to get the number of columns of a given matrix or `intmat`.

Example:

```

ring R;
matrix M[2][3];
nrows(M);
↳ 2
nrows(freemodule(4));
↳ 4
module m=[0,0,1];
nrows(m);
↳ 3
nrows([0,x,0]);
↳ 2

```

See Section 5.1.51 [gen], page 188; Section 4.12 [matrix], page 106; Section 4.13 [module], page 110; Section 5.1.103 [ncols], page 227; Section 4.20 [smatrix], page 127; Section 4.22 [vector], page 131.

5.1.107 numerator

Syntax: `numerator (number_expression)`

Type: number

Purpose: returns the numerator of a number.

Example:

```

ring r = 0, x, dp;
number n = 3/2;
numerator(n);
↳ 3

```

See Section 5.1.9 [cleardenom], page 161; [content], page 798; Section 5.1.22 [denominator], page 170.

5.1.108 nvars

Syntax: `nvars (ring_name)`

Type: int

Purpose: returns the number of variables of a ring.

Example:

```

ring r=(23,t,v),(x,a(1..7)),ls;
// the variables are x,a(1),...,a(7)
nvars(r);
↳ 8

```

See Section 5.1.104 [npars], page 227; Section 4.19 [ring], page 124; Section 5.1.163 [var], page 279; Section 5.1.165 [varstr], page 280.

5.1.109 open

Syntax: `open (link_expression)`

Type: none

Purpose: opens a link.

Example:

```

link l="ssi:tcp localhost:"+system("Singular");
open(l); // start SINGULAR "server" on localhost in batchmode
close(l); // shut down SINGULAR server

```

See Section 5.1.10 [close], page 161; Section 4.9 [link], page 94.

5.1.110 option

Syntax: `option ()`

Type: string

Purpose: lists all defined options.

Syntax: `option (option_name)`

Type: none

Purpose: sets an option.

Note: To disable an option, use the prefix `no`.

Syntax: `option (get)`

Type: intvec

Purpose: dumps the state of all options to an intvec.

Syntax: `option (set, intvec_expression)`

Type: none

Purpose: restores the state of all options from an intvec (produced by `option(get)`).

Values: The following options are used to manipulate the behavior of computations and act like boolean switches. Use the prefix `no` to disable an option. Notice that some options are ring dependent and reset to their default values on a change of the current basering.

`none` turns off all options (including the `prompt` option).

`warn` be aware of pitfalls. See Section 3.9.7 [option(warn)], page 71.

`returnSB` the functions `syz`, `intersect` (2 arguments), `quotient` return a standard base instead of a generating set if `returnSB` is set. This option should not be used for `lift`.

`fastHC` tries to find the highest corner of the staircase (HC) as fast as possible during a standard basis computation (only used for local orderings).

- infRedTail** local normal form computations will not use the ecart to avoid possibly infinite tail reductions: should only be used with extreme care. By default, it is only set in the case of a zero-dimensional ideal.
- intStrategy** avoids division of coefficients during standard basis computations. This option is ring dependent. By default, it is set for rings with characteristic 0 and not set for all other rings.
- lazy** uses a more lazy approach in std computations, which was used in SINGULAR version before 2-0 (and which may lead to faster or slower computations, depending on the example)
- length** select shorter reducers in std computations,
- notRegularity** disables the regularity bound for **res** and **mres** (see Section 5.1.130 [regularity], page 247).
- notSugar** turns off sugar strategy during standard basis computation and reduction.
- notBuckets** disables the bucket representation of polynomials during standard basis computations. This option usually decreases the memory consumption but increases the computation time. It should only be set for memory-critical standard basis computations.
- prot** shows protocol information indicating the progress during the following computations: **facstd**, **fglm**, **groebner**, **intersect**, **lres**, **mres**, **minres**, **mstd**, **res**, **slimgb**, **sres**, **std**, **stdfglm**, **stdhilb**, **syz**. See below for more details.
- qringNF** simplifies modulo the current **qring** in all assignments.
- redSB** computes a reduced standard basis in any standard basis computation (in rings with global or local orderings, See Section 5.1.64 [interred], page 199 for the discussion of reduced for local orderings))
- redTail** reduction of the tails of polynomials during standard basis computations. This option is ring dependent. By default, it is set for rings with global degree orderings and not set for all other rings. This option changes the reduction strategy and may decrease/increase time and memory consumption - it does not ensure tail reduction on the result - use **redSB** for that.
- redThrough** for inhomogeneous input, polynomial reductions during standard basis computations are never postponed, but always finished through. This option is ring dependent. By default, it is set for rings with global degree orderings and not set for all other rings. This option changes the reduction strategy and may decrease/increase time and memory consumption.
- sugarCrit** uses criteria similar to the homogeneous case to keep more pairs which would be excluded by other criteria but which may be useful for downstream computations. This option changes the strategy for criteria and selection and may decrease/increase time and memory consumption.

- weightM** automatically computes suitable weights for the weighted ecart and the weighted sugar method.
- cancelunit**
avoids to divide polynomials by non-constant units in `std` in the local case. Should usually not be used.
- contentSB**
avoids to divide by the content of a polynomial in `std` and related algorithms. Should usually not be used.
- intersectElim**
prefers elimination to compute intersections (experimental, will be removed in the next release). Should usually not be used.
- intersectSyz**
prefers syzygy methods to compute intersections (experimental, will be removed in the next release). Should usually not be used.

The following options, which also control computations, are special, since they are not manipulated by the `option` command but by a direct assignment of a value. Reset the option by assigning the value 0; the command `option(none)` will not reset them! If there is a non-zero value assigned, the command `option()` prints the option.

- multBound**
a multiplicity bound is set (see Section 5.3.4 [multBound], page 298).
- degBound** a degree bound is set (see Section 5.3.1 [degBound], page 297).

The last set of options controls the output of SINGULAR:

- Imap** shows the mapping of variables with the `fetch` and `imap` commands.
- debugLib** warns about syntax errors when loading a library.
- defRes** shows the names of the syzygy modules while converting `resolution` to `list`
- loadLib** shows loading of libraries (set by default).
- loadProc** shows loading of procedures from libraries.
- mem** shows memory usage in square brackets (see Section 5.1.89 [memory], page 216).
- notWarnSB**
do not warn about using a generating set instead of a standard basis.
- prompt** shows prompt (`>`, resp. `.`) if ready for input (default).
- reading** shows the number of characters read from a file.
- redefine** warns about variable redefinitions (set by default).
- usage** shows correct usage in error messages (set by default).

Example:

```
option(prot);
option();
```

```

↳ //options: prot redefine usage prompt
  option(notSugar);
  option();
↳ //options: prot notSugar redefine usage prompt
  option(noprot);
  option();
↳ //options: notSugar redefine usage prompt
  option(none);
  option();
↳ //options: none
  ring r=0,x,dp;
  degBound=22;
  option();
↳ //options: degBound redTail redThrough intStrategy
  intvec i=option(get);
  option(none);
  option(set,i);
  option();
↳ //options: degBound redTail redThrough intStrategy

```

The output reported on `option(prot)` has the following meaning:

(command)	(character)	(meaning)
facstd	F	found a new factor
		all other characters: like the output of <code>std</code> and <code>reduce</code>
fglm	.	basis monomial found
	+	edge monomial found
	-	border monomial found
groebner		all characters: like the output of <code>std/slimgb</code>
lres	.	minimal syzygy found
	n	slanted degree, i.e., row of Betti matrix
	(mn)	calculate in module n
	g	pair found giving reductum and syzygy
mres	[d]	computations of the d-th syzygy module
		all other characters: like the output of <code>std</code>
minres	[d]	minimizing of the d-th syzygy module
mstd		all characters: like the output of <code>std</code>
reduce	r	reduced a leading term
	t	reduced a non-leading term
res	[d]	computations of the d-th syzygy module
		all other characters: like the output of <code>std</code>
slimgb	M[n,m]	parallel reduction of n elements with m non-zero output elements

	v	candidate for postponing, need to canonicalize
	.	postponed a reduction of a pair/S-polynomial
	b	exchange of a reductor by a 'better' one
	e	a new reductor with non-minimal leading term
	r	redTail reduction
	n	no redTail reduction
	B	resort pairs
	C	slimgb_alg::cleanDegs
	(n)	n critical pairs are still to be reduced
	d	the maximal degree of the leading terms is currently d
sres	.	syzygy found
	(n)	n elements remaining
	[n]	finished module n
std	[m:n]	internal ring change to polynomial representation with exponent bound m and n words in exponent vector
	s	found a new element of the standard basis
	-	reduced a pair/S-polynomial to 0
	.	postponed a reduction of a pair/S-polynomial
	h	used Hilbert series criterion
	H(d)	found a 'highest corner' of degree d, no need to consider higher degrees
	(n)	n critical pairs are still to be reduced
	(S:n)	doing complete reduction of n elements
	d	the degree of the leading terms is currently d
stdfglm		all characters in first part: like the output of <code>std</code> all characters in second part: like the output of <code>fglm</code>
stdhilb		all characters: like the output of <code>std</code>
syz		all characters: like the output of <code>std</code>

See Section 5.3.1 [degBound], page 297; Section 5.3.4 [multBound], page 298; Section 5.1.149 [std], page 266.

5.1.111 ord

Syntax: `ord (poly_expression)`
`ord (vector_expression)`

Type: int

Purpose: returns the (weighted) degree of the initial term of a polynomial or a vector; the weights are the weights used for the first block of the ring ordering.

Note: `ord(0)` is -1.
In a global degree ordering `ord` is the same as `deg`.

Example:

```
ring r=7,(x,y),wp(2,3);
ord(0);
↳ -1
```

```

    poly f=x2+y3; // weight on y is 3
    ord(f),deg(f);
    ↪ 9 9
    ring R=7,(x,y),ws(2,3);
    poly f=x2+y3;
    ord(f),deg(f);
    ↪ 4 9
    vector v=[x2,y];
    ord(v),deg(v);
    ↪ 3 4

```

See Section 5.1.19 [deg], page 168; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.112 ordstr

Syntax: ordstr (ring_name)

Type: string

Purpose: returns the description of the monomial ordering of the ring.

Example:

```

    ring r=7,(x,y),wp(2,3);
    ordstr(r);
    ↪ wp(2,3),C

```

See Section 5.1.7 [charstr], page 160; Section 5.1.115 [parstr], page 236; Section 4.19 [ring], page 124; Section 5.1.165 [varstr], page 280.

5.1.113 par

Syntax: par (int_expression)

Type: number

Purpose: par(n); returns the n-th parameter of the basering.

Example:

```

    ring r=(0,a,b,c),(x,y,z),dp;
    char(r); // char to get the characteristic
    ↪ 0
    par(2); // par to get the n-th parameter
    ↪ (b)

```

See Section 5.1.5 [char], page 159; Section 5.1.104 [npars], page 227; Section 5.1.115 [parstr], page 236; Section 4.19 [ring], page 124; Section 5.1.163 [var], page 279.

5.1.114 pardeg

Syntax: pardeg (number_expression)

Type: int

Purpose: returns the degree of a number considered as a polynomial in the ring parameters.

Example:

```

ring r=(0,a,b,c),(x,y,z),dp;
pardeg(a^2*b);
↳ 3

```

See Section 5.1.19 [deg], page 168; Section 4.14 [number], page 113; Section 4.19 [ring], page 124; Section 5.1.163 [var], page 279.

5.1.115 parstr

Syntax: parstr (ring_name)
parstr (int_expression)
parstr (ring_name, int_expression)

Type: string

Purpose: returns the list of parameters of the ring as a string or the name of the n-th parameter where n is given by the int_expression.
If the ring_name is omitted, the basering is used, thus parstr(n) is equivalent to parstr(basering,n).

Example:

```

ring r=(7,a,b,c),(x,y),wp(2,3);
parstr(r);
↳ a,b,c
parstr(2);
↳ b
parstr(r,3);
↳ c

```

See Section 5.1.7 [charstr], page 160; Section 5.1.104 [npars], page 227; Section 5.1.112 [ordstr], page 235; Section 4.19 [ring], page 124; Section 5.1.165 [varstr], page 280.

5.1.116 preimage

Syntax: preimage (map)
preimage (ring_name, map_name, ideal_name)
preimage (ring_name, ideal_expression, ideal_name)

Type: ring
ideal

Purpose: returns the source ring of a map (in the first case) or returns the preimage of an ideal under a given map.
The second argument has to be a map from the basering to the given ring (or an ideal defining such a map), and the ideal has to be an ideal in the given ring.

Note: As preimage is handling ideals (not polynomials), the result of a preimage calculation of a principal ideal is (the closure of) the preimage of the ideal, not that of the polynomial.

Example:

```

ring r1=32003,(x,y,z,w),lp;
ring r=32003,(x,y,z),dp;
ideal i=x,y,z;
ideal i1=x,y;
ideal i0=0;
map f=r1,i;

```

```

        nameof (preimage (f));
    ↪ r1
        setring r1;
        ideal i1=preimage(r,f,i1);
        i1;
    ↪ i1[1]=w
    ↪ i1[2]=y
    ↪ i1[3]=x
        // the kernel of f
        preimage(r,f,i0);
    ↪ _[1]=w
        // or, use:
        kernel(r,f);
    ↪ _[1]=w

```

See Section 4.5 [ideal], page 78; Section 5.1.70 [kernel], page 203; Section 4.11 [map], page 103; Section 4.19 [ring], page 124.

5.1.117 prime

Syntax: prime (int_expression)

Type: int

Purpose: returns the largest prime less than or equal to the argument; returns 2 for all arguments smaller than 3.

Example:

```

        prime(320000);
    ↪ 319993
        prime(32004);
    ↪ 32003
        prime(0);
    ↪ 2
        prime(-1);
    ↪ 2

```

See Section D.2.3 [general.lib], page 790; Section 4.6 [int], page 82.

5.1.118 primefactors

Syntax: primefactors (int/bigint/number_expression)
 primefactors (int/bigint/number_expression , int_expression)

Type: list

Purpose: returns the prime factorisation up to an optionally given bound, b, on the prime factors. When called with int(s)/bigint(s), no ring needs to be active. When called with numbers these are assumed to be integers in a polynomial ring over Q.

The method finds all prime factors of an integer n. n' will contain the sign, be zero, or the rest (when a bound is given) respectively. The returned list contains the following information: The returned list contains the following information:

L[1][i] = i-th prime factor (in ascending order),
 L[2][i] = multiplicity of L[1][i],
 L[3] = n'

Example:

```

bigint n = bigint(7)^12 * bigint(37)^6 * 121;
primefactors(n);
↳ [1]:
↳ [1]:
↳ 7
↳ [2]:
↳ 11
↳ [3]:
↳ 37
↳ [2]:
↳ [1]:
↳ 12
↳ [2]:
↳ 2
↳ [3]:
↳ 6
↳ [3]:
↳ 1
primefactors(n,25);
↳ [1]:
↳ [1]:
↳ 7
↳ [2]:
↳ 11
↳ [2]:
↳ [1]:
↳ 12
↳ [2]:
↳ 2
↳ [3]:
↳ 2565726409

```

See Section 5.1.117 [prime], page 237.

5.1.119 print

Syntax: `print (expression)`
`print (expression, "betti")`
`print (expression, format_string)`

Type: string

Purpose: The first form prints the expression.
The second form prints the graded Betti numbers from a matrix. The Betti numbers are printed in a matrix-like format where the entry d in row i and column j is the minimal number of generators in degree $i + j$ of the j -th syzygy module of R^n/M (the 0th and 1st syzygy module of R^n/M is R^n and M , resp.).
The last form returns the printed output as a string depending on the format string `iw` which determines the format to use to generate the string.

The following format strings are supported:

`"%s"` returns `string(expression)`,

"%2s"	similar to "%s", except that newlines are inserted after every comma and at the end,
"%1"	similar to "%s", except that each object is embraced by its type such that it can be directly used for "cutting and pasting",
"%21"	similar to "%1", except that newlines are inserted after every comma and at the end,
"%;"	returns the string equivalent to typing <code>expression</code> ;
"%t"	returns the string equivalent to typing <code>type expression</code> ;
"%p"	returns the string equivalent to typing <code>print(expression)</code> ;
"%b"	returns the string equivalent to typing <code>print(expression, "betti")</code> ;
"betti"	is not a format string.

Example:

```

ring r=0,(x,y,z),dp;
module m=[1,y],[0,x+z];
m;
↳ m[1]=y*gen(2)+gen(1)
↳ m[2]=x*gen(2)+z*gen(2)
print(m); // the columns generate m
↳ 1,0,
↳ y,x+z
string s=print(m,"%s"); s;
↳ y*gen(2)+gen(1),x*gen(2)+z*gen(2)
s=print(m,"%2s"); s;
↳ y*gen(2)+gen(1),
↳ x*gen(2)+z*gen(2)
↳
s=print(m,"%1"); s;
↳ module(y*gen(2)+gen(1),x*gen(2)+z*gen(2))
s=print(m,"%;"); s;
↳ m[1]=y*gen(2)+gen(1)
↳ m[2]=x*gen(2)+z*gen(2)
↳
s=print(m,"%t"); s;
↳ // m module , rk 2
↳ m[1]=y*gen(2)+gen(1)
↳ m[2]=x*gen(2)+z*gen(2)
s=print(m,"%p"); s;
↳ 1,0,
↳ y,x+z
intmat M=betti(mres(m,0));
print(M,"betti");
↳
0 1
↳ -----
0: 1 1
↳ -----
↳ total: 1 1
↳
list l=r,M;
s=print(l,"%s"); s;

```

```

↳ (QQ), (x,y,z), (dp(3),C), 1,1
   s=print(1,"%2s"); s;
↳ (QQ), (x,y,z), (dp(3),C),
↳ 1,1
↳
   s=print(1,"%1"); s;
↳ list("(QQ), (x,y,z), (dp(3),C)", intmat(intvec(1,1 ), 1,2))

```

See Section 3.5.5 [Type conversion and casting], page 46; Section 5.1.4 [betti], page 157; Section 5.1.17 [dbprint], page 167; [fprintf], page 785; [printf], page 785; Section 5.3.7 [short], page 300; [sprintf], page 785; Section 4.21.3 [string type cast], page 128; Section 5.1.158 [type], page 277.

5.1.120 printf

Procedure from library `standard.lib` (see Section D.1 [standard.lib], page 785).

Syntax: `printf (string_expression [, any_expressions])`

Return: none

Purpose: `printf(fmt, ...)`; performs output formatting. The first argument is a format control string. Additional arguments may be required, depending on the content of the control string. A series of output characters is generated as directed by the control string; these characters are displayed (i.e., printed to standard out).

The control string `fmt` is simply text to be copied, except that the string may contain conversion specifications.

Type `help print`; for a listing of valid conversion specifications. As an addition to the conversions of `print`, the `%n` and `%2` conversion specification does not consume an additional argument, but simply generates a newline character.

Note: If one of the additional arguments is a list, then it should be enclosed once more into a `list()` command, since passing a list as an argument flattens the list by one level.

Example:

```

ring r=0, (x,y,z), dp;
module m=[1,y], [0,x+z];
intmat M=betti(mres(m,0));
list l=r,m,matrix(M);
printf("s:%s,l:%1",1,2);
↳ s:1,l:int(2)
printf("s:%s",l);
↳ s:(QQ), (x,y,z), (dp(3),C)
printf("s:%s",list(l));
↳ s:(QQ), (x,y,z), (dp(3),C), y*gen(2)+gen(1), x*gen(2)+z*gen(2), 1,1
printf("2l:%2l",list(l));
↳ 2l:list("(QQ), (x,y,z), (dp(3),C)",
↳ module(y*gen(2)+gen(1),
↳ x*gen(2)+z*gen(2)),
↳ matrix(ideal(1,
↳ 1),1,2))
↳
printf("%p",matrix(M));
↳ 1,1
printf(";",matrix(M));
↳ _[1,1]=1

```

```

↳ _[1,2]=1
↳
printf("%b",M);
↳          0      1
↳ -----
↳    0:    1      1
↳ -----
↳ total:   1      1
↳

```

See also: [fprintf], page 785; Section 5.1.119 [print], page 238; [sprintf], page 785; Section 4.21 [string], page 127.

5.1.121 prune

Syntax: `prune (module_expression)`

Type: `module`

Purpose: returns the module minimally embedded in a free module such that the corresponding factor modules are isomorphic.

Note: If for the input module the attribute "isHomog" is set, `prune` also sets the attribute "isHomog".

For non-global orderings, only reduction steps with constant units are performed. Hence, the returned module does not need to be minimal.

Example:

```

ring r=0,(x,y,z),dp;
module m=gen(1),gen(3),[x,y,0,z],[x+y,0,0,0,1];
print(m);
↳ 1,0,x,x+y,
↳ 0,0,y,0,
↳ 0,1,0,0,
↳ 0,0,z,0,
↳ 0,0,0,1
print(prune(m));
↳ y,
↳ z

```

See Section 4.13 [module], page 110.

5.1.122 qhweight

Syntax: `qhweight (ideal_expression)`

Type: `intvec`

Purpose: computes the weight vector of the variables for a quasihomogeneous ideal. If the input is not weighted homogeneous, an intvec of zeros is returned.

Example:

```

ring h1=32003,(t,x,y,z),dp;
ideal i=x4+y3+z2;
qhweight(i);
↳ 0,3,4,6

```

See Section 4.5 [ideal], page 78; Section 4.8 [intvec], page 91; Section 5.1.170 [weight], page 283.

5.1.123 qrds

Syntax: `qrds (matrix_expression, number_expression, number_expression, number_expression)`

Type: list

Purpose: computes all eigenvalues with multiplicities of the given matrix by performing the numeric QR double shift algorithm involving Hessenberg form and householder transformations.

This method expects the ground field to be the complex numbers, and all matrix entries to be real numbers, i.e., elements of this ground field with the imaginary part equal to zero.

If the algorithm works, then it returns a list with two entries which are again lists of the same size:

`_[1][i]` is the i -th mutually distinct eigenvalue that was found,

`_[2][i]` is the (int) multiplicity of `_[1][i]`.

If the algorithm does not work (due to an ill-posed matrix), a list with the single entry (int)0 is returned.

The first number argument is used for detection of deflation in the actual QR double shift algorithm. The second number argument is used for ending Heron's iteration whenever square roots are being computed. And the third number argument is used to distinguish between distinct eigenvalues: When the Euclidean distance between two computed eigenvalues is less than this number, then they will be regarded equal, resulting in a higher multiplicity of the corresponding eigenvalue. (A good choice for all three number arguments is a small value like e.g. 10^{-100} .)

Example:

```
ring r=(complex,50),(dummy),dp;
matrix A[3][3]=-10,37,-5,-14,51,-10,-29,99,-18;
bigint b = bigint(10)^100; number t = 1/b;
list L=qrds(A,t,t,t); L;
⇒ [1]:
⇒ [1]:
⇒ (3+i*2)
⇒ [2]:
⇒ (3-i*2)
⇒ [3]:
⇒ 17
⇒ [2]:
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ 1
⇒ [3]:
⇒ 1
```

5.1.124 quote

Syntax: `quote (expression)`

Type: none

Purpose: prevents expressions from evaluation. Used only in connections with write to ssi links, prevents evaluation of an expression before sending it to an other SINGULAR process.

Within a quoted expression, the quote can be "undone" by an `eval` (i.e., each `eval` "undoes" the effect of exactly one quote).

Example:

```

link l="ssi:w example.ssi";
ring r=0,(x,y,z),ds;
ideal i=maxideal(3);
ideal j=x7,x2,z;
option(prot);
// compute i+j before writing, but not std
write (l, quote(std(eval(i+j))));
close(l);
// now read it in again and evaluate:
read(l);
↳ [65535:1]1(12)s2(11)s3(10)--s(7)s(6)-----7-
↳ product criterion:4 chain criterion:0
↳ _[1]=z
↳ _[2]=x2
↳ _[3]=xy2
↳ _[4]=y3
close(l);

```

See Section 4.9.5 [Ssi links], page 96; Section 5.1.29 [eval], page 174; Section 5.1.172 [write], page 284.

5.1.125 quotient

Syntax: `quotient (ideal_expression, ideal_expression)`
`quotient (module_expression, module_expression)`

Type: ideal

Syntax: `quotient (module_expression, ideal_expression)`

Type: module

Purpose: computes the ideal quotient, resp. module quotient. Let R be the basering, I, J ideals and M a module in R^n . Then

$$\text{quotient}(I, J) = \{a \in R \mid aJ \subset I\},$$

$$\text{quotient}(M, J) = \{b \in R^n \mid bJ \subset M\}.$$

Example:

```

ring r=181,(x,y,z),(c,ls);
ideal id1=maxideal(3);
ideal id2=x2+xyz,y2-z3y,z3+y5xz;
ideal id6=quotient(id1,id2);
id6;
↳ id6[1]=z
↳ id6[2]=y
↳ id6[3]=x
quotient(id2,id1);
↳ _[1]=z2
↳ _[2]=yz
↳ _[3]=y2
↳ _[4]=xz
↳ _[5]=xy

```

```

↳ _[6]=x2
module m=x*freemodule(3),y*freemodule(2);
ideal id3=x,y;
quotient(m,id3);
↳ _[1]=[1]
↳ _[2]=[0,1]
↳ _[3]=[0,0,x]

```

See Section 5.1.40 [fglmquot], page 182; Section 4.5 [ideal], page 78; [modQuotient], page 818; Section 4.13 [module], page 110.

5.1.126 random

Syntax: `random (int_expression , int_expression)`

Type: `int`

Purpose: returns a random integer between the integer given by the first `int_expression` and the one given by the second `int_expression`.

Syntax: `random (int_expression , int_expression , int_expression)`

Type: `intmat`

Purpose: returns a random `intmat` where the size is given by the second (number of rows) and third argument (number of columns). The absolute value of the entries of the matrix is smaller than or equal to the integer given as the first argument.

Note: The random generator can be set to a startvalue with the function `system`, resp. by a command line option. The current value of the random generator is `system("random")`.

Internally a random generator with values in 1 to $2^{31} - 2$ and a full period is used, max-min may not be larger than $2^{31}-2$.

Example:

```

random(1,1000);
↳ 35
random(1,2,3);
↳ 0,0,0,
↳ 1,1,-1
system("random",210); // start random generator with 210
random(-1000,1000);
↳ 707
random(-1000,1000);
↳ 284
system("random",210);
random(-1000,1000); // the same random values again
↳ 707

```

See Section 3.1.6 [Command line options], page 19; Section 4.6 [int], page 82; Section 4.7 [intmat], page 88; Section 5.1.153 [system], page 270.

5.1.127 rank

Syntax: `rank (matrix_expression) #*rank (matrix_expression , 1)`

Type: int

Purpose: returns the rank of a given matrix which is filled with elements of the ground field. The first variant uses a LU-decomposition, the second a row-echelon form.

Note: The function works by computing the row echelon form of the matrix using the same algorithm as for `ludecomp`.

Example:

```
ring s = 0, x, dp;
matrix A[100][100];
int i; int j; int r;
for (i = 1; i <= 100; i++)
{
  for (j = 1; j <= 100; j++)
  {
    A[i, j] = random(-10, 10);
  }
}
r = rank(A); r;
↳ 100
```

See Section 5.1.84 [`ludecomp`], page 213.

5.1.128 read

Syntax: `read (link_expression)`
for DBM links:
`read (link_expression)`
`read (link_expression, string_expression)`

Type: any

Purpose: reads data from a link.

For ASCII links, the content of the entire file is returned as a string. If the ASCII link is the empty string, `read` reads from standard input.

For ssi links, one expression is read from the link and returned after evaluation. See Section 4.9.5 [Ssi links], page 96.

For ssi links the `read` command blocks as long as there is no data to be read from the link. The `status` command can be used to check whether or not there is data to be read.

For DBM links, a `read` with one argument returns the value of the next entry in the data base, and a `read` with two arguments returns the value to the key given as the second argument from the data base. See Section 4.9.7 [DBM links], page 99.

Example:

```
ring r=32003,(x,y,z),dp;
ideal i=x+y,z3+22y;
// write the ideal i to the file save_i
write(":w save_i",i);
ring r0=0,(x,y,z),Dp;
// create an ideal k equal to the content
// of the file save_i
string s="ideal k="+read("save_i")+";";
execute(s);
```

```

k;
↪ k[1]=x+y
↪ k[2]=z3+22y

```

See Section 5.1.32 [execute], page 176; Section 5.1.52 [getdump], page 189; Section 4.9 [link], page 94; Section 5.1.148 [status], page 265; Section 5.1.172 [write], page 284.

5.1.129 reduce

Syntax:

```

reduce ( poly_expression, ideal_expression )
reduce ( poly_expression, ideal_expression, int_expression )
reduce ( poly_expression, poly_expression, ideal_expression )
reduce ( vector_expression, ideal_expression )
reduce ( vector_expression, ideal_expression, int_expression )
reduce ( vector_expression, module_expression )
reduce ( vector_expression, module_expression, int_expression )
reduce ( vector_expression, poly_expression, module_expression )
reduce ( ideal_expression, ideal_expression )
reduce ( ideal_expression, ideal_expression, int_expression )
reduce ( ideal_expression, matrix_expression, ideal_expression )
reduce ( module_expression, ideal_expression )
reduce ( module_expression, ideal_expression, int_expression )
reduce ( module_expression, module_expression )
reduce ( module_expression, module_expression, int_expression )
reduce ( module_expression, matrix_expression, module_expression )
reduce ( poly/vector/ideal/module, ideal/module, int, intvec )
reduce ( ideal, matrix, ideal, int )
reduce ( poly, poly, ideal, int )
reduce ( poly, poly, ideal, int, intvec )

```

Type: the type of the first argument

Purpose: reduces a polynomial, vector, ideal or module to its normal form with respect to an ideal or module represented by a standard basis. Returns 0 if and only if the polynomial (resp. vector, ideal, module) is an element (resp. subideal, submodule) of the ideal (resp. module). The result may have no meaning if the second argument is not a standard basis.

The third (optional) argument of type int modifies the behavior:

0 default

1 consider only the leading term and do no tail reduction.

2 tail reduction:n the local/mixed ordering case: reduce also with bad ecart

4 reduce without division, return possibly a non-zero constant multiple of the remainder

If a second argument u of type poly or matrix is given, the first argument p is replaced by p/u . This works only for zero dimensional ideals (resp. modules) in the third argument and gives, even in a local ring, a reduced normal form which is the projection to the quotient by the ideal (resp. module). One may give a degree bound in the fourth argument with respect to a weight vector in the fifth argument in order to have a finite computation. If some of the weights are zero, the procedure may not terminate!

Note: The commands `reduce` and `NF` are synonymous.

Example:

```

ring r1 = 0,(z,y,x),ds;
poly s1=2x5y+7x2y4+3x2yz3;
poly s2=1x2y2z2+3z8;
poly s3=4xy5+2x2y2z3+11x10;
ideal i=s1,s2,s3;
ideal j=std(i);
reduce(3z3yx2+7y4x2+yx5+z12y2x2,j);
↳ -yx5+2401/81y14x2+2744/81y11x5+392/27y8x8+224/81y5x11+16/81y2x14
reduce(3z3yx2+7y4x2+yx5+z12y2x2,j,1);
↳ -yx5+z12y2x2
// 4 arguments:
ring rs=0,x,ds;
// normalform of 1/(1+x) w.r.t. (x3) up to degree 5
reduce(poly(1),1+x,ideal(x3),5);
↳ // ** _ is no standard basis
↳ 1-x+x2

```

See Section 5.1.26 [division], page 172; Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 4.16.3 [poly operations], page 119; Section 5.1.149 [std], page 266; Section 4.22 [vector], page 131.

5.1.130 regularity

Syntax: regularity (list_expression)
regularity (resolution_expression)

Type: int

Purpose: computes the regularity of a homogeneous ideal, resp. module, from a minimal resolution given by the argument.

Let $0 \rightarrow \bigoplus_a K[x]e_{a,n} \rightarrow \dots \rightarrow \bigoplus_a K[x]e_{a,0} \rightarrow I \rightarrow 0$ be a minimal resolution of I considered with homogeneous maps of degree 0. The regularity is the smallest number s with the property $\deg(e_{a,i}) \leq s + i$ for all i .

Note: If applied to a non minimal resolution only an upper bound is returned.
If the input to the commands `res` and `mres` is homogeneous the regularity is computed and used as a degree bound during the computation unless `option(notRegularity)`; is given.

Example:

```

ring rh3=32003,(w,x,y,z),(dp,C);
poly f=x11+y10+z9+x5y2+x2y2z3+xy3*(y2+x)^2;
ideal j=homog(jacob(f),w);
def jr=res(j,0);
regularity(jr);
↳ 25
// example for upper bound behaviour:
list jj=jr;
regularity(jj);
↳ 25
jj=nres(j,0);
regularity(jj);
↳ 27

```

```

    jj=minres(jj);
    regularity(jj);
    ↪ 25

```

See Section 5.1.48 [fres], page 186; Section 4.10 [list], page 101; Section 5.1.93 [minres], page 220; Section 5.1.98 [mres], page 222; Section 5.1.110 [option], page 230; [res], page 785; Section 4.18 [resolution], page 123; Section 5.1.147 [sres], page 264.

5.1.131 repart

Syntax: `repart (number_expression)`

Type: number

Purpose: returns the real part of a number from a complex ground field, returns its argument otherwise.

Example:

```

    ring r=(complex,i),x,dp;
    repart(1+2*i);
    ↪ 1

```

See Section 5.1.60 [impart], page 196.

5.1.132 res

Procedure from library `standard.lib` (see Section D.1 [standard_lib], page 785).

Syntax: `res (ideal_expression, int_expression [, any_expression])`
`res (module_expression, int_expression [, any_expression])`

Type: resolution

Purpose: computes a (possibly minimal) free resolution of an ideal or module using a heuristically chosen method.

The second (int) argument (say `k`) specifies the length of the resolution. If it is not positive then `k` is assumed to be the number of variables of the basering.

If a third argument is given, the returned resolution is minimized.

Depending on the input, the returned resolution is computed using the following methods:

quotient rings:

`nres` (classical method using syzygies), see Section 5.1.105 [nres], page 228.

homogeneous ideals and `k=0`:

`lres` (La'Scala's method), see Section 5.1.83 [lres], page 212.

not minimized resolution and (homogeneous input with `k` not 0, or local rings):

`sres` (Schreyer's method), see Section 5.1.147 [sres], page 264.

all other inputs:

`mres` (classical method), see Section 5.1.98 [mres], page 222.

Note: Accessing single elements of a resolution may require some partial computations to be finished and may therefore take some time.

See also Section 5.1.4 [beti], page 157; Section 5.1.48 [fres], page 186; Section 5.1.58 [hres], page 195; Section 4.5 [ideal], page 78; Section 5.1.83 [lres], page 212; Section 5.1.93 [minres], page 220;

Section 4.13 [module], page 110; Section 5.1.98 [mres], page 222; Section 5.1.105 [nres], page 228; Section 4.18 [resolution], page 123; Section 5.1.147 [sres], page 264.

Example:

```

ring r=0,(x,y,z),dp;
ideal i=xz,yz,x3-y3;
def l=res(i,0); // homogeneous ideal: uses lres
l;
↳ 1      3      2
↳ r <--  r <--  r
↳
↳ 0      1      2
↳
print(betti(l), "betti"); // input to betti may be of type resolution
↳          0      1      2
↳ -----
↳  0:      1      -      -
↳  1:      -      2      1
↳  2:      -      1      1
↳ -----
↳ total:    1      3      2
↳
l[2];          // element access may take some time
↳ _[1]=-x*gen(1)+y*gen(2)
↳ _[2]=-x2*gen(2)+y2*gen(1)+z*gen(3)
i=i,x+1;
l=res(i,0);    // inhomogeneous ideal: uses mres
l;
↳ 1      3      3      1
↳ r <--  r <--  r <--  r
↳
↳ 0      1      2      3
↳ resolution not minimized yet
↳
ring rs=0,(x,y,z),ds;
ideal i=imap(r,i);
def l=res(i,0); // local ring not minimized: uses sres
l;
↳ 1      1
↳ rs <--  rs
↳
↳ 0      1
↳ resolution not minimized yet
↳
res(i,0,0);    // local ring and minimized: uses mres
↳ 1      1
↳ rs <--  rs
↳
↳ 0      1
↳

```

5.1.133 reservedName

Syntax: reservedName ()

Type: none

Syntax: reservedName (string_expression)

Type: int

Purpose: prints a list of all reserved identifiers (first form) or tests whether the string is a reserved identifier (second form). This includes blackbox/newstruct types.

Example:

```
reservedName();
↳ ... // output skipped
reservedName("ring");
↳ 1
reservedName("xyz");
↳ 0
```

See Section 5.1.102 [names], page 225; Section 4.21 [string], page 127.

5.1.134 resultant

Syntax: resultant (poly_expression, poly_expression, ring_variable)

Type: poly

Purpose: computes the resultant of the first and second argument with respect to the variable given as the third argument.

Example:

```
ring r=32003,(x,y,z),dp;
poly f=3*(x+2)^3+y;
poly g=x+y+z;
resultant(f,g,x);
↳ 3y3+9y2z+9yz2+3z3-18y2-36yz-18z2+35y+36z-24
```

See Section 4.16 [poly], page 117; Section 4.19 [ring], page 124.

5.1.135 ringlist

Syntax: ringlist (ring_expression)

Type: list

Purpose: decomposes a ring/qring into a list of 4 (or 6 in the non-commutative case, see Section 7.3.24 [ringlist (plural)], page 352) components. It is identical to `ring_list` with the exception of the first list entry.

1. the field description in the following format:

for \mathbb{Q} , \mathbb{Z}/p : the characteristic, type int (0 or prime number)

for real, complex: a list of:

the characteristic, type int (always 0)

the precision, type list (2 integers: external, internal precision)

the name of the imaginary unit, type string

for transcendental or algebraic extensions: described as a ringlist (that is, as list L with 4 entries: L[1] the characteristic, L[2] the names of the parameters, L[3] the monomial ordering for the ring of parameters (default: lp), L[4] the minimal polynomial (as ideal))

for Z , Z/n , Z/n^m a list ["integer", [n, m]] with:
 the base n is of type int or bigint (if not given $n = 0$, $Z/0 = Z$)
 the exponent m is of type int (if not given $m = 1$)

2. the names of the variables (a list L of strings: $L[i]$ is the name of the i -th variable)
3. the monomial ordering (a list L of lists): each block $L[i]$ consists of
 - the name of the ordering (string)
 - parameters specifying the ordering and the size of the block (intvec : typically the weights for the variables [default: 1])
4. the quotient ideal.

From a list of such structure, a new ring may be defined by the command `ring` (see the following example). If the attribute "maxExp" of the ring is different from the default 32767, it is also set for the list.

Note: All data which depends on a ring belong to the current ring, not to a ring which can be constructed from a modified list. These data will be mapped via `fetch` to the ring to be constructed.

Example:

```

ring r = 0, (x(1..3)), dp;
list l = ringlist(r);
1;
↳ [1]:
↳ 0
↳ [2]:
↳ [1]:
↳ x(1)
↳ [2]:
↳ x(2)
↳ [3]:
↳ x(3)
↳ [3]:
↳ [1]:
↳ [1]:
↳ dp
↳ [2]:
↳ 1,1,1
↳ [2]:
↳ [1]:
↳ C
↳ [2]:
↳ 0
↳ [4]:
↳ _[1]=0
// Now change l and create a new ring, by
//- changing the base field to the function field with parameter a,
//- introducing one extra variable y,
//- defining the block ordering (dp(2),wp(3,4)).
//- define the minpoly after creating the function field
l[1]=list(0,list("a"),list(list("lp",1)),ideal(0));
l[2][size(l[2])+1]="y";
l[3][3]=l[3][2]; // save the module ordering

```

```

l[3][1]=list("dp",intvec(1,1));
l[3][2]=list("wp",intvec(3,4));
attrib(l,"maxExp",100); // and lower the limit for exponents to 100
def ra = ring(l);      //creates the newring
ra; setring ra;
↳ // coefficients: QQ(a)
↳ // number of vars : 4
↳ //      block  1 : ordering dp
↳ //                : names  x(1) x(2)
↳ //      block  2 : ordering wp
↳ //                : names  x(3) y
↳ //                : weights 3 4
↳ //      block  3 : ordering C
  attrib(ra,"maxExp");
↳ 65535
  list lra = ringlist(ra);
  lra[1][4]=ideal(a2+1);
  def Ra = ring(lra);
  setring Ra; Ra;
↳ // coefficients: QQ[a]/(a^2+1)
↳ // number of vars : 4
↳ //      block  1 : ordering dp
↳ //                : names  x(1) x(2)
↳ //      block  2 : ordering wp
↳ //                : names  x(3) y
↳ //                : weights 3 4
↳ //      block  3 : ordering C

```

See Section 4.19.1 [qring], page 124; Section 4.19 [ring], page 124; Section 5.1.136 [ring_list], page 252.

5.1.136 ring_list

Syntax: ring_list (ring-expression)

Type: list

Purpose: decomposes a ring/qring into a list of 4 (or 6 in the non-commutative case, see Section 7.3.24 [ringlist (plural)], page 352) components. It is identical to ringlist with the exception of the first list entry.

1. the field description as **cring**
2. the names of the variables (a list L of strings: L[i] is the name of the i-th variable)
3. the monomial ordering (a list L of lists): each block L[i] consists of
 - the name of the ordering (string)
 - parameters specifying the ordering and the size of the block (intvec : typically the weights for the variables [default: 1])
4. the quotient ideal.

From a list of such structure, a new ring may be defined by the command ring (see the following example).

Note: All data which depends on a ring belong to the current ring, not to a ring which can be constructed from a modified list. These data will be mapped via fetch to the ring to be constructed.

Example:

```

ring r = 0, (x(1..3)), dp;
list l = ring_list(r);
l;
↳ [1]:
↳ QQ
↳ [2]:
↳ [1]:
↳ x(1)
↳ [2]:
↳ x(2)
↳ [3]:
↳ x(3)
↳ [3]:
↳ [1]:
↳ [1]:
↳ dp
↳ [2]:
↳ 1,1,1
↳ [2]:
↳ [1]:
↳ C
↳ [2]:
↳ 0
↳ [4]:
↳ _[1]=0
// Now change l and create a new ring, by
//- changing the base field to ZZ/32003
//- introducing one extra variable y,
//- defining the block ordering (dp(2),wp(3,4)).
//- define the minpoly after creating the function field
l[1]=ZZ/32003;
l[2][size(l[2])+1]="y";
l[3][3]=l[3][2]; // save the module ordering
l[3][1]=list("dp",intvec(1,1));
l[3][2]=list("wp",intvec(3,4));
def ra = ring(l); //creates the newring
ra; setring ra;
↳ // coefficients: ZZ/32003
↳ // number of vars : 4
↳ //      block 1 : ordering dp
↳ //      : names x(1) x(2)
↳ //      block 2 : ordering wp
↳ //      : names x(3) y
↳ //      : weights 3 4
↳ //      block 3 : ordering C

```

See Section 4.19.1 [qring], page 124; Section 4.19 [ring], page 124; Section 5.1.135 [ringlist], page 250.

5.1.137 rvar

Syntax: `rvar (name)`
`rvar (poly_expression)`
`rvar (string_expression)`

Type: `int`

Purpose: returns the number of the variable if the name/polynomial is a ring variable of the basering or if the string is the name of a ring variable of the basering; returns 0 if not. Hence the return value of `rvar` can also be used in a boolean context to check whether the variable exists.

Example:

```

ring r=29,(x,y,z),lp;
rvar(x);
↳ 1
rvar(r);
↳ 0
rvar(y);
↳ 2
rvar(var(3));
↳ 3
rvar("x");
↳ 1

```

See Section 5.1.18 [defined], page 167; Section 4.19 [ring], page 124; Section 5.1.163 [var], page 279; Section 5.1.165 [varstr], page 280.

5.1.138 `sba`

Syntax: `sba (ideal_expression)`
`sba (ideal_expression, int_expression, int_expression)`

Type: `ideal`

Purpose: returns a standard basis of an ideal with respect to the monomial ordering of the basering. A standard basis is a set of generators such that the leading terms generate the leading ideal, resp. module.

Use optional second and third arguments of type `int` to determine the respective variant of the signature-based standard basis algorithm:

The second argument specifies the internal module order `sba` uses:

- 0: induced Schreyer order on the signatures, non-incremental computation of the basis
- 1: position over term order, incremental computation of the basis
- 2: term over position order, non-incremental computation
- 3: Schreyer-weighted degree over index over leading term

The third argument specifies the rewrite order `sba` uses:

- 0: using the rewrite order described in <http://dx.doi.org/10.1016/j.jsc.2010.06.019>
- 1: using the rewrite order described in <http://dx.doi.org/10.1016/j.jsc.2011.05.004>

The standard call of `sba(i)` corresponds to `sba(i,0,1)`.

Note: The standard basis is computed with an optimized version of known signature-based algorithms like Faugere's F5 Algorithm. Whereas the correctness of the algorithms is only guaranteed for global orderings, timings for pure lexicographical orderings can be slow. In this situation you should try to compute the basis w.r.t. the graded reverse-lexicographic ordering and then convert to a basis for the lexicographical ordering using other methods (see Section 5.1.39 [fglm], page 181 and see Section D.4.9 [grwalk.lib], page 815). If the algorithms tend to use too much memory, you should try the other implemented standard basis algorithms (see Section 5.1.149 [std], page 266, see [groebner], page 785, and see Section 5.1.143 [slimgb], page 260). Note that the behaviour of `sba` on an example can be rather different depending on which variant you choose (second and third argument).

Example:

```
// incremental F5 computation
ring r=32003,(x,y,z),dp;
poly s1=1x2y+151xyz10+169y21;
poly s2=1xz14+6x2y4+3z24;
poly s3=5y10z10x+2y20z10+y10z20+11x3;
ideal i=s1,s2,s3;
ideal j=sba(i,1,0);
// non-incremental F5 computation
ring rhom=32003,(x,y,z,h),dp;
ideal i=homog(imap(r,i),h);
ideal j=sba(i,0,0);
// non-incremental signature-based computation
ring whom=32003,(x,y,z),dp;
ideal i=fetch(r,i);
ideal j=sba(i);
```

See Section 5.1.39 [fglm], page 181; [groebner], page 785; Section 4.5 [ideal], page 78; Section 4.19 [ring], page 124; Section 5.1.143 [slimgb], page 260; Section 5.1.149 [std], page 266.

5.1.139 setring

Syntax: `setring ring_name`

Type: none

Purpose: changes the basering to another (already defined) ring.

Example:

```
ring r1=0,(x,y),lp;
// the basering is r1
ring r2=32003,(a(1..8)),ds;
// the basering is r2
setring r1;
// the basering is again r1
nameof(basering);
↳ r1
  listvar();
↳ // r2                                [0] ring
↳ // r1                                [0] *ring
```

Use in procedures:

All changes of the basering by a definition of a new ring or a `setring` command in a procedure are local to this procedure. Use `keepring` to move a ring, which is local to a procedure, up by one nesting level.

See Section 5.2.11 [keepring], page 293; Section 4.19.1 [qring], page 124; Section 4.19 [ring], page 124.

5.1.140 simplex

Syntax: `simplex (matrix_expression, int_expression, int_expression, int_expression, int_expression, int_expression)`

Type: list

Purpose: perform the simplex algorithm for the tableau given by the input, e.g. `simplex (M, m, n, m1, m2, m3)`:

M matrix of numbers :

first row describing the objective function (maximize problem), the remaining rows describing constraints;

m, n, m1, m2, m3 int :

n = number of variables; m = total number of constraints; m1 = number of inequalities "`<=`" (rows 2 ... m1+1 of M); m2 = number of inequalities "`>=`" (rows m1+2 ... m1+m2+1 of M); m3 = number of equalities.

The following assumptions are made:

- * ground field is of type `(real,N)`, $N \geq 4$;
- * the matrix M is of size $m \times n$;
- * $m = m1 + m2 + m3$;
- * the entries $M[2,1], \dots, M[m+1,1]$ are non-negative;
- * the variables $x(i)$ are non-negative;
- * a row $b, a(1), \dots, a(n)$ corresponds to $b + a(1)x(1) + \dots + a(n)x(n)$;
- * for a `<=`, `>=`, or `==` constraint: add "in mind" `>=0`, `<=0`, or `==0`.

The output is a list L with

* L[1] = matrix

* L[2] = int:

0 = finite solution found; 1 = unbounded; -1 = no solution; -2 = error occurred;

* L[3] = intvec :

L[3][k] = number of variable which corresponds to row k+1 of L[1];

* L[4] = intvec :

L[4][j] = number of variable which is represented by column j+1 of L[1] ("non-basis variable");

* L[5] = int :

number of constraints (= m);

* L[6] = int :

number of variables (= n).

The solution can be read off the first column of L[1] as it is done by the procedure [simplexOut], page 878 in `solve.lib`.

Example:

```

ring r = (real,10),(x),lp;

// consider the max. problem:
//
//   maximize  x(1) + x(2) + 3*x(3) - 0.5*x(4)
//
// with constraints:  x(1) +          2*x(3)          <= 740
//                   2*x(2)          - 7*x(4) <= 0
//                   x(2) - x(3) + 2*x(4) >= 0.5
//                   x(1) + x(2) + x(3) + x(4) = 9
//
matrix sm[5][5]=( 0, 1, 1, 3,-0.5,
                  740,-1, 0,-2, 0,
                  0, 0,-2, 0, 7,
                  0.5, 0,-1, 1,-2,
                  9,-1,-1,-1,-1);

int n = 4; // number of constraints
int m = 4; // number of variables
int m1= 2; // number of <= constraints
int m2= 1; // number of >= constraints
int m3= 1; // number of == constraints
simplex(sm, n, m, m1, m2, m3);
↳ [1]:
↳   _[1,1]=17.025
↳   _[1,2]=-0.95
↳   _[1,3]=-0.05
↳   _[1,4]=1.95
↳   _[1,5]=-1.05
↳   _[2,1]=730.55
↳   _[2,2]=0.1
↳   _[2,3]=-0.1
↳   _[2,4]=-1.1
↳   _[2,5]=0.9
↳   _[3,1]=3.325
↳   _[3,2]=-0.35
↳   _[3,3]=-0.15
↳   _[3,4]=0.35
↳   _[3,5]=0.35
↳   _[4,1]=0.95
↳   _[4,2]=-0.1
↳   _[4,3]=0.1
↳   _[4,4]=0.1
↳   _[4,5]=0.1
↳   _[5,1]=4.725
↳   _[5,2]=-0.55
↳   _[5,3]=0.05
↳   _[5,4]=0.55
↳   _[5,5]=-0.45
↳ [2]:
↳ 0
↳ [3]:

```

```

↳      5,2,4,3
↳ [4]:
↳      1,6,8,7
↳ [5]:
↳      4
↳ [6]:
↳      4

```

See [simplexOut], page 878.

5.1.141 simplify

Syntax: `simplify (poly_expression , int_expression)`
`simplify (vector_expression , int_expression)`
`simplify (ideal_expression , int_expression)`
`simplify (module_expression , int_expression)`

Type: the type of the first argument

Purpose: returns the "simplified" first argument depending on the simplification rules specified by the second argument. The simplification rules are the following functions:

- | | |
|----|--|
| 1 | normalize (divide by leading coefficient if this is a unit of the ground field/ring). |
| 2 | erase zero generators/columns. |
| 4 | erase copies of earlier listed generators/columns. |
| 8 | erase generators/columns which a scalar multiples (w.r.t. ground field/ring) of earlier listed generators/columns. |
| 16 | erase generators/columns whose leading monomials are copies of leading monomials of earlier listed generators/columns such that the coefficients of both leading terms are units in the ground field/ring. |
| 32 | erase generators/columns whose leading terms are divisible by leading terms of other (not necessarily earlier) listed generators/columns. |
| 64 | normalize each coefficient of every monomial (of every polynomial) |

Example:

```

ring r=0,(x,y,z),(c,dp);
ideal i=0,2x,2x,4x,3x+y,5x2;
simplify(i,1);
↳ _[1]=0
↳ _[2]=x
↳ _[3]=x
↳ _[4]=x
↳ _[5]=x+1/3y
↳ _[6]=x2
simplify(i,2);
↳ _[1]=2x
↳ _[2]=2x
↳ _[3]=4x
↳ _[4]=3x+y
↳ _[5]=5x2
simplify(i,4);

```

```

↳ _[1]=0
↳ _[2]=2x
↳ _[3]=0
↳ _[4]=4x
↳ _[5]=3x+y
↳ _[6]=5x2
simplify(i,8);
↳ _[1]=0
↳ _[2]=2x
↳ _[3]=0
↳ _[4]=0
↳ _[5]=3x+y
↳ _[6]=5x2
simplify(i,16);
↳ _[1]=0
↳ _[2]=2x
↳ _[3]=0
↳ _[4]=0
↳ _[5]=0
↳ _[6]=5x2
simplify(i,32);
↳ _[1]=0
↳ _[2]=2x
↳ _[3]=0
↳ _[4]=0
↳ _[5]=0
↳ _[6]=0
simplify(i,32+2+1);
↳ _[1]=x
matrix A[2][3]=x,0,2x,y,0,2y;
simplify(A,2+8); // by automatic conversion to module
↳ _[1]=[x,y]

```

See Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

5.1.142 size

Syntax: `size (string-expression)`
 `size (bigint-expression)`
 `size (number-expression)`
 `size (intvec-expression)`
 `size (intmat-expression)`
 `size (poly-expression)`
 `size (vector-expression)`
 `size (ideal-expression)`
 `size (module-expression)`
 `size (matrix-expression)`
 `size (list-expression)`
 `size (resolution-expression)`
 `size (ring-expression)`

Type: `int`

Purpose: depends on the type of argument:

- ideal or module
returns the number of (non-zero) generators.
- string, intvec, list or resolution
returns the length, i.e., the number of characters, entries or elements.
- poly or vector
returns the number of monomials.
- matrix or intmat
returns the number of entries (rows*columns).
- ring
returns the number of elements in the ground field (for Z/p and algebraic extensions) or -1
- number or bigint
returns 0 for 0 or the number of words

Example:

```

string s="hello";
size(s);
↳ 5
intvec iv=1,2;
size(iv);
↳ 2
ring r=0,(x,y,z),lp;
poly f=x+y+z;
size(f);
↳ 3
vector v=[x+y,0,0,1];
size(v);
↳ 3
ideal i=f,y;
size(i);
↳ 2
module m=v,[0,1],[0,0,1],2*v;
size(m);
↳ 4
matrix mm[2][2];
size(mm);
↳ 4
ring r1=(2,a),x,dp;
minpoly=a4+a+1;
size(r1);
↳ 16

```

See Section 4.5 [ideal], page 78; Section 4.7 [intmat], page 88; Section 4.8 [intvec], page 91; Section 4.13 [module], page 110; Section 5.1.103 [ncols], page 227; Section 5.1.106 [nrows], page 228; Section 4.16 [poly], page 117; Section 4.21 [string], page 127; Section 4.22 [vector], page 131.

5.1.143 slimgb

Syntax:

```

slimgb ( ideal_expression )
slimgb ( module_expression )

```

Type: ideal or module

Purpose: Section A.2.3 [slim Groebner bases], page 706

Returns a Groebner basis of an ideal or module with respect to the monomial ordering of the basering (which has to be global).

Note: The algorithm is designed to keep polynomials slim (short with small coefficients). For details see https://www.singular.uni-kl.de/reports/35/paper_35_full.ps.gz. A reduced Groebner basis is returned if option(redSB) is set (see [option(redSB)], page 231). To view the progress of long running computations, use option(prot) (see [option(prot)], page 231).

Warning: Groebner basis computations with inexact coefficients can not be trusted due to rounding errors.

Example:

```
ring r=2,(x,y,z),lp;
poly s1=z*(x*y+1);
poly s2=x2+x;
poly s3=y2+y;
ideal i=s1,s2,s3;
slingb(i);
↳ _[1]=y2+y
↳ _[2]=x2+x
↳ _[3]=yz+z
↳ _[4]=xz+z
```

See [groebner], page 785; Section 4.5 [ideal], page 78; Section 5.1.110 [option], page 230; Section 4.19 [ring], page 124; Section 5.1.149 [std], page 266.

5.1.144 sortvec

Syntax: sortvec (ideal_expression)
sortvec (module_expression)

Type: intvec

Purpose: computes the permutation v which orders the ideal, resp. module, I by its initial terms, starting with the smallest, that is, $I(v[i]) < I(v[i+1])$ for all i .

Example:

```
ring r=0,(x,y,z),dp;
ideal I=y,z,x,x3,xz;
sortvec(I);
↳ 2,1,3,5,4
```

See Section D.2.3 [general_lib], page 790.

5.1.145 sqrfree

Syntax: sqrfree (poly_expression)
sqrfree (poly_expression, 0)
sqrfree (poly_expression, 2)

Type: list of ideal and intvec

Syntax: `sqrfree (poly_expression, 1)`

Type: ideal

Syntax: `sqrfree (poly_expression, 3)`

Type: poly

Purpose: computes the squarefree factors (as an ideal) of the polynomial together with or without the multiplicities (as an intvec) depending on the second argument:

0: returns factors and multiplicities, first factor is a constant.

May also be written with only one argument.

1: returns non-constant factors (no multiplicities).

2: returns non-constant factors and multiplicities.

3: returns the product of non-constant factors, i.e. squarefree part

Note: Not implemented for the coefficient fields real and finite fields of type (p^n, a) .

Example:

```

ring r=3,(x,y,z),dp;
poly f=(x-y)^3*(x+z)*(y-z);
sqrfree(f);
↳ [1]:
↳  _[1]=1
↳  _[2]=-xy+xz-yz+z2
↳  _[3]=-x+y
↳ [2]:
↳  1,1,3
sqrfree(f,1);
↳ _[1]=-xy+xz-yz+z2
↳ _[2]=-x+y
sqrfree(f,2);
↳ [1]:
↳  _[1]=-xy+xz-yz+z2
↳  _[2]=-x+y
↳ [2]:
↳  1,3
sqrfree(f,3);
↳ x2y-xy2-x2z-xyz-y2z-xz2+yz2

```

See Section 5.1.36 [factorize], page 178.

5.1.146 sprintf

Procedure from library `standard.lib` (see Section D.1 [standard.lib], page 785).

Syntax: `sprintf (string_expression [, any_expressions])`

Return: string

Purpose: `sprintf(fmt,...)`; performs output formatting. The first argument is a format control string. Additional arguments may be required, depending on the content of the control string. A series of output characters is generated as directed by the control string; these characters are returned as a string.

The control string `fmt` is simply text to be copied, except that the string may contain conversion specifications.

Type `help print`; for a listing of valid conversion specifications. As an addition to the conversions of `print`, the `%n` and `%2` conversion specification does not consume an additional argument, but simply generates a newline character.

Note: If one of the additional arguments is a list, then it should be wrapped in an additional `list()` command, since passing a list as an argument flattens the list by one level.

Example:

```

ring r=0,(x,y,z),dp;
module m=[1,y],[0,x+z];
intmat M=betti(mres(m,0));
list l = r, m, M;
string s = sprintf("s:%s,%n l:%l", 1, 2); s;
↳ s:1,
↳ l:int(2)
s = sprintf("s:%n%s", 1); s;
↳ s:
↳ (QQ),(x,y,z),(dp(3),C)
s = sprintf("s:%2%s", list(1)); s;
↳ s:
↳ (QQ),(x,y,z),(dp(3),C),y*gen(2)+gen(1),x*gen(2)+z*gen(2),1,1
s = sprintf("2l:%n%2l", list(1)); s;
↳ 2l:
↳ list("(QQ),(x,y,z),(dp(3),C)",
↳ module(y*gen(2)+gen(1),
↳ x*gen(2)+z*gen(2)),
↳ intmat(intvec(1,1),1,2))
↳
s = sprintf("%p", list(1)); s;
↳ [1]:
↳ // coefficients: QQ
↳ // number of vars : 3
↳ //          block  1 : ordering dp
↳ //          : names  x y z
↳ //          block  2 : ordering C
↳ [2]:
↳   _[1]=y*gen(2)+gen(1)
↳   _[2]=x*gen(2)+z*gen(2)
↳ [3]:
↳   1,1
s = sprintf("%;", list(1)); s;
↳ [1]:
↳ // coefficients: QQ
↳ // number of vars : 3
↳ //          block  1 : ordering dp
↳ //          : names  x y z
↳ //          block  2 : ordering C
↳ [2]:
↳   _[1]=y*gen(2)+gen(1)
↳   _[2]=x*gen(2)+z*gen(2)
↳ [3]:
↳   1,1
↳
s = sprintf("%b", M); s;

```

```

↳          0      1
↳ -----
↳      0:      1      1
↳ -----
↳ total:      1      1
↳

```

See also: [fprintf], page 785; Section 5.1.119 [print], page 238; [printf], page 785; Section 4.21 [string], page 127.

5.1.147 sres

Syntax: `sres (ideal_expression, int_expression)`
`sres (module_expression, int_expression)`

Type: resolution

Purpose: computes a free resolution of an ideal or module with Schreyer's method. The ideal, resp. module, has to be a standard basis. More precisely, let M be given by a standard basis and $A_1 = \text{matrix}(M)$. Then `sres` computes a free resolution of $\text{coker}(A_1) = F_0/M$

$$\dots \longrightarrow F_2 \xrightarrow{A_2} F_1 \xrightarrow{A_1} F_0 \longrightarrow F_0/M \longrightarrow 0.$$

If the int expression k is not zero then the computation stops after k steps and returns a list of modules (given by standard bases) $M_i = \text{module}(A_i)$, $i=1..k$.

`sres(M,0)` returns a list of n modules where n is the number of variables of the basering.

Even if `sres` does not compute a minimal resolution, the `betti` command gives the true betti numbers! In many cases of interest `sres` is much faster than any other known method. Let `list L=sres(M,0)`; then `L[1]=M` is identical to the input, `L[2]` is a standard basis with respect to the Schreyer ordering of the first syzygy module of `L[1]`, etc. (`L[i] = M_i` in the notations from above.)

Note: Accessing single elements of a resolution may require some partial computations to be finished and may therefore take some time.

Example:

```

ring r=31991,(t,x,y,z,w),ls;
ideal M=t2x2+tx2y+x2yz,t2y2+ty2z+y2zw,
      t2z2+tz2w+xz2w,t2w2+txw2+xyw2;
M=std(M);
resolution L=sres(M,0);
L;
↳ 1      35      141      209      141      43      4
↳ r <--  r <--  r <--  r <--  r <--  r <--  r
↳
↳ 0      1      2      3      4      5      6
↳ resolution not minimized yet
↳
print(betti(L),"betti");
↳          0      1      2      3      4      5
↳ -----
↳      0:      1      -      -      -      -      -
↳      1:      -      -      -      -      -      -
↳      2:      -      -      -      -      -      -
↳      3:      -      4      -      -      -      -

```

```

↳      4:      -      -      -      -      -      -
↳      5:      -      -      -      -      -      -
↳      6:      -      -      6      -      -      -
↳      7:      -      -      9      16     2      -
↳      8:      -      -      -      2      5      1
↳ -----
↳ total:      1      4      15     18     7      1
↳

```

See Section 5.1.4 [betti], page 157; Section 5.1.48 [fres], page 186; Section 5.1.58 [hres], page 195; Section 4.5 [ideal], page 78; Section 4.6 [int], page 82; Section 5.1.83 [lres], page 212; Section 5.1.93 [minres], page 220; Section 4.13 [module], page 110; Section 5.1.98 [mres], page 222; [res], page 785; Section 5.1.154 [syz], page 275.

5.1.148 status

Syntax: status (link_expression , string_expression)

Type: string

Syntax: status (link_expression , string_expression , string_expression)

Type: int

Purpose: returns the status of the link as asked for by the second argument. If a third argument is given, the result of the comparison to the status string is returned: (status(l,s1)==s2) is equivalent to status(l,s1,s2).

The following string expressions are allowed:

"name" the name string given by the definition of the link (usually the filename)

"type" returns "ASCII", "DBM" or "ssi"

"open" returns "yes" or "no"

"openread" returns "yes" or "no"

"openwrite" returns "yes" or "no"

"read" returns "ready" or "not ready"

"write" returns "ready" or "not ready"

"mode" returns (depending on the type of the link and its status) "", "w", "a", "r" or "rw"

"exists" returns "yes" or "no": existence of the filename for ASCII/ssi links

Syntax: status (list_expression , int_expression)

Type: int

Purpose: the list should be a list L of links, the second argument a timeout in 1/10 seconds. Returns

-2 select returns an error

-1 all links are closed/at eof

0 timeout

>0 (at least) $L[i]$ is ready

Example:

```

link l=":w example.txt";
status(l,"write");
↳ not ready
open(l);
status(l,"write","ready");
↳ 1
close(l);

```

See Section 4.9 [link], page 94; Section 5.1.109 [open], page 230; Section 5.1.128 [read], page 245; Section 5.1.172 [write], page 284.

5.1.149 std

Syntax: `std (ideal_expression)`
`std (module_expression)`
`std (smatrix_expression)`
`std (ideal_expression, intvec_expression)`
`std (module_expression, intvec_expression)`
`std (ideal_expression, intvec_expression, intvec_expression)`
`std (module_expression, intvec_expression, intvec_expression)`
`std (ideal_expression, poly_expression)`
`std (module_expression, vector_expression)`
`std (ideal_expression, ideal_expression)`
`std (module_expression, module_expression)`
`std (ideal_expression, poly_expression, intvec_expression, intvec_expression)`
`std (module_expression, poly_expression, intvec_expression, intvec_expression)`

Type: ideal, module or smatrix

Purpose: returns a standard basis of an ideal or module with respect to the monomial ordering of the basering. For Letterplace rings, a twosided Groebner basis is computed. A standard basis is a set of generators such that the leading terms generate the leading ideal, resp. module.

Use an optional second argument of type intvec as Hilbert series (result of `hilb(i,1)`, see Section 5.1.56 [hilb], page 193), if the ideal, resp. module, is homogeneous (Hilbert driven standard basis computation, [stdhilb], page 785). If the ideal is quasihomogeneous with some weights w and if the Hilbert series is computed w.r.t. to these weights, then use w as third argument.

Use an optional second argument of type poly/vector/ideal, resp. module, to construct the standard basis from an already computed one (given as the first argument) and additional generator(s) (the second argument).

4 arguments G, p, hv, w are the combination of the above: standard basis G , additional generator p , hilbert function hv w.r.t. weights w .

Warning: Groebner basis computations with inexact coefficients can not be trusted due to rounding errors.

Note: The standard basis is computed with a (more or less) straight-forward implementation of the classical Buchberger (resp. Mora) algorithm. For global orderings, use the `groebner` command instead (see [groebner], page 785), which heuristically chooses the "best" algorithm to compute a Groebner basis.

To view the progress of long running computations, use `option(prot)` (see `[option(prot)]`, page 231).

Example:

```

// local computation
ring r=32003,(x,y,z),ds;
poly s1=1x2y+151xyz10+169y21;
poly s2=1xz14+6x2y4+3z24;
poly s3=5y10z10x+2y20z10+y10z20+11x3;
ideal i=s1,s2,s3;
ideal j=std(i);
degree(j);
↳ // dimension (local) = 0
↳ // multiplicity = 1512
// Hilbert driven elimination (standard)
ring rhom=32003,(x,y,z,h),dp;
ideal i=homog(imap(r,i),h);
ideal j=std(i);
intvec iv=hilb(j,1);
ring rlex=32003,(x,y,z,h),lp;
ideal i=fetch(rhom,i);
ideal j=std(i,iv);
j=subst(j,h,1);
j[1];
↳ z64
// Hilbert driven elimination (ideal is quasihomogeneous)
intvec w=10,1,1;
ring whom=32003,(x,y,z),wp(w);
ideal i=fetch(r,i);
ideal j=std(i);
intvec iw=hilb(j,1,w);
ring wlex=32003,(x,y,z),lp;
ideal i=fetch(whom,i);
ideal j=std(i,iw,w);
j[1];
↳ z64

```

See Section 5.1.34 `[facstd]`, page 177; Section 5.1.39 `[fglm]`, page 181; `[groebner]`, page 785; Section 4.5 `[ideal]`, page 78; Section 4.13 `[module]`, page 110; Section 5.1.99 `[mstd]`, page 223; Section 5.1.110 `[option]`, page 230; Section 4.19 `[ring]`, page 124; Section 4.20 `[smatrix]`, page 127; `[stdfglm]`, page 785; `[stdhilb]`, page 785.

5.1.150 `stdfglm`

Procedure from library `standard.lib` (see Section D.1 `[standard.lib]`, page 785).

Syntax: `stdfglm (ideal-expression)`
`stdfglm (ideal-expression, string-expression)`

Type: ideal

Purpose: computes the standard basis of the ideal in the basering via `fglm` from the ordering given as the second argument to the ordering of the basering. If no second argument is given, "dp" is used. The standard basis for the given ordering (resp. for "dp") is computed via the command `groebner` except if a further argument "std" or "slimgb" is given in which case `std` resp. `slimgb` is used.

Example:

```

ring r = 0,(x,y,z),lp;
ideal i = y3+x2,x2y+x2,x3-x2,z4-x2-y;
stdfglm(i); //uses fglm from "dp" (with groebner) to "lp"
↳ _[1]=z12
↳ _[2]=yz4-z8
↳ _[3]=y2+y-z8-z4
↳ _[4]=xy-xz4-y+z4
↳ _[5]=x2+y-z4
stdfglm(i,"std"); //uses fglm from "dp" (with std) to "lp"
↳ _[1]=z12
↳ _[2]=yz4-z8
↳ _[3]=y2+y-z8-z4
↳ _[4]=xy-xz4-y+z4
↳ _[5]=x2+y-z4
ring s = (0,x),(y,z,u,v),lp;
minpoly = x2+1;
ideal i = u5-v4,zv-u2,zu3-v3,z2u-v2,z3-uv,yv-zu,yu-z2,yz-v,y2-u,u-xy2;
weight(i);
↳ 2,3,4,5
stdfglm(i,"(a(2,3,4,5),dp)"); //uses fglm from "(a(2,3,4,5),dp)" to "lp"
↳ _[1]=v2
↳ _[2]=u
↳ _[3]=zv
↳ _[4]=z2
↳ _[5]=yv
↳ _[6]=yz-v
↳ _[7]=y2

```

See also: Section 5.1.39 [fglm], page 181; [groebner], page 785; Section 5.1.143 [slimgb], page 260; Section 5.1.149 [std], page 266; [stdhilb], page 785.

5.1.151 stdhilb

Procedure from library `standard.lib` (see Section D.1 [standard.lib], page 785).

Syntax: `stdhilb (ideal_expression)`
`stdhilb (module_expression)`
`stdhilb (ideal_expression, intvec_expression)`
`stdhilb (module_expression, intvec_expression)`
`stdhilb (ideal_expression, list of string-expressions, and intvec_expression)`

Type: type of the first argument

Purpose: Compute a Groebner basis of the ideal/module in the basering by using the Hilbert driven Groebner basis algorithm. If an argument of type string, stating "`std`" resp. "`slimgb`", is given, the standard basis computation uses `std` or `slimgb`, otherwise a heuristically chosen method (default)
If an optional second argument `w` of type `intvec` is given, `w` is used as variable weights. If `w` is not given, it is computed as $w[i] = \deg(\text{var}(i))$. If the ideal is homogeneous w.r.t. `w` then the Hilbert series is computed w.r.t. to these weights.

Theory: If the ideal is not homogeneous compute first a Groebner basis of the homogenization [w.r.t. the weights `w`] of the ideal/module, then the Hilbert function and, finally, a

Groebner basis in the original ring by using the computed Hilbert function. If the given w does not coincide with the variable weights of the basering, the result may not be a groebner basis in the original ring.

Note: 'Homogeneous' means weighted homogeneous with respect to the weights $w[i]$ of the variables $\text{var}(i)$ of the basering. Parameters are not converted to variables.

Example:

```

ring r = 0, (x,y,z), lp;
ideal i = y3+x2,x2y+x2z2,x3-z9,z4-y2-xz;
ideal j = stdhilb(i); j;
↳ j[1]=z10
↳ j[2]=yz9
↳ j[3]=2y2z4-z8
↳ j[4]=2y3z3-2y2z5-yz7
↳ j[5]=y4+y3z2
↳ j[6]=xz+y2-z4
↳ j[7]=xy2-xz4-y3z
↳ j[8]=x2+y3
ring r1 = 0, (x,y,z), wp(3,2,1);
ideal i = y3+x2,x2y+x2z2,x3-z9,z4-y2-xz; //ideal is homogeneous
ideal j = stdhilb(i,"std"); j;
↳ j[1]=y2+xz-z4
↳ j[2]=x2-xyz+yz4
↳ j[3]=2xz5-z8
↳ j[4]=2xyz4-yz7+z9
↳ j[5]=z10
↳ j[6]=2yz9+z11
//this is equivalent to:
intvec v = hilb(std(i),1);
ideal j1 = std(i,v,intvec(3,2,1)); j1;
↳ j1[1]=y2+xz-z4
↳ j1[2]=x2-xyz+yz4
↳ j1[3]=2xz5-z8
↳ j1[4]=2xyz4-yz7+z9
↳ j1[5]=z10
↳ j1[6]=yz9
size(NF(j,j1))+size(NF(j1,j)); //j and j1 define the same ideal
↳ 0

```

See also: [groebner], page 785; Section 5.1.143 [slimgb], page 260; Section 5.1.149 [std], page 266; [stdfglm], page 785.

5.1.152 subst

Syntax: `subst (poly_expression, variable, poly_expression)`
`subst (poly_expression, variable, poly_expression , ... variable, poly_expression)`
`subst (vector_expression, variable, poly_expression)`
`subst (ideal_expression, variable, poly_expression)`
`subst (module_expression, variable, poly_expression)`

Type: poly, vector, ideal or module (corresponding to the first argument)

Purpose: substitutes one or more ring variable(s)/parameter variable(s) by (a) polynomial(s). Note that in the case of more than one substitution pair, the substitutions will be

performed sequentially and not simultaneously. The below examples illustrate this behaviour.

Note, that the coefficients must be polynomial when substituting a parameter.

Example:

```
ring r=0,(x,y,z),dp;
poly f=x2+y2+z2+x+y+z;
subst(f,x,y,y,z); // first substitute x by y, then y by z
↳ 3z2+3z
subst(f,y,z,x,y); // first substitute y by z, then x by y
↳ y2+2z2+y+2z
```

See Section 4.5 [ideal], page 78; Section 4.11 [map], page 103; Section 4.13 [module], page 110; Section 4.16 [poly], page 117; [substitute], page 799; Section 4.22 [vector], page 131.

5.1.153 system

Syntax: `system (string-expression)`
`system (string-expression, expression)`

Type: depends on the desired function, may be none

Purpose: interface to internal data and the operating system. The string-expression determines the command to execute. Some commands require an additional argument (second form) where the type of the argument depends on the command. See below for a list of all possible commands.

Note: Not all functions work on every platform.

Functions:

```
system("alarm", int )
    abort the Singular process after computing for that many seconds (system+user cpu time).
```

```
system("absFact", poly )
    absolute factorization of the polynomial (from a polynomial ring over a transcendental extension) Returns a list of the ideal of the factors, intvec of multiplicities, ideal of minimal polynomials and the number of factors.
```

```
system("blackbox")
    list all blackbox data types.
```

```
system("browsers");
    returns a string about available help browsers. See Section 3.1.3 [The online help system], page 15.
```

```
system("bracket", poly, poly )
    returns the Lie bracket [p,q].
```

```
system("complexNearZero", number-expression )
    checks for a small value for floating point numbers
```

```
system("contributors")
    returns names of people who contributed to the SINGULAR kernel as string.
```

```
system("cpu")
    returns the number of cpus as int (for creating multiple threads/processes).
    (see system("--cpus")).
```


`system("denom_list")`
 returns the list of denominators (number) which occurred in the latest std computation(s). Is reset to the empty list at ring changes or by this system call.

`system("eigenvals", matrix)`
 returns the list of the eigenvalues of the matrix (as ideal, intvec). (see `system("hessenberg")`).

`system("env", ring)`
 returns the enveloping algebra (i.e. $R \text{ tensor } R^{\text{opp}}$) See `system("opp")`.

`system("executable", string)`
 returns the path of the command given as argument or the empty string (for: not found) See `system("Singular")`. See `system("getenv", "PATH")`.

`system("getenv", string-expression)`
 returns the value of the shell environment variable given as the second argument. The return type is string.

`system("getPrecDigits")`
 returns the precision for floating point numbers

`system("gmsnf", ideal, ideal, matrix,int, int)`
 Gauss-Manin system: for `gmspoly.lib`, `gmssing.lib`

`system("HC")`
 returns the degree of the "highest corner" from the last std computation (or 0).

`system("hessenberg", matrix)`
 returns the Hessenberg matrix (via QR algorithm).

`system("install", s1, s2, p3, i4)`
 install a new method p3 for s2 for the newstruct type s1. s2 must be a reserved operator with i4 operands (i4 may be 1,2,3; use 4 for more than 3 or a varying number of arguments) See Section 4.23.4 [Commands for user defined types], page 135.

`system("LLL", B)`
 B must be a matrix or an intmat. Interface to NTLs LLL (Exact Arithmetic Variant over ZZ). Returns the same type as the input.
 B is an $m \times n$ matrix, viewed as m rows of n -vectors. m may be less than, equal to, or greater than n , and the rows need not be linearly independent. B is transformed into an LLL-reduced basis. The first $m - \text{rank}(B)$ rows of B are zero.
 More specifically, elementary row transformations are performed on B so that the non-zero rows of new-B form an LLL-reduced basis for the lattice spanned by the rows of old-B.

`system("nblocks")` or `system("nblocks", ring_name)`
 returns the number of blocks of the given ring, or of the current basering, if no second argument is given. The return type is int.

`system("nc_hilb", ideal, int, [...])`
 internal support for `ncHilb.lib`, return nothing

`system("neworder", ideal)`
 string of the ring variables in an heurically good order for `char_series`

`system("newstruct")`
 list all newstruct data types.

`system("opp", ring)`
 returns the opposite ring.

`system("oppose", ring R, poly p)`
 returns the opposite polynomial of p from R.

`system("pcvLAddL", list, list)`
`system("pcvPMuL", poly, list)`
`system("pcvMinDeg", poly)`
`system("pcvP2CV", list, int, int)`
`system("pcvCV2P", list, int, int)`
`system("pcvDim", int, int)`
`system("pcvBasis", int, int)` internal for `mondromy.lib`

`system("pid")`
 returns the process number as int (for creating unique names).

`system("random")` or `system("random", int)`
 returns or sets the seed of the random generator.

`system("reduce_bound", poly, ideal, int)`
 or `system("reduce_bound", ideal, ideal, int)`
 or `system("reduce_bound", vector, module, int)`
 or `system("reduce_bound", module, module, int)` returns the normal-form of the first argument wrt. the second up to the given degree bound (wrt. total degree)

`system("reserve", int)`
 reserve a port and listen with the given backlog. (see `system("reservedLink")`).

`system("reservedLink")`
 accept a connect at the reserved port and return a (write-only) link to it. (see `system("reserve")`).

`system("semaphore", string, int)`
 operations for semaphores: string may be "init", "exists", "acquire", "try_acquire", "release", "get_value", and int is the number of the semaphore. Returns -2 for wrong command, -1 for error or the result of the command.

`system("semic", list, list)`
 or `system("semic", list, list, int)` computes from list of spectrum numbers and list of spectrum numbers the semicontinuity index (qh, if 3rd argument is 1).

`system("setenv", string_expression, string_expression)`
 sets the shell environment variable given as the second argument to the value given as the third argument. Returns the third argument. Might not be available on all platforms.

`system("sh", string_expression)`
 shell escape, returns the return code of the shell as int. The string is sent literally to the shell.

`system("shrinktest", poly, i2)`
 internal for shift algebra (with i2 variables): shrink the poly

`system("Singular")`
 returns the absolute (path) name of the running SINGULAR as string.

`system("SingularLib")`
 returns the colon separated library search path name as string.

`system("spadd", list, list)`
 or `system("spadd", list, list, int)` computes from list of spectrum numbers and list of spectrum numbers the sum of the lists.

`system("spectrum", poly)`
 or `system("spectrum", poly, int)`

`system("spmul", list, int)`
 or `system("spmul", list, list, int)` computes from list of spectrum numbers the multiple of it.

`system("std_syz", module, int)`
 compute a partial groebner base of a module, stop after the given column

`system("tensorModuleMult", int, module)`
 internal for sheafcoh.lib (see `id_TensorModuleMult`)

`system("twostd", ideal)`
 returns the two-sided standard basis of the two-sided ideal.

`system("uname")`
 returns a string identifying the architecture for which SINGULAR was compiled.

`system("verifyGB", ideal_expression/module_expression)`
 checks, if an ideal/module is a Groebner base

`system("version")`
 returns the version number of SINGULAR as int. (Version a-b-c-d returns $a*10000+b*1000+c*100+d$)

`system("with")`
 without an argument: returns a string describing the current version of SINGULAR, its build options, the used path names and other configurations
 with a string argument: test for that feature and return an int.

`system("--cpus")`
 returns the number of available cpu cores as int (for using multiple cores). (see `system("cpu")`).

`system("-")`
 prints the values of all options.

`system("-long_option_name")`
 returns the value of the (command-line) option `long_option_name`. The type of the returned value is either string or int. See Section 3.1.6 [Command line options], page 19, for more info.

`system("-long_option_name", expression)`
 sets the value of the (command-line) option `long_option_name` to the value given by the expression. Type of the expression must be string, or int. See Section 3.1.6 [Command line options], page 19, for more info. Among others, this can be used for setting the seed of the random number generator, the used help browser, the minimal display time, or the timer resolution.

Example:

```
// a listing of the current directory:
system("sh","ls");
// execute a shell, return to SINGULAR with exit:
system("sh","sh");
string unique_name="/tmp/xx"+string(system("pid"));
unique_name;
↳ /tmp/xx4711
system("uname")
↳ ix86-Linux
system("getenv","PATH");
↳ /bin:/usr/bin:/usr/local/bin
system("Singular");
↳ /usr/local/bin/Singular
// report value of all options
system("--");
↳ // --batch          0
↳ // --execute
↳ // --sdb            0
↳ // --echo           1
↳ // --profile        0
↳ // --quiet          1
↳ // --sort           0
↳ // --random         12345678
↳ // --no-tty         1
↳ // --user-option
↳ // --allow-net      0
↳ // --browser
↳ // --cntrlc
↳ // --emacs          0
↳ // --no-stdlib      0
↳ // --no-rc          1
↳ // --no-warn        0
↳ // --no-out         0
↳ // --no-shell       0
↳ // --min-time       "0.5"
↳ // --cpus           4
↳ // --threads        4
↳ // --flint-threads  1
↳ // --MPport
↳ // --MPhost
↳ // --link
↳ // --ticks-per-sec  1
// set minimal display time to 0.02 seconds
system("--min-time", "0.02");
// set timer resolution to 0.01 seconds
```

```

system("--ticks-per-sec", 100);
// re-seed random number generator
system("--random", 12345678);
// allow your web browser to access HTML pages from the net
system("--allow-net", 1);
// and set help browser to firefox
system("--browser", "firefox");
↳ // ** Could not get 'DataDir'.
↳ // ** Either set environment variable 'SINGULAR_DATA_DIR' to 'DataDir',
↳ // ** or make sure that 'DataDir' is at "/home/hannes/test/Singular/./s
  are/"
↳ // ** Could not get 'IdxFile'.
↳ // ** Either set environment variable 'SINGULAR_IDX_FILE' to 'IdxFile',
↳ // ** Could not get 'DataDir'.
↳ // ** Either set environment variable 'SINGULAR_DATA_DIR' to 'DataDir',
↳ // ** or make sure that 'DataDir' is at "/home/hannes/test/Singular/./s
  are/"
↳ // ** or make sure that 'IdxFile' is at "%D/singular/singular.idx"
↳ // ** resource 'x' not found
↳ // ** Setting help browser to 'dummy'.

```

5.1.154 syz

Syntax: syz (ideal_expression)
 syz (module_expression)
 syz (ideal_expression, string_expression)
 syz (module_expression, string_expression)

Type: module

Purpose: computes the first syzygy (i.e., the module of relations of the given generators) of the ideal, resp. module.

An optional second argument specifies the Groebner base algorithm to use. Possible values are "std"(default) and "slimgb".

Only for use of "std": If option(returnSB) is set, a standard basis is returned, otherwise a generating set.

Example:

```

ring R=0,(x,y),(c,dp);
ideal i=x,y;
syz(i);
↳ _[1]=[y,-x]

```

See Section 5.1.48 [fres], page 186; Section 5.1.58 [hres], page 195; Section 4.5 [ideal], page 78; Section 5.1.80 [lift], page 209; Section 5.1.81 [liftstd], page 209; Section 5.1.83 [ires], page 212; Section 4.13 [module], page 110; Section 5.1.98 [mres], page 222; Section D.4.21 [nfmodsyz.lib], page 823; Section 5.1.105 [nres], page 228; Section 5.1.110 [option], page 230; [res], page 785; Section 5.1.147 [sres], page 264.

5.1.155 tensor

Syntax: tensor (matrix_expression , matrix_expression)
 tensor (module_expression , module_expression)
 tensor (smatrix_expression , smatrix_expression)

Type: same as the first argument

Purpose: computes the tensor product (Kronecker product) of A and B

Example:

```

ring r=32003,(x,y,z),(c,ds);
matrix A[3][3]=1,2,3,4,5,6,7,8,9;
matrix B[2][2]=x,y,0,z;
print(A);
↳ 1,2,3,
↳ 4,5,6,
↳ 7,8,9
print(B);
↳ x,y,
↳ 0,z
print(tensor(A,B));
↳ x, y, 2x,2y,3x,3y,
↳ 0, z, 0, 2z,0, 3z,
↳ 4x,4y,5x,5y,6x,6y,
↳ 0, 4z,0, 5z,0, 6z,
↳ 7x,7y,8x,8y,9x,9y,
↳ 0, 7z,0, 8z,0, 9z

```

See Section 4.12 [matrix], page 106; Section 4.13 [module], page 110.

5.1.156 trace

Syntax: trace (intmat_expression)
 trace (matrix_expression)

Type: int, if the argument is an intmat, resp.
 poly, if the argument is a matrix

Purpose: returns the trace of an intmat, resp. matrix.

Example:

```

intmat m[2][2]=1,2,3,4;
print(m);
↳      1      2
↳      3      4
trace(m);
↳ 5

```

See Section 4.7 [intmat], page 88; Section 4.12 [matrix], page 106.

5.1.157 transpose

Syntax: transpose (intmat_expression)
 transpose (matrix_expression)
 transpose (smatrix_expression)
 transpose (module_expression)

Type: intmat, matrix, or module, corresponding to the argument

Purpose: transposes a matrix.

Example:

```

ring R=0,x,dp;
matrix m[2][3]=1,2,3,4,5,6;
print(m);
↳ 1,2,3,
↳ 4,5,6
print(transpose(m));
↳ 1,4,
↳ 2,5,
↳ 3,6

```

See Section 4.7 [intmat], page 88; Section 4.12 [matrix], page 106; Section 4.13 [module], page 110; Section 4.20 [smatrix], page 127.

5.1.158 type

Syntax: type name ;
type (name) ;

Type: none

Purpose: prints the name, level, type and value of a variable. To display the value of an expression, it is sufficient to type the expression followed by ;.

Example:

```

int i=3;
i;
↳ 3
type(i);
↳ // i int 3

```

See Chapter 4 [Data types], page 72; Section 5.1.82 [listvar], page 210; Section 5.1.119 [print], page 238.

5.1.159 typeof

Syntax: typeof (expression)

Type: string

Purpose: returns the type of an expression as string.

Returns the type of the first list element if the expression is an expression list.

Possible types are: "ideal", "int", "intmat", "intvec", "list", "map", "matrix", "module", "number", "none", "poly", "proc", "qring", "resolution", "ring", "string", "vector".

For internal use only is the type "?unknown type?".

Example:

```

int i=9; i;
↳ 9
typeof(_);
↳ int
print(i);
↳ 9
typeof(_);
↳ string

```

```

    type i;
    ↪ // i int 9
    typeof(_);
    ↪ string
    string s=typeof(i);
    s;
    ↪ int
    typeof(s);
    ↪ string
    proc p() { "hello"; return();}
    p();
    ↪ hello
    typeof(_);
    ↪ ?unknown type?

```

See Chapter 4 [Data types], page 72; Section 5.1.158 [type], page 277.

5.1.160 univariate

Syntax: univariate (poly_expression)

Type: int

Purpose: returns 0 for not univariate, -1 for a constant or the number of the variable of the univariate polynomial.

Example:

```

    ring r=0,(x,y,z),dp;
    univariate(x2+1);
    ↪ 1
    univariate(x2+y+1);
    ↪ 0
    univariate(1);
    ↪ -1
    univariate(var(2));
    ↪ 2
    var(univariate(z));
    ↪ z

```

See Section 5.1.77 [leadexp], page 207; Section 5.1.163 [var], page 279.

5.1.161 uresolve

Syntax: uresolve (ideal_expression, int_expression, int_expression, int_expression)

Type: list

Purpose: computes all complex roots of a zerodimensional ideal. Makes either use of the multipolynomial resultant of Macaulay (second argument = 1), which works only for homogeneous ideals, or uses the sparse resultant of Gelfand, Kapranov and Zelevinsky (second argument = 0). The sparse resultant algorithm uses a mixed polyhedral subdivision of the Minkowski sum of the Newton polytopes in order to construct the sparse resultant matrix. Its determinant is a nonzero multiple of the sparse resultant. The u-resultant of B.\ L. van der Waerden and Laguerre's algorithm are used to determine the complex roots.

The third argument defines the precision of the fractional part if the ground field is the field of rational numbers, otherwise it will be ignored.

The fourth argument (can be 0, 1 or 2) gives the number of extra runs of Laguerre's algorithm (with corrupted roots), leading to better results.

Note: If the ground field is the field of complex numbers, the elements of the list are of type number, otherwise of type string.

See Section 5.1.74 [laguerre], page 205; Section 5.1.97 [mpresmat], page 222.

5.1.162 vandermonde

Syntax: `vandermonde (ideal_expression, ideal_expression, int_expression)`

Type: poly

Purpose: `vandermonde(p,v,d)` computes the (unique) polynomial of degree `d` with prescribed values `v[1], ..., v[N]` at the points `p0, ..., pN-1`, `N=(d+1)n`, `n` the number of ring variables.

The returned polynomial is $\sum c_{\alpha_1 \dots \alpha_n} \cdot x_1^{\alpha_1} \cdot \dots \cdot x_n^{\alpha_n}$, where the coefficients $c_{\alpha_1 \dots \alpha_n}$ are the solution of the (transposed) Vandermonde system of linear equations

$$\sum_{\alpha_1 + \dots + \alpha_n \leq d} c_{\alpha_1 \dots \alpha_n} \cdot p_1^{(k-1)\alpha_1} \cdot \dots \cdot p_n^{(k-1)\alpha_n} = v[k], \quad k = 1, \dots, N.$$

Note: the ground field has to be the field of rational numbers. Moreover, `ncols(p)==n`, the number of variables in the basering, and all the given generators have to be numbers different from 0,1 or -1. Finally, `ncols(v)==(d+1)n`, and all given generators have to be numbers.

Example:

```
ring r=0,(x,y),dp;
// determine f with deg(f)=2 and with given values v of f
// at 9 points: (2,3)^0=(1,1), ..., (2,3)^8=(2^8,3^8)
// valuation point: (2,3)
ideal p=2,3;
ideal v=1,2,3,4,5,6,7,8,9;
poly ip=vandermonde(p,v,2);
ip[1..5]; // the 5 first terms of ip:
↳ -1/9797760x2y2-595/85536x2y+55/396576xy2+935/384x2-1309/3240xy
// compute value of ip at the point 2^8,3^8, result must be 9
subst(subst(ip,x,2^8),y,3^8);
↳ 9
```

5.1.163 var

Syntax: `var (int_expression)`

Type: poly

Purpose: `var(n)` returns the `n`-th ring variable.

Example:

```
ring r=0,(x,y,z),dp;
var(2);
↳ y
```

See Section 4.6 [int], page 82; Section 5.1.108 [nvars], page 229; Section 4.19 [ring], page 124; Section 5.1.137 [rvar], page 253; Section 5.1.160 [univariate], page 278; Section 5.1.165 [varstr], page 280.

5.1.164 variables

Syntax: `variables (poly_expression)`
`variables (ideal_expression)`
`variables (matrix_expression)`

Type: ideal

Purpose: `variables(p)` returns the list of all ring variables the argument depends on.

Example:

```

ring r=0,(x,y,z),dp;
variables(2);
↳ _[1]=0
variables(x+y2);
↳ _[1]=x
↳ _[2]=y
variables(ideal(x+y2,x3y,z));
↳ _[1]=x
↳ _[2]=y
↳ _[3]=z
string(variables(ideal(x+y2,x3y,z)));
↳ x,y,z

```

See Section 5.1.77 [leadexp], page 207; Section 5.1.108 [nvars], page 229; Section 5.1.160 [univariate], page 278; Section 5.1.163 [var], page 279; Section 5.1.165 [varstr], page 280.

5.1.165 varstr

Syntax: `varstr (ring_name)`
`varstr (int_expression)`
`varstr (ring_name, int_expression)`

Type: string

Purpose: returns the list of the names of the ring variables as a string or the name of the n-th ring variable, where n is given by the int_expression. If the ring name is omitted, the basering is used, thus `varstr(n)` is equivalent to `varstr(basing,n)`.

Example:

```

ring r=0,(x,y,z),dp;
varstr(r);
↳ x,y,z
varstr(r,1);
↳ x
varstr(2);
↳ y

```

See Section 5.1.7 [charstr], page 160; Section 4.6 [int], page 82; Section 5.1.108 [nvars], page 229; Section 5.1.112 [ordstr], page 235; Section 5.1.115 [parstr], page 236; Section 4.19 [ring], page 124; Section 5.1.163 [var], page 279.

5.1.166 vdim

Syntax: vdim (ideal_expression)
vdim (module_expression)

Type: int

Purpose: computes the vector space dimension of the ring, resp. free module, modulo the ideal, resp. module, generated by the initial terms of the given generators. If the generators form a standard basis, this is the same as the vector space dimension of the ring, resp. free module, modulo the ideal, resp. module.
If the ideal, resp. module, is not zero-dimensional, -1 is returned.

Example:

```
ring r=0,(x,y),ds;
ideal i=x2+y2,x2-y2;
ideal j=std(i);
vdim(j);
↳ 4
```

See [codim], page 868; Section 5.1.20 [degree], page 169; Section 5.1.25 [dim], page 171; Section 4.5 [ideal], page 78; Section 5.1.69 [kbase], page 203; Section 5.1.100 [mult], page 224; Section 5.1.149 [std], page 266.

5.1.167 waitall

Syntax: waitall (list_expression)
waitall (list_expression , int_expression)

Type: int

Purpose: Expects a list of open links (of mode ssi:fork, ssi:tcp) and waits until all of them are finished, i.e., are ready to be read.

In the first case, the command waits for all links to finish (or to crash, see below) and may therefore run forever.

In the second case, a timeout in milliseconds can be provided, forcing the command to terminate after the specified time. If the given timeout is 0, the command checks whether all links are finished or not, but does not wait for any link (polling).

Return values are:

-1: The read state of all links is "eof", see Section 4.9 [link], page 94, Section 5.1.148 [status], page 265. This might happen if all the links crashed.

0: timeout (or polling): None of the links is ready.

1: All links are ready. (Note: There might be links whose read state is "eof", but at least one link is ready.)

Example:

```
link l1 = "ssi:fork"; open(l1);
link l2 = "ssi:fork"; open(l2);
link l3 = "ssi:fork"; open(l3);
list l = list(l1,l2,l3);
write(l1, quote(system("sh", "sleep 15")));
write(l2, quote(system("sh", "sleep 10")));
write(l3, quote(system("sh", "sleep 11")));
waitall(l, 5000); // terminates after 5sec with result 0
↳ 0
```

```

        waitall(1);          // terminates after 10 more sec
    ↪ 1
        close(11);
        close(12);
        close(13);

```

See Section 5.1.168 [waitfirst], page 282.

5.1.168 waitfirst

Syntax: waitfirst (list_expression)
 waitfirst (list_expression , int_expression)

Type: int

Purpose: Expects a list of open links (of mode ssi:fork, ssi:tcp) and waits until the first of them is finished, i.e., is ready to be read.

In the first case, the command waits until the first link is finished (or all of them crashed, see below) and may therefore run forever.

In the second case, a timeout in milliseconds can be provided, forcing the command to terminate after the specified time. If the given timeout is 0, the command checks whether one of the links is finished or not, but does not wait for any link (polling).

Return values are:

-1: The read state of all links is "eof", see Section 4.9 [link], page 94, Section 5.1.148 [status], page 265. This might happen if all the links crashed.

0: timeout (or polling): None of the links is ready.

i>1: At least the link whose list index is i is ready.

Example:

```

        link l1 = "ssi:fork"; open(l1);
        link l2 = "ssi:fork"; open(l2);
        link l3 = "ssi:fork"; open(l3);
        list l = list(l1,l2,l3);
        write(l1, quote(system("sh", "sleep 15")));
        write(l2, quote(system("sh", "sleep 13")));
        write(l3, quote(system("sh", "sleep 11")));
        waitfirst(l, 5000); // terminates after 5sec with result 0
    ↪ 0
        waitfirst(1);      // terminates after 6 more sec with result 3
    ↪ 3
        close(l1);
        close(l2);
        close(l3);

```

See Section 5.1.167 [waitall], page 281.

5.1.169 wedge

Syntax: wedge (matrix_expression , int_expression)

Type: matrix

Purpose: wedge(M,n) computes the n-th exterior power of the matrix M.

Example:

```

ring r;
matrix m[2][3]=x,y,y,z,z,x;
print(m);
↳ x,y,y,
↳ z,z,x
print(wedge(m,2));
↳ xz-yz,-x2+yz,xy-yz

```

See Section 4.6 [int], page 82; Section 4.12 [matrix], page 106; Section 5.1.92 [minor], page 218.

5.1.170 weight

Syntax: `weight (ideal_expression)`
`weight (module_expression)`

Type: intvec

Purpose: computes an "optimal" weight vector for an ideal, resp. module, which may be used as weight vector for the variables in order to speed up the standard basis algorithm. If the input is weighted homogeneous, a weight vector for which the input is weighted homogeneous is found.

Example:

```

ring h1=32003,(t,x,y,z),dp;
ideal i=
9x8+y7t3z4+5x4y2t2+2xy2z3t2,
9y8+7xy6t+2x5y4t2+2x2yz3t2,
9z8+3x2y3z2t4;
intvec e=weight(i);
e;
↳ 5,7,5,7
ring r=32003,(a,b,c,d),wp(e);
map f=h1,a,b,c,d;
ideal i0=std(f(i));

```

See Section 4.5 [ideal], page 78; Section 4.8 [intvec], page 91; Section 5.1.122 [qhweight], page 241.

5.1.171 weightKB

Procedure from library `standard.lib` (see Section D.1 [standard.lib], page 785).

Syntax: `weightKB (module_expression, int_expression, list_expression)`
`weightKB (ideal_expression, int_expression, list_expression)`

Return: the same as the input type of the first argument

Purpose: If `I,d,wim` denotes the three arguments then `weightKB` computes the weighted degree-d part of a vector space basis (consisting of monomials) of the quotient ring, resp. of the quotient module, modulo `I` w.r.t. weights given by `wim`. The information about the weights is given as a list of two intvec: `wim[1]` weights for all variables (positive), `wim[2]` weights for the module generators.

Note: This is a generalization of the command `kbasis` with the same first two arguments.

Example:

```

ring R=0, (x,y), wp(1,2);
weightKB(ideal(0),3,intvec(1,2));
↪ _[1]=x3
↪ _[2]=xy

```

See also: Section 5.1.69 [kbase], page 203.

5.1.172 write

Syntax: `write (link_expression, expression_list)`
for DBM links:
`write (link, string_expression, string_expression)`
`write (link, string_expression)`

Type: none

Purpose: writes data to a link.
If the link is of type ASCII, all expressions are converted to strings (and separated by a newline character) before they are written. As a consequence, only such values which can be converted to a string can be written to an ASCII link.
For ssi links, ring-dependent expressions are written together with a ring description. To prevent an evaluation of the expression before it is written, the `quote` command (possibly together with `eval`) can be used. A `write` blocks (i.e., does not return to the prompt), as long as a ssi link is not ready for writing.
For DBM links, `write` with three arguments inserts the first string as key and the second string as value into the dbm data base.
Called with two arguments, it deletes the entry with the key specified by the string from the data base.

Example:

```

// write the values of the variables f and i as strings into
// the file "outfile" (overwrite it, if it exists)
write(":w outfile",f,i);

// now append the string "that was f,i" (without the quotes)
// at the end of the file "outfile"
write(":a outfile","that was f,i");
// alternatively, links could be used:
link l=":a outfile"; l;
// type : ASCII
// mode : a
// name : outfile
// open : no
// read : not ready
// write: not ready
write(l," that was f,i");
// saving and retrieving data (ASCII format):
ring r=32003,(x,y,z),dp;
ideal i=x+y,z3+22y;
write(":w save_i",i);// this writes x+y,z3+22y to the file save_i
ring r=32003,(x,y,z),dp;
string s=read("save_i"); //creates the string x+y,z3+22y
execute("ideal k="+s+""); // this defines an ideal k which
// is equal to i.

```

```
// for large objects, the ssi format and ssi links are better:
write("ssi:w save_i.ssi",i);
def j=read("ssi:r save_i.ssi");
```

See Chapter 4 [Data types], page 72; Section 5.1.27 [dump], page 173; Section 5.1.29 [eval], page 174; Section 4.9 [link], page 94; Section 5.1.119 [print], page 238; [printf], page 785; Section 5.1.124 [quote], page 242; Section 5.1.128 [read], page 245; Section 5.3.7 [short], page 300.

5.2 Control structures

A sequence of commands surrounded by curly brackets (`{` and `}`) is a so-called block. Blocks are used in SINGULAR in order to define procedures and to collect commands belonging to `if`, `else`, `for` and `while` statements and to the `example` part in libraries. Even if the sequence of statements consists of only a single command it has to be surrounded by curly brackets! Variables which are defined inside a block are not local to that block. Note that there need not be an ending semicolon at the end of the block.

Example:

```
if ( i>j )
{
  // This is the block
  int temp;
  temp=i;
  i=j;
  j=temp;
  kill temp;
}
```

5.2.1 apply

Syntax: `apply(expression , function);`

Purpose: applies the function to all elements of the first argument. The first argument must be of type `intvec`, `intmat`, or `list`. The result will be an expression list, its type and format will be set by the following assign. The function must be a kernel command or a procedure which takes one argument and returns a value.

Example:

```
proc p(int x) {return(x^2);}
intvec v=1,2,3;
apply(v,p);
↳ 1 4 9
intvec vv=apply(v,p);vv;
↳ 1,4,9
list ll=apply(v,p);ll;
↳ [1]:
↳ 1
↳ [2]:
↳ 4
↳ [3]:
↳ 9
```

5.2.2 break

Syntax: `break;`

Purpose: leaves the innermost `for` or `while` block.

Example:

```
while (1)
{
    ...
    if ( ... )
    {
        break; // leave the while block
    }
}
```

See Section 5.2 [Control structures], page 285; Section 5.2.8 [for], page 290; Section 5.2.15 [while], page 296.

5.2.3 breakpoint

Syntax: `breakpoint(proc_name);`
`breakpoint(proc_name, line_no);`

Purpose: sets a breakpoint at the beginning of the specified procedure or at the given line.

Note: Line number 1 is the first line of a library (for procedures from libraries), resp. the line with the `{`.

A line number of -1 removes all breakpoint from that procedure.

Example:

```
breakpoint(groebner);
↳ breakpoint 1, at line 838 in groebner
breakpoint(groebner, 176);
↳ breakpoint 2, at line 176 in groebner
breakpoint(groebner, -1);
↳ breakpoints in groebner deleted(0x6)
```

See Section 3.9.3 [Source code debugger], page 68; Section 5.2.16 [~], page 297.

5.2.4 continue

Syntax: `continue;`

Purpose: skips the rest of the innermost `for` or `while` loop und jumps to the beginning of the block. This command is only valid inside a `for` or a `while` construction.

Note: Unlike the C-construct it **does not execute the increment statement**. The command `continue` is mainly for internal use.

Example:

```
for (int i = 1 ; i<=10; i=i+1)
{
    ...
    if (i==3) { i=8;continue; }
    // skip the rest if i is 3 and
    // continue with the next i: 8
```



```

        i;
    }
    ↪ 1
    ↪ 2
    ↪ 8
    ↪ 9
    ↪ 10

```

See Section 5.2 [Control structures], page 285; Section 5.2.8 [for], page 290; Section 5.2.15 [while], page 296.

5.2.5 else

Syntax: `if (boolean_expression) true_block else false_block`

Purpose: executes `false_block` if the `boolean_expression` of the `if` statement is false. This command is only valid in combination with an `if` command.

Example:

```

    int i=3;
    if ( i > 5)
    {
        "i is bigger than 5";
    }
    else
    {
        "i is smaller than 6";
    }
    ↪ i is smaller than 6

```

See Section 5.2 [Control structures], page 285; Section 4.6.5 [boolean expressions], page 86; Section 5.2.9 [if], page 291.

5.2.6 export

Syntax: `export name ;`
`export list_of_names ;`

Purpose: converts a local variable of a procedure to a global one, that is the identifier is not removed at the end of the procedure. However, the package the variable belongs to is not changed.

Note: Objects defined in a ring are not automatically exported when exporting the ring.

Example:

```

    proc p1
    {
        int i,j;
        export(i);
        intmat m;
        listvar();
        export(m);
    }
    p1();
    ↪ // m

```

[1] intmat 1 x 1

```

⇒ // j                [1] int 0
⇒ // i                [0] int 0
listvar();
⇒ // m                [0] intmat 1 x 1
⇒ // i                [0] int 0

```

See Section 5.2.7 [exportto], page 288; Section 5.2.10 [importfrom], page 291; Section 5.2.11 [keeping], page 293.

5.2.7 exportto

Syntax: `exportto(package_name , name);`
`exportto(package_name , list_of_names);`

Purpose: transfers an identifier in the current package into the one specified by `package_name`. `package_name` can be `Current`, `Top` or any other identifier of type package.

Note: Objects defined in a ring are not automatically exported when exporting the ring.

Warning: The identifier is transferred to the other package. It does no longer exist in the current package. If the identifier should only be copied, Section 5.2.10 [importfrom], page 291 should be used instead.

Example:

```

proc p1
{
  int i,j;
  exportto(Current,i);
  intmat m;
  listvar(Current);
  exportto(Top,m);
}
p1();
⇒ // Top                [0] package Top (T)
⇒ // ::m                [1] intmat 1 x 1
⇒ // ::i                [0] int 0
⇒ // ::j                [1] int 0
⇒ // ::#                [1] list, size: 0
⇒ // ::p1               [0] proc
⇒ // ::mathicgb_prOrder [0] proc from singmathic.so (C)
⇒ // ::mathicgb         [0] proc from singmathic.so (C)
⇒ // ::create_ring      [0] proc from standard.lib
⇒ // ::min              [0] proc from standard.lib
⇒ // ::max              [0] proc from standard.lib
⇒ // ::datetime         [0] proc from standard.lib
⇒ // ::weightKB         [0] proc from standard.lib
⇒ // ::fprintf          [0] proc from standard.lib
⇒ // ::printf           [0] proc from standard.lib
⇒ // ::sprintf          [0] proc from standard.lib
⇒ // ::quotient4        [0] proc from standard.lib
⇒ // ::quotient5        [0] proc from standard.lib
⇒ // ::quotient3        [0] proc from standard.lib
⇒ // ::quotient2        [0] proc from standard.lib
⇒ // ::quotient1        [0] proc from standard.lib
⇒ // ::quot             [0] proc from standard.lib

```

```

⇒ // ::res [0] proc from standard.lib
⇒ // ::groebner [0] proc from standard.lib
⇒ // ::qslimgb [0] proc from standard.lib
⇒ // ::hilbRing [0] proc from standard.lib
⇒ // ::par2varRing [0] proc from standard.lib
⇒ // ::quotientList [0] proc from standard.lib
⇒ // ::stdhilb [0] proc from standard.lib
⇒ // ::stdfglm [0] proc from standard.lib
⇒ // ::Float [0] proc from kernel (C)
⇒ // ::crossprod [0] proc from kernel (C)
package Test1;
exportto(Test1,p1);
listvar(Top);
⇒ // Top [0] package Top (T)
⇒ // ::m [0] intmat 1 x 1
⇒ // ::i [0] int 0
⇒ // ::mathicgb_prOrder [0] proc from singmathic.so (C)
⇒ // ::mathicgb [0] proc from singmathic.so (C)
⇒ // ::create_ring [0] proc from standard.lib
⇒ // ::min [0] proc from standard.lib
⇒ // ::max [0] proc from standard.lib
⇒ // ::datetime [0] proc from standard.lib
⇒ // ::weightKB [0] proc from standard.lib
⇒ // ::fprintf [0] proc from standard.lib
⇒ // ::printf [0] proc from standard.lib
⇒ // ::sprintf [0] proc from standard.lib
⇒ // ::quotient4 [0] proc from standard.lib
⇒ // ::quotient5 [0] proc from standard.lib
⇒ // ::quotient3 [0] proc from standard.lib
⇒ // ::quotient2 [0] proc from standard.lib
⇒ // ::quotient1 [0] proc from standard.lib
⇒ // ::quot [0] proc from standard.lib
⇒ // ::res [0] proc from standard.lib
⇒ // ::groebner [0] proc from standard.lib
⇒ // ::qslimgb [0] proc from standard.lib
⇒ // ::hilbRing [0] proc from standard.lib
⇒ // ::par2varRing [0] proc from standard.lib
⇒ // ::quotientList [0] proc from standard.lib
⇒ // ::stdhilb [0] proc from standard.lib
⇒ // ::stdfglm [0] proc from standard.lib
⇒ // ::Float [0] proc from kernel (C)
⇒ // ::crossprod [0] proc from kernel (C)
listvar(Test1);
⇒ // Test1 [0] package Test1 (N)
⇒ // ::p1 [0] proc
Test1::p1();
⇒ // Test1 [0] package Test1 (N)
⇒ // ::m [1] intmat 1 x 1
⇒ // ::i [0] int 0
⇒ // ::j [1] int 0
⇒ // ::# [1] list, size: 0
⇒ // ::p1 [0] proc
⇒ // ** redefining m ( exportto(Top,m);)

```

```

listvar(Top);
↳ // Top [0] package Top (T)
↳ // ::m [0] intmat 1 x 1
↳ // ::i [0] int 0
↳ // ::mathicgb_prOrder [0] proc from singmathic.so (C)
↳ // ::mathicgb [0] proc from singmathic.so (C)
↳ // ::create_ring [0] proc from standard.lib
↳ // ::min [0] proc from standard.lib
↳ // ::max [0] proc from standard.lib
↳ // ::datetime [0] proc from standard.lib
↳ // ::weightKB [0] proc from standard.lib
↳ // ::fprintf [0] proc from standard.lib
↳ // ::printf [0] proc from standard.lib
↳ // ::sprintf [0] proc from standard.lib
↳ // ::quotient4 [0] proc from standard.lib
↳ // ::quotient5 [0] proc from standard.lib
↳ // ::quotient3 [0] proc from standard.lib
↳ // ::quotient2 [0] proc from standard.lib
↳ // ::quotient1 [0] proc from standard.lib
↳ // ::quot [0] proc from standard.lib
↳ // ::res [0] proc from standard.lib
↳ // ::groebner [0] proc from standard.lib
↳ // ::qslimgb [0] proc from standard.lib
↳ // ::hilbRing [0] proc from standard.lib
↳ // ::par2varRing [0] proc from standard.lib
↳ // ::quotientList [0] proc from standard.lib
↳ // ::stdhilb [0] proc from standard.lib
↳ // ::stdfglm [0] proc from standard.lib
↳ // ::Float [0] proc from kernel (C)
↳ // ::crossprod [0] proc from kernel (C)
listvar(Test1);
↳ // Test1 [0] package Test1 (N)
↳ // ::i [0] int 0
↳ // ::p1 [0] proc

```

See Section 5.2.6 [export], page 287; Section 5.2.10 [importfrom], page 291; Section 5.2.11 [keeping], page 293.

5.2.8 for

Syntax: for (init_command; boolean_expression; iterate_commands) block

Purpose: repetitive, conditional execution of a command block.

The command `init_command` is executed first. Then `boolean_expression` is evaluated. If its value is `TRUE` the block is executed, otherwise the `for` statement is complete. After each execution of the block, the command `iterate_command` is executed and `boolean_expression` is evaluated. This is repeated until `boolean_expression` evaluates to `FALSE`.

The command `break;` leaves the innermost `for` construct.

Example:

```

// sum of 1 to 10:
int s=0;
for (int i=1; i<=10; i=i+1)

```

```

{
    s=s+i;
}
s;
↳ 55

```

See Section 5.2 [Control structures], page 285; Section 4.6.5 [boolean expressions], page 86; Section 5.2.2 [break], page 286; Section 5.2.4 [continue], page 286; Section 5.2.9 [if], page 291; Section 5.2.15 [while], page 296.

5.2.9 if

Syntax: `if (boolean_expression) true_block`
`if (boolean_expression) true_block else false_block`

Purpose: executes `true_block` if the boolean condition is true. If the `if` statement is followed by an `else` statement and the boolean condition is false, then `false_block` is executed.

Example:

```

int i = 9;
matrix m[i][i];
if ( i > 5 and typeof(m) == "matrix" )
{
    m[i][i] = i;
}

```

See Section 5.2 [Control structures], page 285; Section 4.6.5 [boolean expressions], page 86; Section 5.2.2 [break], page 286; Section 5.2.5 [else], page 287.

5.2.10 importfrom

Syntax: `importfrom(package_name , name);`
`importfrom(package_name , list_of_names);`

Purpose: creates a new identifier in the current package which is a copy of the one specified by `name` in the package `package_name`. `package_name` can be `Top` or any other identifier of type package.

Note: Objects defined in a ring are not automatically imported when importing the ring.

Warning: The identifier is copied to the current package. It does still exist (independently) in the package `package_name`. If the identifier should be erased in the package from which it originates, Section 5.2.7 [exportto], page 288 should be used instead.

Example:

```

listvar(Top);
↳ // Top [0] package Top (T)
↳ // ::mathicgb_prOrder [0] proc from singmathic.so (C)
↳ // ::mathicgb [0] proc from singmathic.so (C)
↳ // ::create_ring [0] proc from standard.lib
↳ // ::min [0] proc from standard.lib
↳ // ::max [0] proc from standard.lib
↳ // ::datetime [0] proc from standard.lib
↳ // ::weightKB [0] proc from standard.lib
↳ // ::fprintf [0] proc from standard.lib
↳ // ::printf [0] proc from standard.lib

```

```

⇒ // ::sprintf [0] proc from standard.lib
⇒ // ::quotient4 [0] proc from standard.lib
⇒ // ::quotient5 [0] proc from standard.lib
⇒ // ::quotient3 [0] proc from standard.lib
⇒ // ::quotient2 [0] proc from standard.lib
⇒ // ::quotient1 [0] proc from standard.lib
⇒ // ::quot [0] proc from standard.lib
⇒ // ::res [0] proc from standard.lib
⇒ // ::groebner [0] proc from standard.lib
⇒ // ::qslimb [0] proc from standard.lib
⇒ // ::hilbRing [0] proc from standard.lib
⇒ // ::par2varRing [0] proc from standard.lib
⇒ // ::quotientList [0] proc from standard.lib
⇒ // ::stdhilb [0] proc from standard.lib
⇒ // ::stdfglm [0] proc from standard.lib
⇒ // ::Float [0] proc from kernel (C)
⇒ // ::crossprod [0] proc from kernel (C)
load("inout.lib");
listvar(Top);
⇒ // Top [0] package Top (T)
⇒ // ::mathicgb_prOrder [0] proc from singmathic.so (C)
⇒ // ::mathicgb [0] proc from singmathic.so (C)
⇒ // ::create_ring [0] proc from standard.lib
⇒ // ::min [0] proc from standard.lib
⇒ // ::max [0] proc from standard.lib
⇒ // ::datetime [0] proc from standard.lib
⇒ // ::weightKB [0] proc from standard.lib
⇒ // ::fprintf [0] proc from standard.lib
⇒ // ::printf [0] proc from standard.lib
⇒ // ::sprintf [0] proc from standard.lib
⇒ // ::quotient4 [0] proc from standard.lib
⇒ // ::quotient5 [0] proc from standard.lib
⇒ // ::quotient3 [0] proc from standard.lib
⇒ // ::quotient2 [0] proc from standard.lib
⇒ // ::quotient1 [0] proc from standard.lib
⇒ // ::quot [0] proc from standard.lib
⇒ // ::res [0] proc from standard.lib
⇒ // ::groebner [0] proc from standard.lib
⇒ // ::qslimb [0] proc from standard.lib
⇒ // ::hilbRing [0] proc from standard.lib
⇒ // ::par2varRing [0] proc from standard.lib
⇒ // ::quotientList [0] proc from standard.lib
⇒ // ::stdhilb [0] proc from standard.lib
⇒ // ::stdfglm [0] proc from standard.lib
⇒ // ::Float [0] proc from kernel (C)
⇒ // ::crossprod [0] proc from kernel (C)
importfrom(Inout,pause);
listvar(Top);
⇒ // Top [0] package Top (T)
⇒ // ::pause [0] proc from inout.lib
⇒ // ::mathicgb_prOrder [0] proc from singmathic.so (C)
⇒ // ::mathicgb [0] proc from singmathic.so (C)
⇒ // ::create_ring [0] proc from standard.lib

```

```

⇒ // ::min [0] proc from standard.lib
⇒ // ::max [0] proc from standard.lib
⇒ // ::datetime [0] proc from standard.lib
⇒ // ::weightKB [0] proc from standard.lib
⇒ // ::fprintf [0] proc from standard.lib
⇒ // ::printf [0] proc from standard.lib
⇒ // ::sprintf [0] proc from standard.lib
⇒ // ::quotient4 [0] proc from standard.lib
⇒ // ::quotient5 [0] proc from standard.lib
⇒ // ::quotient3 [0] proc from standard.lib
⇒ // ::quotient2 [0] proc from standard.lib
⇒ // ::quotient1 [0] proc from standard.lib
⇒ // ::quot [0] proc from standard.lib
⇒ // ::res [0] proc from standard.lib
⇒ // ::groebner [0] proc from standard.lib
⇒ // ::qslimb [0] proc from standard.lib
⇒ // ::hilbRing [0] proc from standard.lib
⇒ // ::par2varRing [0] proc from standard.lib
⇒ // ::quotientList [0] proc from standard.lib
⇒ // ::stdhilb [0] proc from standard.lib
⇒ // ::stdfglm [0] proc from standard.lib
⇒ // ::Float [0] proc from kernel (C)
⇒ // ::crossprod [0] proc from kernel (C)

```

See Section 5.2.6 [export], page 287; Section 5.2.7 [exportto], page 288; Section 5.2.11 [keepring], page 293.

5.2.11 keepring

Syntax: keepring name ;

Warning: This command is obsolete. Instead the respective identifiers in the ring should be exported and the ring itself should subsequently be returned. The command is only included for backward compatibility and may be removed in future releases.

Purpose: moves the specified ring to the next (upper) level. This command can only be used inside of procedures and it should be the last command before the `return` statement. There it provides the possibility to keep a ring which is local to the procedure (and its objects) accessible after the procedure ended without making the ring global.

Example:

```

proc P1
{
  ring r=0,x,dp;
  keepring r;
}
proc P2
{
  "inside P2: " + nameof(basering);
  P1();
  "inside P2, after call of P1: " + nameof(basering);
}
ring r1= 0,y,dp;
P2();

```

```

↳ inside P2: r1
↳ inside P2, after call of P1: r
"at top level: " + nameof(basering);
↳ at top level: r1

```

See Section 4.19 [ring], page 124.

5.2.12 load

Syntax: `load(string_expression);`
`load(string_expression , string_expression);`

Type: none

Purpose: reads a library of procedures from a file. In contrast to the command `LIB` (see note below), the command `load` does not add the procedures of the library to the package `Top`, but only to the package corresponding to the library. If the given filename does not start with `.` or `/`, the following directories are searched for it (in the given order): the current directory, the directories given in the environment variable `SINGULARPATH`, some default directories relative to the location of the `SINGULAR` executable program, and finally some default absolute directories. You can view the search path which `SINGULAR` uses to locate its libraries, by starting up `SINGULAR` with the option `-v`, or by issuing the command `system("with");`.

The second string selections options for loading.

Note: `load(<string_expr>,"with")` is equivalent to
`LIB <string_expr>`.

Note: `load(<string_expr>,"try")` is equivalent to
`LIB <string_expr>` which never fails - test the package name to distinguish.

All loaded libraries are displayed by the `listvar(package);` command:

```

option(loadLib); // show loading of libraries;
                // standard.lib is loaded

listvar(package);
↳ // Singmathic [0] package Singmathic (C,singmathic.s\
o)
↳ // Standard [0] package Standard (S,standard.lib)
↳ // Top [0] package Top (T)
// the names of the procedures of inout.lib
load("inout.lib"); // are now known to Singular
↳ // ** loaded inout.lib (4.1.2.0, Feb_2019)
listvar(package);
↳ // Inout [0] package Inout (S,inout.lib)
↳ // Singmathic [0] package Singmathic (C,singmathic.s\
o)
↳ // Standard [0] package Standard (S,standard.lib)
↳ // Top [0] package Top (T)
load("blabla.lib","try");
listvar(package);
↳ // Inout [0] package Inout (S,inout.lib)
↳ // Singmathic [0] package Singmathic (C,singmathic.s\
o)
↳ // Standard [0] package Standard (S,standard.lib)
↳ // Top [0] package Top (T)

```



```

option(noloadLib); // do not show loading of libraries;
load("matrix.lib","try");
listvar(package);
⇒ // Elim [0] package Elim (S,elim.lib)
⇒ // Triang [0] package Triang (S,triang.lib)
⇒ // Absfact [0] package Absfact (S,absfact.lib)
⇒ // Primdec [0] package Primdec (S,primdec.lib)
⇒ // Ring [0] package Ring (S,ring.lib)
⇒ // Random [0] package Random (S,random.lib)
⇒ // General [0] package General (S,general.lib)
⇒ // Polylib [0] package Polylib (S,polylib.lib)
⇒ // Nctools [0] package Nctools (S,nctools.lib)
⇒ // Matrix [0] package Matrix (S,matrix.lib)
⇒ // Inout [0] package Inout (S,inout.lib)
⇒ // Singmathic [0] package Singmathic (C,singmathic.s\
o)
⇒ // Standard [0] package Standard (S,standard.lib)
⇒ // Top [0] package Top (T)

```

Each time a library (Section 3.8 [Libraries], page 54) / dynamic module (Section 3.10 [Dynamic loading], page 71) is loaded, the corresponding package is created, if it does not already exist.

The name of a package corresponding to a SINGULAR library is derived from the name of the library file. The first letter is capitalized and everything to right of the left-most dot is dropped. For a dynamic module the packagename is hard-coded in the binary file.

Only the names of the procedures in the library are loaded, the body of the procedures is read during the first call of this procedure. This minimizes memory consumption by unused procedures. When SINGULAR is started with the `-q` or `--quiet` option, no message about the loading of a library is displayed.

```

option(loadLib); // show loading of libraries; standard.lib is loaded
// the names of the procedures of inout.lib
load("inout.lib"); // are now known to Singular
⇒ // ** loaded inout.lib (4.1.2.0, Feb_2019)
listvar();

```

See Section 3.1.6 [Command line options], page 19; Section A.1.9 [Dynamic modules], page 701; Section 5.1.79 [LIB], page 208; Section 2.3.3 [Procedures and libraries], page 10; Appendix D [SINGULAR libraries], page 785; Section 5.2.7 [exportto], page 288; Section 5.2.10 [importfrom], page 291; Section 4.15 [package], page 117; Section 4.17 [proc], page 121; Section D.1 [standard.lib], page 785; Section 4.21 [string], page 127; Section 5.1.153 [system], page 270.

5.2.13 quit

Syntax: `exit;`
`quit;`

Purpose: quits SINGULAR; works also from inside a procedure or from an interrupt. Instead of `quit`, the synonymous command `exit` may be used.

Example:

```
quit;
```

5.2.14 return

Syntax: `return (expression_list);`
`return ();`

Type: any

Purpose: returns the result(s) of a procedure and can only be used inside a procedure. Note that the brackets are required even if no return value is given.

Example:

```
proc p2
{
  int i,j;
  for(i=1;i<=10;i++)
  {
    j=j+i;
  }
  return(j);
}
// can also return an expression list, i.e., more than one value
proc tworeturn ()
{ return (1,2); }
int i,j = tworeturn();
// return type may even depend on the input
proc type_return (int i)
{
  if (i > 0) {return (i);}
  else {return (list(i));}
}
// then we need def type (or list) to collect value
def t1 = type_return(1);
def t2 = type_return(-1);
```

See Chapter 4 [Data types], page 72; Section 4.17 [proc], page 121.

5.2.15 while

Syntax: `while (boolean_expression) block`

Purpose: repetitive, conditional execution of block.
 The `boolean_expression` is evaluated and if its value is `TRUE`, the block gets executed. This is repeated until `boolean_expression` evaluates to `FALSE`. The command `break` leaves the innermost `while` construction.

Example:

```
int i = 9;
while (i>0)
{
  // ... // do something for i=9, 8, ..., 1
  i = i - 1;
}
while (1)
{
  // ... // do something forever
  if (i == -5) // but leave the loop if i is -5
  {
```

```

        break;
    }
}

```

See Section 5.2 [Control structures], page 285; Section 4.6.5 [boolean expressions], page 86; Section 5.2.2 [break], page 286.

5.2.16 ~ (break point)

Syntax: ~;

Purpose: sets a break point. Whenever SINGULAR reaches the command ~; in a sequence of commands it prompts for input. The user may now input lines of SINGULAR commands. The line length cannot exceed 80 characters. SINGULAR proceeds with the execution of the command following ~; as soon as it receives an empty line. Furthermore, the debug mode will be activated: See Section 3.9.3 [Source code debugger], page 68.

Example:

```

proc t
{
  int i=2;
  ~;
  return(i+1);
}
t();
↳ -- break point in t --
↳ -- 0: called from STDIN --
// here local variables of the procedure can be accessed
i;
↳ 2
↳ -- break point in t --

↳ 3

```

See Section 3.9.4 [Break points], page 69.

5.3 System variables

5.3.1 degBound

Type: int

Purpose: The standard basis computation is stopped if the total (weighted) degree exceeds degBound. degBound should not be used for a global ordering with inhomogeneous input, if the ordering is not dp or Dp. (Remark: elimination requires always an elimination ordering).
Reset this bound by setting degBound to 0.
The exact meaning of "degree" depends on the ring ordering and the command: slimgb uses always the total degree with weights 1, std does so for block orderings, only.

Example:

```

degBound = 7;
option();
↳ //options for 'std'-command: degBound
ideal j=std(i);
degBound;
↳ 7
degBound = 0; //resets degree bound to infinity

```

See Section 5.1.19 [deg], page 168; Section 4.6 [int], page 82; Section 5.1.110 [option], page 230; Section 5.1.149 [std], page 266.

5.3.2 echo

Type: int

Purpose: input is echoed if `echo >= voice`.
`echo` is a local setting for a procedure and defaulted to 0.
`echo` does not affect the output of commands.

Example:

```

echo = 1;
int i = echo;
↳ int i = echo;

```

See Section 4.6 [int], page 82; Section 5.3.11 [voice], page 303.

5.3.3 minpoly

Type: number

Purpose: describes the coefficient field of the current basering as an algebraic extension with the minimal polynomial equal to `minpoly`. Setting the `minpoly` should be the first command after defining the ring.

Note: The minimal polynomial has to be specified in the syntax of a polynomial. Its variable is not one of the ring variables, but the algebraic element which is being adjoined to the field. Algebraic extensions in SINGULAR are only possible over the rational numbers or over \mathbb{Z}/p , p a prime number.
SINGULAR does not check whether the given polynomial is irreducible! It can be checked in advance with the function `factorize` (see Section 5.1.36 [factorize], page 178).

Example:

```

//(Q[i]/(i^2+1))[x,y,z]:
ring Cxyz=(0,i),(x,y,z),dp;
minpoly=i^2+1;
i2; //this is a number, not a poly
↳ -1

```

See Section 5.1.36 [factorize], page 178; Section 4.19 [ring], page 124.

5.3.4 multBound

Type: int

Purpose: The standard basis computation is stopped if the ideal is zero-dimensional in a ring with local ordering and its multiplicity (`mult`) is lower than `multBound`.
Reset this bound by setting `multBound` to 0.

Example:

```

ring r=0,(x,y,z),ds;
ideal i,j;
i=x7+y7+z6,x6+y8+z7,x7+y5+z8,
x2y3+y2z3+x3z2,x3y2+y3z2+x2z3;
multBound=100;
j=std(i);
degree(j);
↳ // dimension (local) = 0
↳ // multiplicity = 98
multBound=0; //disables multBound
j=std(i);
degree(j);
↳ // dimension (local) = 0
↳ // multiplicity = 86

```

See Section 4.6 [int], page 82; Section 5.1.100 [mult], page 224; Section 5.1.110 [option], page 230; Section 5.1.149 [std], page 266.

5.3.5 noether

Type: poly

Purpose: The standard basis computation in local rings cuts off all monomials above (in the sense of the monomial ordering) the monomial `noether` during the computation. Reset `noether` by setting `noether` to 0.

Example:

```

ring R=32003,(x,y,z),ds;
ideal i=x2+y12,y13;
std(i);
↳ _[1]=x2+y12
↳ _[2]=y13
noether=x11;
std(i);
↳ _[1]=x2
noether=0; //disables noether

```

See Section 4.16 [poly], page 117; Section 5.1.149 [std], page 266.

5.3.6 printlevel

Type: int

Purpose: sets the debug level for `dbprint`. If `printlevel` \geq `voice` then `dbprint` is equivalent to `print`, otherwise nothing is printed.

Note: See Section 3.8.6 [Procedures in a library], page 57, for a small example about how this is used for the display of comments while procedures are executed.

Example:

```

voice;
↳ 1
printlevel=0;
dbprint(1);

```

```

    printlevel=voice;
    dbprint(1);
    ↪ 1

```

See Section 5.1.17 [dbprint], page 167; Section 4.6 [int], page 82; Section 5.3.11 [voice], page 303.

5.3.7 short

Type: int

Purpose: the output of monomials is done in the short manner, if `short` is non-zero. A C-like notion is used, if `short` is zero. Both notations may be used as input. The default depends on the names of the ring variables (0 if there are names of variables longer than 1 character, 1 otherwise). Every change of the basering sets `short` to the previous value for that ring. In other words, the value of the variable `short` is "ring-local".
If the names are long, or the ring non-commutative, `short` can not be schaged to 1.

Example:

```

    ring r=23,x,dp;
    int save=short;
    short=1;
    2x2,x2;
    ↪ 2x2 x2
    short=0;
    2x2,x2;
    ↪ 2*x^2 x^2
    short=save; //resets short to the previous value

```

See Section 4.6 [int], page 82.

5.3.8 timer

Type: int

Purpose:

1. the CPU time (i.e, user and system time) used for each command is printed if `timer > 0`, if this time is bigger than a (customizable) minimal time and if `printlevel+1 >= voice` (which is by default true on the SINGULAR top level, but not true while procedures are executed).
2. yields the CPU time used since the start-up of SINGULAR in a (customizable) resolution.

The default setting of `timer` is 0, the default minimal time is 0.5 seconds, and the default timer resolution is 1 (i.e., the default unit of time is one second). The minimal time and timer resolution can be set using the command line options `--min-time` and `--ticks-per-sec` and can be checked using `system("--min-time")` and `system("--ticks-per-sec")`.

How to use `timer` in order to measure the time for a sequence of commands, see example below.

Note for Windows95/98:

The value of the `timer` cannot be used (resp. trusted) when SINGULAR is run under Windows95/98 (this is due to the shortcomings of the Windows95/98 operating system). Use Section 5.3.10 [rtimer], page 303, instead.

Example:

```

timer=1; // The time of each command is printed
int t=timer; // initialize t by timer
ring r=0,(x,y,z),dp;
poly p=(x+2y+3z+4xy+5xz+6yz)^20;
// timer as int_expression:
t=timer-t;
t; // yields the time in ticks-per-sec (default 1)
↳ 0
    // since t was initialized by timer
int tps=system("--ticks-per-sec");
t div tps; // yields the time in seconds truncated to int
↳ 0
timer=0;
system("--ticks-per-sec",1000); // set timer resolution to ms
t=timer; // initialize t by timer
p=(x+2y+3z+4xy+5xz+6yz)^20;
timer-t; // time in ms
↳ 20

```

See Section 3.1.6 [Command line options], page 19; Section 5.3.6 [printlevel], page 299; Section 5.3.10 [rtimer], page 303; Section 5.1.153 [system], page 270; Section 5.3.11 [voice], page 303.

5.3.9 TRACE

Type: int

Purpose: sets level of debugging.

```

TRACE=0   No debugging messages are printed.
TRACE=1   Messages about entering and leaving of procedures are displayed.
TRACE=3   Messages about entering and leaving of procedures together with line numbers are displayed.
TRACE=4   Each line is echoed and the interpretation of commands in this line is suspended until the user presses RETURN.
TRACE=8   (debug version only:) show basering for all levels
TRACE=128
           show all calls to kernel routines
TRACE=256
           show all assigns
TRACE=512
           show all automatic type conversions
TRACE=1024
           profiling: print line numbers to smon.out

```

TRACE is defaulted to 0.

TRACE does not affect the output of commands.

Example:

```

TRACE=1;
LIB "general.lib";
sum(1..100);
↳ entering sum (level 0)
↳ entering   lsum (level 1)
↳ entering   lsum (level 2)
↳ entering   lsum (level 3)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ leaving    lsum (level 3)
↳ entering   lsum (level 3)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ leaving    lsum (level 3)
↳ leaving    lsum (level 2)
↳ entering   lsum (level 2)
↳ entering   lsum (level 3)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ leaving    lsum (level 3)
↳ entering   lsum (level 3)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ leaving    lsum (level 3)
↳ leaving    lsum (level 2)
↳ leaving    lsum (level 1)
↳ entering   lsum (level 1)
↳ entering   lsum (level 2)
↳ entering   lsum (level 3)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ leaving    lsum (level 3)
↳ entering   lsum (level 3)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)
↳ leaving    lsum (level 3)
↳ leaving    lsum (level 2)
↳ entering   lsum (level 2)
↳ entering   lsum (level 3)
↳ entering   lsum (level 4)
↳ leaving    lsum (level 4)

```



```

↳ entering      lsum (level 4)
↳ leaving       lsum (level 4)
↳ leaving       lsum (level 3)
↳ entering      lsum (level 3)
↳ entering      lsum (level 4)
↳ leaving       lsum (level 4)
↳ entering      lsum (level 4)
↳ leaving       lsum (level 4)
↳ leaving       lsum (level 3)
↳ leaving       lsum (level 2)
↳ leaving       lsum (level 1)
↳ leaving       sum (level 0)
↳ 5050

```

See Section 4.6 [int], page 82.

5.3.10 rtimer

Type: int

Purpose: identical to `timer` (see Section 5.3.8 [timer], page 300), except that real times (i.e., wall-clock) times are reported, instead of CPU times. This can be trusted on all operating systems (including Windows95/98).

5.3.11 voice

Type: int

Purpose: shows the nesting level of procedures.

Note: See Section 3.8 [Libraries], page 54, for a small example how this is used for the display of comments while procedures are executed.

Example:

```

    voice;
↳ 1
proc p
{
    voice;
};
p();
↳ 2

```

See Section 5.1.17 [dbprint], page 167; Section 5.1.82 [listvar], page 210; Section 5.3.6 [printlevel], page 299.

6 Tricks and pitfalls

6.1 Limitations

SINGULAR has the following limitations:

- the characteristic of a prime field must be less than or equal to 2147483647 (2^{31})
(the characteristic of a prime field in the factory routines must be less than 536870912 (2^{29}))
(the characteristic of a prime field in the NTL routines must be less than NTL_SP_BOUND (2^{30}) on 32bit machines - This is always the case since currently, only factory uses NTL.)
- the number of elements in $GF(p,n)$ must be less than 65536
- the (weighted) degree of a monomial must be less or equal than 2147483647
- the rank of any free module must be less or equal than 2147483647
- the maximal allowed exponent of a ring variable depends on the ordering of the ring and is at least 32767. See also Section B.2 [Monomial orderings], page 760 for setting other limits.
- the precision of long floating point numbers (for ground field `real`) must be less or equal than 32767
- integers (of type `int`) have the limited range from -2147483648 to 2147483647
- floating point numbers (type `number` from field `real`) have a limited range which is machine dependent. A typical range is -1.0e-38 to 1.0e+38. The string representation of overflow and underflow is machine dependent, as well. For example "Inf" on Linux, or "+. +00e+00" on HPUX.
Their input syntax is given by `scanf`, but must start with a digit.
- floating point numbers (type `number` from field `real` with a precision `p` larger then 3) use internally `mpf_set_default_prec(3.5*p+1)`.
Their input syntax is given by `mpf_set_str` from GMP, but must start with a digit.
- the length of an identifier is unlimited but `listvar` displays only the first 20 characters
- statements may not contain more than 10000 tokens
- tokens (i.e. strings, numbers, ...) may not be longer than 16382 characters
- All input to SINGULAR must be 7-bit clean, i.e. special characters like the the German Umlaute (ä, ö, etc.), or the French accent characters may neither appear as input to SINGULAR, nor in libraries or procedure definitions.

6.2 System dependent limitations

Ports of SINGULAR to different systems do not always implement all possible parts of SINGULAR:

- dynamic modules are implemented for
 - unix systems with ELF format for executables (Linux, Solaris, FreeBSD)

6.3 Major differences to the C programming language

Although many constructs from SINGULAR's programming language are similar to those from the C programming language, there are some subtle differences. Most notably:

6.3.1 No rvalue of increments and assignments

The increment operator `++` (resp. decrement operator `--`) has no rvalue, i.e., cannot be used on the right-hand sides of assignments. So, instead of

```
j = i++; // WRONG!!!
```

(which results in an error), it must be written

```
i++; j = i;
```

Likewise, an assignment expression does not have a result. Therefore, compound assignments like `i = j = k;` are not allowed and result in an error.

6.3.2 Evaluation of logical expressions

All arguments of a logical expression are first evaluated and then the value of the logical expression is determined. For example, the logical expressions `(a || b)` is evaluated by first evaluating `a` and `b`, even though the value of `b` has no influence on the value of `(a || b)`, if `a` evaluates to true.

Note, that this evaluation is different from the left-to-right, conditional evaluation of logical expressions (as found in most programming languages). For example, in these other languages, the value of `(1 || b)` is determined without ever evaluating `b`. This causes some problems with boolean tests on variables, which might not be defined at evaluation time. For example, the following results in an error, if the variable `i` is undefined:

```
if (defined(i) && i > 0) {} // WRONG!!!
```

This must be written instead as:

```
if (defined(i))
{
  if (i > 0) {}
}
```

However, there are several short work-arounds for this problem:

1. If a variable (say, `i`) is only to be used as a boolean flag, then define (value is TRUE) and undefine (value is FALSE) `i` instead of assigning a value. Using this scheme, it is sufficient to simply write

```
if (defined(i))
```

in order to check whether `i` is TRUE. Use the command `kill` to undefine a variable, i.e. to assign it a FALSE value (see Section 5.1.71 [kill], page 204).

2. If a variable can have more than two values, then define it, if necessary, before it is used for the first time. For example, if the following is used within a procedure

```
if (! defined(DEBUG)) { int DEBUG = 1;}
...
if (DEBUG == 3) {...}
if (DEBUG == 2) {...}
...
```

then a user of this procedure does not need to care about the existence of the `DEBUG` variable – this remains hidden from the user. However, if `DEBUG` exists globally, then its local default value is overwritten by its global one.

6.3.3 No case or switch statement

SINGULAR does not offer a `case` (or `switch`) statement. However, it can be imitated in the following way:

```

while (1)
{
    if (choice == choice_1) { ...; break;}
    ...
    if (choice == choice_n) { ...; break;}
    // default case
    ...; break;
}

```

6.3.4 Usage of commas

In SINGULAR, a comma separates list elements and the value of a comma expression is a list. Hence, commas cannot be used to combine several expressions into a single expression. For example, instead of writing

```
for (i=1, j=5; i<5 || j<10; i++, j++) {...} // WRONG!!!!!!
```

one has to write

```
for (i,j = 1,5; i<5 || j<10; i++, j++) {...}
```

6.3.5 Usage of brackets

In SINGULAR, curly brackets (`{ }`) **must always** be used to enclose the statement body following such constructs like `if`, `else`, `for`, or `while`, even if this block consists of only a single statement. Similarly, in the return statement of a procedure, parentheses (`()`) **must always** be used to enclose the return value. Even if there is no value to return, parentheses have to be used after a return statement (i.e., `return()`). For example,

```
if (i == 1) return i; // WRONG!!!!!!
```

results in an error. Instead, it must be written as

```
if (i == 1) { return (i); }.
```

6.3.6 Behavior of continue

SINGULAR's `continue` construct is only valid inside the body of a `for` or `while` construct. It skips the rest of the loop-body and jumps to the beginning of the block. Unlike the C-construct SINGULAR's `continue` **does not execute the increment statement**. For example,

```

for (int i = 1 ; i<=10; i=i+1)
{
    ...
    if (i==3) { i=8;continue; }
    // skip the rest if i is 3 and
    // continue with the next i: 8
    i;
}
↳ 1
↳ 2
↳ 8
↳ 9
↳ 10

```

6.3.7 Return type of procedures

Although the SINGULAR language is a strongly typed programming language, the type of the return value of a procedure does not need to be specified. As a consequence, the return type of a procedure may vary, i.e., may, for example, depend on the input. However, the return value of such a procedure may then only be assigned to a variable of type `def`.

```
proc type_return (int i)
{
  if (i > 0) {return (i);}
  else {return (list(i));}
}
def t1 = type_return(1);
def t2 = type_return(-1);
typeof(t1); typeof(t2);
↳ int
↳ list
```

Furthermore, it is mandatory to assign the return value of a procedure to a variable of type `def`, if a procedure changes the current ring using the `keepring` command (see Section 5.2.11 [keepring], page 293) and returns a ring-dependent value (like a polynomial or module).

```
proc def_return
{
  ring r=0,(x,y),dp;
  poly p = x;
  keepring r;
  return (x);
}
def p = def_return();
// poly p = def_return(); would be WRONG!!!
typeof(p);
↳ poly
```

On the other hand, more than one value can be returned by a single `return` statement. For example,

```
proc tworeturn () { return (1,2); }
int i,j = tworeturn();
```

6.3.8 First index is 1

Although the SINGULAR language is C like, the indices of all objects which may have an index start at 1.

```
ring r;
ideal i=1,x,z;
i[2];
↳ x
intvec v=1,2,3;
v[1];
↳ 1
poly p=x+y+z;
p[2];
↳ y
vector h=[x+y,x,z];
h[1];
```

```

↳ x+y
  h[1][1];
↳ x

```

6.4 Miscellaneous oddities

1. integer division

If two numerical constants (i.e., two sequences of digits) are divided using the / operator, the surrounding whitespace determines which division to use: if there is no space between the constants and the / operator (e.g., "3/2"), both numerical constants are treated as of type `number` and the current ring division is used. If there is at least one space surrounding the / operator (e.g., "3 / 2"), both numerical constants are treated as of type `int` and an integer division is performed. To avoid confusion, use the `div` operator instead of / for integer division and an explicit type cast to `number` for ring division. Note, that this problem does only occur for divisions of numerical constants. It also applies for large numerical constants which are of type `bigint`.

```

ring r=32002,x,dp;
↳ // ** 32002 is invalid as characteristic of the ground field. 32003 is us\
  ed.
  3/2; // ring division
↳ -16000
  3 / 2; // integer division
↳ // ** int division with '/': use 'div' instead in line >> 3 / 2; // int\
  eger division<<
↳ 1
  3 div 2;
↳ 1
  number(3) / number(2);
↳ -16000
  number a=3;
  number b=2;
  a/b;
↳ -16000
  int c=3;
  int d=2;
  c / d;
↳ // ** int division with '/': use 'div' instead in line >> c / d;<<
↳ 1

```

2. monomials and precedence

The formation of a monomial has precedence over all operators (a monomial is here an optional coefficient followed by any sequence of ring variables (possibly followed by an exponent) which only consist of letters, digits and (over the rationals) / without any whitespace):

```

ring r=0,(x,y),dp;
2xy^2 == (2*x*y)^2;
↳ 1
2xy^2 == 2x*y^2;
↳ 0
2x*y^2 == 2*x * (y^2);
↳ 1

```

During that formation no operator is involved: in the non-commutative case, we have

```

LIB "nctools.lib";
ring r = 0,(x,y),dp;
def S = superCommutative();
xy == yx;
↳ 1
x*y == y*x;
↳ 1
x*y, y*x;
↳ xy xy

```

3. meaning of `mult`

For an arbitrary ideal or module `i`, `mult(i)` returns the multiplicity of the ideal generated by the leading monomials of the given generators of `i`, hence depends on the monomial ordering! A standard mistake is to interpret `degree(i)` or `mult(i)` for an inhomogeneous ideal `i` as the degree of the homogenization or as something like the 'degree of the affine part'. For the ordering `dp` (degree reverse lexicographical) the converse is true: if `i` is given by a standard basis, `mult(i)` is the degree of the homogeneous ideal obtained by homogenization of `i` and then putting the homogenizing variable to 0, hence it is the degree of the part at infinity (this can also be checked by looking at the initial ideal).

4. size of ideals

`size` counts the non-zero entries of an ideal or module. Use `ncols` to determine the actual number of entries in the ideal or module.

5. computations in `qring`

In order to speed up computations in quotient rings, SINGULAR usually does not reduce polynomials w.r.t. the quotient ideal; rather the given representative is used as long as possible during computations. If it is necessary, reduction is done during standard base computations. To reduce a polynomial `f` by hand w.r.t. the current quotient ideal use the command `reduce(f,std(0))` (see Section 5.1.129 [reduce], page 246).

6. degree of a polynomial

`degBound`

The exact meaning of "degree" depends on the ring ordering and the command: `slimgb` uses always the total degree with weights 1, `std` does so only for block orderings.

`hilb`

the degree is the total degree with weights 1 unless a weight vector is given

`kbase`

the degree is the total degree with weights 1 (to use another weight vector see [weightKB], page 785)

7. substring selection

To extract substrings from a `string`, square brackets are used, enclosing either two comma-separated `ints` or an `intvec`. Although two comma-separated `ints` represent an `intvec`, they mean different things in substring access. Square brackets enclosing two `ints` (e.g. `s[2,6]`) return a substring where the first integer denotes the starting position and the second integer denotes the length of the substring. The result is returned as a `string`. Square brackets enclosing an `intvec` (e.g. `s[intvec(2,6)]`) return the characters of the string at the position given by the values of the `intvec`. The result is returned as an expression list of strings.

```

string s = "one-word";
s[2,6];    // a substring starting at the second char
↳ ne-wor
size(_);

```

```

↳ 6
  intvec v = 2,6;
  s[v];      // the second and the sixth char
↳ n o
  string st = s[v]; // stick together by an assignment
  st;
↳ no
  size(_);
↳ 2
  v = 2,6,8;
  s[v];
↳ n o d

```

8. packages and indexed variables

See example

```

package K;
string varok; exportto(K,varok);
string work(1); exportto(K,work(1));
int i(1..3); exportto(K,i(1..3));
// Toplevel does not contain i(1..3)
listvar();
// i(1..3) are stored in Package 'K'
listvar(K);
↳ // K                               [0] package K (N)
↳ // ::i(3)                           [0] int 0
↳ // ::i(2)                           [0] int 0
↳ // ::i(1)                           [0] int 0
↳ // ::work(1)                         [0] string
↳ // ::varok                           [0] string

```

6.5 Identifier resolution

In SINGULAR, an identifier (i.e., a "word") is resolved in the following way and order: It is checked for

1. a reserved name (like `ring`, `std`, ...),
2. a local variable (w.r.t. a procedure),
3. a local ring variable (w.r.t. the current basering locally set in a procedure),
4. a global ring variable (w.r.t. the current basering)
5. a global variable,
6. a monomial consisting of local ring variables written without operators,
7. a monomial consisting of global ring variables written without operators.

Consequently, it is allowed to have general variables with the same name as ring variables. However, the above identifier resolution order must be kept in mind. Otherwise, surprising results may come up.

```

ring r=0,(x,y),dp;
int x;
x*y; // resolved product int*poly, i.e., 0*y
↳ 0
xy; // "xy" is one identifier and resolved to monomial xy
↳ xy

```


For these reasons, we strongly recommend not to use variables which have the same name(s) as ring variables.

Moreover, we strongly recommend not to use ring variables whose name is fully contained in (i.e., is a substring of) another name of a ring variable. Otherwise, effects like the following might occur:

```
ring r=0,(x, x1),dp; // name x is substring of name x1 !!!!!!!!
x;x1; // resolved polynomial x
↳ x
↳ x1
short=0; 2x1; // resolved to monomial 2*x^1 !!!!!!!
↳ 2*x
2*x1; // resolved to product 2 times x1
↳ 2*x1
```

7 Non-commutative subsystem

SINGULAR has three non-commutative subsystems, handling various classes of associative non-commutative rings: PLURAL, SCA and LETTERPLACE.

7.1 PLURAL

What is and what does PLURAL?

PLURAL is a kernel extension of SINGULAR, providing many algorithms for computations within non-commutative G - and GR - algebras (see Section 7.4 [Mathematical background (plural)], page 360 for detailed information on algebras and algorithms).

It uses the same data structures as SINGULAR, sometimes interpreting them in a different way and/or modifying them for its own purposes. In spite of such a difference, one can always transfer objects between commutative rings of SINGULAR and non-commutative rings of PLURAL.

With PLURAL, one can set up a non-commutative G -algebra, say A , with a Poincaré-Birkhoff-Witt (PBW) basis, (see Section 7.4.1 [G-algebras], page 360 for step-by-step building instructions and also Section 7.5 [PLURAL libraries], page 365 for procedures for setting many important algebras easily). Afterwards, one can proceed to the factor-algebra of A modulo a two-sided ideal (see Section 7.3.29 [twostd (plural)], page 358), thus obtaining a GR -algebra (see Section 7.2.5 [qring (plural)], page 324 type).

Functionalities of PLURAL (enlisted in Section 7.3 [Functions (plural)], page 329) are accessible as soon as the basering becomes non-commutative (see Section 7.3.16 [nc_algebra], page 343 and the library Section 7.5.10 [ncalg.lib], page 458 with many readily predefined algebras).

One can perform various computations with polynomials and ideals in A and with vectors and submodules of a free module A^n .

What PLURAL does not:

PLURAL does not perform computations in the free algebra or in its general factor algebras (instead, these computations can be possibly done by Section 7.7 [LETTERPLACE], page 613).

In PLURAL one can only work with G -algebras and with their factor-algebras by two-sided ideals (GR -algebras).

PLURAL requires a global monomial ordering (see Section B.2.2 [General definitions for orderings], page 760). However, SCA (Section 7.6 [Graded commutative algebras (SCA)], page 611) provides the possibility of computations in a tensor product of a non-commutative graded commutative algebra (equipped with a global ordering) with a commutative algebra (equipped with any ordering).

PLURAL does not handle non-commutative parameters, i.e. the elements of the coefficient field (or a ring) mutually commute with all variables. Defining parameters, one **cannot** impose non-commutative relations on them. Moreover, it is impossible to introduce parameters which do not commute with variables. However, Section 7.5.21 [olga.lib], page 578 offers a rich functionality for working within Ore localizations of G -algebras and Section 7.5.25 [ratgb.lib], page 609 provides Groebner bases for so-called rational localizations of G -algebras.

PLURAL does not yet support rings like Z as coefficients.

PLURAL conventions

***-multiplication (plural)**

in the non-commutative case, the correct multiplication of y by x must be written as $y*x$.

Both expressions yx and xy are equal, since they are interpreted as commutative expressions. See example in Section 7.2.4.2 [poly expressions (plural)], page 323.

Note, that PLURAL output consists only of standard monomials, even when the signs $*$ are omitted.

ideal (plural)

Unless stated otherwise, an expression of type **ideal** as understood by PLURAL as a list of generators of a **left** ideal. For more information see Section 7.2.1 [ideal (plural)], page 313.

For a **two-sided ideal** T , use the command Section 7.3.29 [twostd (plural)], page 358 for computing the two-sided Groebner basis of T .

For a **right ideal** I , use Section 7.5.20.10 [rightStd], page 567 from `nctools_lib` for computing the right Groebner basis of I .

module (plural)

Unless stated otherwise, a **module** as understood by PLURAL is **either** a finitely generated **left** submodule of a free module (of finite rank)

or a factor module of a free module (of finite rank) by its left submodule (see Section 7.2.3 [module (plural)], page 320 for details). The concrete interpretation left to a function.

qring (plural)

It is only possible to build factor-algebras modulo **two-sided** ideals (see Section 7.2.5 [qring (plural)], page 324), which have to be given via their two-sided Groebner basis (see Section 7.3.29 [twostd (plural)], page 358).

7.2 Data types (plural)

This chapter explains all data types of PLURAL in alphabetical order. For every type, there is a description of the declaration syntax

as well as information about how to build expressions of certain types.

The term "expression list" in PLURAL refers to any comma separated list of expressions.

For the general syntax of a declaration see Section 3.5.1 [General command syntax], page 41.

7.2.1 ideal (plural)

For PLURAL ideals are **left** ideals, unless stated otherwise.

Ideals are represented as lists of polynomials which are interpreted as left generators of the ideal.

For the operations with two-sided ideals see Section 7.3.29 [twostd (plural)], page 358.

Like polynomials, ideals can only be defined or accessed with respect to a basering.

Note: `size` counts only the non-zero generators of an ideal whereas `ncols` counts all generators.

7.2.1.1 ideal declarations (plural)

Syntax: `ideal name = list_of_poly_and_ideal_expressions ;`
 `ideal name = ideal_expression ;`

Purpose: defines a left ideal.

Default: 0

Example:

```

ring r=0,(x,y,z),dp;
def R=nc_algebra(-1,0); // an anti-commutative algebra
setring R;
poly s1 = x2;
poly s2 = y3;
poly s3 = z;
ideal i = s1, s2-s1, 0,s3*s2, s3^4;
i;
↳ i[1]=x2
↳ i[2]=y3-x2
↳ i[3]=0
↳ i[4]=-y3z
↳ i[5]=z4
size(i);
↳ 4
ncols(i);
↳ 5

```

7.2.1.2 ideal expressions (plural)

An ideal expression is:

1. an identifier of type ideal
2. a function returning an ideal
3. a combination of ideal expressions by the arithmetic operations + or *
4. a power of an ideal expression (operator ^ or **)

Note that the computation of the product $i*i$ involves all products of generators of i while i^2 involves only the different ones, and is therefore faster.

5. a type cast to ideal

Example:

```

ring r=0,(x,y,z),dp;
def R=nc_algebra(-1,0); // an anticommutative algebra
setring R;
ideal m = maxideal(1);
m;
↳ m[1]=x
↳ m[2]=y
↳ m[3]=z
poly f = x2;
poly g = y3;
ideal i = x*y*z , f-g, g*(x-y) + f^4 ,0, 2x-z2y;
ideal M = i + maxideal(10);
i = M*M;
ncols(i);
↳ 598
i = M^2;
ncols(i);
↳ 690
i[ncols(i)];
↳ x20

```

```

vector v = [x,y-z,x2,y-x,x2yz2-y];
ideal j = ideal(v);
j;
↳ j[1]=x
↳ j[2]=y-z
↳ j[3]=x2
↳ j[4]=-x+y
↳ j[5]=x2yz2-y

```

7.2.1.3 ideal operations (plural)

- + addition (concatenation of the generators and simplification)
- * multiplication (with ideal, poly, vector, module; in case of multiplication with ideal or module, the result will be simplified)
- ^ exponentiation (by a non-negative integer)

ideal_expression [intvec_expression]

are polynomial generators of the ideal, index 1 gives the first generator.

Note: For simplification of an ideal, see also Section 5.1.141 [simplify], page 258.

Example:

```

ring r=0,(x,y,z),dp;
matrix D[3][3];
D[1,2]=-z; D[1,3]=y; D[2,3]=x;
def R=nc_algebra(1,D); // this algebra is U(so_3)
setring R;
ideal I = 0,x,0,1;
I;
↳ I[1]=0
↳ I[2]=x
↳ I[3]=0
↳ I[4]=1
I + 0; // simplification
↳ _[1]=1
I*x;
↳ _[1]=0
↳ _[2]=x2
↳ _[3]=0
↳ _[4]=x
ideal J = I,0,x,x-z;
I * J; // multiplication with simplification
↳ _[1]=1
vector V = [x,y,z];
print(I*V);
↳ 0,x2,0,x,
↳ 0,xy,0,y,
↳ 0,xz,0,z
ideal m = maxideal(1);
m^2;
↳ _[1]=x2
↳ _[2]=xy

```

```

↳ _[3]=xz
↳ _[4]=y2
↳ _[5]=yz
↳ _[6]=z2
ideal II = I[2..4];
II;
↳ II[1]=x
↳ II[2]=0
↳ II[3]=1

```

7.2.1.4 ideal related functions (plural)

dim	Gelfand-Kirillov dimension of basering modulo the ideal of leading terms (see Section 7.3.3 [dim (plural)], page 331)
eliminate	elimination of variables (see Section 7.3.5 [eliminate (plural)], page 333)
intersect	ideal intersection (see Section 7.3.9 [intersect (plural)], page 337)
kbase	vector space basis of basering modulo the leading ideal (see Section 7.3.10 [kbase (plural)], page 337)
lead	leading terms of a set of generators (see Section 5.1.75 [lead], page 206)
lift	lift-matrix (see Section 7.3.11 [lift (plural)], page 338)
liftstd	left Groebner basis and transformation matrix computation (see Section 7.3.12 [liftstd (plural)], page 339)
maxideal	generators of a power of the maximal ideal at 0 (see Section 5.1.88 [maxideal], page 216)
modulo	represents $(h1 + h2)/h1 \cong h2/(h1 \cap h2)$ (see Section 7.3.14 [modulo (plural)], page 341)
mres	minimal free resolution of an ideal and a minimal set of generators of the given ideal (see Section 7.3.15 [mres (plural)], page 342)
ncols	number of columns (see Section 5.1.103 [ncols], page 227)
nres	computes a free resolution of an ideal resp. module M which is minimized from the second free module on (see Section 7.3.18 [nres (plural)], page 345)
oppose	creates an opposite ideal of a given ideal from the given ring into a basering (see Section 7.3.19 [oppose], page 347)
preimage	preimage under a ring map (see Section 7.3.21 [preimage (plural)], page 349)
quotient	ideal quotient (see Section 7.3.22 [quotient (plural)], page 350)
reduce	left normal form with respect to a left Groebner basis (see Section 7.3.23 [reduce (plural)], page 351)
simplify	simplify a set of polynomials (see Section 5.1.141 [simplify], page 258)
size	number of non-zero generators (see Section 5.1.142 [size], page 259)
slimgb	left Groebner basis computation with slim technique (see Section 7.3.25 [slimgb (plural)], page 354)
std	left Groebner basis computation (see Section 7.3.26 [std (plural)], page 355)

subst	substitute a ring variable (see Section 7.3.27 [subst (plural)], page 357)
syz	computation of the first syzygy module (see Section 7.3.28 [syz (plural)], page 357)
twostd	two-sided Groebner basis computation (see Section 7.3.29 [twostd (plural)], page 358)
vdim	vector space dimension of basering modulo the leading ideal (see Section 7.3.30 [vdim (plural)], page 359)

7.2.2 map (plural)

Maps are ring maps from a preimage ring (source) into the basering (target), defined by specifying images for source variables in the target ring.

Note:

- the target of a map is **ALWAYS** the actual basering
- the preimage ring has to be stored "by its name", that means, maps can only be used in such contexts, where the name of the preimage ring can be resolved (this has to be considered in subprocedures). See also Section 6.5 [Identifier resolution], page 310, Section 3.7.4 [Names in procedures], page 54.

Maps between rings with different coefficient fields are possible and listed below.

Canonically realized are

- $Q \rightarrow Q(a, \dots)$ (Q : the rational numbers)
- $Q \rightarrow R$ (R : the real numbers)
- $Q \rightarrow C$ (C : the complex numbers)
- $Z/p \rightarrow (Z/p)(a, \dots)$ (Z : the integers)
- $Z/p \rightarrow GF(p^n)$ (GF : the Galois field)
- $Z/p \rightarrow R$
- $R \rightarrow C$

Possible are furthermore

- $Z/p \rightarrow Q$, $[i]_p \mapsto i \in [-p/2, p/2] \subseteq Z$
- $Z/p \rightarrow Z/p'$, $[i]_p \mapsto i \in [-p/2, p/2] \subseteq Z$, $i \mapsto [i]_{p'} \in Z/p'$
- $C \rightarrow R$, by taking the real part

Finally, in PLURAL we allow the mapping from rings with coefficient field Q to rings whose ground fields have finite characteristic:

- $Q \rightarrow Z/p$
- $Q \rightarrow (Z/p)(a, \dots)$

Note: In these cases the denominator and the numerator of a number are mapped separately by the usual map from Z to Z/p , and the image of the number is built again afterwards by division. It is thus not allowed to map numbers whose denominator is divisible by the characteristic of the target ground field, or objects containing such numbers. We, therefore, strongly recommend to use such maps only to map objects with integer coefficients.

Note that - in contrast to the commutative case - maps between non-commutative rings easily fail to be a morphism.

7.2.2.1 map declarations (plural)

Syntax: `map name = preimage_ring_name , ideal_expression ;`
`map name = preimage_ring_name , list_of_poly_and_ideal_expressions ;`
`map name = map_expression ;`

Purpose: defines a ring map from `preimage_ring` to basering.
 Maps the variables of the `preimage_ring` to the generators of the ideal.
 If the ideal contains less elements than the number of variables in the `preimage_ring`, the remaining variables are mapped to 0.
 If the ideal contains more elements, extra elements are ignored.
 The image ring is always the current basering. For the mapping of coefficients from different fields see Section 7.2.2 [map (plural)], page 317.

Default: none

Note: There are standard mappings for maps which are close to the identity map: `fetch (plural)` and `imap (plural)`.

The name of a map serves as the function which maps objects from the `preimage_ring` into the basering. These objects must be defined by names (no evaluation in the `preimage_ring` is possible).

Example:

```
// an easy example
ring r1 = 0,(a,b),dp; // a commutative ring
poly P = a^2+ab+b^3;
ring r2 = 0,(x,y),dp;
def W=nc_algebra(1,-1); // a Weyl algebra
setring W;
map M = r1, x^2, -y^3;
// note: M is just a map and not a morphism of K-algebras
M(P);
↳ -y9-x2y3+x4
// now, a more involved example
LIB "ncalg.lib";
def Usl2 = makeUsl2();
// this algebra is U(sl_2), generated by e,f,h
setring Usl2;
poly P = 4*e*f+h^2-2*h; // the central el-t of Usl2
poly Q = e^3*f-h^4; // some polynomial
ring W1 = 0,(D,X),dp;
def W2=nc_algebra(1,-1);
setring W2; // this is the opposite Weyl algebra
map F = Usl2, -X, D*D*X, 2*D*X;
F(P); // 0, because P is in the kernel of F
↳ 0
F(Q);
↳ -16D4X4+96D3X3-D2X4-112D2X2+6DX3+16DX-6X2
```

See Section 7.3.7 [fetch (plural)], page 335; Section 7.2.1.2 [ideal expressions (plural)], page 314; Section 7.3.8 [imap (plural)], page 336; Section 7.2.2 [map (plural)], page 317; Section 7.2.7 [ring (plural)], page 327.

7.2.2.2 map expressions (plural)

A map expression is:

1. an identifier of type map
2. a function returning map
3. a composition of maps using parentheses, e.g. $f(g)$

7.2.2.3 map (plural) operations

() composition of maps. If, for example, f and g are maps, then $f(g)$ is a map expression giving the composition $f \circ g$ of f and g , provided the target ring of g is the basering of f .

map_expression [int_expressions]
is a map entry (the image of the corresponding variable)

Example:

```
LIB "ncalg.lib";
def Us12 = makeUs12(); // this algebra is U(sl_2)
setring Us12;
map F = Us12, f, e, -h; // involutive endomorphism of U(sl_2)
F;
  ↪ F[1]=f
  ↪ F[2]=e
  ↪ F[3]=-h
map G = F(F);
G;
  ↪ G[1]=e
  ↪ G[2]=f
  ↪ G[3]=h
poly p = (f+e*h)^2 + 3*h-e;
P;
  ↪ e2h2+2e2h+2efh-2ef+f2-h2-e+3h
F(p);
  ↪ f2h2-2efh-2f2h+e2-2ef+h2-f-h
G(p);
  ↪ e2h2+2e2h+2efh-2ef+f2-h2-e+3h
(G(p) == p); // G is the identity
  ↪ 1
```

7.2.2.4 map related functions (plural)

fetch (plural)

the identity map between rings and q Rings (see Section 7.3.7 [fetch (plural)], page 335)

imap (plural)

a convenient map procedure for inclusions and projections of rings (see Section 7.3.8 [imap (plural)], page 336)

preimage (plural)

preimage under a ring map (see Section 7.3.21 [preimage (plural)], page 349)

subst substitute a ring variable (see Section 7.3.27 [subst (plural)], page 357)

See also Section 7.5.19 [ncpreim.lib], page 551 for the advanced preimage algorithm.

7.2.3 module (plural)

Modules are **left** submodules of a free module over the basering with basis `gen(1)`, `gen(2)`, \dots , `gen(n)` for some natural number `n`.

They are represented by lists of vectors, which generate the left submodule. Like vectors, they can only be defined or accessed with respect to a basering.

If M is a left submodule of R^n (where R is the basering) generated by vectors v_1, \dots, v_k , then these generators may be considered as the generators of relations of R^n/M between the canonical generators `gen(1)`, \dots , `gen(n)`. Hence, any finitely generated R -module can be represented in PLURAL by its module of relations. This is the so-called Coker-representation.

The assignments `module M=v1, ..., vk`; `matrix A=M`; create the presentation matrix of size $n \times k$, with the columns of A being the vectors v_1, \dots, v_k which generate M .

7.2.3.1 module declarations (plural)

Syntax: `module name = list_of_vector_expressions` (which are interpreted as left generators of the module) ;

`module name = module_expression` ;

Purpose: defines a left module.

Default: `[0]`

Example:

```
ring r=0,(x,y,z),(c,dp);
matrix D[3][3];
D[1,2]=-z; D[1,3]=y; D[2,3]=x;
def R=nc_algebra(1,D); // this algebra is U(so_3)
setring R;
vector s1 = [x2,y3,z];
vector s2 = [xy,1,0];
vector s3 = [0,x2-y2,z];
poly f = -x*y;
module m = s1, s2-s1,f*(s3-s1);
m;
↳ m[1]=[x2,y3,z]
↳ m[2]=[-x2+xy,-y3+1,-z]
↳ m[3]=[x3y-2x2z-xy,xy4-x3y+xy3+2x2z+xy]
// show m in matrix format (columns generate m)
print(m);
↳ x2,-x2+xy,x3y-2x2z-xy,
↳ y3,-y3+1, xy4-x3y+xy3+2x2z+xy,
↳ z, -z, 0
```

7.2.3.2 module expressions (plural)

A module expression is:

1. an identifier of type module
2. a function returning module
3. module expressions combined by the arithmetic operation `+`
4. multiplication of a module expression with an ideal or a poly expression: `*`
5. a type cast to module

7.2.3.3 module operations (plural)

+ addition (concatenation of the generators and simplification) Note that "-" implicitly converts a module into a matrix; see below example.

***** right or left multiplication with number, ideal, or poly (but not 'module' * 'module'!)

`module_expression [int_expression , int_expression]`

is a module entry, where the first index indicates the row and the second the column

`module_expressions [int_expression]`

is a vector, where the index indicates the column (generator)

Example:

```
ring A=0,(x,y,z),Dp;
matrix D[3][3];
D[1,2]=-z; D[1,3]=y; D[2,3]=x; // this algebra is U(so_3)
def B=nc_algebra(1,D);
setring B;
module M = [x,y],[0,0,x*z];
module N = matrix((x+y-z)*M) - matrix(M*(x+y-z)); // no - for type module
print(N);
↳ -y-z,0,
↳ -x+z,0,
↳ 0, -x2-xy-yz-z2
```

7.2.3.4 module related functions (plural)

eliminate

elimination of variables (see Section 7.3.5 [eliminate (plural)], page 333)

freemodule

the free module of given rank (see Section 5.1.47 [freemodule], page 186)

intersect

module intersection (see Section 7.3.9 [intersect (plural)], page 337)

kbase

vector space basis of free module over the basering modulo the module of leading terms (see Section 7.3.10 [kbase (plural)], page 337)

lead

initial module (see Section 5.1.75 [lead], page 206)

lift

lift-matrix (see Section 7.3.11 [lift (plural)], page 338)

liftstd

left Groebner basis and transformation matrix computation (see Section 7.3.12 [liftstd (plural)], page 339)

modulo

represents $(h_1 + h_2)/h_1 \cong h_2/(h_1 \cap h_2)$ (see Section 7.3.14 [modulo (plural)], page 341)

mres

minimal free resolution of a module and a minimal set of generators of the given ideal module (see Section 7.3.15 [mres (plural)], page 342)

ncols

number of columns (see Section 5.1.103 [ncols], page 227)

nres

computes a free resolution of an ideal resp. module M which is minimized from the second free module on (see Section 7.3.18 [nres (plural)], page 345)

nrows

number of rows (see Section 5.1.106 [nrows], page 228)

oppose	creates an opposite module of a given module from the given ring into a basering (see Section 7.3.19 [oppose], page 347)
print	nice print format (see Section 5.1.119 [print], page 238)
prune	minimize the embedding into a free module (see Section 5.1.121 [prune], page 241)
quotient	module quotient (see Section 7.3.22 [quotient (plural)], page 350)
reduce	left normal form with respect to a left Groebner basis (see Section 7.3.23 [reduce (plural)], page 351)
simplify	simplify a set of vectors (see Section 5.1.141 [simplify], page 258)
size	number of non-zero generators (see Section 5.1.142 [size], page 259)
std	left Groebner basis computation (see Section 7.3.26 [std (plural)], page 355)
subst	substitute a ring variable (see Section 7.3.27 [subst (plural)], page 357)
syz	computation of the first syzygy module (see Section 7.3.28 [syz (plural)], page 357)
vdim	vector space dimension of free module over the basering modulo module of leading terms (see Section 7.3.30 [vdim (plural)], page 359)

7.2.4 poly (plural)

Polynomials and vectors are the basic data for all main algorithms in PLURAL. Polynomials consist of finitely many terms (coefficient*monomial) which are combined by the usual polynomial operations (see Section 7.2.4.2 [poly expressions (plural)], page 323). Polynomials can only be defined or accessed with respect to a basering which determines the coefficient type, the names of the indeterminants and the monomial ordering.

Example:

```
ring r=32003,(x,y,z),dp;
poly f=x3+y5+z2;
```

Remark: Remember the conventions on polynomial multiplication we follow (*-multiplication in Section 7.1 [PLURAL], page 312).

7.2.4.1 poly declarations (plural)

Syntax: poly name = poly_expression ;

Purpose: defines a polynomial.

Default: 0

Example:

```
ring r = 32003,(x,y,z),dp;
def R=nc_algebra(-1,1);
setring R;
// ring of some differential-like operators
R;
↳ // coefficients: ZZ/32003
↳ // number of vars : 3
↳ //          block  1 : ordering dp
↳ //                   : names   x y z
↳ //          block  2 : ordering C
↳ // noncommutative relations:
```

```

↳ //    yx=-xy+1
↳ //    zx=-xz+1
↳ //    zy=-yz+1
yx;      // not correct input
↳ xy
y*x;     // correct input
↳ -xy+1
poly s1  = x3y2+151x5y+186xy6+169y9;
poly s2  = 1*x^2*y^2*z^2+3z8;
poly s3  = 5/4x4y2+4/5*x*y^5+2x2y2z3+y7+11x10;
int a,b,c,t=37,5,4,1;
poly f=3*x^a+x*y^(b+c)+t*x^a*y^b*z^c;
f;
↳ x37y5z4+3x37+xy9
short = 0;
f;
↳ x^37*y^5*z^4+3*x^37+xy^9

```

7.2.4.2 poly expressions (plural)

A polynomial expression is (optional parts in square brackets):

1. a monomial (there are NO spaces allowed inside a monomial)

```
[coefficient] ring_variable [exponent] [ring_variable [exponent] ...]
```

monomials which contain an indexed ring variable must be built from `ring_variable` and `coefficient` with the operations `*` and `^`

2. an identifier of type `poly`
3. a function returning `poly`
4. polynomial expressions combined by the arithmetic operations `+`, `-`, `*`, `/`, or `^`.
5. a type cast to `poly`

Example:

```

ring r=0,(x,y),dp;
def R=nc_algebra(1,1); // make it a Weyl algebra
setring R;
R;
↳ // coefficients: QQ
↳ // number of vars : 2
↳ //          block  1 : ordering dp
↳ //          : names  x y
↳ //          block  2 : ordering C
↳ // noncommutative relations:
↳ //    yx=xy+1
yx;      // not correct input
↳ xy
y*x;     // correct input
↳ xy+1
poly f = 10x2*y3 + 2y2*x^2 - 2*x*y + y - x + 2;
lead(f);
↳ 10x2y3
leadmonom(f);

```

```

↳ x2y3
simplify(f,1); // normalize leading coefficient
↳ x2y3+1/5x2y2+3/5xy-1/10x+1/10y+3/5
cleardenom(f);
↳ 10x2y3+2x2y2+6xy-x+y+6

```

7.2.4.3 poly operations (plural)

+ addition
- negation or subtraction
***** multiplication
/ commutative division by a monomial, non divisible terms yield 0
^, ** power by a positive integer
<, <=, >, >=, ==, <>
 comparison (of leading monomials w.r.t. monomial ordering)
poly_expression [intvec_expression]
 the sum of monomials at the indicated places w.r.t. the monomial ordering

7.2.4.4 poly related functions (plural)

bracket computes the (iterated) Lie bracket of two polynomials (see Section 7.3.2 [bracket], page 330)
lead leading term (see Section 5.1.75 [lead], page 206)
leadcoef coefficient of the leading term (see Section 5.1.76 [leadcoef], page 207)
leadexp the exponent vector of the leading monomial (see Section 5.1.77 [leadexp], page 207)
leadmonom
 leading monomial (see Section 5.1.78 [leadmonom], page 208)
oppose creates an opposite polynomial of a given polynomial from the given ring into a basering (see Section 7.3.19 [oppose], page 347)
reduce left normal form with respect to a left Groebner basis (see Section 7.3.23 [reduce (plural)], page 351)
simplify normalize a polynomial (see Section 5.1.141 [simplify], page 258)
size number of monomials (see Section 5.1.142 [size], page 259)
subst substitute a ring variable (see Section 7.3.27 [subst (plural)], page 357)
var the indicated variable of the ring (see Section 5.1.163 [var], page 279)

7.2.5 qring (plural)

PLURAL offers the possibility to compute within factor-rings modulo two-sided ideals. The ideal has to be given as a two-sided Groebner basis (see Section 7.3.29 [twostd (plural)], page 358 command).

For a detailed description of the concept of rings and quotient rings see Section 3.3 [Rings and orderings], page 30.

Note: we highly recommend to turn on `option(redSB); option(redTail);` while computing in qrings. Otherwise results may have a difficult interpretation.

7.2.5.1 qring declaration (plural)

Syntax: `qring name = ideal_expression ;`

Default: none

Purpose: declares a quotient ring as the basering modulo an `ideal_expression` and sets it as current basering.

Note: reports error if an ideal is not a two-sided Groebner basis.

Example:

```
ring r=0,(z,u,v,w),dp;
def R=nc_algebra(-1,0); // an anticommutative algebra
setring R;
option(redSB);
option(redTail);
ideal i=z^2,u^2,v^2,w^2, zuv-w;
qring Q = i; // incorrect call produces error
↳ // ** i is no standard basis
↳ // ** i is no twosided standard basis
kill Q;
setring R; // go back to the ring R
qring q=twostd(i); // now it is an exterior algebra modulo <zuv-w>
q;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ //          block 1 : ordering dp
↳ //          : names  z u v w
↳ //          block 2 : ordering C
↳ // noncommutative relations:
↳ //    uz=-zu
↳ //    vz=-zv
↳ //    wz=-zw
↳ //    vu=-uv
↳ //    wu=-uw
↳ //    wv=-vw
↳ // quotient ring from ideal
↳ _[1]=w2
↳ _[2]=vw
↳ _[3]=uw
↳ _[4]=zw
↳ _[5]=v2
↳ _[6]=u2
↳ _[7]=z2
↳ _[8]=zuv-w
poly k = (v-u)*(zv+u-w);
k; // the output is not yet totally reduced
↳ zuv-uv+uw-vw
poly ek=reduce(k,std(0));
ek; // the reduced form
↳ -uv+w
```

7.2.5.2 qring related functions (plural)

envelope enveloping ring (see Section 7.3.6 [envelope], page 334)
nvars number of ring variables (see Section 5.1.108 [nvars], page 229)
opposite opposite ring (see Section 7.3.20 [opposite], page 348)
setring set a new basering (see Section 5.1.139 [setring], page 255)

7.2.6 resolution (plural)

The type resolution is intended as an intermediate representation which internally retains additional information obtained during computation of resolutions. It furthermore enables the use of partial results to compute, for example, Betti numbers or minimal resolutions. Like ideals and modules, a resolution can only be defined w.r.t. a basering.

Note: to access the elements of a resolution, it has to be assigned to a list. This assignment also completes computations and may therefore take time, (resp. an access directly with the brackets [,] causes implicitly a cast to a list).

7.2.6.1 resolution declarations (plural)

Syntax: resolution name = resolution_expression ;

Purpose: defines a resolution.

Default: none

Example:

```
ring r=0,(x,y,z),dp;
matrix D[3][3];
D[1,2]=z;
def R=nc_algebra(1,D); // it is a Heisenberg algebra
setring R;
ideal i=z2+z,x+y;
resolution re=nres(i,0);
re;
↳ 1      2      1
↳ R <--  R <--  R
↳
↳ 0      1      2
↳ resolution not minimized yet
↳
list l = re;
l;
↳ [1]:
↳   _[1]=z2+z
↳   _[2]=x+y
↳ [2]:
↳   _[1]=z2*gen(2)-x*gen(1)-y*gen(1)+z*gen(2)
↳ [3]:
↳   _[1]=0
print(matrix(1[2]));
↳ -x-y,
↳ z2+z
print(module(transpose(matrix(1[2]))*transpose(matrix(1[1])))); // check
↳ 0
```


7.2.6.2 resolution expressions (plural)

A resolution expression is:

1. an identifier of type resolution
2. a function returning a resolution
3. a type cast to resolution from a list of ideals, resp. modules.

7.2.6.3 resolution related functions (plural)

betti	Betti numbers of a resolution (see Section 7.3.1 [betti (plural)], page 329)
minres	minimizes a free resolution (see Section 7.3.13 [minres (plural)], page 340)
mres	computes a minimal free resolution of an ideal resp. module and a minimal set of generators of the given ideal resp. module (see Section 7.3.15 [mres (plural)], page 342)
nres	computes a free resolution of an ideal resp. module M which is minimized from the second module on (see Section 7.3.18 [nres (plural)], page 345)

7.2.7 ring (plural)

Rings are used to describe properties of polynomials, ideals etc. Almost all computations in PLURAL require a basering. For a detailed description of the concept of rings see Section 3.3 [Rings and orderings], page 30.

Note: PLURAL usually works with global orderings (see Section 7.1 [PLURAL], page 312) but one can use certain local once when graded commutative rings are being used.

7.2.7.1 ring declarations (plural)

Syntax: `ring name = (coefficient_field), (names_of_ring_variables), (ordering);`

Default: `32003, (x,y,z), (dp,C);`

Purpose: declares a ring and sets it as the actual basering.

The `coefficient_field` is given by one of the following:

1. a non-negative `int_expression` less or equal 2147483647.
2. an `expression_list` of an `int_expression` and one or more names.
3. the name `real`.
4. an `expression_list` of the name `real` and an `int_expression`.
5. an `expression_list` of the name `complex`, an optional `int_expression` and a name.

'`names_of_ring_variables`' must be a list of names or indexed names.

'`ordering`' is a list of block orderings where each block ordering is either

1. `lp`, `dp`, `Dp`, optionally followed by a size parameter in parentheses.
2. `wp`, `Wp`, or `a` followed by a weight vector given as an `intvec_expression` in parentheses.
3. `M` followed by an `intmat_expression` in parentheses.
4. `c` or `C`.

As long as all non-commuting variables are global, any ordering may be used. In graded commutative algebras, one may also use `ls`, `ds`, `Ds`, `ws`, and `Ws`.

If one of `coefficient_field`, `names_of_ring_variables`, and `ordering` consists of only one entry, the parentheses around this entry may be omitted.

In order to create a non-commutative structure over a commutative ring, use Section 7.3.16 [`nc_algebra`], page 343.

7.2.7.2 ring operations (plural)

+ construct a tensor product $C = A \otimes_{\mathbf{K}} B$ of two G -algebras A and B over the ground field. Let, e.g.,

$$A = k_1 \langle x_1, \dots, x_n \mid \{x_j x_i = c_{ij} \cdot x_i x_j + d_{ij}\}, 1 \leq i < j \leq n \rangle, \text{ and } B = k_2 \langle y_1, \dots, y_m \mid \{y_j y_i = q_{ij} \cdot y_i y_j + r_{ij}\}, 1 \leq i < j \leq m \rangle$$

be two G -algebras, then C is defined to be the algebra

$$C = K \langle x_1, \dots, x_n, y_1, \dots, y_m \mid \{x_j x_i = c_{ij} \cdot x_i x_j + d_{ij}, 1 \leq i < j \leq n\}, \{y_j y_i = q_{ij} \cdot y_i y_j + r_{ij}, 1 \leq i < j \leq m\}, \{y_j x_i = x_i y_j, 1 \leq j \leq m, 1 \leq i \leq n\} \rangle.$$

Concerning the ground fields k_1 resp. k_2 of A resp. B , take the following guidelines for $A \otimes_{\mathbf{K}} B$ into consideration:

- Neither k_1 nor k_2 may be R or C .
- If the characteristic of k_1 and k_2 differs, then one of them must be Q .
- At most one of k_1 and k_2 may have parameters.
- If one of k_1 and k_2 is an algebraic extension of Z/p it may not be defined by a `charstr` of type (p^n, a) .

One can create a ring using `ring(list)`, see also `ringlist`.

Example:

```
LIB "ncalg.lib";
def a = makeUs12();           // U(sl_2) in e,f,h presentation
ring W0 = 0, (x,d), dp;
def W = Weyl();              // 1st Weyl algebra in x,d
def S = a+W;
setring S;
S;
⇨ // coefficients: QQ
⇨ // number of vars : 5
⇨ //      block 1 : ordering dp
⇨ //                : names e f h
⇨ //      block 2 : ordering dp
⇨ //                : names x d
⇨ //      block 3 : ordering C
⇨ // noncommutative relations:
⇨ //      fe=ef-h
⇨ //      he=eh+2e
⇨ //      hf=fh-2f
⇨ //      dx=xd+1
```

7.2.7.3 ring related functions (plural)

`charstr` description of the coefficient field of a ring (see Section 5.1.7 [`charstr`], page 160)

envelope	enveloping ring (see Section 7.3.6 [envelope], page 334)
npars	number of ring parameters (see Section 5.1.104 [npars], page 227)
nvars	number of ring variables (see Section 5.1.108 [nvars], page 229)
opposite	opposite ring (see Section 7.3.20 [opposite], page 348)
ordstr	monomial ordering of a ring (see Section 5.1.112 [ordstr], page 235)
parstr	names of all ring parameters or the name of the n-th ring parameter (see Section 5.1.115 [parstr], page 236)
qring	quotient ring (see Section 7.2.5 [qring (plural)], page 324)
ringlist	decomposes a ring into a list of its components (see Section 7.3.24 [ringlist (plural)], page 352)
setring	set a new basering (see Section 5.1.139 [setring], page 255)
varstr	names of all ring variables or the name of the n-th ring variable (see Section 5.1.165 [varstr], page 280)

7.3 Functions (plural)

This chapter gives a complete reference of all functions and commands of the PLURAL kernel, i.e. all built-in commands (for the PLURAL libraries see Section 7.5 [PLURAL libraries], page 365).

The general syntax of a function is

```
[target =] function_name (<arguments>);
```

Note, that both **Control structures** and **System variables** of PLURAL are the same as of SINGULAR (see Section 5.2 [Control structures], page 285, Section 5.3 [System variables], page 297).

7.3.1 betti (plural)

Syntax: `betti (list_expression)`
`betti (resolution_expression)`
`betti (list_expression , int_expression)`
`betti (resolution_expression , int_expression)`

Type: intmat

Note: in the non-commutative case, computing Betti numbers makes sense only if the basering R has homogeneous relations. The output of the command can be pretty-printed using `print(, 'betti')`, i.e., with "betti" as second argument; see below example.

Purpose: with 1 argument: computes the graded Betti numbers of a minimal resolution of R^n/M , if R denotes the basering and M a homogeneous submodule of R^n and the argument represents a resolution of R^n/M .

The entry d of the intmat at place (i, j) is the minimal number of generators in degree $i+j$ of the j -th syzygy module (= module of relations) of R^n/M (the 0th (resp. 1st) syzygy module of R^n/M is R^n (resp. M)). The argument is considered to be the result of a `mres` or `nres` command. This implies that a zero is only allowed (and counted) as a generator in the first module.

For the computation `betti` uses only the initial monomials. This could lead to confusing results for a non-homogeneous input.

If the optional second argument is non-zero, the Betti numbers will be minimized.

Example:

```

int i;int N=2;
ring r=0,(x(1..N),d(1..N),q(1..N)),Dp;
matrix D[3*N][3*N];
for (i=1;i<=N;i++)
{ D[i,N+i]=q(i)^2; }
def W=nc_algebra(1,D); setring W;
// this algebra is a kind of homogenized Weyl algebra
W;
↳ // coefficients: QQ
↳ // number of vars : 6
↳ //      block 1 : ordering Dp
↳ //      : names x(1) x(2) d(1) d(2) q(1) q(2)
↳ //      block 2 : ordering C
↳ // noncommutative relations:
↳ //      d(1)x(1)=x(1)*d(1)+q(1)^2
↳ //      d(2)x(2)=x(2)*d(2)+q(2)^2
ideal I = x(1),x(2),d(1),d(2),q(1),q(2);
option(redSB);
option(redTail);
resolution R = mres(I,0);
// thus R will be the full length minimal resolution
print(betti(R),"betti");
↳
↳      0      1      2      3      4      5      6
↳ -----
↳ 0:      1      6     15     20     15      6      1
↳ -----
↳ total:   1      6     15     20     15      6      1
↳

```

7.3.2 bracket

Syntax: bracket (poly_expression, poly_expression)
 bracket (poly_expression, poly_expression, int_expression)

Type: poly

Purpose: Computes the Lie bracket $[p, q] = pq - qp$ of the first polynomial with the second. Uses special routines, based on the Leibniz rule. If the third argument is $N > 1$, then the right normed bracket $[a, [\dots [a, b]]]$ will be computed.

Note: effective both with PLURAL and LETTERPLACE rings.

Example:

```

ring r=(0,Q),(x,y,z),Dp; // first, let us do a Plural example
minpoly=Q^2-Q+1;
matrix C[3][3]; matrix D[3][3];
C[1,2]=Q2; C[1,3]=1/Q2; C[2,3]=Q2;
D[1,2]=-Q*z; D[1,3]=1/Q*y; D[2,3]=-Q*x;
def R=nc_algebra(C,D); setring R; R;
↳ // coefficients: QQ[Q]/(Q2-Q+1)
↳ // number of vars : 3
↳ //      block 1 : ordering Dp
↳ //      : names x y z

```

```

↳ //          block 2 : ordering C
↳ // noncommutative relations:
↳ //    yx=(Q-1)*xy+(-Q)*z
↳ //    zx=(-Q)*xz+(-Q+1)*y
↳ //    zy=(Q-1)*yz+(-Q)*x
// this is a quantum deformation of U(so_3),
// where Q is a 6th root of unity
poly p=Q^4*x2+y2+Q^4*z2+Q*(1-Q^4)*x*y*z;
// p is the central element of the algebra
p=p^3; // any power of a central element is central
poly q=(x+Q*y+Q^2*z)^4;
// take q to be some big noncentral element
size(q); // check how many monomials are in big polynomial q
↳ 28
bracket(p,q); // check p*q=q*p
↳ 0
// a more common behaviour of the bracket follows:
bracket(x+Q*y+Q^2*z,z);
↳ (Q+1)*xz+(Q+1)*yz+(Q-1)*x+(Q-1)*y
kill R; setring r; // Now consider an example for Letterplace
LIB "freegb.lib";
ring R = freeAlgebra(r,5); // F<x,y,z> with deg left lex ordering
bracket(x,y);
↳ x*y-y*x
bracket(x,y,2);
↳ x*x*y-2*x*y*x+y*x*x
bracket(x,y,3);
↳ x*x*x*y-3*x*x*y*x+3*x*y*x*x-y*x*x*x
bracket(z^2,x+Q*y,2);
↳ x*z*z*z*z+(Q)*y*z*z*z*z-2*z*z*x*z*z+(-2*Q)*z*z*y*z*z+z*z*z*z*x+(Q)*z*z*
z*y

```

7.3.3 dim (plural)

Syntax: dim (ideal_expression)
dim (module_expression)

Type: int

Purpose: computes the Gelfand-Kirillov dimension of the ideal, resp. module, generated by the leading monomials of the given generators of the ideal, resp. module. This is also the dimension of the ideal resp. submodule, if it is represented by a left Groebner basis.

Note: The dimension of a submodule of a free module is defined to be the Gelfand-Kirillov dimension of the left module with the presentation via given submodule.
The computed Gelfand-Kirillov dimension is taken relative to the ground field. In order to compute the complete Gelfand-Kirillov dimension, one has to add the transcendence degree of the ground field over its prime field.

Example:

```

ring r=0,(x,y,Dx,Dy),dp;
matrix M[4][4]; M[1,3]=1;M[2,4]=1;
def R = nc_algebra(1,M); // 2nd Weyl algebra
setring R;

```

```

    dim(std(0)); // the GK dimension of the ring itself
    ↪ 4
    ideal I=x*Dy^2-2*y*Dy^2+2*Dy, Dx^3+3*Dy^2;
    dim(std(I)); // the GK dimension of the module R/I
    ↪ 2
    module T = (x*Dx -2)*gen(1), Dx^3*gen(1), (y*Dy +3)*gen(2);
    dim(std(T)); // the GK dimension of the module R^2/T
    ↪ 3

```

See Section 4.5 [ideal], page 78; Section 4.13 [module], page 110; Section 5.1.149 [std], page 266; Section 5.1.166 [vdim], page 281.

7.3.4 division (plural)

Syntax: `division (ideal-expression, ideal-expression)`
`division (module-expression, module-expression)`
`division (ideal-expression, ideal-expression, int-expression)`
`division (module-expression, module-expression, int-expression)`
`division (ideal-expression, ideal-expression, int-expression, intvec-expression)`
`division (module-expression, module-expression, int-expression, intvec-expression)`

Type: list

Purpose: `division` computes a left division with remainder. For two left ideals resp. modules M (first argument) and N (second argument), it returns a list T,R,U where T is a matrix, R is a left ideal resp. a module, and U is a diagonal matrix of units such that $\text{transpose}(U)*\text{transpose}(\text{matrix}(M))=\text{transpose}(T)*\text{transpose}(\text{matrix}(N)) + \text{transpose}(\text{matrix}(R))$. From this data one gets a left standard representation for the left normal form R of M with respect to a left Groebner basis of N . `division` uses different algorithms depending on whether N is represented by a Groebner basis. For a GR-algebra, the matrix U is the identity matrix. A matrix T as above is also computed by `lift`.

For additional arguments n (third argument) and w (fourth argument), `division` returns a list T,R as above such that $\text{transpose}(\text{matrix}(M))=\text{transpose}(T)*\text{transpose}(\text{matrix}(N)) + \text{transpose}(\text{matrix}(R))$ is a left standard representation for the left normal form R of M with respect to N up to weighted degree n with respect to the weight vector w . The weighted degree of T and R respect to w is at most n . If the weight vector w is not given, `division` uses the standard weight vector $w=1, \dots, 1$.

Example:

```

LIB "dmod.lib";
ring r = 0,(x,y),dp;
poly f = x^3+xy;
def S = Sannfs(f); setring S; // compute the annihilator of f^s
LD; // is not a Groebner basis yet!
↪ LD[1]=3*x^2*Dy-x*Dx+y*Dy
↪ LD[2]=x*Dx+2*y*Dy-3*s
poly f = imap(r,f);
poly P = f*Dx-s*diff(f,x);
division(P,LD); // so P is in the ideal via the cofactors in _[1]
↪ [1]:

```

```

↳   _[1,1]=-2/3*y
↳   _[2,1]=x^2+1/3*y
↳ [2]:
↳   _[1]=0
↳ [3]:
↳   _[1,1]=1
ideal I = LD, f; // consider a bigger ideal
list L = division(s^2, I); // the normal form is -2s-1
L;
↳ [1]:
↳   _[1,1]=2/3*x^2*Dy-1/3*x*Dx+2/3*s+1/3
↳   _[2,1]=2/3*x^2*Dy-1/3*x*Dx-1/3*s-2/3
↳   _[3,1]=-2*x*Dy^2+Dx*Dy
↳ [2]:
↳   _[1]=-2*s-1
↳ [3]:
↳   _[1,1]=1
// now we show that the formula above holds
matrix M[1][1] = s^2; matrix N = matrix(I);
matrix T = matrix(L[1]); matrix R = matrix(L[2]); matrix U = matrix(L[3])
// the formula must return zero:
transpose(U)*transpose(M) - transpose(T)*transpose(N) - transpose(R);
↳ _[1,1]=0

```

See Section 4.5 [ideal], page 78; Section 5.1.80 [lift], page 209; Section 4.13 [module], page 110; Section 4.16 [poly], page 117; Section 4.22 [vector], page 131.

7.3.5 eliminate (plural)

Syntax: `eliminate (ideal_expression, product_of_ring_variables)`
`eliminate (module_expression, product_of_ring_variables)`

Type: the same as the type of the first argument

Purpose: eliminates variables occurring as factors of the second argument from an ideal (resp. a submodule of a free module), by intersecting it (resp. each component of the submodule) with the subring not containing these variables.

Note: `eliminate` neither needs a special ordering on the basering nor a Groebner basis as input. Moreover, `eliminate` does not work in non-commutative quotients.

Remark: in a non-commutative algebra, not every subset of a set of variables generates a proper subalgebra. But if it is so, there may be cases, when no elimination (by means of Groebner bases) is possible; in such situations error messages will be reported. See also Section 7.5.19 [ncpreim.lib], page 551 for the advanced algorithm for elimination and preimage.

Example:

```

ring r=0,(e,f,h,a),Dp;
matrix d[4][4];
d[1,2]=-h; d[1,3]=2*e; d[2,3]=-2*f;
def R=nc_algebra(1,d); setring R;
// this algebra is U(sl_2), tensored with K[a] over K
option(redSB);
option(redTail);

```

```

poly p = 4*e*f+h^2-2*h - a;
// p is a central element with parameter
ideal I = e^3, f^3, h^3-4*h, p; // take this ideal
// and intersect I with the ring K[a]
ideal J = eliminate(I,e*f*h);
// if we want substitute 'a' with a value,
// it has to be a root of this polynomial
J;
↳ J[1]=a3-32a2+192a
// now we try to eliminate h,
// that is we intersect I with the subalgebra S,
// generated by e and f.
// But S is not closed in itself, since f*e-e*f=-h !
// the next command will definitely produce an error
eliminate(I,h);
↳ ? no elimination is possible: subalgebra is not admissible
↳ ? error occurred in or before ./examples/eliminate_(plural).sing 1\
   ine 21: 'eliminate(I,h)';
// since a commutes with e,f,h, we can eliminate it:
eliminate(I,a);
↳ _[1]=h3-4h
↳ _[2]=fh2-2fh
↳ _[3]=f3
↳ _[4]=eh2+2eh
↳ _[5]=2efh-h2-2h
↳ _[6]=e3

```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.2.3 [module (plural)], page 320; Section 7.3.26 [std (plural)], page 355.

7.3.6 envelope

Syntax: envelope (ring_name)

Type: ring

Purpose: creates an enveloping algebra of a given algebra, that is $A^{env} = A \otimes_K A^{opp}$, where A^{opp} is the opposite algebra of A .

Remark: You have to activate the ring with the `setring` command. For the presentation, see explanation of opposite in Section 7.3.20 [opposite], page 348.

```

LIB "ncalg.lib";
def A = makeUsl2();
setring A; A;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ //          block  1 : ordering dp
↳ //                   : names    e f h
↳ //          block  2 : ordering C
↳ // noncommutative relations:
↳ //    fe=ef-h
↳ //    he=eh+2e
↳ //    hf=fh-2f
def Aenv = envelope(A);

```



```

setring Aenv;
Aenv;
↳ // coefficients: QQ
↳ // number of vars : 6
↳ //      block  1 : ordering dp
↳ //                : names   e f h
↳ //      block  2 : ordering a
↳ //                : names   H F E
↳ //                : weights 1 1 1
↳ //      block  3 : ordering ls
↳ //                : names   H F E
↳ //      block  4 : ordering C
↳ // noncommutative relations:
↳ //      fe=ef-h
↳ //      he=eh+2e
↳ //      hf=fh-2f
↳ //      FH=HF-2F
↳ //      EH=HE+2E
↳ //      EF=FE-H

```

See Section 7.3.19 [oppose], page 347; Section 7.3.20 [opposite], page 348.

7.3.7 fetch (plural)

Syntax: `fetch (ring_name, name)`

Type: number, poly, vector, ideal, module, matrix or list (the same type as the second argument)

Purpose: maps objects between rings. `fetch` is the identity map between rings and qings, the i -th variable of the source ring is mapped to the i -th variable of the basering. The coefficient fields must be compatible. (See Section 7.2.2 [map (plural)], page 317 for a description of possible mappings between different ground fields). `fetch` offers a convenient way to change variable names or orderings, or to map objects from a ring to a quotient ring of that ring or vice versa.

Note: Compared with `imap`, `fetch` uses the position of the ring variables, not their names.

Example:

```

LIB "ncalg.lib";
def Us12 = makeUs12(); // this algebra is U(sl_2)
setring Us12;
option(redSB);
option(redTail);
poly C = 4*e*f+h^2-2*h; // the central element of Us12
ideal I = e^3,f^3,h^3-4*h;
ideal J = twostd(I);
// print a compact presentation of J:
print(matrix(ideal(J[1..5]))); // first 5 generators
↳ h3-4h,fh2-2fh,eh2+2eh,f2h-2f2,2efh-h2-2h
print(matrix(ideal(J[6..size(J)]))); // last generators
↳ e2h+2e2,f3,ef2-fh,e2f-eh-2e,e3
ideal QC = twostd(C-8);
qring Q = QC;
ideal QJ = fetch(Us12,J);

```

```

QJ = std(QJ);
// thus QJ is the image of I in the factor-algebra QC
print(matrix(QJ)); // print QJ compactly
↳ h3-4h,fh2-2fh,eh2+2eh,f2h-2f2,e2h+2e2,f3,e3

```

See Section 7.3.8 [imap (plural)], page 336; Section 7.2.2 [map (plural)], page 317; Section 7.2.5 [qring (plural)], page 324; Section 7.2.7 [ring (plural)], page 327.

7.3.8 imap (plural)

Syntax: `imap (ring_name, name)`

Type: number, poly, vector, ideal, module, matrix or list (the same type as the second argument)

Purpose: identity map on common subrings. `imap` is the map between rings and qrings with compatible ground fields which is the identity on variables and parameters of the same name and 0 otherwise. (See Section 7.2.2 [map (plural)], page 317 for a description of possible mappings between different ground fields). Useful for mappings from a homogenized ring to the original ring or for mappings from/to rings with/without parameters. Compared with `fetch`, `imap` uses the names of variables and parameters. **Unlike map and fetch, imap can map parameters to variables.**

Example:

```

LIB "ncalg.lib";
ring ABP=0,(p4,p5,a,b),dp; // a commutative ring
def Usl3 = makeUsl(3);
def BIG = Usl3+ABP;
setring BIG;
poly P4 = 3*x(1)*y(1)+3*x(2)*y(2)+3*x(3)*y(3);
P4 = P4 +h(1)^2+h(1)*h(2)+h(2)^2-3*h(1)-3*h(2);
// P4 is a central element of Usl3 of degree 2
poly P5 = 4*x(1)*y(1) + h(1)^2 - 2*h(1);
// P5 is a central element of the subalgebra of U(sl_3),
// generated by x(1),y(1),h(1)
ideal J = x(1),x(2),h(1)-a,h(2)-b;
// we are interested in the module U(sl_3)/J,
// which depends on parameters a,b
ideal I = p4-P4, p5-P5;
ideal K = I, J;
ideal E = eliminate(K,x(1)*x(2)*x(3)*y(1)*y(2)*y(3)*h(1)*h(2));
E; // this is the ideal of central characters in ABP
↳ E[1]=a*b+b^2-p4+p5+a+3*b
↳ E[2]=a^2-p5+2*a
↳ E[3]=b^3+p4*a-p5*a-a^2-p4*b+3*b^2
// what are the characters on nonzero a,b?
ring abP = (0,a,b),(p4,p5),dp;
ideal abE = imap(BIG, E);
option(redSB);
option(redTail);
abE = std(abE);
// here come characters (indeed, we have only one)
// that is a maximal ideal in K[p4,p5]
abE;

```

```

↳ abE[1]=p5+(-a^2-2*a)
↳ abE[2]=p4+(-a^2-a*b-3*a-b^2-3*b)

```

See Section 7.3.7 [fetch (plural)], page 335; Section 7.2.2 [map (plural)], page 317; Section 7.2.5 [qring (plural)], page 324; Section 7.2.7 [ring (plural)], page 327.

7.3.9 intersect (plural)

Syntax: `intersect (expression_list of ideal_expression)`
 `intersect (expression_list of module_expression)`

Type: ideal, resp. module

Purpose: computes the intersection of ideals, resp. modules.

Example:

```

ring r=0,(x,y),dp;
def R=nc_algebra(-1,0); //anti-commutative algebra
setring R;
module M=[x,x],[y,0];
module N=[0,y^2],[y,x];
option(redSB);
module Res;
Res=intersect(M,N);
print(Res);
↳ y2, 0,
↳ -xy,xy2
kill r,R;
//-----
LIB "ncalg.lib";
ring r=0,(x,d),dp;
def RR=Weyl(); // make r into Weyl algebra
setring RR;
ideal I = x+d^2;
ideal J = d-1;
ideal H = intersect(I,J);
H;
↳ H[1]=d4+xd2-2d3-2xd+d2+x+2d-2
↳ H[2]=xd3+x2d-xd2+d3-x2+xd-2d2-x+1

```

7.3.10 kbase (plural)

Syntax: `kbase (ideal_expression)`
 `kbase (module_expression)`
 `kbase (ideal_expression, int_expression)`
 `kbase (module_expression, int_expression)`

Type: the same as the input type of the first argument

Purpose: with one argument: computes the vector space basis of the factor-module that equals ring (resp. free module) modulo the ideal (resp. submodule), generated by the initial terms of the given generators.

If the factor-module is not of finite dimension, -1 is returned.

If the generators form a Groebner basis, this is the same as the vector space basis of the factor-module.

when called with two arguments: computes the part of a vector space basis of the respective quotient with degree (of monomials) equal to the second argument. Here, the quotient does not need to be finite dimensional.

Note: in the non-commutative case, a ring modulo an ideal has a ring structure if and only if the ideal is two-sided.

Also, `kbase` respects module grading given by the `isHomog` attribute of input modules.

Example:

```
ring r=0,(x,y,z),dp;
matrix d[3][3];
d[1,2]=-z; d[1,3]=2x; d[2,3]=-2y;
def R=nc_algebra(1,d); // this algebra is U(sl_2)
setring R;
ideal i=x^2,y^2,z^2-1;
i=std(i);
print(matrix(i)); // print a compact presentation of i
↳ z^2-1,yz-y,xz+x,y^2,2xy-z-1,x^2
kbase(i);
↳ _[1]=z
↳ _[2]=y
↳ _[3]=x
↳ _[4]=1
vdim(i);
↳ 4
ideal j=x,z-1;
j=std(j);
kbase(j,3);
↳ _[1]=y^3
```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.2.3 [module (plural)], page 320; Section 7.3.30 [vdim (plural)], page 359.

7.3.11 lift (plural)

Syntax: `lift (ideal_expression, subideal_expression)`
`lift (module_expression, submodule_expression)`

Type: matrix

Purpose: computes the (left) transformation matrix which expresses the (left) generators of a submodule in terms of the (left) generators of a module. Uses different algorithms for modules which are (resp. are not) represented by Groebner bases.

More precisely, if `m` is the module, `sm` the submodule, and `T` the transformation matrix returned by `lift`, then `transpose(matrix(sm)) = transpose(T)*transpose(matrix(m))`.

If `m` and `sm` are ideals, `ideal(sm) = ideal(transpose(T)*transpose(matrix(m)))`.

Note: Gives a warning if `sm` is not a submodule.

Example:

```
ring r = (0,a),(e,f,h),(c,dp);
matrix D[3][3];
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
```

```

def R=nc_algebra(1,D); // this algebra is a parametric U(sl_2)
setring R;
ideal I = e,h-a; // consider this parametric ideal
I = std(I); // left Groebner basis
print(matrix(I)); // print a compact presentation of I
↳ h+(-a),e
poly Z = 4*e*f+h^2-2*h; // a central element in R
Z = Z - NF(Z,I); // a central character
ideal j = std(Z);
j;
↳ j[1]=4*ef+h2-2*h+(-a2-2a)
matrix T = lift(I,j);
print(T);
↳ h+(a+2),
↳ 4*f
ideal tj = ideal(transpose(T)*transpose(matrix(I)));
size(ideal(matrix(j)-matrix(tj))); // test for 0
↳ 0

```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.3.12 [liftstd (plural)], page 339; Section 7.2.3 [module (plural)], page 320.

7.3.12 liftstd (plural)

Syntax: liftstd (ideal_expression, matrix_name)
liftstd (module_expression, matrix_name)
liftstd (ideal_expression, matrix_name, module_name)
liftstd (module_expression, matrix_name, module_name)

Type: ideal or module

Purpose: returns a left Groebner basis of an ideal or module and a left transformation matrix from the given ideal, resp. module, to the Groebner basis.

That is, if m is the ideal or module, sm is the left Groebner basis of m , returned by `liftstd`, and T is a left transformation matrix, then $sm=module(transpose(transpose(T)*transpose(matrix(m))))$.

If m is an ideal, $sm=ideal(transpose(T)*transpose(matrix(m)))$.

In an optional third argument the left syzygy module will be returned.

Example:

```

LIB "ncalg.lib";
def A = makeUsl2();
setring A; // this algebra is U(sl_2)
ideal i = e2,f;
option(redSB);
option(redTail);
matrix T;
ideal j = liftstd(i,T);
// the Groebner basis in a compact form:
print(matrix(j));
↳ f,2h2+2h,2eh+2e,e2
print(T); // the transformation matrix
↳ 0,f2, -f,1,
↳ 1,-e2f+4eh+8e,e2,0

```

```

ideal tj = ideal(transpose(T)*transpose(matrix(i)));
size(ideal(matrix(j)-matrix(tj))); // test for 0
↳ 0
module S; ideal k = liftstd(i,T,S); // the third argument
S = std(S); print(S); // the syzygy module
↳ -ef-2h+6,-f3,
↳ e3,      e2f2-6efh-6ef+6h2+18h+12

```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.2.7 [ring (plural)], page 327; Section 7.3.26 [std (plural)], page 355.

7.3.13 minres (plural)

Syntax: minres (list_expression)

Type: list

Syntax: minres (resolution_expression)

Type: resolution

Purpose: minimizes a free resolution of an ideal or module given by the list_expression, resp. resolution_expression.

Example:

```

LIB "ncalg.lib";
def A = makeUs12();
setring A; // this algebra is U(sl_2)
ideal i=e,f,h;
i=std(i);
resolution F=nres(i,0); F;
↳ 1      3      3      1
↳ A <--  A <--  A <--  A
↳
↳ 0      1      2      3
↳ resolution not minimized yet
↳
list lF = F; lF;
↳ [1]:
↳  _[1]=h
↳  _[2]=f
↳  _[3]=e
↳ [2]:
↳  _[1]=f*gen(1)-h*gen(2)-2*gen(2)
↳  _[2]=e*gen(1)-h*gen(3)+2*gen(3)
↳  _[3]=e*gen(2)-f*gen(3)-gen(1)
↳ [3]:
↳  _[1]=e*gen(1)-f*gen(2)+h*gen(3)
print(betti(lF), "betti");
↳          0      1      2      3
↳ -----
↳  0:      1      -      3      1
↳ -----
↳ total:   1      0      3      1
↳
resolution MF=minres(F); MF;

```

```

↳ 1      2      2      1
↳ A <--  A <--  A <--  A
↳
↳ 0      1      2      3
↳
list LMF = F; LMF;
↳ [1]:
↳   _[1]=f
↳   _[2]=e
↳ [2]:
↳   _[1]=-ef*gen(1)+f2*gen(2)+2h*gen(1)+2*gen(1)
↳   _[2]=-e2*gen(1)+ef*gen(2)+h*gen(2)-2*gen(2)
↳ [3]:
↳   _[1]=e*gen(1)-f*gen(2)
print(betti(LMF), "betti");
↳           0      1      2      3
↳ -----
↳    0:      1      -      -      -
↳    1:      -      -      2      1
↳ -----
↳ total:      1      0      2      1
↳

```

See Section 7.3.15 [mres (plural)], page 342; Section 7.3.18 [nres (plural)], page 345.

7.3.14 modulo (plural)

Syntax: modulo (ideal-expression, ideal-expression)
 modulo (module-expression, module-expression)

Type: module

Purpose: modulo(h1,h2) represents $h_1/(h_1 \cap h_2) \cong (h_1 + h_2)/h_2$, where h_1 and h_2 are considered as submodules of the same free module R^l ($l=1$ for ideals).
 Let H_1 (resp. H_2) be the matrix of size $l \times k$ (resp. $l \times m$), having the generators of h_1 (resp. h_2) as columns.
 Then $h_1/(h_1 \cap h_2) \cong R^k / \ker(\overline{H}_1)$, where $\overline{H}_1 : R^k \rightarrow R^l / \text{Im}(H_2) = R^l / h_2$ is the induced map given by H_1 .
 modulo(h1,h2) returns generators of the kernel of this induced map.

Note: If, for at least one of h_1 or h_2 , the attribute isHomog is st, then modulo(h1,h2) also sets this attribute (if the weights are compatible).

Example:

```

LIB "ncalg.lib";
def A = makeUs12();
setring A; // this algebra is U(sl_2)
option(redSB);
option(redTail);
ideal H2 = e2,f2,h2-1;
H2 = twostd(H2);
print(matrix(H2)); // print H2 in a compact form
↳ h2-1,fh-f,eh+e,f2,2ef-h-1,e2
ideal H1 = std(e);

```

```

ideal T = modulo(H1,H2);
T = NF(std(H2+T),H2);
T = std(T);
T;
↳ T[1]=h-1
↳ T[2]=e

```

See also Section 7.3.28 [syz (plural)], page 357.

7.3.15 mres (plural)

Syntax: `mres (ideal_expression, int_expression)`
`mres (module_expression, int_expression)`

Type: resolution

Purpose: computes a minimal free resolution of an ideal or module M with the Groebner basis method. More precisely, let $A = \text{matrix}(M)$, then `mres` computes a free resolution of $\text{coker}(A) = F_0/M$

$$\dots \longrightarrow F_2 \xrightarrow{A_2} F_1 \xrightarrow{A_1} F_0 \longrightarrow F_0/M \longrightarrow 0,$$

where the columns of the matrix A_1 are a (possibly) minimal set of generators of M . If the int expression k is not zero, then the computation stops after k steps and returns a resolution consisting of modules $M_i = \text{module}(A_i)$, $i = 1 \dots k$.

`mres(M,0)` returns a resolution consisting of at most $n+2$ modules, where n is the number of variables of the basering. Let `list L=mres(M,0)`; then `L[1]` consists of a minimal set of generators M , `L[2]` consists of a minimal set of generators for the first syzygy module of `L[1]`, etc., until `L[p+1]`, such that `L[i] ≠ 0` for $i \leq p$, but `L[p+1]` (the first syzygy module of `L[p]`) is 0 (if the basering is not a qring).

Note: Accessing single elements of a resolution may require that some partial computations have to be finished and may therefore take some time. Hence, assigning right away to a list is the recommended way to do it.

Example:

```

LIB "ncalg.lib";
def A = makeUs12();
setring A; // this algebra is U(sl_2)
option(redSB);
option(redTail);
ideal i = e,f,h;
i = std(i);
resolution M=mres(i,0);
M;
↳ 1      2      2      1
↳ A <--  A <--  A <--  A
↳
↳ 0      1      2      3
↳
list l = M; l;
↳ [1]:
↳   _[1]=f
↳   _[2]=e

```



```

↳ [2]:
↳   _[1]=ef*gen(1)-f2*gen(2)-2h*gen(1)-2*gen(1)
↳   _[2]=e2*gen(1)-ef*gen(2)-h*gen(2)+2*gen(2)
↳ [3]:
↳   _[1]=e*gen(1)-f*gen(2)
// see the exactness at this point
size(ideal(transpose(matrix(1[2]))*transpose(matrix(1[1]))));
↳ 0
print(matrix(M[3]));
↳ e,
↳ -f
// see the exactness at this point
size(ideal(transpose(matrix(1[3]))*transpose(matrix(1[2]))));
↳ 0

```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.3.13 [minres (plural)], page 340; Section 7.2.3 [module (plural)], page 320; Section 7.3.18 [nres (plural)], page 345.

7.3.16 nc_algebra

Syntax:

```

nc_algebra( matrix_expression C, matrix_expression D )
nc_algebra( number_expression n, matrix_expression D )
nc_algebra( matrix_expression C, poly_expression p )
nc_algebra( number_expression n, poly_expression p )

```

Type: ring

Purpose: Executed in the basering \mathbf{r} , say, in k variables x_1, \dots, x_k , `nc_algebra` creates and returns the non-commutative extension of \mathbf{r} subject to relations $\{x_j x_i = c_{ij} \cdot x_i x_j + d_{ij}, 1 \leq i < j \leq k\}$, where c_{ij} and d_{ij} must be put into two strictly upper triangular matrices C with entries c_{ij} from the ground field of \mathbf{r} and D with (commutative) polynomial entries d_{ij} from \mathbf{r} . See all the details in Section 7.4.1 [G-algebras], page 360.
 If $\forall i < j, c_{ij} = n$, one can input the number n instead of matrix C .
 If $\forall i < j, d_{ij} = p$, one can input the polynomial p instead of matrix D .

Note: The returned ring should be activated afterwards, using the command `setring`.

Remark: At present, `PLURAL` does not check the non-degeneracy conditions (see Section 7.4.1 [G-algebras], page 360) while setting an algebra.

Example:

```

LIB "nctools.lib";
// ----- first example: C, D are matrices -----
ring r1 = (0,Q),(x,y,z),Dp;
minpoly = rootofUnity(6);
matrix C[3][3];
matrix D[3][3];
C[1,2]=Q2; C[1,3]=1/Q2; C[2,3]=Q2;
D[1,2]=-Q*z; D[1,3]=1/Q*y; D[2,3]=-Q*x;
def S=nc_algebra(C,D);
// this algebra is a quantum deformation U'_q(so_3),
// where Q is a 6th root of unity
setring S;S;
↳ // coefficients: QQ[Q]/(Q2-Q+1)

```

```

↳ // number of vars : 3
↳ //          block 1 : ordering Dp
↳ //          : names  x y z
↳ //          block 2 : ordering C
↳ // noncommutative relations:
↳ //   yx=(Q-1)*xy+(-Q)*z
↳ //   zx=(-Q)*xz+(-Q+1)*y
↳ //   zy=(Q-1)*yz+(-Q)*x
kill r1,S;
// ----- second example: number n=1, D is a matrix
ring r2=0,(Xa,Xb,Xc,Ya,Yb,Yc,Ha,Hb),dp;
matrix d[8][8];
d[1,2]=-Xc; d[1,4]=-Ha; d[1,6]=Yb; d[1,7]=2*Xa;
d[1,8]=-Xa; d[2,5]=-Hb; d[2,6]=-Ya; d[2,7]=-Xb;
d[2,8]=2*Xb; d[3,4]=Xb; d[3,5]=-Xa; d[3,6]=-Ha-Hb;
d[3,7]=Xc; d[3,8]=Xc; d[4,5]=Yc; d[4,7]=-2*Ya;
d[4,8]=Ya; d[5,7]=Yb; d[5,8]=-2*Yb;
d[6,7]=-Yc; d[6,8]=-Yc;
def S=nc_algebra(1,d); // this algebra is U(sl_3)
setring S;S;
↳ // coefficients: QQ
↳ // number of vars : 8
↳ //          block 1 : ordering dp
↳ //          : names  Xa Xb Xc Ya Yb Yc Ha Hb
↳ //          block 2 : ordering C
↳ // noncommutative relations:
↳ //   XbXa=Xa*Xb-Xc
↳ //   YaXa=Xa*Ya-Ha
↳ //   YcXa=Xa*Yc+Yb
↳ //   HaXa=Xa*Ha+2*Xa
↳ //   HbXa=Xa*Hb-Xa
↳ //   YbXb=Xb*Yb-Hb
↳ //   YcXb=Xb*Yc-Ya
↳ //   HaXb=Xb*Ha-Xb
↳ //   HbXb=Xb*Hb+2*Xb
↳ //   YaXc=Xc*Ya+Xb
↳ //   YbXc=Xc*Yb-Xa
↳ //   YcXc=Xc*Yc-Ha-Hb
↳ //   HaXc=Xc*Ha+Xc
↳ //   HbXc=Xc*Hb+Xc
↳ //   YbYa=Ya*Yb+Yc
↳ //   HaYa=Ya*Ha-2*Ya
↳ //   HbYa=Ya*Hb+Ya
↳ //   HaYb=Yb*Ha+Yb
↳ //   HbYb=Yb*Hb-2*Yb
↳ //   HaYc=Yc*Ha-Yc
↳ //   HbYc=Yc*Hb-Yc
kill r2,S;
// ---- third example: C is a matrix, p=0 is a poly
ring r3=0,(a,b,c,d),lp;
matrix c[4][4];
c[1,2]=1; c[1,3]=3; c[1,4]=-2;
c[2,3]=-1; c[2,4]=-3; c[3,4]=1;

```

```

def S=nc_algebra(c,0); // it is a quasi--commutative algebra
setring S;S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ //           block 1 : ordering lp
↳ //           : names  a b c d
↳ //           block 2 : ordering C
↳ // noncommutative relations:
↳ //   ca=3ac
↳ //   da=-2ad
↳ //   cb=-bc
↳ //   db=-3bd
kill r3,S;
// -- fourth example : number n = -1, poly p = 3w
ring r4=0,(u,v,w),dp;
def S=nc_algebra(-1,3w);
setring S;S;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ //           block 1 : ordering dp
↳ //           : names  u v w
↳ //           block 2 : ordering C
↳ // noncommutative relations:
↳ //   vu=-uv+3w
↳ //   wu=-uw+3w
↳ //   wv=-vw+3w
kill r4,S;

```

See also Section 7.5.10 [ncalg_lib], page 458; Section 7.5.20 [nctools_lib], page 559; Section 7.5.24 [qmatrix_lib], page 606.

7.3.17 ncalgebra

Syntax:

```

ncalgebra( matrix_expression C, matrix_expression D )
ncalgebra( number_expression n, matrix_expression D )
ncalgebra( matrix_expression C, poly_expression p )
ncalgebra( number_expression n, poly_expression p )

```

Type: none

Purpose: Works like Section 7.3.16 [nc_algebra], page 343 but changes the basering.

Remark: This function is **deprecated** and should be substituted by `nc_algebra`, since it violates the general SINGULAR policy: only Section 4.19 [ring], page 124 and Section 5.1.139 [setring], page 255 can change the basering. More concretely, replace by `def A = nc_algebra(C, D); setring A;` which will additionally introduce a new ring. Afterwards, some objects may have to be mapped into the new ring.

See also Section 7.3.16 [nc_algebra], page 343; Section 7.5.20 [nctools_lib], page 559.

7.3.18 nres (plural)

Syntax: `nres (ideal_expression, int_expression)`
`nres (module_expression, int_expression)`

Type: resolution

Purpose: computes a free resolution of an ideal or module which is minimized from the second module on (by the Groebner basis method).

Note: Assigning a resolution to a list is the best choice of usage. The resolution may be minimized by using the command `minres`. Use the command `betti` to compute Betti numbers.

Example:

```
LIB "ncalg.lib";
def A = makeUs12();
setring A; // this algebra is U(sl_2)
option(redSB);
option(redTail);
ideal i = e,f,h;
i = std(i);
resolution F=nres(i,0); F;
↳ 1      3      3      1
↳ A <--  A <--  A <--  A
↳
↳ 0      1      2      3
↳ resolution not minimized yet
↳
list l = F; l;
↳ [1]:
↳  _[1]=h
↳  _[2]=f
↳  _[3]=e
↳ [2]:
↳  _[1]=f*gen(1)-h*gen(2)-2*gen(2)
↳  _[2]=e*gen(1)-h*gen(3)+2*gen(3)
↳  _[3]=e*gen(2)-f*gen(3)-gen(1)
↳ [3]:
↳  _[1]=e*gen(1)-f*gen(2)+h*gen(3)
// see the exactness at this point:
size(ideal(transpose(matrix(l[2]))*transpose(matrix(l[1]))));
↳ 0
// see the exactness at this point:
size(ideal(transpose(matrix(l[3]))*transpose(matrix(l[2]))));
↳ 0
print(betti(l), "betti");
↳          0      1      2      3
↳ -----
↳ 0:      1      -      3      1
↳ -----
↳ total:  1      0      3      1
↳
print(betti(minres(l)), "betti");
↳          0      1      2      3
↳ -----
↳ 0:      1      -      -      -
↳ 1:      -      -      2      1
↳ -----
```

```

↳ total:      1      0      2      1
↳

```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.3.13 [minres (plural)], page 340; Section 7.2.3 [module (plural)], page 320; Section 7.3.15 [mres (plural)], page 342.

7.3.19 oppose

Syntax: `oppose (ring_name, name)`

Type: poly, vector, ideal, module or matrix (the same type as the second argument)

Purpose: for a given object in the given ring, creates its opposite object in the opposite (Section 7.3.20 [opposite], page 348) ring (the last one is assumed to be the current ring).

Remark: for any object O , $(O^{opp})^{opp} = O$.

```

LIB "ncalg.lib";
def r = makeUs12();
setring r;
matrix m[3][4];
poly p = (h^2-1)*f*e;
vector v = [1,e*h,0,p];
ideal i = h*e, f^2*e,h*f*e;
m      = e,f,h,1,0,h^2, p,0,0,1,e^2,e*f*h+1;
module mm = module(m);
def b    = opposite(r);
setring b; b;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ //      block  1 : ordering a
↳ //                : names   H F E
↳ //                : weights 1 1 1
↳ //      block  2 : ordering ls
↳ //                : names   H F E
↳ //      block  3 : ordering C
↳ // noncommutative relations:
↳ //      FH=HF-2F
↳ //      EH=HE+2E
↳ //      EF=FE-H
// we will oppose these objects: p,v,i,m,mm
poly P    = oppose(r,p);
vector V  = oppose(r,v);
ideal I   = oppose(r,i);
matrix M  = oppose(r,m);
module MM = oppose(r,mm);
def c = opposite(b);
setring c; // now let's check the correctness:
// print compact presentations of objects
print(oppose(b,P)-imap(r,p));
↳ 0
print(oppose(b,V)-imap(r,v));
↳ [0]
print(matrix(oppose(b,I))-imap(r,i));
↳ 0,0,0
print(matrix(oppose(b,M))-imap(r,m));

```

```

↳ 0,0,0,0,
↳ 0,0,0,0,
↳ 0,0,0,0
print(matrix(oppose(b,MM))-imap(r,mm));
↳ 0,0,0,0,
↳ 0,0,0,0,
↳ 0,0,0,0

```

See Section 7.3.6 [envelope], page 334; Section 7.3.20 [opposite], page 348.

7.3.20 opposite

Syntax: `opposite (ring_name)`

Type: ring

Purpose: creates an opposite algebra of a given algebra.

Note: activate the ring with the `setring` command.

An opposite algebra of a given algebra $(A, \#)$ is an algebra $(A, *)$ with the same vector space but with the opposite multiplication, i.e.

$\forall f, g \in A^{opp}$, a new multiplication $*$ on A^{opp} is defined to be $f * g := g \# f$.

This is an identity functor on commutative algebras.

Remark: Starting from the variables x_1, \dots, x_N and the ordering $<$ of the given algebra, an opposite algebra will have variables X_N, \dots, X_1 (where the case and the position are reverted). Moreover, it is equipped with an opposed ordering $<_{opp}$ (it is given by the matrix, obtained from the matrix ordering of $<$ with the reverse order of columns). Currently not implemented for non-global orderings.

```

LIB "ncalg.lib";
def B = makeQso3(3);
// this algebra is a quantum deformation of U(so_3),
// where the quantum parameter is a 6th root of unity
setring B; B;
↳ // coefficients: QQ[Q]/(Q2-Q+1)
↳ // number of vars : 3
↳ //      block  1 : ordering dp
↳ //                : names  x y z
↳ //      block  2 : ordering C
↳ // noncommutative relations:
↳ //      yx=(Q-1)*xy+(-Q)*z
↳ //      zx=(-Q)*xz+(-Q+1)*y
↳ //      zy=(Q-1)*yz+(-Q)*x
def Bopp = opposite(B);
setring Bopp;
Bopp;
↳ // coefficients: QQ[Q]/(Q2-Q+1)
↳ // number of vars : 3
↳ //      block  1 : ordering a
↳ //                : names  Z Y X
↳ //                : weights 1 1 1
↳ //      block  2 : ordering ls
↳ //                : names  Z Y X
↳ //      block  3 : ordering C

```

```

↳ // noncommutative relations:
↳ //   YZ=(Q-1)*ZY+(-Q)*X
↳ //   XZ=(-Q)*ZX+(-Q+1)*Y
↳ //   XY=(Q-1)*YX+(-Q)*Z
def Bcheck = opposite(Bopp);
setring Bcheck; Bcheck; // check that (B-opp)-opp = B
↳ // coefficients: QQ[Q]/(Q2-Q+1)
↳ // number of vars : 3
↳ //       block  1 : ordering wp
↳ //                   : names   x y z
↳ //                   : weights 1 1 1
↳ //       block  2 : ordering C
↳ //       block  3 : ordering C
↳ // noncommutative relations:
↳ //   yx=(Q-1)*xy+(-Q)*z
↳ //   zx=(-Q)*xz+(-Q+1)*y
↳ //   zy=(Q-1)*yz+(-Q)*x

```

See Section B.2.6 [Matrix orderings], page 762; Section 7.3.6 [envelope], page 334; Section 7.3.19 [oppose], page 347.

7.3.21 preimage (plural)

Syntax: `preimage (ring_name, map_name, ideal_name)`
`preimage (ring_name, ideal_expression, ideal_name)`

Type: ideal

Purpose: returns the preimage of an ideal under a given map. The second argument has to be a map from the basering to the given ring (or an ideal defining such a map), and the ideal has to be an ideal in the given ring.

Note: To compute the kernel of a map, the preimage of zero has to be determined. Hence there is no special command for computing the kernel of a map in PLURAL.

Remark: In the non-commutative case, the command `preimage` is implemented only for maps $A \rightarrow B$, where A is a commutative ring. See Section 7.5.19 [ncpreim.lib], page 551 for the most general available implementation.

Example:

```

LIB "ncalg.lib";
ring R = 0,a,dp;
def Usl2 = makeUsl2();
setring Usl2;
poly C = 4*e*f+h^2-2*h;
// C is a central element of U(s12)
ideal I = e^3, f^3, h^3-4*h;
ideal Z = 0; // zero
ideal J = twostd(I); // two-sided GB
ideal K = std(I); // left GB
map Phi = R,C; // phi maps a (in R) to C (in U(s12))
setring R;
ideal PreJ = preimage(Usl2,Phi,J);
// the central character of J
PreJ;

```

```

↳ PreJ[1]=a2-8a
factorize(PreJ[1],1);
↳ _[1]=a
↳ _[2]=a-8
// hence, there are two simple characters for J
ideal PreK = preimage(Us12,Phi,K);
// the central character of K
PreK;
↳ PreK[1]=a3-32a2+192a
factorize(PreK[1],1);
↳ _[1]=a
↳ _[2]=a-24
↳ _[3]=a-8
// hence, there are three simple characters for K
preimage(Us12, Phi, Z); // kernel of phi
↳ _[1]=0

```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.2.2 [map (plural)], page 317; Section 7.2.7 [ring (plural)], page 327.

7.3.22 quotient (plural)

Syntax: `quotient (ideal_expression, ideal_expression)`
 `quotient (module_expression, module_expression)`

Type: ideal

Syntax: `quotient (module_expression, ideal_expression)`

Type: module

Purpose: computes the ideal quotient, resp. module quotient. Let R be the basering, I, J ideals and M, N submodules in R^n . Then

$$\begin{aligned} \text{quotient}(I, J) &= \{a \in R \mid aJ \subset I\}, \\ \text{quotient}(M, J) &= \{b \in R^n \mid bJ \subset M\}. \end{aligned}$$

Note: It can only be used for two-sided ideals (bimodules) in the second argument, otherwise the result may have no meaning.

Example:

```

//----- a very simple example -----
ring r=(0,q),(x,y),Dp;
def R=nc_algebra(q,0); // this algebra is a quantum plane
setring R;
option(returnSB);
poly f1 = x^3+2*x*y^2+2*x^2*y;
poly f2 = y;
poly f3 = x^2;
poly f4 = x+y;
ideal i = f1,f2;
ideal I = twostd(i);
ideal j = f3,f4;
ideal J = twostd(j);
quotient(I,J);

```



```

↳ _[1]=y
↳ _[2]=x2
module M = x*freemodule(3), y*freemodule(2);
quotient(M, ideal(x,y));
↳ _[1]=gen(1)
↳ _[2]=gen(2)
↳ _[3]=x*gen(3)
kill r,R;
//----- a bit more involved example
LIB "ncalg.lib";
def Usl2 = makeUsl2();
// this algebra is U(sl_2)
setring Usl2;
ideal i = e3,f3,h3-4*h;
ideal I = std(i);
poly C = 4*e*f+h^2-2*h;
ideal H = twostd(C-8);
option(returnSB);
ideal Q = quotient(I,H);
// print a compact presentation of Q:
print(matrix(Q));
↳ h,f3,ef2-4f,e2f-6e,e3

```

See Section 7.2.1 [ideal (plural)], page 313; Section 7.2.3 [module (plural)], page 320.

7.3.23 reduce (plural)

Syntax:

```

reduce ( poly_expression, ideal_expression )
reduce ( poly_expression, ideal_expression, int_expression )
reduce ( vector_expression, ideal_expression )
reduce ( vector_expression, ideal_expression, int_expression )
reduce ( vector_expression, module_expression )
reduce ( vector_expression, module_expression, int_expression )
reduce ( ideal_expression, ideal_expression )
reduce ( ideal_expression, ideal_expression, int_expression )
reduce ( module_expression, ideal_expression )
reduce ( module_expression, ideal_expression, int_expression )
reduce ( module_expression, module_expression )
reduce ( module_expression, module_expression, int_expression )

```

Type: the type of the first argument

Purpose: reduces a polynomial, vector, ideal or module to its **left** normal form with respect to an ideal or module represented by a left Groebner basis, if the second argument is a left Groebner basis.

returns 0 if and only if the polynomial (resp. vector, ideal, module) is an element (resp. subideal, submodule) of the ideal (resp. module).

Otherwise, the result may have no meaning.

The third (optional) argument 1 of type int forces a reduction which considers only the leading term and does no tail reduction.

Note: The commands `reduce` and `NF` are synonymous.

Example:

```

ring r=(0,a),(e,f,h),Dp;
matrix d[3][3];
d[1,2]=-h; d[1,3]=2e; d[2,3]=-2f;
def R=nc_algebra(1,d);
setring R;
// this algebra is U(sl_2) over Q(a)
ideal I = e2, f2, h2-1;
I = std(I);
// print a compact presentation of I
print(matrix(I));
↳ h2-1,fh-f,f2,eh+e,2*ef-h2-h,e2
ideal J = e, h-a;
J = std(J);
// print a compact presentation of J
print(matrix(J));
↳ h+(-a),e
poly z=4*e*f+h^2-2*h;
// z is the central element of U(sl_2)
reduce(z,I); // the central character of I:
↳ 3
reduce(z,J); // the central character of J:
↳ (a2+2a)
poly nz = z - NF(z,J); // nz will belong to J
reduce(nz,J);
↳ 0
reduce(I,J);
↳ _[1]=(a2-1)
↳ _[2]=(a-1)*f
↳ _[3]=f2
↳ _[4]=0
↳ _[5]=(-a2+a)
↳ _[6]=0

```

See also Section 7.2.1 [ideal (plural)], page 313; Section 7.2.3 [module (plural)], page 320; Section 7.3.26 [std (plural)], page 355.

7.3.24 ringlist (plural)

Syntax: ringlist (ring-expression)
ringlist (qring-expression)

Type: list

Purpose: decomposes a ring/qring into a list of 6 (or 4 in the commutative case) components. The first 4 components are common both for the commutative and for the non-commutative cases, the 5th and the 6th appear only in the non-commutative case.

5. upper triangle square matrix with nonzero upper triangle, containing structural coefficients of a G-algebra (this corresponds to the matrix C from the definition of Section 7.4.1 [G-algebras], page 360)
6. square matrix, containing structural polynomials of a G-algebra (this corresponds to the matrix D from the definition of Section 7.4.1 [G-algebras], page 360)

Note: After modifying a list acquired with `ringlist`, one can construct a corresponding ring with `ring(list)`.

Example:

```
// consider the quantized Weyl algebra
ring r = (0,q),(x,d),Dp;
def RS=nc_algebra(q,1);
setring RS; RS;
↳ // coefficients: QQ(q)
↳ // number of vars : 2
↳ //      block 1 : ordering Dp
↳ //      : names x d
↳ //      block 2 : ordering C
↳ // noncommutative relations:
↳ // dx=(q)*xd+1
list l = ringlist(RS);
l;
↳ [1]:
↳ [1]:
↳ 0
↳ [2]:
↳ [1]:
↳ q
↳ [3]:
↳ [1]:
↳ lp
↳ [2]:
↳ 1
↳ [4]:
↳ _[1]=0
↳ [2]:
↳ [1]:
↳ x
↳ [2]:
↳ d
↳ [3]:
↳ [1]:
↳ [1]:
↳ Dp
↳ [2]:
↳ 1,1
↳ [2]:
↳ [1]:
↳ C
↳ [2]:
↳ 0
↳ [4]:
↳ _[1]=0
↳ [5]:
↳ _[1,1]=0
↳ _[1,2]=(q)
↳ _[2,1]=0
```

```

↳   _[2,2]=0
↳ [6]:
↳   _[1,1]=0
↳   _[1,2]=1
↳   _[2,1]=0
↳   _[2,2]=0
// now, change the relation d*x = q*x*d +1
// into the relation d*x=(q2+1)*x*d + q*d + 1
matrix S = l[5]; // matrix of coefficients
S[1,2] = q^2+1;
l[5] = S;
matrix T = l[6]; // matrix of polynomials
T[1,2] = q*d+1;
l[6] = T;
def rr = ring(l);
setring rr; rr;
↳ // coefficients: QQ(q)
↳ // number of vars : 2
↳ //      block  1 : ordering Dp
↳ //              : names   x d
↳ //      block  2 : ordering C
↳ // noncommutative relations:
↳ //      dx=(q2+1)*xd+(q)*d+1

```

See also Section 7.2.7 [ring (plural)], page 327; Section 5.1.135 [ringlist], page 250.

7.3.25 `slimgb` (plural)

Syntax: `slimgb (ideal_expression)`
`slimgb (module_expression)`

Type: same type as argument

Purpose: returns a left Groebner basis of a left ideal or module with respect to the global monomial ordering of the basering.

Note: The commutative algorithm is described in the diploma thesis of Michael Brickenstein "Neue Varianten zur Berechnung von Groebnerbasen", written 2004 under supervision of G.-M. Greuel in Kaiserslautern.

It is designed to keep polynomials or vectors slim (short with small coefficients). Currently best results are examples over function fields (parameters).

The current implementation may not be optimal for weighted degree orderings.

The program only supports the options `prot`, which will give protocol output and `redSB` for returning a reduced Groebner basis. The protocol messages of `slimgb` mean the following:

`M[n,m]` means a parallel reduction of `n` elements with `m` non-zero output elements,
`b` notices an exchange trick described in the thesis and
`e` adds a reductor with non-minimal leading term.

`slimgb` works for grade commutative algebras but not for general GR-algebras. Please use `qslimgb` instead.

For a detailed commutative example see Section A.2.3 [slim Groebner bases], page 706.

Example:

```

LIB "nctools.lib";
LIB "ncalg.lib";
def U = makeUsl(2); setring U;
// U is the U(sl_2) algebra
ideal I = e^3, f^3, h^3-4*h;
option(redSB);
ideal J = slingb(I);
J;
↳ J[1]=h3-4h
↳ J[2]=fh2-2fh
↳ J[3]=eh2+2eh
↳ J[4]=2efh-h2-2h
↳ J[5]=f3
↳ J[6]=e3
// compare slingb with std:
ideal K = std(I);
print(matrix(NF(K,J)));
↳ 0,0,0,0,0,0
print(matrix(NF(J,K)));
↳ 0,0,0,0,0,0
// hence both Groebner bases are equal.
// Another example for exterior algebras
ring r;
def E = Exterior(); setring E; E;
↳ // coefficients: ZZ/32003
↳ // number of vars : 3
↳ //      block 1 : ordering dp
↳ //      : names x y z
↳ //      block 2 : ordering C
↳ // noncommutative relations:
↳ //   yx=-xy
↳ //   zx=-xz
↳ //   zy=-yz
↳ // quotient ring from ideal
↳ _[1]=z2
↳ _[2]=y2
↳ _[3]=x2
slingb(xy+z);
↳ _[1]=yz
↳ _[2]=xz
↳ _[3]=xy+z

```

See Section 5.1.110 [option], page 230; Section 7.3.26 [std (plural)], page 355.

7.3.26 std (plural)

Syntax: std (ideal_expression)
 std (module_expression)
 std (ideal_expression, poly_expression)
 std (module_expression, vector_expression)

Type: ideal or module

Purpose: returns a left Groebner basis (see Section 7.4.2 [Groebner bases in G-algebras], page 361 for a definition) of an ideal or module with respect to the monomial ordering of the basering.

Use an optional second argument of type poly, resp. vector, to construct the Groebner basis from an already computed one (given as the first argument) and one additional generator (the second argument).

Note: To view the progress of long running computations, use `option(prot)`. (see Section 5.1.110 [option], page 230(prot)).

Example:

```
LIB "ncalg.lib";
option(prot);
def R = makeUs12();
// this algebra is U(sl_2)
setring R;
ideal I = e2, f2, h2-1;
I=std(I);
↳ 2(2)s
↳ s
↳ s
↳ 3s
↳ (3)2(2)s
↳ s
↳ (4)(3)(2)3s
↳ 2(4)(3)(2)32product criterion:6 chain criterion:3
I;
↳ I[1]=h2-1
↳ I[2]=fh-f
↳ I[3]=eh+e
↳ I[4]=f2
↳ I[5]=2ef-h-1
↳ I[6]=e2
kill R;
//-----
def RQ = makeQso3(3);
// this algebra is U'_q(so_3),
// where Q is a 6th root of unity
setring RQ;
RQ;
↳ // coefficients: QQ[Q]/(Q2-Q+1)
↳ // number of vars : 3
↳ //      block 1 : ordering dp
↳ //      : names x y z
↳ //      block 2 : ordering C
↳ // noncommutative relations:
↳ //      yx=(Q-1)*xy+(-Q)*z
↳ //      zx=(-Q)*xz+(-Q+1)*y
↳ //      zy=(Q-1)*yz+(-Q)*x
ideal J=x2, y2, z2;
J=std(J);
↳ 2(2)s
↳ s
```

```

↳ s
↳ 3s
↳ (4)s
↳ 2(3)s
↳ (5)s
↳ (6)s
↳ 1(8)s
↳ (7)(5)s
↳ (3)(2)product criterion:0 chain criterion:17
J;
↳ J[1]=z
↳ J[2]=y
↳ J[3]=x

```

See also Section 7.2.1 [ideal (plural)], page 313; Section 7.2.7 [ring (plural)], page 327.

7.3.27 subst (plural)

Syntax: subst (poly_expression, ring_variable, poly_expression)
 subst (vector_expression, ring_variable, poly_expression)
 subst (ideal_expression, ring_variable, poly_expression)
 subst (module_expression, ring_variable, poly_expression)

Type: poly, vector, ideal or module (corresponding to the first argument)

Purpose: substitutes a ring variable by a polynomial.

Example:

```

LIB "ncalg.lib";
def R = makeUs12();
// this algebra is U(s1_2)
setring R;
poly C = e*f*h;
poly C1 = subst(C,e,h^3);
C1;
↳ fh4-6fh3+12fh2-8fh
poly C2 = subst(C,f,e+f);
C2;
↳ e2h+efh

```

See also Section 7.2.2 [map (plural)], page 317.

7.3.28 syz (plural)

Syntax: syz (ideal_expression)
 syz (module_expression)

Type: module

Purpose: computes the first syzygy (i.e., the module of relations of the given generators) of the ideal, resp. module.

Note: if S is a matrix of a left syzygy module of left submodule given by matrix M , then $\text{transpose}(S) * \text{transpose}(M) = 0$.

Example:

```

LIB "ncalg.lib";
def R = makeQso3(3); setring R;
// we wish to have completely reduced bases:
option(redSB); option(redTail);
ideal tst;
ideal J = x3+x,x*y*z;
print(syz(J));
↳ -yz,
↳ x2+1
ideal K = x+y+z,y+z,z;
module S = syz(K);
print(S);
↳ (Q-1),      (-Q+1)*z,   (-Q)*y,
↳ (Q)*z+(-Q+1), (Q-1)*z+(Q), -x+(Q)*y,
↳ y+(-Q)*z,    x+(-Q),    x+(-Q+1)
tst = ideal(transpose(S)*transpose(K));
// check the property of a syzygy module (tst==0):
size(tst);
↳ 0
// Now compute the Groebner basis of K ...
K = std(K);
// ... print a matrix presentation of K ...
print(matrix(K));
↳ z,y,x
S = syz(K); // ... and its syzygy module
print(S);
↳ y,    x,      (Q-1),
↳ (Q)*z,(Q),    x,
↳ (Q-1),(-Q+1)*z,(Q)*y
tst = ideal(transpose(S)*transpose(K));
// check the property of a syzygy module (tst==0):
size(tst);
↳ 0
// Note the "commutative" (not transposed) syzygy property does not hold
size(ideal(matrix(K)*matrix(S)));
↳ 3

```

See also Section 7.2.1 [ideal (plural)], page 313; Section 7.3.13 [minres (plural)], page 340; Section 7.2.3 [module (plural)], page 320; Section 7.3.15 [mres (plural)], page 342; Section 7.3.18 [nres (plural)], page 345.

7.3.29 twostd (plural)

Syntax: twostd(ideal.expression);

Type: ideal

Purpose: returns a two-sided Groebner basis of an input

Note: Treating the input as a set of

two-sided generators of a two-sided ideal T , two-sided Groebner basis is a left (and a right) Groebner basis of T . (see Section 5.1.149 [std], page 266).

Remark: There are algebras with no two-sided ideals except 0 and the whole algebra (like Weyl algebras).

Example:

```

LIB "ncalg.lib";
def U = makeUsl2(); // this algebra is U(sl_2)
setring U;
ideal i= e^3, f^3, h^3 - 4*h;
option(redSB);
option(redTail);
ideal I = std(i);
print(matrix(I)); // print a compact presentation of I
↳ h3-4h,fh2-2fh,eh2+2eh,2efh-h2-2h,f3,e3
ideal J = twostd(i);
// print a compact presentation of J:
print(matrix(ideal(J[1..6]))); // first 6 gen's
↳ h3-4h,fh2-2fh,eh2+2eh,f2h-2f2,2efh-h2-2h,e2h+2e2
print(matrix(ideal(J[7..size(J)]))); // the rest of gen's
↳ f3,ef2-fh,e2f-eh-2e,e3
// compute the set of elements present in J but not in I
ideal K = NF(J,I);
K = K+0; // simplify K
print(matrix(K));
↳ f2h-2f2,e2h+2e2,ef2-fh,e2f-eh-2e

```

7.3.30 vdim (plural)

Syntax: vdim (ideal_expression)
vdim (module_expression)

Type: int

Purpose: computes the vector space dimension of the factor-module that equals ring (resp. free module) modulo the ideal (resp. submodule), generated by the leading terms of the given generators.

If the factor-module is not of finite dimension, -1 is returned.

If the generators form a left Groebner basis, this is the same as the vector space dimension of the left factor module.

Note: In the non-commutative case, a ring modulo an ideal has a ring structure if and only if the ideal is two-sided.

Example:

```

ring R=0,(x,y,z),dp;
matrix d[3][3];
d[1,2]=-z; d[1,3]=2x; d[2,3]=-2y;
def RS=nc_algebra(1,d); //U(sl_2)
setring RS;
option(redSB); option(redTail);
ideal I=x3,y3,z3-z;
I=std(I);
I;
↳ I[1]=z3-z
↳ I[2]=y3
↳ I[3]=x3
↳ I[4]=y2z2-y2z
↳ I[5]=x2z2+x2z

```

```

↳ I [6]=x2y2z-2xyz2-2xyz+2z2+2z
vdim(I);
↳ 21

```

See also Section 7.2.1 [ideal (plural)], page 313; Section 7.3.10 [kbase (plural)], page 337; Section 7.3.26 [std (plural)], page 355.

7.4 Mathematical background (plural)

This section introduces some of the mathematical notions and definitions used throughout the PLURAL manual. For details, please, refer to appropriate articles or text books (see Section 7.4.4 [References (plural)], page 364). A detailed discussion of the subjects in this section can be found in the doctoral thesis [LV] of V. Levandovskyy (see Section 7.4.4 [References (plural)], page 364).

All algebras are assumed to be associative K -algebras for some field K .

7.4.1 G-algebras

Definition (PBW basis)

Let K be a field, and let a K -algebra A be generated by variables x_1, \dots, x_n subject to some relations. We call A an algebra with **PBW basis** (Poincaré-Birkhoff-Witt basis), if a K -basis of A is $\text{Mon}(x_1, \dots, x_n) = \{x_1^{a_1} x_2^{a_2} \dots x_n^{a_n} \mid a_i \in \mathbb{N} \cup \{0\}\}$, where a power-product $x_1^{a_1} x_2^{a_2} \dots x_n^{a_n}$ (in this particular order) is called a **monomial**. For example, $x_1 x_2$ is a monomial, while $x_2 x_1$ is, in general, not a monomial.

Definition (G-algebra)

Let K be a field, and let a K -algebra A be given in terms of generators subject to the following relations:

$$A = K\langle x_1, \dots, x_n \mid \{x_j x_i = c_{ij} \cdot x_i x_j + d_{ij}\}, 1 \leq i < j \leq n \rangle, \text{ where } c_{ij} \in K^*, d_{ij} \in K[x_1, \dots, x_n].$$

A is called a **G-algebra**, if the following conditions hold:

- there is a monomial well-ordering $<$ on $K[x_1, x_2, \dots, x_n]$ such that $\forall i < j \text{ LM}(d_{ij}) < x_i x_j$,
- **non-degeneracy conditions**: $\forall 1 \leq i < j < k \leq n : \mathcal{NDC}_{ijk} = 0$, where

$$\mathcal{NDC}_{ijk} = c_{ik} c_{jk} \cdot d_{ij} x_k - x_k d_{ij} + c_{jk} \cdot x_j d_{ik} - c_{ij} \cdot d_{ik} x_j + d_{jk} x_i - c_{ij} c_{ik} \cdot x_i d_{jk}.$$

Note: Note that non-degeneracy conditions ensure associativity of multiplication, defined by the relations. It is also proved, that they are necessary and sufficient to guarantee the PBW property of an algebra, defined via C_{ij} and D_{ij} as above.

Theorem (properties of G-algebras)

Let A be a G -algebra. Then

- A has a PBW (Poincaré-Birkhoff-Witt) basis,
- A is left and right noetherian,
- A is an integral domain.

Setting up a G-algebra

In order to set up a G -algebra one has to do the following steps:

- define a commutative ring $R = K[x_1, \dots, x_n]$, equipped with a monomial ordering $<$ (see Section 7.2.7.1 [ring declarations (plural)], page 327).
This provides us with the information on a field K (together with its parameters), variables $\{x_i\}$ and an ordering $<$.
From the sequence of variables we will build a G -algebra with the Poincaré-Birkhoff-Witt (PBW) basis $\{x_1^{a_1} x_2^{a_2} \dots x_n^{a_n}\}$.
- define strictly $n \times n$ upper triangular matrices (of type `matrix`)
 1. $C = \{c_{ij}, i < j\}$, with nonzero entries c_{ij} of type number (c_{ij} for $i \geq j$ will be ignored).
 2. $D = \{d_{ij}, i < j\}$, with polynomial entries d_{ij} from R (d_{ij} for $i \geq j$ will be ignored).
- Call the initialization function `nc_algebra(C,D)` (see Section 7.3.16 [nc_algebra], page 343) with the data C and D .

PLURAL does not check automatically whether the non-degeneracy conditions hold but it provides a procedure Section 7.5.20.3 [ndcond], page 561 from the library Section 7.5.20 [nctools_lib], page 559 to check this.

7.4.2 Groebner bases in G-algebras

We follow the notations, used in the SINGULAR Manual (e.g. in Section C.1 [Standard bases], page 766).

For a G -algebra A , we denote by ${}_A\langle g_1, \dots, g_s \rangle$ the left submodule of a free module A^r , generated by elements $\{g_1, \dots, g_s\} \subset A^r$.

Let $<$ be a fixed monomial well-ordering on the G -algebra A with the Poincaré-Birkhoff-Witt (PBW) basis $\{x^\alpha = x_1^{a_1} x_2^{a_2} \dots x_n^{a_n}\}$. For a given free module A^r with the basis $\{e_1, \dots, e_r\}$, $<$ denotes also a fixed module ordering on the set of monomials $\{x^\alpha e_i \mid \alpha \in \mathbf{N}^n, 1 \leq i \leq r\}$.

Definition

For a set $S \subset A^r$, define $L(S)$ to be the K -vector space, spanned on the leading monomials of elements of S , $L(S) = \oplus \{Kx^\alpha e_i \mid \exists s \in S, \text{LM}(s) = x^\alpha e_i\}$.

We call $L(S)$ the **span of leading monomials of S** .

Let $I \subset A^r$ be a left A -submodule. A finite set $G \subset I$ is called a **left Groebner basis** of I if and only if $L(G) = L(I)$, that is for any $f \in I \setminus \{0\}$ there exists a $g \in G$ satisfying $\text{LM}(g) \mid \text{LM}(f)$, i.e., if $\text{LM}(f) = x^\alpha e_i$, then $\text{LM}(f) = x^\beta e_i$ with $\beta_j \leq \alpha_j$, $1 \leq j \leq n$.

Remark: In general non-commutative algorithms are working with global well-orderings only (see Section 7.1 [PLURAL], page 312, Section B.2 [Monomial orderings], page 760 and Section 3.3.3 [Term orderings], page 34), unless we deal with graded commutative algebras via Section 7.6 [Graded commutative algebras (SCA)], page 611.

A Groebner basis $G \subset A^r$ is called **minimal** (or **reduced**) if $0 \notin G$ and if $\text{LM}(g) \notin L(G \setminus \{g\})$ for all $g \in G$. Note, that any Groebner basis can be made minimal by deleting successively those g with $\text{LM}(h) \mid \text{LM}(g)$ for some $h \in G \setminus \{g\}$.

For $f \in A^r$ and $G \subset A^r$ we say that f is **completely reduced with respect to G** if no monomial of f is contained in $L(G)$.

Left Normal Form

A map $\text{NF} : A^r \times \{G \mid G \text{ a (left) Groebner basis}\} \rightarrow A^r, (f|G) \mapsto \text{NF}(f|G)$, is called a **(left) normal form** on A^r if for any $f \in A^r$ and any left Groebner basis G the following holds:

- (i) $\text{NF}(0|G) = 0$,
- (ii) if $\text{NF}(f|G) \neq 0$ then $\text{LM}(g)$ does not divide $\text{LM}(\text{NF}(f|G))$ for all $g \in G$,
- (iii) $f - \text{NF}(f|G) \in {}_A\langle G \rangle$.

$\text{NF}(f|G)$ is called a **left normal form of f with respect to G** (note that such a map is not unique).

Remark: As we have already mentioned in the definitions **ideal** and **module** (see Section 7.1 [PLURAL], page 312), by **NF** (or **reduce**) PLURAL understands a left normal form. Note, that **rightNF** from Section 7.5.20 [nctools_lib], page 559 allows to compute a right normal form.

Left ideal membership (plural)

For a left Groebner basis G of I the following holds: $f \in I$ if and only if the left normal form $\text{NF}(f|G) = 0$.

For computing a left Groebner basis \mathbf{G} of \mathbf{I} , use Section 7.3.26 [std (plural)], page 355.

For computing a left normal form of \mathbf{f} with respect to \mathbf{G} , use Section 7.3.23 [reduce (plural)], page 351.

Right ideal membership (plural)

The right ideal membership is analogous to the left one:

for computing a right Groebner basis \mathbf{G} of \mathbf{I} , use Section 7.5.20.10 [rightStd], page 567 from Section 7.5.20 [nctools_lib], page 559,

for computing a right normal form of \mathbf{f} with respect to \mathbf{G} , use Section 7.5.20.11 [rightNF], page 568 from Section 7.5.20 [nctools_lib], page 559.

Two-sided ideal membership (plural)

Let J be a two-sided ideal and T be a two-sided Groebner basis of J .

Then $f \in J$ if and only if the left normal form $\text{NF}(f|T) = 0$.

For computing a two-sided Groebner basis \mathbf{T} of \mathbf{J} , use Section 7.3.29 [twostd (plural)], page 358,

for computing a normal form of \mathbf{f} with respect to \mathbf{T} , use Section 7.3.23 [reduce (plural)], page 351.

7.4.3 Syzygies and resolutions (plural)

Syzygies

Let A be a GR-algebra. A **left** (resp. **right**) **syzygy** between k elements $\{f_1, \dots, f_k\} \subset A^r$ is a k -tuple $(g_1, \dots, g_k) \in A^k$ satisfying

$$\sum_{i=1}^k g_i f_i = 0 \quad \text{resp.} \quad \sum_{i=1}^k f_i g_i = 0.$$

The set of all left (resp. right) syzygies between $\{f_1, \dots, f_k\}$ is a left (resp. right) submodule S of A^k .

Remark: With respect to the definitions of `ideal` and `module` (see Section 7.1 [PLURAL], page 312), by `syz PLURAL` understands an inquiry to compute the left syzygy module.

Note, that `rightModulo(M, std(0))` from Section 7.5.20 [nctools.lib], page 559 computes the right syzygy module of M .

If S is a matrix of a left syzygy module of left submodule given by matrix M , then `transpose(S)*transpose(M) = 0` (but, in general, $M \cdot S \neq 0$).

Note, that the syzygy modules of I depend on a choice of generators $\{g_1, \dots, g_s\}$, but one can show that they depend on I uniquely up to direct summands.

Free resolutions

Let $I = \langle g_1, \dots, g_s \rangle \subseteq A^r$ and $M = A^r/I$. A **free resolution of M** is a long exact sequence

$$\dots \longrightarrow F_2 \xrightarrow{B_2} F_1 \xrightarrow{B_1} F_0 \longrightarrow M \longrightarrow 0,$$

with `transpose(B_{i+1}) \cdot transpose(B_i) = 0`

and where the columns of the matrix B_1 generate I . Note, that resolutions over factor-algebras need not to be of finite length.

Generalized Hilbert Syzygy Theorem

For a G -algebra A , generated by n variables, there exists a free resolution of length smaller or equal than n .

Example:

```
ring R=0,(x,y,z),dp;
matrix d[3][3];
d[1,2]=-z; d[1,3]=2x; d[2,3]=-2y;
def U=nc_algebra(1,d); // this algebra is U(sl_2)
setring U;
option(redSB); option(redTail);
ideal I=x^3,y^3,z^3-z;
I=std(I);
I;
↳ I[1]=z^3-z
↳ I[2]=y^3
↳ I[3]=x^3
↳ I[4]=y^2z^2-y^2z
↳ I[5]=x^2z^2+x^2z
↳ I[6]=x^2y^2z-2xyz^2-2xyz+2z^2+2z
resolution resI = mres(I,0);
resI;
↳ 1      5      7      3
↳ U <--  U <--  U <--  U
↳
↳ 0      1      2      3
↳
list l = resI;
// The matrix A_1 is given by
print(matrix(l[1]));
```

```

↳ z3-z,y3,x3,y2z2-y2z,x2z2+x2z
// We see that the columns of A_1 generate I.
// The matrix A_2 is given by
print(matrix(1[2]));
↳ 0,      0,      y2,  x2,  6yz,      -36xy+18z+24,-6xz,
↳ z2+11z+30,0,      0,  0,  2x2z+12x2,  2x3,      0,
↳ 0,      z2-11z+30,0,  0,  0,      -2y3,      2y2z-12y2,
↳ -y,      0,      -z-5,0,  x2y-6xz-30x,9x2,      x3,
↳ 0,      -x,      0,  -z+5,-y3,      -9y2,      -xy2-4yz+28y
ideal tst; // now let us show that the resolution is exact
matrix TST;
TST = transpose(matrix(1[3]))*transpose(matrix(1[2])); // 2nd term
size(ideal(TST));
↳ 0
TST = transpose(matrix(1[2]))*transpose(matrix(1[1])); // 1st term
size(ideal(TST));
↳ 0

```

7.4.4 References (plural)

The Centre for Computer Algebra Kaiserslautern publishes a series of preprints which are electronically available at <https://www.mathematik.uni-kl.de/organisation/zca/reports-on-ca/>. Other sources to check are the following books and articles:

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7.5 PLURAL libraries

The content of libraries, created for PLURAL is described in the following subsections.

Use the LIB command for loading of single libraries.

Note: For any computation in PLURAL, the monomial ordering must be a global ordering.

See also Section D.11.3 [jacobson_lib], page 889 for the diagonalization of matrices over Ore Euclidean domains.

7.5.1 bimodules_lib

Library: bimodules.lib

Purpose: Tools for handling bimodules

Authors: Ann Christina Foldenauer, Christina.Foldenauer@rwth-aachen.de
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Overview:

The main purpose of this library is the handling of bimodules which will help e.g. to determine weak normal forms of representation matrices and total divisors within non-commutative, non-simple G -algebras. We will use modules homomorphisms between a G -algebra and its enveloping algebra in order to work left Groebner basis theory on bimodules. Assume we have defined a (non-commutative) G -algebra A over the field K , and an (A,A) -bimodule M . Instead of working with M over A , we define the enveloping algebra $A^{\text{env}} = A \otimes_K A^{\text{opp}}$ (this can be done with command `envelope(A)`) and embed M into A^{env} via `imap()`. Thus we obtain the left A^{env} -module $M \otimes 1$ in A^{env} . This has a lot of advantages, because left module theory has much more commands that are already implemented in SINGULAR:PLURAL. Two important procedures that we can use are `std()` which computes the left Groebner basis, and `NF()` which computes the left normal form. With the help of this method we are also able to determine the set of bisyzygies of a bimodule.

A built-in command `twostd` in PLURAL computes the two-sided Groebner basis of an ideal by using the right completion algorithm of [2]. `bistd` from this library uses very different approach, which is often superior to the right completion.

References:

- The procedure `bistd()` is the implementation of an algorithm M. del Socorro Garcia Roman presented in [1](page 66-78).
- [1] Maria del Socorro Garcia Roman, Effective methods in Algebras with PBW bases: G -algebras and Yang-Baxter Algebras, Ph.D. thesis, Universidad de La Laguna, 2005.
 - [2] Viktor Levandovskyy, Non-commutative Computer Algebra for polynomial Algebras: Groebner Bases, Applications and Implementations, Ph.D. thesis, Kaiserslautern, 2005.
 - [3] N. Jacobson, The theory of rings, AMS, 1943.
 - [4] P. M. Cohn, Free Rings and their Relations, Academic Press Inc. (London) Ltd., 1971.

Procedures: See also: Section 7.5.10 [`ncalg.lib`], page 458; Section 7.5.20 [`nctools.lib`], page 559.

7.5.1.1 bistd

Procedure from library `bimodules.lib` (see Section 7.5.1 [`bimodules.lib`], page 365).

Usage: `bistd(M)`; M is (two-sided) ideal/module

Return: ideal or module (same type as the argument)

Purpose: Computes the two-sided Groebner basis of an ideal/module with the help the enveloping algebra of the basering, alternative to `twostd()` for ideals.

Example:


```

LIB "bimodules.lib";
ring w = 0,(x,s),Dp;
def W=nc_algebra(1,s); // 1st shift algebra
setring W;
matrix m[3][3]=[s^2,s+1,0],[s+1,0,s^3-x^2*s],[2*s+1,s^3+s^2,s^2];
print(m);
  ↪ s2,    s+1,    s+1,
  ↪ 0,     -x2s+s3,2s+1,
  ↪ s3+s2,s2,     0
module L = m; module M2 = bistd(L);
print(M2);
  ↪ 1,1,s+1,
  ↪ 0,1,0,
  ↪ 0,0,s2

```

7.5.1.2 bitrinity

Procedure from library `bimodules.lib` (see Section 7.5.1 [bimodules_lib], page 365).

Usage: `bitrinity(M)`; M is (two-sided) ideal/module

Return: ring, the enveloping algebra of the basering, with objects in it. additionally it exports a list $L = \text{Coeff}, \text{BiSyz}$.

Theory: Let ψ_s be the epimorphism of left $R(X) R^{\text{opp}}$ modules:
 $\psi_s(s(X)_K t) = \text{smt} := (s_1 m t_1, \dots, s_s m t_s) = (\psi(s_1(X) t_1), \dots, \psi(s_s(X) t_s))$ in R^s .

Then $\psi_s(A) := (\psi_s(a_{ij}))$ for every matrix A in $\text{Mat}(n \times m, R)$.

For a two-sided ideal $I = \langle f_1, \dots, f_j \rangle$ with Groebner basis $G = \{g_1, \dots, g_k\}$ in R , Coeff is the Coefficient-Matrix and BiSyz a bisyzygy matrix.

Let C be the submatrix of Coeff , where C is Coeff without the first row. Then $(g_1, \dots, g_k) = \psi_s(C^T * (f_1 \dots f_j)^T)$ and $(0, \dots, 0) = \psi_s(\text{BiSyz}^T * (f_1 \dots f_j)^T)$. The first row of Coeff ($G_1 \dots G_n$) corresponds to the image of the Groebner basis of I : $\psi_s((G_1 \dots G_n)) = G = \{g_1 \dots g_k\}$.

For a (R,R) -bimodule M with Groebner basis $G = \{g_1, \dots, g_k\}$ in R^r , Coeff is the coefficient matrix and BiSyz a bisyzygy matrix.

Let C be the submatrix of Coeff , where C is Coeff without the first r rows. Then $(g_1 \dots g_k) = \psi_s(C^T * (f_1 \dots f_j)^T)$ and $(0 \dots 0) = \psi_s(\text{BiSyz}^T * (f_1 \dots f_j)^T)$.

The first r rows of $\text{Coeff} = (G_1 \dots G_n)$ (Here G_i denotes to the i -th column of the first r rows) corresponds to the image of the Groebner basis of M : $\psi_s((G_1 \dots G_n)) = G = \{g_1 \dots g_k\}$.

Purpose: This procedure returns a coefficient matrix in the enveloping algebra of the basering R , that gives implicitly the two-sided Groebner basis of a (R,R) -bimodule M and the coefficients that produce the Groebner basis with the help of the originally used generators of M . Additionally it calculates the bisyzygies of M as left-module of the enveloping algebra of R .

Auxiliary procedures:

Note: To get list $L = \text{Coeff}, \text{BiSyz}$, we set: `def G = bitrinity(); setring G; L;` or `$L[1]; L[2];`

Example:

```

LIB "bimodules.lib";
ring r = 0,(x,s),dp;

```

```

def R = nc_algebra(1,s); setring R; // 1st shift algebra
poly f = x*s + s^2; // only one generator
ideal I = f; // note, two sided Groebner basis of I is xs, s^2
def G = bitrinity(I);
setring G;
print(L[1]); // Coeff
↳ S2, SX,
↳ s-S, -s+S+1
//the first row shows the Groebnerbasis of I consists of
// psi_s(SX) = xs , phi(S^2) = s^2:
// remember phi(a (X) b - c (X) d) = psi_s(a (X) b) - phi(c (X) d) := ab - cd in R.
// psi_s((-s+S+1)*(x*s + s^2)) = psi_s(-xs2-s3+xsS+xs+s2S)
// = -xs^2-s^3+xs^2+xs+s^3 = xs
// psi_s((s-S)*(x*s + s^2)) = psi_s(xs2+s3-xsS-s2S+s2) = s^2
print(L[2]); //Biszygies
↳ sX+sS-2s-SX-S2,x+s-X-S+1,s2-2sS+S2
// e.g. psi_s((x2-2sS+s-X2+2S2+2X+S-1)(x*s + s^2))
// = psi_s(x3s+x2s2-2xs2S+xs2-2s3S+s3-xsX2+2xsS2+2xsX+xsS-xs-s2X2+2s2S2+2s2X-s2S)
// = x^3s+x^2s^2-2xs^3+xs^2-2s^4+s^3-xsx^2+2xs^3+2xsx+xs^2-xs-s^2x^2+2s^4+2s^2x-s^3
// = 0 in R

```

7.5.1.3 liftenvelope

Procedure from library `bimodules.lib` (see Section 7.5.1 [`bimodules.lib`], page 365).

Usage: `liftenvelope(M,g)`; M ideal/module, g poly

Return: ring, the enveloping algebra of the basering R.

Given a two-sided ideal M in R and a polynomial g in R this procedure returns the enveloping algebra of R. Additionally it exports a list $l = C, B$; where B is the left Groebner basis of the left-syzygies of $M \otimes 1$ and C is a vector of coefficients in the enveloping algebra of R such that $\text{psi}_s(C^T * (f_1 \dots f_n)) = g$.

psi_s is an epimorphism of left R (X) R^{opp} modules:

$\text{psi}_s(s(X)_K t) = \text{smt} := (s_1 m t_1, \dots, s_s m t_s) = (\psi(s_1(X) t_1), \dots, \psi(s_s(X) t_s))$ in R^s .

Then $\text{psi}_s(A) := (\text{psi}_s(a_{ij}))$ for every matrix A in $\text{Mat}(n \times m, R)$.

Assume: The second component has to be an element of the first component.

Purpose: This procedure is used for computing total divisors. Let $\{f_1, \dots, f_n\}$ be the generators of the first component and let the second component be called g. Then the returned list $l = C, B = (b_1, \dots, b_n)$; defines an affine set $A = C + \sum_i a_i b_i$ with (a_1, \dots, a_n) in the enveloping algebra of the basering R such that $\text{psi}_s(a^T * (f_1 \dots f_n)) = g$ for all a in A. For certain rings R, we can find pure tensors within this set A, and if we do, `liftenvelope()` helps us to decide whether f is a total divisor of g.

Note: To get list $l = C, B$. we set: `def G = liftenvelope(); setring G; l`; or `l[1]; l[2];`.

Example:

```

LIB "bimodules.lib";
ring r = 0, (x,s), dp;
def R = nc_algebra(1,s); setring R;
ideal I = x*s;
poly p = s*x*s*x; // = (s (x) x) * x*s = (sX) * x*s
p;

```

```

↳ x2s2+3xs2+2s2
def J = liftenvelope(I,p);
setring J;
print(l[1]);
↳ 0
//2s+SX = (2s (x) 1) + (1 (x) sx)
print(l[2]);
↳ sX-2s-SX,x-X+1,s2-2sS+S2
// Groebnerbasis of BiSyz(I) as LeftSyz in R^{env}
// We get : 2s+SX + ( sX - 2s -SX) = sX - a pure tensor!!!!

```

7.5.1.4 CompDecomp

Procedure from library `bimodules.lib` (see Section 7.5.1 [bimodules_lib], page 365).

Usage: `CompDecomp(p)`; p poly

Note: This procedure only works if the basering is an enveloping algebra A^{env} of a (non-commutative) ring A . Thus also the polynomial in the argument has to be in A^{env} .

Return: Returns an ideal I in A^{env} , where the sum of all terms of the argument with the same right side (of the tensor summands) are stored as a generator of I .
Let $b \neq c$, then for $p = (a(X) b) + (c(X) b) + (a(X) c)$ the ideal $I := \text{CompDecomp}(p)$ is given by: $I[1] = (a(X) b) + (c(X) b)$; $I[2] = a(X) c$.

Purpose: By decomposing the polynomial we can easily check whether the given polynomial is a pure tensor.

Example:

```

LIB "bimodules.lib";
ring r = 0,(x,s),dp;
def R = nc_algebra(1,s); setring R; //1st shift algebra
def Re = envelope(R); setring Re; //basing is now R^{env} = R (X) R^{opp}
poly f = X*S*x^2+5*x*S*X+S*X; f;
↳ x2SX+x2S+5xSX+SX
ideal I = CompDecomp(f);
print(matrix(I)); // what means that f = (x2+5x+1)*SX + x2*S
↳ x2SX+5xSX+SX,x2S
poly p = x*S+X^2*S+2*s+x*X^2*s+5*x*s; p;
↳ xsX2+5xs+xS+2s+SX2+2SX+S
ideal Q = CompDecomp(p);
print(matrix(Q));
↳ xsX2,5xs+2s,xS+S,SX2,2SX

```

7.5.1.5 isPureTensor

Procedure from library `bimodules.lib` (see Section 7.5.1 [bimodules_lib], page 365).

Usage: `isPureTensor(g)`; g poly

Note: This procedure only works if the basering is an enveloping algebra A^{env} of a (non-commutative) ring A . Thus also the polynomial in the argument has to be in A^{env} .

Return: Returns 0 if g is not a pure tensor and if g is a pure tensor then `isPureTensor()` returns a vector v with $v = a*\text{gen}(1)+b*\text{gen}(2) = (a,b)^T$ with $a(X) b = g$.

Purpose: Checks whether a given polynomial in \mathbb{A}^{env} is a pure tensor. This is also an auxiliary procedure for checking total divisibility.

Example:

```
LIB "bimodules.lib";
ring r = 0,(x,s),dp;
def R = nc_algebra(1,s); setring R; //1st shift algebra
def Re = envelope(R); setring Re; //basing is now  $R^{\text{env}} = R(X) R^{\text{opp}}$ 
poly p = x*(x*s)*x + s^2*x; p;
  ↳ x3s+x2s+xs2+2s2
// p is of the form q(X)1, a pure tensor indeed:
isPureTensor(p);
  ↳ x3s*gen(1)+x2s*gen(1)+xs2*gen(1)+2s2*gen(1)+gen(2)
// v = transpose( x3s+x2s+xs2+2s2 1 ) i.e. p = x3s+x2s+xs2+2s2 (X) 1
poly g = S*X+ x*s*X+ S^2*x;
g;
  ↳ xsX+xS2+SX
isPureTensor(g); // indeed g is not a pure tensor
  ↳ 0
poly d = x*X+s*X+x*S*X+s*S*X;d;
  ↳ xSX+xX+sSX+sX
isPureTensor(d); // d is a pure tensor indeed
  ↳ x*gen(1)+s*gen(1)+SX*gen(2)+X*gen(2)
// v = transpose( x+s S*X+X ) i.e. d = x+s (X) s*x+x
// remember that * denotes to the opposite multiplication s*x = xs in R.
```

7.5.1.6 isTwoSidedGB

Procedure from library `bimodules.lib` (see Section 7.5.1 [bimodules_lib], page 365).

Usage: `isTwoSidedGB(I)`; I ideal

Return: Returns 0 if the generators of a given ideal are not two-sided, 1 if they are.

Note: This procedure should only be used for non-commutative rings, as every element is two-sided in a commutative ring.

Purpose: Auxiliary procedure for diagonal forms. Let R be a non-commutative ring (e.g. G -algebra), and p in R , this program checks whether p is two-sided i.e. $Rp = pR$.

Example:

```
LIB "bimodules.lib";
ring r = 0,(x,s),dp;
def R = nc_algebra(1,s); setring R; //1st shift algebra
ideal I = s^2, x*s, s^2 + 3*x*s;
isTwoSidedGB(I); // I is two-sided
  ↳ 1
ideal J = s^2+x;
isTwoSidedGB(J); // J is not two-sided; twostd(J) = s,x;
  ↳ 0
```

7.5.2 bfun_lib

Library: `bfun.lib`

Purpose: Algorithms for b-functions and Bernstein-Sato polynomial

Authors: Daniel Andres, daniel.andres@math.rwth-aachen.de
 Viktor Levandovskyy, levandov@math.rwth-aachen.de

Overview: Given a polynomial ring $R = K[x_1, \dots, x_n]$ and a polynomial F in R , one is interested in the global b-function (also known as Bernstein-Sato polynomial) $b(s)$ in $K[s]$, defined to be the non-zero monic polynomial of minimal degree, satisfying a functional identity $L * F^{s+1} = b(s) F^s$, for some operator L in $D[s]$ (* stands for the action of differential operator)

By D one denotes the n -th Weyl algebra
 $K\langle x_1, \dots, x_n, d_1, \dots, d_n \mid d_j x_j = x_j d_j + 1 \rangle$.

One is interested in the following data:

- Bernstein-Sato polynomial $b(s)$ in $K[s]$,
- the list of its roots, which are known to be rational
- the multiplicities of the roots.

There is a constructive definition of a b-function of a holonomic ideal I in D (that is, an ideal I in a Weyl algebra D , such that D/I is holonomic module) with respect to the given weight vector w : For a polynomial p in D , its initial form w.r.t. $(-w, w)$ is defined as the sum of all terms of p which have maximal weighted total degree where the weight of x_i is $-w_i$ and the weight of d_i is w_i . Let J be the initial ideal of I w.r.t. $(-w, w)$, i.e. the K -vector space generated by all initial forms w.r.t. $(-w, w)$ of elements of I . Put $s = w_1 x_1 d_1 + \dots + w_n x_n d_n$. Then the monic generator $b_w(s)$ of the intersection of J with the PID $K[s]$ is called the b-function of I w.r.t. w . Unlike Bernstein-Sato polynomial, general b-function with respect to arbitrary weights need not have rational roots at all. However, b-function of a holonomic ideal is known to be non-zero as well.

References:

- [SST] Saito, Sturmfels, Takayama: Groebner Deformations of Hypergeometric Differential Equations (2000),
 Noro: An Efficient Modular Algorithm for Computing the Global b-function, (2002).

Procedures: See also: Section 7.5.4 [dmod.lib], page 395; Section 7.5.5 [dmodapp.lib], page 414; Section 7.5.7 [dmodvar.lib], page 446; Section D.6.13 [gmssing.lib], page 862.

7.5.2.1 bfct

Procedure from library `bfun.lib` (see Section 7.5.2 [bfun.lib], page 370).

Usage: `bfct(f [,s,t,v]);` f a poly, s, t optional ints, v an optional intvec

Return: list of ideal and intvec

Purpose: computes the roots of the Bernstein-Sato polynomial $b(s)$ for the hypersurface defined by f .

Assume: The basering is commutative and of characteristic 0.

Background:

In this proc, the initial Malgrange ideal is computed according to the algorithm by Masayuki Noro and then a system of linear equations is solved by linear reductions.

Note: In the output list, the ideal contains all the roots and the intvec their multiplicities.

If $s <> 0$, `std` is used for GB computations, otherwise, and by default, `slimgb` is used.
 If $t <> 0$, a matrix ordering is used for Groebner basis computations, otherwise, and by default, a block ordering is used.
 If v is a positive weight vector, v is used for homogenization computations, otherwise and by default, no weights are used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "bfun.lib";
ring r = 0, (x,y), dp;
poly f = x^2+y^3+x*y^2;
bfct(f);
⇒ [1]:
⇒  _[1]=-5/6
⇒  _[2]=-1
⇒  _[3]=-7/6
⇒ [2]:
⇒  1,1,1
intvec v = 3,2;
bfct(f,1,0,v);
⇒ [1]:
⇒  _[1]=-5/6
⇒  _[2]=-1
⇒  _[3]=-7/6
⇒ [2]:
⇒  1,1,1
```

7.5.2.2 bfctSyz

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `bfctSyz(f [,r,s,t,u,v]);` f poly, r,s,t,u optional ints, v opt. intvec

Return: list of ideal and intvec

Purpose: computes the roots of the Bernstein-Sato polynomial $b(s)$ for the hypersurface defined by f

Assume: The basering is commutative and of characteristic 0.

Background:

In this proc, the initial Malgrange ideal is computed according to the algorithm by Masayuki Noro and then a system of linear equations is solved by computing syzygies.

Note: In the output list, the ideal contains all the roots and the intvec their multiplicities.

If $r <> 0$, `std` is used for GB computations in characteristic 0, otherwise, and by default, `slimgb` is used.

If $s <> 0$, a matrix ordering is used for GB computations, otherwise, and by default, a block ordering is used.

If $t <> 0$, the computation of the intersection is solely performed over characteristic 0, otherwise and by default, a modular method is used.

If $u < 0$ and by default, `std` is used for GB computations in characteristic > 0 , otherwise, `slimgb` is used.

If v is a positive weight vector, v is used for homogenization computations, otherwise and by default, no weights are used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "bfun.lib";
ring r = 0, (x,y), dp;
poly f = x^2+y^3+x*y^2;
bfctSyz(f);
↳ [1]:
↳   _[1]==-5/6
↳   _[2]==-1
↳   _[3]==-7/6
↳ [2]:
↳   1,1,1
intvec v = 3,2;
bfctSyz(f,0,1,1,0,v);
↳ [1]:
↳   _[1]==-5/6
↳   _[2]==-1
↳   _[3]==-7/6
↳ [2]:
↳   1,1,1
```

7.5.2.3 bfctAnn

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `bfctAnn(f [,a,b,c]);` f a poly, a , b , c optional ints

Return: list of ideal and intvec

Purpose: computes the roots of the Bernstein-Sato polynomial $b(s)$ for the hypersurface defined by f .

Assume: The basering is commutative and of characteristic 0.

Background:

In this proc, $\text{Ann}(f^s)$ is computed and then a system of linear equations is solved by linear reductions.

Note: In the output list, the ideal contains all the roots and the intvec their multiplicities.

If $a < 0$, only f is appended to $\text{Ann}(f^s)$, otherwise, and by default, f and all its partial derivatives are appended.

If $b < 0$, `std` is used for GB computations, otherwise, and by default, `slimgb` is used.

If $c < 0$, `std` is used for Groebner basis computations of ideals $\langle I+J \rangle$ when I is already a Groebner basis of $\langle I \rangle$.

Otherwise, and by default the engine determined by the switch b is used.

Note that in the case $c < 0$, the choice for b will be overwritten only for the types of ideals mentioned above.

This means that if $b < 0$, specifying c has no effect.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "bfun.lib";
ring r = 0, (x,y), dp;
poly f = x^2+y^3+x*y^2;
bfctAnn(f);
↳ [1]:
↳  _[1]=-5/6
↳  _[2]=-1
↳  _[3]=-7/6
↳ [2]:
↳  1,1,1
def R = reiffen(4,5); setring R;
RC; // the Reiffen curve in 4,5
↳ xy4+y5+x4
bfctAnn(RC,0,1);
↳ [1]:
↳  _[1]=-9/20
↳  _[2]=-11/20
↳  _[3]=-13/20
↳  _[4]=-7/10
↳  _[5]=-17/20
↳  _[6]=-9/10
↳  _[7]=-19/20
↳  _[8]=-1
↳  _[9]=-21/20
↳  _[10]=-11/10
↳  _[11]=-23/20
↳  _[12]=-13/10
↳  _[13]=-27/20
↳ [2]:
↳  1,1,1,1,1,1,1,1,1,1,1,1,1
```

7.5.2.4 bfctOneGB

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `bfctOneGB(f [,s,t]);` `f` a poly, `s,t` optional ints

Return: list of ideal and intvec

Purpose: computes the roots of the Bernstein-Sato polynomial $b(s)$ for the hypersurface defined by `f`, using only one GB computation

Assume: The basering is commutative and of characteristic 0.

Background:

In this proc, the initial Malgrange ideal is computed based on the algorithm by Masayuki Noro and combined with an elimination ordering.

Note: In the output list, the ideal contains all the roots and the intvec their multiplicities.

If `s<>0`, `std` is used for the GB computation, otherwise, and by default, `slimgb` is used.

If $t < 0$, a matrix ordering is used for GB computations, otherwise, and by default, a block ordering is used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel ≥ 2`, all the debug messages will be printed.

Example:

```
LIB "bfun.lib";
ring r = 0, (x,y), dp;
poly f = x^2+y^3+x*y^2;
bfctOneGB(f);
↳ [1]:
↳   _[1] = -5/6
↳   _[2] = -1
↳   _[3] = -7/6
↳ [2]:
↳   1, 1, 1
bfctOneGB(f, 1, 1);
↳ [1]:
↳   _[1] = -5/6
↳   _[2] = -1
↳   _[3] = -7/6
↳ [2]:
↳   1, 1, 1
```

7.5.2.5 bfctIdeal

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `bfctIdeal(I, w[s,t]);` I an ideal, w an intvec, s,t optional ints

Return: list of ideal and intvec

Purpose: computes the roots of the global b-function of I w.r.t. the weight $(-w, w)$.

Assume: The basering is the n -th Weyl algebra in characteristic 0 and for all $1 ≤ i ≤ n$ the identity $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$ holds, i.e. the sequence of variables is given by $x(1), \dots, x(n), D(1), \dots, D(n)$, where $D(i)$ is the differential operator belonging to $x(i)$. Further we assume that I is holonomic.

Background:

In this proc, the initial ideal of I is computed according to the algorithm by Masayuki Noro and then a system of linear equations is solved by linear reductions.

Note: In the output list, say L,

- L[1] of type ideal contains all the rational roots of a b-function,
- L[2] of type intvec contains the multiplicities of above roots,
- optional L[3] of type string is the part of b-function without rational roots.

Note, that a b-function of degree 0 is encoded via $L[1][1]=0$, $L[2]=0$ and $L[3]$ is 1 (for nonzero constant) or 0 (for zero b-function).

If $s < 0$, `std` is used for GB computations in characteristic 0, otherwise, and by default, `slimgb` is used.

If $t < 0$, a matrix ordering is used for GB computations, otherwise, and by default, a block ordering is used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "bfun.lib";
ring @D = 0, (x,y,Dx,Dy), dp;
def D = Weyl();
setring D;
ideal I = 3*x^2*Dy+2*y*Dx, 2*x*Dx+3*y*Dy+6; I = std(I);
intvec w1 = 0,1;
intvec w2 = 2,3;
bfctIdeal(I,w1);
⇒ [1]:
⇒  _[1]=0
⇒  _[2]=-2/3
⇒  _[3]=-4/3
⇒ [2]:
⇒  1,1,1
bfctIdeal(I,w2,0,1);
⇒ [1]:
⇒  _[1]=-6
⇒ [2]:
⇒  1
ideal J = I[size(I)]; // J is not holonomic by construction
bfctIdeal(J,w1); // b-function of D/J w.r.t. w1 is non-zero
⇒ WARNING: given ideal is not holonomic
⇒ ... setting bound for degree of b-function to 10 and proceeding
⇒ [1]:
⇒  _[1]=0
⇒  _[2]=-2/3
⇒  _[3]=-4/3
⇒ [2]:
⇒  1,1,1
bfctIdeal(J,w2); // b-function of D/J w.r.t. w2 is zero
⇒ WARNING: given ideal is not holonomic
⇒ ... setting bound for degree of b-function to 10 and proceeding
⇒ // Intersection is zero
⇒ [1]:
⇒  _[1]=0
⇒ [2]:
⇒  0
⇒ [3]:
⇒  0
```

7.5.2.6 pIntersect

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `pIntersect(f, I [,s]);` `f` a poly, `I` an ideal, `s` an optional int

Return: vector, coefficient vector of the monic polynomial

Purpose: compute the intersection of ideal `I` with the subalgebra `K[f]`

Assume: `I` is given as Groebner basis, basering is not a qring.

Note: If the intersection is zero, this proc might not terminate.
If $s > 0$ is given, it is searched for the generator of the intersection only up to degree s . Otherwise (and by default), no bound is assumed.

Display: If `printlevel=1`, progress debug messages will be printed,
if `printlevel ≥ 2`, all the debug messages will be printed.

Example:

```
LIB "bfun.lib";
ring r = 0, (x,y), dp;
poly f = x^2+y^3+xy^2;
def D = initialMalgrange(f);
setring D;
inF;
↳ inF[1]=x*Dt
↳ inF[2]=2*x*y*Dx+3*y^2*Dx-y^2*Dy-2*x*Dy
↳ inF[3]=2*x^2*Dx+xy*Dx+xy*Dy+18*t*Dt+9*x*Dx-x*Dy+6*y*Dy+4*x+18
↳ inF[4]=18*t*Dt^2+6*y*Dt*Dy-y*Dt+27*Dt
↳ inF[5]=y^2*Dt
↳ inF[6]=2*t*y*Dt+2*x*y*Dx+2*y^2*Dx-6*t*Dt-3*x*Dx-x*Dy-2*y*Dy+2*y-6
↳ inF[7]=x*y^2+y^3+x^2
↳ inF[8]=2*y^3*Dx-2*y^3*Dy-3*y^2*Dx-2*x*y*Dy+y^2*Dy-4*y^2+36*t*Dt+18*x*Dx+1\
2*y*Dy+36
pIntersect(t*Dt, inF);
↳ gen(4)-1/36*gen(2)
pIntersect(t*Dt, inF, 1);
↳ // Try a bound of at least 2
↳ 0
```

7.5.2.7 pIntersectSyz

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `pIntersectSyz(f, I [,p,s,t]);` f poly, I ideal, p, t optial ints, p prime

Return: vector, coefficient vector of the monic polynomial

Purpose: compute the intersection of an ideal I with the subalgebra $K[f]$

Assume: I is given as Groebner basis.

Note: If the intersection is zero, this procedure might not terminate.
If $p > 0$ is given, this proc computes the generator of the intersection in char p first and then only searches for a generator of the obtained degree in the basering. Otherwise, it searches for all degrees by computing syzygies.
If $s < > 0$, `std` is used for Groebner basis computations in char 0, otherwise, and by default, `slimgb` is used.
If $t < > 0$ and by default, `std` is used for Groebner basis computations in char > 0 , otherwise, `slimgb` is used.

Display: If `printlevel=1`, progress debug messages will be printed,
if `printlevel ≥ 2`, all the debug messages will be printed.

Example:

```

LIB "bfun.lib";
ring r = 0, (x,y), dp;
poly f = x^2+y^3+x*y^2;
def D = initialMalgrange(f);
setring D;
inF;
↳ inF[1]=x*Dt
↳ inF[2]=2*x*y*Dx+3*y^2*Dx-y^2*Dy-2*x*Dy
↳ inF[3]=2*x^2*Dx+x*y*Dx+x*y*Dy+18*t*Dt+9*x*Dx-x*Dy+6*y*Dy+4*x+18
↳ inF[4]=18*t*Dt^2+6*y*Dt*Dy-y*Dt+27*Dt
↳ inF[5]=y^2*Dt
↳ inF[6]=2*t*y*Dt+2*x*y*Dx+2*y^2*Dx-6*t*Dt-3*x*Dx-x*Dy-2*y*Dy+2*y-6
↳ inF[7]=x*y^2+y^3+x^2
↳ inF[8]=2*y^3*Dx-2*y^3*Dy-3*y^2*Dx-2*x*y*Dy+y^2*Dy-4*y^2+36*t*Dt+18*x*Dx+1\
    2*y*Dy+36
poly s = t*Dt;
pIntersectSyz(s, inF);
↳ gen(4)-1/36*gen(2)
int p = prime(20000);
pIntersectSyz(s, inF, p, 0, 0);
↳ gen(4)-1/36*gen(2)

```

7.5.2.8 linReduce

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `linReduce(f, I [,s,t,u]);` f a poly, I an ideal, s,t,u optional ints

Return: poly or list, linear reductum (over field) of f by elements from I

Purpose: reduce a polynomial only by linear reductions (no monomial multiplications)

Note: If $s < 0$, a list consisting of the reduced polynomial and the coefficient vector of the used reductions is returned, otherwise (and by default) only reduced polynomial is returned.

If $t < 0$ (and by default) all monomials are reduced (if possible), otherwise, only leading monomials are reduced.

If $u < 0$ (and by default), the ideal is linearly pre-reduced, i.e. instead of the given ideal, the output of `linReduceIdeal` is used.

If u is set to 0 and the given ideal does not equal the output of `linReduceIdeal`, the result might not be as expected.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "bfun.lib";
ring r = 0, (x,y), dp;
ideal I = 1,y,xy;
poly f = 5xy+7y+3;
poly g = 7x+5y+3;
linReduce(g,I); // reduces tails
↳ 7x
linReduce(g,I,0,0); // no reductions of tails
↳ 7x+5y+3

```

```

linReduce(f,I,1); // reduces tails and shows reductions used
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ -5*gen(3)-7*gen(2)-3*gen(1)
f = x3+y2+x2+y+x;
I = x3-y3, y3-x2,x3-y2,x2-y,y2-x;
list l = linReduce(f,I,1);
l;
⇒ [1]:
⇒ 5y
⇒ [2]:
⇒ gen(5)-4*gen(4)+2*gen(3)-3*gen(2)-3*gen(1)
module M = I;
f - (l[1]-(M*l[2])[1,1]);
⇒ 0

```

7.5.2.9 linReduceIdeal

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `linReduceIdeal(I [,s,t,u]);` I an ideal, s,t,u optional ints

Return: ideal or list, linear reductum (over field) of I by its elements

Purpose: reduces a list of polys only by linear reductions (no monomial multiplications)

Note: If $s < 0$, a list consisting of the reduced ideal and the coefficient vectors of the used reductions given as module is returned. Otherwise (and by default), only the reduced ideal is returned. If $t < 0$ (and by default) all monomials are reduced (if possible), otherwise, only leading monomials are reduced. If $u < 0$ (and by default), the ideal is first sorted in increasing order. If u is set to 0 and the given ideal is not sorted in the way described, the result might not be as expected.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "bfun.lib";
ring r = 0, (x,y), dp;
ideal I = 3,x+9,y4+5x,2y4+7x+2;
linReduceIdeal(I); // reduces tails
⇒ _[1]=0
⇒ _[2]=3
⇒ _[3]=x
⇒ _[4]=y4
linReduceIdeal(I,0,0); // no reductions of tails
⇒ _[1]=0
⇒ _[2]=3
⇒ _[3]=x+9
⇒ _[4]=y4+5x
list l = linReduceIdeal(I,1); // reduces tails and shows reductions used

```

```

1;
↳ [1] :
↳  _[1]=0
↳  _[2]=3
↳  _[3]=x
↳  _[4]=y4
↳ [2] :
↳  _[1]=gen(4)-2*gen(3)+3*gen(2)-29/3*gen(1)
↳  _[2]=gen(1)
↳  _[3]=gen(2)-3*gen(1)
↳  _[4]=gen(3)-5*gen(2)+15*gen(1)
module M = I;
matrix(l[1]) - matrix(M)*matrix(l[2]);
↳  _[1,1]=0
↳  _[1,2]=0
↳  _[1,3]=0
↳  _[1,4]=0

```

7.5.2.10 linSyzSolve

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

- Usage:** `linSyzSolve(I[,s]);` I an ideal, s an optional int
- Return:** vector, coefficient vector of linear combination of 0 in elements of I
- Purpose:** compute a linear dependency between the elements of an ideal if such one exists
- Note:** If `s<>0`, `std` is used for Groebner basis computations, otherwise, `slimgb` is used.
By default, `slimgb` is used in char 0 and `std` in char >0.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "bfun.lib";
ring r = 0,x,dp;
ideal I = x,2x;
linSyzSolve(I);
↳ gen(2)-2*gen(1)
ideal J = x,x2;
linSyzSolve(J);
↳ 0

```

7.5.2.11 allPositive

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

- Usage:** `allPositive(v);` v an intvec
- Return:** int, 1 if all components of v are positive, or 0 otherwise
- Purpose:** check whether all components of an intvec are positive
- Example:**

```

LIB "bfun.lib";
intvec v = 1,2,3;
allPositive(v);
↳ 1
intvec w = 1,-2,3;
allPositive(w);
↳ 0

```

7.5.2.12 scalarProd

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `scalarProd(v,w)`; v,w intvecs

Return: int, the standard scalar product of v and w

Purpose: computes the scalar product of two intvecs

Assume: the arguments are of the same size

Example:

```

LIB "bfun.lib";
intvec v = 1,2,3;
intvec w = 4,5,6;
scalarProd(v,w);
↳ 32

```

7.5.2.13 vec2poly

Procedure from library `bfun.lib` (see Section 7.5.2 [`bfun.lib`], page 370).

Usage: `vec2poly(v [,i])`; v a vector or an intvec, i an optional int

Return: poly, an univariate polynomial in i -th variable with coefficients given by v

Purpose: constructs an univariate polynomial in $K[\text{var}(i)]$ with given coefficients, such that the coefficient at $\text{var}(i)^{\{j-1\}}$ is $v[j]$.

Note: The optional argument i must be positive, by default i is 1.

Example:

```

LIB "bfun.lib";
ring r = 0,(x,y),dp;
vector v = gen(1) + 3*gen(3) + 22/9*gen(4);
intvec iv = 3,2,1;
vec2poly(v,2);
↳ 22/9y3+3y2+1
vec2poly(iv);
↳ x2+2x+3

```

7.5.3 central_lib

Library: `central.lib`

Purpose: Computation of central elements of GR-algebras

Author: Oleksandr Motsak, U@D, where $U=\{\text{motsak}\}$, $D=\{\text{mathematik.uni-kl.de}\}$

Overview: A library for computing elements of the center and centralizers of sets of elements in GR-algebras.

Procedures:

7.5.3.1 centralizeSet

Procedure from library `central.lib` (see Section 7.5.3 [`central.lib`], page 381).

Usage: `centralizeSet(F, V);` F, V ideals

Input: F, V finite sets of elements of the base algebra

Return: ideal, generated by computed elements

Purpose: computes a vector space basis of the centralizer of the set F in the vector space generated by V over the ground field

Example:

```
LIB "central.lib";
ring A = 0, (e(1..4)), dp;
matrix D[4][4]=0;
D[2,4] = -e(1);
D[3,4] = -e(2);
// This is A_4_1 - the first real Lie algebra of dimension 4.
def A_4_1 = nc_algebra(1,D); setring A_4_1;
ideal F = variablesSorted(); F;
  => F[1]=e(1)
  => F[2]=e(4)
  => F[3]=e(3)
  => F[4]=e(2)
// the center of A_4_1 is generated by
// e(1) and -1/2*e(2)^2+e(1)*e(3)
// therefore one may consider computing it in the following way:
// 1. Compute a PBW basis consisting of
//    monomials with exponent <= (1,2,1,0)
ideal V = PBW_maxMonom( e(1) * e(2)^2 * e(3) );
// 2. Compute the centralizer of F within the vector space
//    spanned by these monomials:
ideal C = centralizeSet( F, V ); C;
  => C[1]=e(1)
  => C[2]=e(2)^2-2*e(1)*e(3)
inCenter(C); // check the result
  => 1
```

See also: Section 7.5.3.7 [`centralizer`], page 386; Section 7.5.3.2 [`centralizerVS`], page 382; Section 7.5.3.11 [`inCentralizer`], page 388.

7.5.3.2 centralizerVS

Procedure from library `central.lib` (see Section 7.5.3 [`central.lib`], page 381).

Usage: `centralizerVS(F, D);` F ideal, D int

Return: ideal, generated by computed elements

Purpose: computes a vector space basis of `centralizer(F)` up to degree D

Note: D must be non-negative

Example:

```
LIB "central.lib";
ring AA = 0, (x,y,z), dp;
```



```

matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
ideal F = x, y;
// find generators of the vector space of elements
// of degree <= 4 commuting with x and y:
ideal C = centralizerVS(F, 4);
C;
⇒ C[1]=4xy+z2-2z
⇒ C[2]=16x2y2+8xyz2+z4-32xyz-4z3-4z2+16z
inCentralizer(C, F); // check the result
⇒ 1

```

See also: Section 7.5.3.4 [centerVS], page 383; Section 7.5.3.7 [centralizer], page 386; Section 7.5.3.11 [inCentralizer], page 388.

7.5.3.3 centralizerRed

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `centralizerRed(F, D[, N]);` F ideal, D int, N optional int

Return: ideal, generated by computed elements

Purpose: computes subalgebra generators of `centralizer(F)` up to degree D.

Note: In general, one cannot compute the whole `centralizer(F)`.
Hence, one has to specify a termination condition via arguments D and/or N.
If D is positive, only centralizing elements up to degree D are computed.
If D is negative, the termination is determined by N only.
If N is given, the computation stops if at least N elements have been found.
Warning: if N is given and bigger than the actual number of generators,
the procedure may not terminate.
Current ordering must be a degree compatible well-ordering.

Example:

```

LIB "central.lib";
ring AA = 0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
ideal F = x, y;
// find subalgebra generators of degree <= 4 of the subalgebra of
// all elements commuting with x and y:
ideal C = centralizerRed(F, 4);
C;
⇒ C[1]=4xy+z2-2z
inCentralizer(C, F); // check the result
⇒ 1

```

See also: Section 7.5.3.5 [centerRed], page 384; Section 7.5.3.7 [centralizer], page 386; Section 7.5.3.2 [centralizerVS], page 382; Section 7.5.3.11 [inCentralizer], page 388.

7.5.3.4 centerVS

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `centerVS(D);` D int

Return: ideal, generated by computed elements

Purpose: computes a vector space basis of the center up to degree D

Note: D must be non-negative

Example:

```
LIB "central.lib";
ring AA = 0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
// find a basis of the vector space of all
// central elements of degree <= 4:
ideal Z = centerVS(4);
Z;
↪ Z[1]=4xy+z2-2z
↪ Z[2]=16x2y2+8xyz2+z4-32xyz-4z3-4z2+16z
// note that the second element is the square of the first
// plus a multiple of the first:
Z[2] - Z[1]^2 + 8*Z[1];
↪ 0
inCenter(Z); // check the result
↪ 1
```

See also: Section 7.5.3.6 [center], page 385; Section 7.5.3.2 [centralizerVS], page 382; Section 7.5.3.10 [inCenter], page 388.

7.5.3.5 centerRed

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `centerRed(D[, N]);` D int, N optional int

Return: ideal, generated by computed elements

Purpose: computes subalgebra generators of the center up to degree D

Note: In general, one cannot compute the whole center.
Hence, one has to specify a termination condition via arguments D and/or N.
If D is positive, only central elements up to degree D will be found.
If D is negative, the termination is determined by N only.
If N is given, the computation stops if at least N elements have been found.
Warning: if N is given and bigger than the actual number of generators,
the procedure may not terminate.
Current ordering must be a degree compatible well-ordering.

Example:

```
LIB "central.lib";
ring AA = 0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=z;
def A = nc_algebra(1,D); setring A; // it is a Heisenberg algebra
// find a basis of the vector space of
// central elements of degree <= 3:
```

```

ideal VSZ = centerVS(3);
// There should be 3 degrees of z.
VSZ;
↪ VSZ[1]=z
↪ VSZ[2]=z2
↪ VSZ[3]=z3
inCenter(VSZ); // check the result
↪ 1
// find "minimal" central elements of degree <= 3
ideal SAZ = centerRed(3);
// Only 'z' must be computed
SAZ;
↪ SAZ[1]=z
inCenter(SAZ); // check the result
↪ 1

```

See also: Section 7.5.3.6 [center], page 385; Section 7.5.3.4 [centerVS], page 383; Section 7.5.3.3 [centralizerRed], page 383; Section 7.5.3.10 [inCenter], page 388.

7.5.3.6 center

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `center(D[, N]);` D int, N optional int

Return: ideal, generated by computed elements

Purpose: computes subalgebra generators of the center up to degree D

Note: In general, one cannot compute the whole center.
Hence, one has to specify a termination condition via arguments D and/or N.
If D is positive, only central elements up to degree D will be found.
If D is negative, the termination is determined by N only.
If N is given, the computation stops if at least N elements have been found.
Warning: if N is given and bigger than the actual number of generators,
the procedure may not terminate.
Current ordering must be a degree compatible well-ordering.

Example:

```

LIB "central.lib";
ring AA = 0, (x,y,z,t), dp;
matrix D[4][4]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2) tensored with K[t]
// find generators of the center of degree <= 3:
ideal Z = center(3);
Z;
↪ Z[1]=t
↪ Z[2]=4xy+z2-2z
inCenter(Z); // check the result
↪ 1
// find at least one generator of the center:
ideal Z2 = center(-1, 1);
Z2;
↪ Z2[1]=t
inCenter(Z2); // check the result

```

↪ 1

See also: Section 7.5.3.7 [centralizer], page 386; Section 7.5.3.10 [inCenter], page 388.

7.5.3.7 centralizer

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `centralizer(F, D[, N]);` F poly/ideal, D int, N optional int

Return: ideal, generated by computed elements

Purpose: computes subalgebra generators of `centralizer(F)` up to degree D

Note: In general, one cannot compute the whole `centralizer(F)`.
Hence, one has to specify a termination condition via arguments D and/or N.
If D is positive, only centralizing elements up to degree D will be found.
If D is negative, the termination is determined by N only.
If N is given, the computation stops if at least N elements have been found.
Warning: if N is given and bigger than the actual number of generators, the procedure may not terminate.
Current ordering must be a degree compatible well-ordering.

Example:

```
LIB "central.lib";
ring AA = 0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
poly f = 4*x*y+z^2-2*z; // a central polynomial
f;
↪ 4xy+z2-2z
// find generators of the centralizer of f of degree <= 2:
ideal c = centralizer(f, 2);
c; // since f is central, the answer consists of generators of A
↪ c[1]=z
↪ c[2]=y
↪ c[3]=x
inCentralizer(c, f); // check the result
↪ 1
// find at least two generators of the centralizer of f:
ideal cc = centralizer(f,-1,2);
cc;
↪ cc[1]=z
↪ cc[2]=y
↪ cc[3]=x
inCentralizer(cc, f); // check the result
↪ 1
poly g = z^2-2*z; // some non-central polynomial
// find generators of the centralizer of g of degree <= 2:
c = centralizer(g, 2);
c;
↪ c[1]=z
↪ c[2]=xy
inCentralizer(c, g); // check the result
↪ 1
```

```

// find at least one generator of the centralizer of g:
centralizer(g,-1,1);
↳ _[1]=z
// find at least two generators of the centralizer of g:
cc = centralizer(g,-1,2);
cc;
↳ cc[1]=z
↳ cc[2]=xy
inCentralizer(cc, g); // check the result
↳ 1

```

See also: Section 7.5.3.6 [center], page 385; Section 7.5.3.11 [inCentralizer], page 388.

7.5.3.8 sa_reduce

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `sa_reduce(V)`; V ideal

Return: ideal, generated by computed elements

Purpose: compute a subalgebra basis of an algebra generated by the elements of V

Note: At the moment the usage of this procedure is limited to G -algebras

Example:

```

LIB "central.lib";
ring AA = 0, (x,y,z), dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
poly f = 4*x*y+z^2-2*z; // a central polynomial
ideal I = f, f*f, f*f*f - 10*f*f, f+3*z^3; I;
↳ I[1]=4xy+z2-2z
↳ I[2]=16x2y2+8xyz2+z4-32xyz-4z3+32xy+4z2
↳ I[3]=64x3y3+48x2y2z2+12xyz4+z6-288x2y2z-96xyz3-6z5+352x2y2+224xyz2+2z4-12\
8xyz+32z3-64xy-40z2
↳ I[4]=3z3+4xy+z2-2z
sa_reduce(I); // should be just f and z^3
↳ _[1]=4xy+z2-2z
↳ _[2]=z3

```

See also: Section 7.5.3.9 [sa_poly_reduce], page 387.

7.5.3.9 sa_poly_reduce

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `sa_poly_reduce(p, V)`; p poly, V ideal

Return: polynomial, a reduction of p w.r.t. V

Purpose: computes a reduction of the polynomial p w.r.t. the subalgebra generated by elements of V

Note: At the moment the usage of this procedure is limited to G -algebras

Example:

```

LIB "central.lib";
ring AA = 0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
poly f = 4*x*y+z^2-2*z; // a central polynomial
sa_poly_reduce(f + 3*f*f + x, ideal(f) ); // should be just 'x'
↳ x

```

See also: Section 7.5.3.8 [sa_reduce], page 387.

7.5.3.10 inCenter

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `inCenter(E)`; E poly/list/ideal

Return: integer, 1 if E is in the center, 0 otherwise

Purpose: check whether the elements of E are central

Example:

```

LIB "central.lib";
ring R=0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z;
D[1,3]=2*x;
D[2,3]=-2*y;
def r = nc_algebra(1,D); setring r; // this is U(sl_2)
poly p=4*x*y+z^2-2*z;
inCenter(p);
↳ 1
poly f=4*x*y;
inCenter(f);
↳ 0
list l= list( 1, p, p^2, p^3);
inCenter(l);
↳ 1
ideal I= p, f;
inCenter(I);
↳ 0

```

7.5.3.11 inCentralizer

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `inCentralizer(E, S)`; E poly/list/ideal, S poly/ideal

Return: integer, 1 if E is in the centralizer(S), 0 otherwise

Purpose: check whether the elements of E are in the centralizer(S)

Example:

```

LIB "central.lib";
ring R = 0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z;

```

```

def r = nc_algebra(1,D); setring r; // the Heisenberg algebra
poly f = x^2;
poly a = z; // 'z' is central => it lies in every centralizer!
poly b = y^2;
inCentralizer(a, f);
↳ 1
inCentralizer(b, f);
↳ 0
list l = list(1, a);
inCentralizer(l, f);
↳ 1
ideal I = a, b;
inCentralizer(I, f);
↳ 0
printlevel = 2;
inCentralizer(a, f); // yes
↳ 1
inCentralizer(b, f); // no
↳ [1]:
↳ POLY: y2 is NOT in the centralizer of polynomial {x2}
↳ 0

```

7.5.3.12 isCartan

Procedure from library `central.lib` (see Section 7.5.3 [`central.lib`], page 381).

Usage: `isCartan(f);` f poly

Purpose: check whether f is a Cartan element.

Return: integer, 1 if f is a Cartan element and 0 otherwise.

Note: f is a Cartan element of the algebra A
if and only if for all g in A there exists C in K such that $[f, g] = C * g$
if and only if for all variables v_i there exist C in K such that $[f, v_i] = C * v_i$.

Example:

```

LIB "central.lib";
ring R=0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z;
D[1,3]=2*x;
D[2,3]=-2*y;
def r = nc_algebra(1,D); setring r; // this is U(sl_2) with cartan - z
isCartan(z); // yes!
↳ 1
poly p=4*x*y+z^2-2*z;
isCartan(p); // central elements are Cartan elements!
↳ 1
poly f=4*x*y;
isCartan(f); // no way!
↳ 0
isCartan( 10 + p + z ); // scalar + central + cartan
↳ 1

```

7.5.3.13 applyAdF

Procedure from library `central.lib` (see Section 7.5.3 [`central.lib`], page 381).

Usage: `applyAdF(B, f)`; B ideal, f poly

Purpose: Apply Ad_f to every element of B

Return: ideal, generated by $\text{Ad}_f(B[i])$, $1 \leq i \leq \text{size}(B)$

Note: $\text{Ad}_f(v) := [f, v] = f*v - v*f$

Example:

```
LIB "central.lib";
ring AA = 0, (e, f, h), dp;
matrix D[3][3]=0;
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
// Let us consider the linear map Ad_{e} from A_2 into A.
// Compute the PBW basis of A_2:
ideal Basis = PBW_maxDeg( 2 ); Basis;
↳ Basis[1]=e
↳ Basis[2]=f
↳ Basis[3]=h
↳ Basis[4]=h^2
↳ Basis[5]=fh
↳ Basis[6]=f^2
↳ Basis[7]=eh
↳ Basis[8]=ef
↳ Basis[9]=e^2
// Compute images of basis elements under the linear map Ad_e:
ideal Image = applyAdF( Basis, e ); Image;
↳ Image[1]=0
↳ Image[2]=h
↳ Image[3]=-2e
↳ Image[4]=-4eh-4e
↳ Image[5]=-2ef+h^2+2h
↳ Image[6]=2fh-2f
↳ Image[7]=-2e^2
↳ Image[8]=eh
↳ Image[9]=0
// Now we have a linear map given by: Basis_i --> Image_i
// Let's compute its kernel K:
// 1. compute syzygy module C:
module C = linearMapKernel( Image ); C;
↳ C[1]=gen(1)
↳ C[2]=gen(8)+1/4*gen(4)-1/2*gen(3)
↳ C[3]=gen(9)
// 2. compute corresponding combinations of basis vectors:
ideal K = linearCombinations(Basis, C); K;
↳ K[1]=e
↳ K[2]=ef+1/4h^2-1/2h
↳ K[3]=e^2
// Let's check that Ad_e(K) is zero:
applyAdF( K, e );
↳ _[1]=0
```



```

↳ _[2]=0
↳ _[3]=0

```

See also: Section 7.5.3.14 [linearMapKernel], page 391.

7.5.3.14 linearMapKernel

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `linearMapKernel(Images);` Images ideal

Purpose: Computes the syzygy module of the linear map given by Images.

Return: syzygy module, or `int(0)` if all images are zeroes

Example:

```

LIB "central.lib";
ring AA = 0, (e, f, h), dp;
matrix D[3][3]=0;
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
def A = nc_algebra(1,D); // this algebra is U(sl_2)
setring A;
// Let us consider the linear map Ad_{e} from A_2 into A.
// Compute the PBW basis of A_2:
ideal Basis = PBW_maxDeg( 2 ); Basis;
↳ Basis[1]=e
↳ Basis[2]=f
↳ Basis[3]=h
↳ Basis[4]=h2
↳ Basis[5]=fh
↳ Basis[6]=f2
↳ Basis[7]=eh
↳ Basis[8]=ef
↳ Basis[9]=e2
// Compute images of basis elements under the linear map Ad_e:
ideal Image = applyAdF( Basis, e ); Image;
↳ Image[1]=0
↳ Image[2]=h
↳ Image[3]=-2e
↳ Image[4]=-4eh-4e
↳ Image[5]=-2ef+h2+2h
↳ Image[6]=2fh-2f
↳ Image[7]=-2e2
↳ Image[8]=eh
↳ Image[9]=0
// Now we have a linear map given by: Basis_i --> Image_i
// Let's compute its kernel K:
// 1. compute syzygy module C:
module C = linearMapKernel( Image ); C;
↳ C[1]=gen(1)
↳ C[2]=gen(8)+1/4*gen(4)-1/2*gen(3)
↳ C[3]=gen(9)
// 2. compute corresponding combinations of basis vectors:
ideal K = linearCombinations(Basis, C); K;
↳ K[1]=e
↳ K[2]=ef+1/4h2-1/2h

```

```

↳ K[3]=e2
// Let's check that Ad_e(K) is zero:
ideal Z = applyAdF( K, e ); Z;
↳ Z[1]=0
↳ Z[2]=0
↳ Z[3]=0
// Now linearMapKernel will return a single integer 0:
def CC = linearMapKernel(Z); typeof(CC); CC;
↳ int
↳ 0

```

See also: Section 7.5.3.13 [applyAdF], page 390; Section 7.5.3.14 [linearMapKernel], page 391.

7.5.3.15 linearCombinations

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `linearCombinations(Basis, C);` Basis ideal, C module

Purpose: forms linear combinations of elements from Basis by replacing `gen(i)` by `Basis[i]` in C

Return: ideal generated by computed linear combinations

Example:

```

LIB "central.lib";
ring AA = 0,(e,f,h),dp;
matrix D[3][3]=0;
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
// Let us consider the linear map Ad_{e} from A_2 into A.
// Compute the PBW basis of A_2:
ideal Basis = PBW_maxDeg( 2 ); Basis;
↳ Basis[1]=e
↳ Basis[2]=f
↳ Basis[3]=h
↳ Basis[4]=h2
↳ Basis[5]=fh
↳ Basis[6]=f2
↳ Basis[7]=eh
↳ Basis[8]=ef
↳ Basis[9]=e2
// Compute images of basis elements under the linear map Ad_e:
ideal Image = applyAdF( Basis, e ); Image;
↳ Image[1]=0
↳ Image[2]=h
↳ Image[3]=-2e
↳ Image[4]=-4eh-4e
↳ Image[5]=-2ef+h2+2h
↳ Image[6]=2fh-2f
↳ Image[7]=-2e2
↳ Image[8]=eh
↳ Image[9]=0
// Now we have a linear map given by: Basis_i --> Image_i
// Let's compute its kernel K:
// 1. compute syzygy module C:
module C = linearMapKernel( Image ); C;

```

```

↳ C[1]=gen(1)
↳ C[2]=gen(8)+1/4*gen(4)-1/2*gen(3)
↳ C[3]=gen(9)
// 2. compute corresponding combinations of basis vectors:
ideal K = linearCombinations(Basis, C); K;
↳ K[1]=e
↳ K[2]=ef+1/4h2-1/2h
↳ K[3]=e2
// Let's check that Ad_e(K) is zero:
applyAdF( K, e );
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0

```

See also: Section 7.5.3.13 [applyAdF], page 390; Section 7.5.3.14 [linearMapKernel], page 391.

7.5.3.16 variablesStandard

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `variablesStandard();`

Return: ideal, generated by algebra variables

Purpose: computes the set of algebra variables taken in their natural order

Example:

```

LIB "central.lib";
ring AA = 0, (x,y,z), dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
// Variables in their natural order:
variablesStandard();
↳ _[1]=x
↳ _[2]=y
↳ _[3]=z

```

See also: Section 7.5.3.17 [variablesSorted], page 393.

7.5.3.17 variablesSorted

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: `variablesSorted();`

Return: ideal, generated by sorted algebra variables

Purpose: computes the set of algebra variables sorted so that Cartan variables go first

Note: This is a heuristics for the computation of the center: it is better to compute centralizers of Cartan variables first since in this case we can omit solving the system of equations.

Example:

```

LIB "central.lib";
ring AA = 0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
// There is only one Cartan variable - z in U(sl_2),
// it must go 1st:
variablesSorted();
  ↪ _[1]=z
  ↪ _[2]=y
  ↪ _[3]=x

```

See also: Section 7.5.3.16 [variablesStandard], page 393.

7.5.3.18 PBW_eqDeg

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: PBW_eqDeg(Deg); Deg int

Purpose: Compute PBW elements of a given degree.

Return: ideal consisting of found elements.

Note: Unit is omitted. Weights are ignored!

Example:

```

LIB "central.lib";
ring AA = 0,(e,f,h),dp;
matrix D[3][3]=0;
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
// PBW Basis of A_2 \ A_1 - monomials of degree == 2:
PBW_eqDeg( 2 );
  ↪ _[1]=h2
  ↪ _[2]=fh
  ↪ _[3]=f2
  ↪ _[4]=eh
  ↪ _[5]=ef
  ↪ _[6]=e2

```

7.5.3.19 PBW_maxDeg

Procedure from library `central.lib` (see Section 7.5.3 [central.lib], page 381).

Usage: PBW_maxDeg(MaxDeg); MaxDeg int

Purpose: Compute PBW elements up to a given maximal degree.

Return: ideal consisting of found elements.

Note: unit is omitted. Weights are ignored!

Example:

```

LIB "central.lib";
ring AA = 0,(e,f,h),dp;
matrix D[3][3]=0;
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;

```

```

def A = nc_algebra(1,D); // this algebra is U(sl_2)
setring A;
// PBW Basis of A_2 - monomials of degree <= 2, without unit:
PBW_maxDeg( 2 );
↳ _[1]=e
↳ _[2]=f
↳ _[3]=h
↳ _[4]=h2
↳ _[5]=fh
↳ _[6]=f2
↳ _[7]=eh
↳ _[8]=ef
↳ _[9]=e2

```

7.5.3.20 PBW_maxMonom

Procedure from library `central.lib` (see Section 7.5.3 [`central.lib`], page 381).

Usage: PBW_maxMonom(m); m poly

Purpose: Compute PBW elements up to a given maximal one.

Return: ideal consisting of found elements.

Note: Unit is omitted. Weights are ignored!

Example:

```

LIB "central.lib";
ring AA = 0,(e,f,h),dp;
matrix D[3][3]=0;
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
def A = nc_algebra(1,D); // this algebra is U(sl_2)
setring A;
// At most 1st degree in e, h and at most 2nd degree in f, unit is omitted:
PBW_maxMonom( e*(f^2)* h );
↳ _[1]=e
↳ _[2]=f
↳ _[3]=ef
↳ _[4]=f2
↳ _[5]=ef2
↳ _[6]=h
↳ _[7]=eh
↳ _[8]=fh
↳ _[9]=efh
↳ _[10]=f2h
↳ _[11]=ef2h

```

7.5.4 dmod_lib

Library: dmod.lib

Purpose: Algorithms for algebraic D-modules

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Overview: Let K be a field of characteristic 0. Given a polynomial ring $R = K[x_1, \dots, x_n]$ and a polynomial F in R , one is interested in the $R[1/F]$ -module of rank one, generated by the symbol F^s for a symbolic discrete variable s . In fact, the module $R[1/F]^*F^s$ has a structure of a $D(R)[s]$ -module, where $D(R)$ is an n -th Weyl algebra $K\langle x_1, \dots, x_n, d_1, \dots, d_n \mid d_j x_j = x_j d_j + 1 \rangle$ and $D(R)[s] = D(R)$ tensored with $K[s]$ over K . Constructively, one needs to find a left ideal $I = I(F^s)$ in $D(R)$, such that $K[x_1, \dots, x_n, 1/F]^*F^s$ is isomorphic to $D(R)/I$ as a $D(R)$ -module. We often write just D for $D(R)$ and $D[s]$ for $D(R)[s]$. One is interested in the following data:

- $\text{Ann } F^s = I = I(F^s)$ in $D(R)[s]$, denoted by LD in the output
- global Bernstein polynomial in $K[s]$, denoted by bs ,
- its minimal integer root s_0 , the list of all roots of bs , which are known to be rational, with their multiplicities, which is denoted by BS
- $\text{Ann } F^{s_0} = I(F^{s_0})$ in $D(R)$, denoted by LD_0 in the output (LD_0 is a holonomic ideal in $D(R)$)
- $\text{Ann}^{(1)} F^s$ in $D(R)[s]$, denoted by LD_1 (logarithmic derivations)
- an operator in $D(R)[s]$, denoted by PS , such that the functional equality $PS^*F^{(s+1)} = bs^*F^s$ holds in $K[x_1, \dots, x_n, 1/F]^*F^s$.

References:

We provide the following implementations of algorithms:

- (OT) the classical $\text{Ann } F^s$ algorithm from Oaku and Takayama (Journal of Pure and Applied Math., 1999),
- (LOT) Levandovskyy's modification of the Oaku-Takayama algorithm (ISSAC 2007)
- (BM) the $\text{Ann } F^s$ algorithm by Briancon and Maisonobe (Remarques sur l'ideal de Bernstein associe a des polynomes, preprint, 2002)
- (LM08) V. Levandovskyy and J. Martin-Morales, ISSAC 2008
- (C) Countinho, A Primer of Algebraic D-Modules,
- (SST) Saito, Sturmfels, Takayama 'Groebner Deformations of Hypergeometric Differential Equations', Springer, 2000

Guide:

- $\text{Ann } F^s = I(F^s) = LD$ in $D(R)[s]$ can be computed by `Sannfs` [`BM`, `OT`, `LOT`]
- $\text{Ann}^{(1)} F^s$ in $D(R)[s]$ can be computed by `Sannfslog`
- global Bernstein polynomial bs in $K[s]$ can be computed by `bernsteinBM`
- $\text{Ann } F^{s_0} = I(F^{s_0}) = LD_0$ in $D(R)$ can be computed by `annfs0`, `annfs`, `annfsBM`, `annfsOT`, `annfsLOT`, `annfs2`, `annfsRB` etc.
- all the relevant data to F^s (LD , LD_0 , bs , PS) are computed by `operatorBM`
- operator PS can be computed via `operatorModulo` or `operatorBM`
- annihilator of $F^{\{s_1\}}$ for a number s_1 is computed with `annfspecial`
- annihilator of $F_1^{s_1} * \dots * F_p^{s_p}$ is computed with `annfsBMI`
- computing the multiplicity of a rational number r in the Bernstein poly of a given ideal goes with `checkRoot`
- check, whether a given univariate polynomial divides the Bernstein poly goes with `checkFactor`

Procedures: See also: Section 7.5.2 [`bfun_lib`], page 370; Section 7.5.5 [`dmodapp_lib`], page 414; Section 7.5.7 [`dmodvar_lib`], page 446; Section D.6.13 [`gmssing_lib`], page 862.

7.5.4.1 annfs

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `annfs(f [,S,eng]);` f a poly, S a string, eng an optional int

Return: ring

Purpose: compute the D-module structure of `basing[1/f]*f^s` with the algorithm given in S and with the Groebner basis engine given in "eng"

Note: activate the output ring with the `setring` command.

String S; S can be one of the following:

'bm' (default) - for the algorithm of Briancon and Maisonobe,

'ot' - for the algorithm of Oaku and Takayama,

'lot' - for the Levandovskyy's modification of the algorithm of OT.

If `eng <> 0`, `std` is used for Groebner basis computations,

otherwise and by default `slimgb` is used.

In the output ring:

- the ideal LD (which is a Groebner basis) is the needed D-module structure,

- the list BS contains roots and multiplicities of a BS-polynomial of f.

Display: If `printlevel=1`, progress debug messages will be printed,
if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y,z), Dp;
poly F = z*x^2+y^3;
def A = annfs(F); // here, the default BM algorithm will be used
setring A; // the Weyl algebra in (x,y,z,Dx,Dy,Dz)
LD; //the annihilator of F^{-1} over A
↳ LD[1]=y*Dy+3*z*Dz+3
↳ LD[2]=x*Dx-2*z*Dz
↳ LD[3]=x^2*Dy-3*y^2*Dz
↳ LD[4]=3*y^2*Dx-2*x*z*Dy
↳ LD[5]=y^3*Dz+x^2*z*Dz+x^2
↳ LD[6]=2*x*z*Dy^2+9*y*z*Dx*Dz+3*y*Dx
↳ LD[7]=9*y*z*Dx^2*Dz+4*z^2*Dy^2*Dz+3*y*Dx^2+2*z*Dy^2
↳ LD[8]=4*z^2*Dy^3*Dz-27*z^2*Dx^2*Dz^2+2*z*Dy^3-54*z*Dx^2*Dz-6*Dx^2
BS; // roots with multiplicities of BS polynomial
↳ [1]:
↳   _[1]=-1
↳   _[2]=-4/3
↳   _[3]=-5/3
↳   _[4]=-5/6
↳   _[5]=-7/6
↳ [2]:
↳   1,1,1,1,1
```

7.5.4.2 annfspecial

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `annfspecial(I,F,mir,n);` I an ideal, F a poly, int mir, number n

Return: ideal

- Purpose:** compute the annihilator ideal of F^n in the Weyl Algebra for the given rational number n
- Assume:** The basering is $D[s]$ and contains 's' explicitly as a variable, the ideal I is the Ann F^s in $D[s]$ (obtained with e.g. SannfsBM), the integer 'mir' is the minimal integer root of the BS polynomial of F , and the number n is rational.
- Note:** We compute the real annihilator for any rational value of n (both generic and exceptional). The implementation goes along the lines of the Algorithm 5.3.15 from Saito-Sturmfels-Takayama.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y), dp;
poly F = x3-y2;
def B = annfs(F); setring B;
minIntRoot(BS[1],0);
↳ -1
// So, the minimal integer root is -1
setring r;
def A = SannfsBM(F);
setring A;
poly F = x3-y2;
annfspecial(LD,F,-1,3/4); // generic root
↳ _[1]=4*x*Dx+6*y*Dy-9
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=18*x*y*Dy^2-8*y*Dx^2-33*x*Dy
↳ _[4]=54*y^2*Dy^3+16*y*Dx^3+66*x*Dx*Dy-9*y^2*Dy+66*Dy
annfspecial(LD,F,-1,-2); // integer but still generic root
↳ _[1]=2*x*Dx+3*y*Dy+12
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=9*x*y*Dy^2-4*y*Dx^2+33*x*Dy
↳ _[4]=27*y^2*Dy^3+8*y*Dx^3-66*x*Dx*Dy+144*y*Dy^2-66*Dy
annfspecial(LD,F,-1,1); // exceptional root
↳ _[1]=Dx*Dy
↳ _[2]=2*x*Dx+3*y*Dy-6
↳ _[3]=Dy^3
↳ _[4]=y*Dy^2-Dy
↳ _[5]=3*x*Dy^2+Dx^2
↳ _[6]=3*x^2*Dy+2*y*Dx
↳ _[7]=Dx^3+3*Dy^2
↳ _[8]=y*Dx^2+3*x*Dy
```

7.5.4.3 Sannfs

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

- Usage:** `Sannfs(f [,S,eng]);` f a poly, S a string, `eng` an optional int
- Return:** ring
- Purpose:** compute the D-module structure of basering[f^s] with the algorithm given in S and with the Groebner basis engine given in `eng`

Note: activate the output ring with the `setring` command.
 The value of a string `S` can be
 'bm' (default) - for the algorithm of Briancon and Maisonobe,
 'lot' - for the Levandovskyy's modification of the algorithm of OT,
 'ot' - for the algorithm of Oaku and Takayama.
 If `eng <> 0`, `std` is used for Groebner basis computations,
 otherwise, and by default `slimgb` is used.
 In the output ring:
 - the ideal `LD` is the needed D-module structure.

Display: If `printlevel=1`, progress debug messages will be printed,
 if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y,z), Dp;
poly F = x^3+y^3+z^3;
printlevel = 0;
def A = Sannfs(F); // here, the default BM algorithm will be used
setring A;
LD;
↳ LD[1]=z^2*Dy-y^2*Dz
↳ LD[2]=x*Dx+y*Dy+z*Dz-3*s
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
```

7.5.4.4 Sannfslog

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `Sannfslog(f [,eng]);` `f` a poly, `eng` an optional int

Return: ring

Purpose: compute the D-module structure of `basing[1/f]*f^s`

Note: activate the output ring with the `setring` command.
 In the output ring `D[s]`, the ideal `LD1` is generated by the elements
 in `Ann F^s` in `D[s]`, coming from logarithmic derivations.
 If `eng <> 0`, `std` is used for Groebner basis computations,
 otherwise, and by default `slimgb` is used.

Display: If `printlevel=1`, progress debug messages will be printed,
 if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y), Dp;
poly F = x4+y5+x*y4;
printlevel = 0;
def A = Sannfslog(F);
setring A;
LD1;
↳ LD1[1]=4*x^2*Dx+5*x*y*Dx+3*x*y*Dy+4*y^2*Dy-16*x*s-20*y*s
↳ LD1[2]=16*x*y^2*Dx+4*y^3*Dx+12*y^3*Dy-125*x*y*Dx-4*x^2*Dy+5*x*y*Dy-100*y^2*Dy-64*y^2*s+500*y*s
```

7.5.4.5 bernsteinBM

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

- Usage:** `bernsteinBM(f [,eng]);` f a poly, `eng` an optional int
- Return:** list (of roots of the Bernstein polynomial b and their multiplicities)
- Purpose:** compute the global Bernstein-Sato polynomial for a hypersurface, defined by f , according to the algorithm by Briancon and Maisonobe
- Note:** If `eng <> 0`, `std` is used for Groebner basis computations, otherwise, and by default `slimgb` is used.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y,z,w), Dp;
poly F = x^3+y^3+z^2*w;
printlevel = 0;
bernsteinBM(F);
⇒ [1]:
⇒   _[1]==-1
⇒   _[2]==-2
⇒   _[3]==-3/2
⇒   _[4]==-5/3
⇒   _[5]==-7/3
⇒   _[6]==-7/6
⇒   _[7]==-11/6
⇒ [2]:
⇒   1,1,1,1,1,1,1
```

7.5.4.6 bernsteinLift

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

- Usage:** `bernsteinLift(I, F [,eng]);` I an ideal, F a poly, `eng` an optional int
- Return:** list
- Purpose:** compute the (multiple of) Bernstein-Sato polynomial with lift-like method, based on the output of `Sannfs`-like procedure
- Note:** the output list contains the roots with multiplicities of the candidate for being Bernstein-Sato polynomial of f .
If `eng <> 0`, `std` is used for Groebner basis computations, otherwise and by default `slimgb` is used.
If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y,z), Dp;
poly F = x^3+y^3+z^3;
printlevel = 0;
def A = Sannfs(F);  setring A;
```

```

LD;
↳ LD[1]=z^2*Dy-y^2*Dz
↳ LD[2]=x*Dx+y*Dy+z*Dz-3*s
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
poly F = imap(r,F);
list L = bernsteinLift(LD,F); L;
↳ [1]:
↳   _[1]=-2
↳   _[2]=-4/3
↳   _[3]=-5/3
↳   _[4]=-1
↳ [2]:
↳   1,1,1,2
poly bs = fl2poly(L,"s"); bs; // the candidate for Bernstein-Sato polynomial
↳ s^5+7*s^4+173/9*s^3+233/9*s^2+154/9*s+40/9

```

7.5.4.7 operatorBM

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `operatorBM(f [,eng]);` f a poly, eng an optional int

Return: ring

Purpose: compute the B-operator and other relevant data for $\text{Ann } F^s$, using e.g. algorithm by Briancon and Maisonobe for $\text{Ann } F^s$ and BS.

Note: activate the output ring with the `setring` command. In the output ring $D[s]$

- the polynomial F is the same as the input,
- the ideal LD is the annihilator of f^s in $Dn[s]$,
- the ideal $LD0$ is the needed D -mod structure, where $LD0 = LD|_{s=0}$,
- the polynomial bs is the global Bernstein polynomial of f in the variable s ,
- the list BS contains all the roots with multiplicities of the global Bernstein polynomial of f ,
- the polynomial PS is an operator in $Dn[s]$ such that $PS*f^{(s+1)} = bs*f^s$.

If $eng < 0$, `std` is used for Groebner basis computations, otherwise and by default `slimgb` is used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel` ≥ 2 , all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
ring r = 0,(x,y,z),Dp;
poly F = x^3+y^3+z^3;
printlevel = 0;
def A = operatorBM(F);
setring A;
F; // the original polynomial itself
↳ x^3+y^3+z^3
LD; // generic annihilator
↳ LD[1]=x*Dx+y*Dy+z*Dz-3*s
↳ LD[2]=z^2*Dy-y^2*Dz
↳ LD[3]=z^2*Dx-x^2*Dz

```

```

↳ LD[4]=y^2*Dx-x^2*Dy
↳ LD[5]=x^3*Dz+y^3*Dz+z^3*Dz-3*z^2*s
↳ LD[6]=x^3*Dy+y^3*Dy+y^2*z*Dz-3*y^2*s
LD0; // annihilator
↳ LD0[1]=x*Dx+y*Dy+z*Dz+6
↳ LD0[2]=z^2*Dy-y^2*Dz
↳ LD0[3]=z^2*Dx-x^2*Dz
↳ LD0[4]=y^2*Dx-x^2*Dy
↳ LD0[5]=x^3*Dz+y^3*Dz+z^3*Dz+6*z^2
↳ LD0[6]=x^3*Dy+y^3*Dy+y^2*z*Dz+6*y^2
bs; // normalized Bernstein poly
↳ s^5+7*s^4+173/9*s^3+233/9*s^2+154/9*s+40/9
BS; // roots and multiplicities of the Bernstein poly
↳ [1]:
↳   _[1]=-2
↳   _[2]=-4/3
↳   _[3]=-5/3
↳   _[4]=-1
↳ [2]:
↳   1,1,1,2
PS; // the operator, s.t. PS*F^{s+1} = bs*F^s mod LD
↳ 2/81*y*z*Dx^3*Dy*Dz-2/81*y*z*Dy^4*Dz-4/81*y^2*Dy^2*Dz^3-2/81*y*z*Dy*Dz^4+\
  2/81*y*Dx^3*Dy*s-2/81*y*Dy^4*s+2/81*z*Dx^3*Dz*s+2/27*z*Dy^3*Dz*s+2/27*y*D\
  y*Dz^3*s-2/81*z*Dz^4*s+2/27*y*Dx^3*Dy-2/27*y*Dy^4+2/27*z*Dx^3*Dz+2/27*z*D\
  y^3*Dz-10/81*y*Dy*Dz^3-2/27*z*Dz^4+1/27*Dx^3*s^2+1/9*Dy^3*s^2+1/9*Dz^3*s^2+\
  2+5/27*Dx^3*s+11/27*Dy^3*s+11/27*Dz^3*s+20/81*Dx^3+8/27*Dy^3+16/81*Dz^3
reduce(PS*F-bs,LD); // check the property of PS
↳ 0

```

7.5.4.8 operatorModulo

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `operatorModulo(f,I,b)`; f a poly, I an ideal, b a poly

Return: poly

Purpose: compute the B-operator from the polynomial f , ideal $I = \text{Ann } f^s$ and Bernstein-Sato polynomial b using modulo i.e. kernel of module homomorphism

Note: The computations take place in the ring, similar to the one returned by `Sannfs` procedure.

Note, that operator is not completely reduced wrt $\text{Ann } f^{s+1}$.

If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
// LIB "dmod.lib"; option(prot); option(mem);
ring r = 0, (x,y), Dp;
poly F = x^3+y^3+x*y^3;
def A = Sannfs(F); // here we get LD = ann f^s
setring A;
poly F = imap(r,F);

```

```

def B = annfs0(LD,F); // to obtain BS polynomial
list BS = imap(B,BS); poly bs = fl2poly(BS,"s");
poly PS = operatorModulo(F,LD,bs);
LD = groebner(LD);
PS = NF(PS,subst(LD,s,s+1)); // reduction modulo Ann s^{s+1}
↳ // ** _ is no standard basis
size(PS);
↳ 56
lead(PS);
↳ -2/243*y^3*Dx*Dy^3
reduce(PS*F-bs,LD); // check the defining property of PS
↳ 0

```

7.5.4.9 annfsParamBM

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `annfsParamBM(f [,eng]);` `f` a poly, `eng` an optional int

Return: ring

Purpose: compute the generic $\text{Ann } F^{\wedge}$ and exceptional parametric constellations of a polynomial with parametric coefficients with the BM algorithm

Note: activate the output ring with the `setring` command. In this ring,
 - the ideal `LD` is the D-module structure of $\text{Ann } F^{\wedge}$
 - the ideal `Param` contains special parameters as entries
 If `eng <> 0`, `std` is used for Groebner basis computations, otherwise, and by default `slingb` is used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
ring r = (0,a,b),(x,y),Dp;
poly F = x^2 - (y-a)*(y-b);
printlevel = 0;
def A = annfsParamBM(F); setring A;
LD;
↳ LD[1]=2*y*Dx+2*x*Dy+(-a-b)*Dx
↳ LD[2]=x^2*Dy-y^2*Dy+(a+b)*y*Dy+2*y*s+(-a*b)*Dy+(-a-b)*s
↳ LD[3]=4*x^2*Dx+4*x*y*Dy+(-2*a-2*b)*x*Dy-8*x*s+(a^2-2*a*b+b^2)*Dx
Param;
↳ Param[1]=(a-b)
setring r;
poly G = x^2-(y-a)^2; // try the exceptional value b=a of parameters
def B = annfsParamBM(G); setring B;
LD;
↳ LD[1]=y*Dx+x*Dy+(-a)*Dx
↳ LD[2]=x*Dx+y*Dy+(-a)*Dy-2*s
↳ LD[3]=x^2*Dy-y^2*Dy+(2*a)*y*Dy+2*y*s+(-a^2)*Dy+(-2*a)*s
Param;
↳ Param[1]=0

```

7.5.4.10 annfsBMI

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `annfsBMI(F [,eng,met,us]);` F an ideal, `eng`, `met`, `us` optional ints

Return: ring

Purpose: compute two kinds of Bernstein-Sato ideals, associated to $f = F[1]^* \dots F[P]$, with the multivariate algorithm by Briancon and Maisonobe.

Note: activate the output ring with the `setring` command. In this ring,
 - the ideal `LD` is the annihilator of $F[1]^{s_1} \dots F[P]^{s_p}$,
 - the list or ideal `BS` is a Bernstein-Sato ideal of a polynomial $f = F[1]^* \dots F[P]$.
 If `eng <> 0`, `std` is used for Groebner basis computations, otherwise, and by default `slimgb` is used.
 If `met < 0`, the B-Sigma ideal (cf. Castro and Ucha, 'On the computation of Bernstein-Sato ideals', 2005) is computed.
 If $0 < \text{met} < P$, then the ideal `B_P` (cf. the paper) is computed.
 Otherwise, and by default, the ideal `B` (cf. the paper) is computed.
 If `us <> 0`, then syzygies-driven method is used.
 If the output ideal happens to be principal, the list of factors with their multiplicities is returned instead of the ideal.
 If `printlevel=1`, progress debug messages will be printed, if `printlevel >= 2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y), Dp;
ideal F = x,y,x+y;
printlevel = 0;
// *1* let us compute the B ideal
def A = annfsBMI(F); setring A;
LD; // annihilator
↳ LD[1]=x*Dx+y*Dy-s(1)-s(2)-s(3)
↳ LD[2]=x*y*Dy+y^2*Dy-x*s(2)-y*s(2)-y*s(3)
↳ LD[3]=y^2*Dx*Dy-y^2*Dy^2+y*Dy*s(1)-y*Dx*s(2)+2*y*Dy*s(2)-y*Dx*s(3)+y*Dy*s(3)-s(1)*s(2)-s(2)^2-s(2)*s(3)-s(2)
BS; // Bernstein-Sato ideal
↳ [1]:
↳ _[1]=s(1)+1
↳ _[2]=s(2)+1
↳ _[3]=s(3)+1
↳ _[4]=s(1)+s(2)+s(3)+2
↳ _[5]=s(1)+s(2)+s(3)+3
↳ _[6]=s(1)+s(2)+s(3)+4
↳ [2]:
↳ 1,1,1,1,1,1
// *2* now, let us compute B-Sigma ideal
setring r;
def Sigma = annfsBMI(F,0,-1); setring Sigma;
print(matrix(lead(LD))); // compact form of leading
↳ x*Dx,x*y*Dy,y^2*Dx*Dy
// monomials from the annihilator
BS; // Bernstein-Sato B-Sigma ideal: it is principal,
```

```

↳ [1]:
↳   _[1]=s(1)+s(2)+s(3)+2
↳ [2]:
↳   1
// so factors and multiplicities are returned
// *3* and now, let us compute B-i ideal
setring r;
def Bi = annfsBMI(F,0,3); // that is F[3]=x+y is taken
setring Bi;
print(matrix(lead(LD))); // compact form of leading
↳ x*Dx,x*y*Dy,y^2*Dx*Dy
// monomials from the annihilator
BS; // the B_3 ideal: it is principal, so factors
↳ [1]:
↳   _[1]=s(3)+1
↳   _[2]=s(1)+s(2)+s(3)+2
↳ [2]:
↳   1,1
// and multiplicities are returned

```

7.5.4.11 checkRoot

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `checkRoot(f,alpha [,S,eng]);` poly `f`, number `alpha`, string `S`, int `eng`

Return: int

Assume: Basing is a commutative ring, `alpha` is a positive rational number.

Purpose: check whether a negative rational number `-alpha` is a root of the global Bernstein-Sato polynomial of `f` and compute its multiplicity, with the algorithm given by `S` and with the Groebner basis engine given by `eng`.

Note: The annihilator of f^s in $D[s]$ is needed and hence it is computed with the algorithm by Briancon and Maisonobe. The value of a string `S` can be `'alg1'` (default) - for the algorithm 1 of [LM08] `'alg2'` - for the algorithm 2 of [LM08]

Depending on the value of `S`, the output of type int is:

`'alg1'`: 1 only if `-alpha` is a root of the global Bernstein-Sato polynomial

`'alg2'`: the multiplicity of `-alpha` as a root of the global Bernstein-Sato polynomial of `f`. If `-alpha` is not a root, the output is 0.

If `eng <> 0`, `std` is used for Groebner basis computations, otherwise (and by default) `slimgb` is used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
printlevel=0;
ring r = 0, (x,y), Dp;
poly F = x^4+y^5+x*y^4;
checkRoot(F,11/20); // -11/20 is a root of bf
↳ 1
poly G = x*y;

```

```
checkRoot(G,1,"alg2"); // -1 is a root of bg with multiplicity 2
↳ 2
```

7.5.4.12 SannfsBFCT

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `SannfsBFCT(f [,a,b,c]);` f a poly, a,b,c optional ints

Return: ring

Purpose: compute a Groebner basis either of $\text{Ann}(f^s) + \langle f \rangle$ or of $\text{Ann}(f^s) + \langle f, f_1, \dots, f_n \rangle$ in $D[s]$

Note: Activate the output ring with the `setring` command.
This procedure, unlike `SannfsBM`, returns the ring $D[s]$ with an anti-elimination ordering for s .

The output ring contains an ideal LD , being a Groebner basis either of $\text{Ann}(f^s) + \langle f \rangle$, if $a=0$ (and by default), or of $\text{Ann}(f^s) + \langle f, f_1, \dots, f_n \rangle$, otherwise.

Here, f_i stands for the i -th partial derivative of f .

If $b < 0$, `std` is used for Groebner basis computations, otherwise, and by default `slimgb` is used.

If $c > 0$, `std` is used for Groebner basis computations of ideals $\langle I+J \rangle$ when I is already a Groebner basis of $\langle I \rangle$.

Otherwise, and by default the engine determined by the switch b is used. Note that in the case $c > 0$, the choice for b will be overwritten only for the types of ideals mentioned above.

This means that if $b < 0$, specifying c has no effect.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y,z,w), Dp;
poly F = x^3+y^3+z^3*w;
// compute Ann(F^s)+<F> using slimgb only
def A = SannfsBFCT(F);
setring A; A;
↳ // coefficients: QQ
↳ // number of vars : 9
↳ //      block  1 : ordering dp
↳ //                : names    s
↳ //      block  2 : ordering dp
↳ //                : names    x y z w Dx Dy Dz Dw
↳ //      block  3 : ordering C
↳ // noncommutative relations:
↳ //      Dxx=x*Dx+1
↳ //      Dyy=y*Dy+1
↳ //      Dzz=z*Dz+1
↳ //      Dww=w*Dw+1
LD;
↳ LD[1]=z*Dz-3*w*Dw
↳ LD[2]=3*s-x*Dx-y*Dy-3*w*Dw
```



```

↳ LD[3]=y^2*Dx-x^2*Dy
↳ LD[4]=z^2*w*Dy-y^2*Dz
↳ LD[5]=z^3*Dy-3*y^2*Dw
↳ LD[6]=z^2*w*Dx-x^2*Dz
↳ LD[7]=z^3*Dx-3*x^2*Dw
↳ LD[8]=z^3*w+x^3+y^3
↳ LD[9]=x^3*Dy+y^3*Dy+3*y^2*w*Dw+3*y^2
↳ LD[10]=x^3*Dx+x^2*y*Dy+3*x^2*w*Dw+3*x^2
↳ LD[11]=3*z*w^2*Dy*Dw-y^2*Dz^2+2*z*w*Dy
↳ LD[12]=3*z*w^2*Dx*Dw-x^2*Dz^2+2*z*w*Dx
↳ LD[13]=3*z^2*w^2*Dw+x^3*Dz+y^3*Dz+3*z^2*w
↳ LD[14]=9*w^3*Dy*Dw^2-y^2*Dz^3+18*w^2*Dy*Dw+2*w*Dy
↳ LD[15]=9*w^3*Dx*Dw^2-x^2*Dz^3+18*w^2*Dx*Dw+2*w*Dx
↳ LD[16]=9*z*w^3*Dw^2+x^3*Dz^2+y^3*Dz^2+24*z*w^2*Dw+6*z*w
↳ LD[17]=27*w^4*Dw^3+x^3*Dz^3+y^3*Dz^3+135*w^3*Dw^2+114*w^2*Dw+6*w
// the Bernstein-Sato poly of F:
vec2poly(pIntersect(s,LD));
↳ s^6+28/3*s^5+320/9*s^4+1910/27*s^3+2093/27*s^2+1198/27*s+280/27
// a fancier example:
def R = reiffen(4,5); setring R;
RC; // the Reiffen curve in 4,5
↳ xy4+y5+x4
// compute Ann(RC^s)+<RC,diff(RC,x),diff(RC,y)>
// using std for GB computations of ideals <I+J>
// where I is already a GB of <I>
// and slimgb for other ideals
def B = SannfsBFCT(RC,1,0,1);
setring B;
// the Bernstein-Sato poly of RC:
(s-1)*vec2poly(pIntersect(s,LD));
↳ s^13+10*s^12+44*s^11+44099/400*s^10+13355001/80000*s^9+22138611/160000*s^8+
1747493/160000*s^7-7874303503/64000000*s^6-4244944536107/25600000000*s^5-
3066298289417/25600000000*s^4-2787777479229/51200000000*s^3-19980507461\
787/128000000000*s^2-663659243177931/25600000000000*s-48839201079669/25\
600000000000

```

7.5.4.13 annfs0

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `annfs0(I, F [,eng]);` I an ideal, F a poly, eng an optional int

Return: ring

Purpose: compute the annihilator ideal of f^s in the Weyl Algebra, based on the output of Sannfs-like procedure

Note: activate the output ring with the `setring` command. In this ring,
- the ideal LD (which is a Groebner basis) is the annihilator of f^s ,
- the list BS contains the roots with multiplicities of BS polynomial of f.
If `eng <> 0`, `std` is used for Groebner basis computations, otherwise and by default `slimgb` is used.
If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
ring r = 0, (x,y,z), Dp;
poly F = x^3+y^3+z^3;
printlevel = 0;
def A = SannfsBM(F);  setring A;
// alternatively, one can use SannfsOT or SannfsLOT
LD;
↳ LD[1]=z^2*Dy-y^2*Dz
↳ LD[2]=x*Dx+y*Dy+z*Dz-3*s
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
poly F = imap(r,F);
def B = annfs0(LD,F);  setring B;
LD;
↳ LD[1]=x*Dx+y*Dy+z*Dz+6
↳ LD[2]=z^2*Dy-y^2*Dz
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
↳ LD[5]=x^3*Dz+y^3*Dz+z^3*Dz+6*z^2
↳ LD[6]=x^3*Dy+y^3*Dy+y^2*z*Dz+6*y^2
BS;
↳ [1]:
↳   _[1]=-2
↳   _[2]=-4/3
↳   _[3]=-5/3
↳   _[4]=-1
↳ [2]:
↳   1,1,1,2

```

7.5.4.14 annfs2

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `annfs2(I, F [,eng]);` I an ideal, F a poly, eng an optional int

Return: ring

Purpose: compute the annihilator ideal of f^s in the Weyl Algebra, based on the output of Sannfs-like procedure
annfs2 uses shorter expressions in the variable s (the idea of Noro).

Note: activate the output ring with the `setring` command. In this ring,
- the ideal LD (which is a Groebner basis) is the annihilator of f^s ,
- the list BS contains the roots with multiplicities of the BS polynomial.
If `eng <> 0`, `std` is used for Groebner basis computations, otherwise and by default `slimgb` is used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
ring r = 0, (x,y,z), Dp;
poly F = x^3+y^3+z^3;
printlevel = 0;

```

```

def A = SannfsBM(F);
setring A;
LD;
↳ LD[1]=z^2*Dy-y^2*Dz
↳ LD[2]=x*Dx+y*Dy+z*Dz-3*s
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
poly F = imap(r,F);
def B = annfs2(LD,F);
setring B;
LD;
↳ LD[1]=x*Dx+y*Dy+z*Dz+6
↳ LD[2]=z^2*Dy-y^2*Dz
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
↳ LD[5]=x^3*Dz+y^3*Dz+z^3*Dz+6*z^2
↳ LD[6]=x^3*Dy+y^3*Dy+y^2*z*Dz+6*y^2
BS;
↳ [1]:
↳   _[1]=-2
↳   _[2]=-5/3
↳   _[3]=-4/3
↳   _[4]=-1
↳ [2]:
↳   1,1,1,2

```

7.5.4.15 annfsRB

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `annfsRB(I, F [,eng]);` I an ideal, F a poly, eng an optional int

Return: ring

Purpose: compute the annihilator ideal of f^s in the Weyl Algebra, based on the output of Sannfs like procedure

Note: activate the output ring with the `setring` command. In this ring,
 - the ideal LD (which is a Groebner basis) is the annihilator of f^s ,
 - the list BS contains the roots with multiplicities of a Bernstein polynomial of f.
 If `eng <> 0`, `std` is used for Groebner basis computations, otherwise and by default `slimgb` is used.
 This procedure uses in addition to F its Jacobian ideal.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
ring r = 0, (x,y,z), Dp;
poly F = x^3+y^3+z^3;
printlevel = 0;
def A = SannfsBM(F); setring A;
LD; // s-parametric ahhinilator
↳ LD[1]=z^2*Dy-y^2*Dz

```

```

↳ LD[2]=x*Dx+y*Dy+z*Dz-3*s
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
poly F = imap(r,F);
def B = annfsRB(LD,F); setring B;
LD;
↳ LD[1]=x*Dx+y*Dy+z*Dz+6
↳ LD[2]=z^2*Dy-y^2*Dz
↳ LD[3]=z^2*Dx-x^2*Dz
↳ LD[4]=y^2*Dx-x^2*Dy
↳ LD[5]=x^3*Dz+y^3*Dz+z^3*Dz+6*z^2
↳ LD[6]=x^3*Dy+y^3*Dy+y^2*z*Dz+6*y^2
BS;
↳ [1]:
↳   _[1]==-2
↳   _[2]==-5/3
↳   _[3]==-4/3
↳   _[4]==-1
↳ [2]:
↳   1,1,1,2

```

7.5.4.16 checkFactor

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `checkFactor(I,f,qs [,eng]);` I an ideal, f a poly, qs a poly, eng an optional int

Assume: `checkFactor` is called from the basering, created by Sannfs-like proc, that is, from the Weyl algebra in $x_1, \dots, x_N, d_1, \dots, d_N$ tensored with $K[s]$. The ideal I is the annihilator of f^{\sim} in $D[s]$, that is the ideal, computed by Sannfs-like procedure (usually called LD there). Moreover, f is a polynomial in $K[x_1, \dots, x_N]$ and qs is a polynomial in $K[s]$.

Return: int, 1 if qs is a factor of the global Bernstein polynomial of f and 0 otherwise

Purpose: check whether a univariate polynomial qs is a factor of the Bernstein-Sato polynomial of f without explicit knowledge of the latter.

Note: If `eng <> 0`, `std` is used for Groebner basis computations, otherwise (and by default) `slingb` is used.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmod.lib";
ring r = 0, (x,y), Dp;
poly F = x^4+y^5+x*y^4;
printlevel = 0;
def A = Sannfs(F);
setring A;
poly F = imap(r,F);
checkFactor(LD,F,20*s+31); // -31/20 is not a root of bs
↳ 0
checkFactor(LD,F,20*s+11); // -11/20 is a root of bs
↳ 1

```

```
checkFactor(LD,F,(20*s+11)^2); // the multiplicity of -11/20 is 1
↳ 0
```

7.5.4.17 arrange

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `arrange(p); int p`

Return: ring

Purpose: set up the polynomial, describing a hyperplane arrangement

Note: must be executed in a commutative ring

Assume: basering is present and it is commutative

Example:

```
LIB "dmod.lib";
ring X = 0, (x,y,z,t), dp;
poly q = arrange(3);
factorize(q,1);
↳ _[1]=x
↳ _[2]=y
↳ _[3]=x+y
↳ _[4]=z
↳ _[5]=x+z
↳ _[6]=y+z
↳ _[7]=x+y+z
```

7.5.4.18 reiffen

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `reiffen(p, q); int p, int q`

Return: ring

Purpose: set up the polynomial, describing a Reiffen curve

Note: activate the output ring with the `setring` command and find the curve as a polynomial RC.

A Reiffen curve is defined as $RC = x^p + y^q + xy^{\{q-1\}}$, $q \geq p+1 \geq 5$

Example:

```
LIB "dmod.lib";
def r = reiffen(4,5);
setring r;
RC;
↳ xy4+y5+x4
```

7.5.4.19 isHolonomic

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `isHolonomic(M); M an ideal/module/matrix`

Return: int, 1 if M is holonomic over the base ring, and 0 otherwise

Assume: basering is a Weyl algebra in characteristic 0

Purpose: check whether M is holonomic over the base ring

Note: M is holonomic if $2 \cdot \dim(M) = \dim(R)$, where R is the base ring; \dim stands for Gelfand-Kirillov dimension

Example:

```
LIB "dmod.lib";
ring R = 0, (x,y), dp;
poly F = x*y*(x+y);
def A = annfsBM(F,0);
setring A;
LD;
↳ LD[1]=x*Dx+y*Dy+3
↳ LD[2]=x*y*Dy+y^2*Dy+x+2*y
↳ LD[3]=y^2*Dx*Dy-y^2*Dy^2+2*y*Dx-4*y*Dy-2
isHolonomic(LD);
↳ 1
ideal I = std(LD[1]);
I;
↳ I[1]=x*Dx+y*Dy+3
isHolonomic(I);
↳ 0
```

7.5.4.20 convloc

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `convloc(L)`; L a list

Return: list

Purpose: convert a ringlist L into another ringlist, where all the 'p' orderings are replaced with the 's' orderings, e.g. `dp` by `ds`.

Assume: L is a result of a ringlist command

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y,z), (Dp(2), dp(1));
list L = ringlist(r);
list N = convloc(L);
def rs = ring(N);
setring rs;
rs;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ //      block  1 : ordering Ds
↳ //                : names  x y
↳ //      block  2 : ordering ds
↳ //                : names  z
↳ //      block  3 : ordering C
```

7.5.4.21 minIntRoot

Procedure from library `dmod.lib` (see Section 7.5.4 [`dmod.lib`], page 395).

Usage: `minIntRoot(P, fact)`; P an ideal, fact an int

Return: int

Purpose: minimal integer root of a maximal ideal P

Note: if fact==1, P is the result of some 'factorize' call,
else P is treated as the result of bernstein::gmssing.lib
in both cases without constants and multiplicities

Example:

```
LIB "dmod.lib";
ring r = 0, (x,y), ds;
poly f1 = x*y*(x+y);
ideal I1 = bernstein(f1)[1]; // a local Bernstein poly
I1;
⇒ I1[1]==-4/3
⇒ I1[2]==-1
⇒ I1[3]==-2/3
minIntRoot(I1,0);
⇒ -1
poly f2 = x2-y3;
ideal I2 = bernstein(f2)[1];
I2;
⇒ I2[1]==-7/6
⇒ I2[2]==-1
⇒ I2[3]==-5/6
minIntRoot(I2,0);
⇒ -1
// now we illustrate the behaviour of factorize
// together with a global ordering
ring r2 = 0,x,dp;
poly f3 = 9*(x+2/3)*(x+1)*(x+4/3); //global b-polynomial of f1=x*y*(x+y)
ideal I3 = factorize(f3,1);
I3;
⇒ I3[1]=x+1
⇒ I3[2]=3x+2
⇒ I3[3]=3x+4
minIntRoot(I3,1);
⇒ -1
// and a more interesting situation
ring s = 0, (x,y,z), ds;
poly f = x3 + y3 + z3;
ideal I = bernstein(f)[1];
I;
⇒ I[1]==-2
⇒ I[2]==-5/3
⇒ I[3]==-4/3
⇒ I[4]==-1
minIntRoot(I,0);
⇒ -2
```

7.5.4.22 isRational

Procedure from library dmod.lib (see Section 7.5.4 [dmod.lib], page 395).

Usage: isRational(n); n number

Return: int

Purpose: determine whether n is a rational number, that is it does not contain parameters.

Assume: ground field is of characteristic 0

Example:

```
LIB "dmod.lib";
ring r = (0,a),(x,y),dp;
number n1 = 11/73;
isRational(n1);
↳ 1
number n2 = (11*a+3)/72;
isRational(n2);
↳ 0
```

7.5.5 dmodapp_lib

Library: dmodapp.lib

Purpose: Applications of algebraic D-modules

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Support: DFG Graduiertenkolleg 1632 'Experimentelle und konstruktive Algebra'

Overview: Let K be a field of characteristic 0, $R = K[x_1, \dots, x_N]$ and D be the Weyl algebra in variables $x_1, \dots, x_N, d_1, \dots, d_N$. In this library there are the following procedures for algebraic D-modules:

- given a cyclic representation D/I of a holonomic module and a polynomial F in R , it is proved that the localization of D/I with respect to the mult. closed set of all powers of F is a holonomic D-module. Thus we aim to compute its cyclic representation D/L for an ideal L in D . The procedures for the localization are `DLoc`, `SDLoc` and `DLoc0`.
- annihilator in D of a given polynomial F from R as well as of a given rational function G/F from $\text{Quot}(R)$. These can be computed via procedures `annPoly` resp. `annRat`.
- Groebner bases with respect to weights (according to (SST)), given an arbitrary integer vector containing weights for variables, one computes the homogenization of a given ideal relative to this vector, then one computes a Groebner basis and returns the dehomogenization of the result), initial forms and initial ideals in Weyl algebras with respect to a given weight vector can be computed with `GBWeight`, `inForm`, `initialMalgrange` and `initialIdealW`.
- restriction and integration of a holonomic module D/I . Suppose I annihilates a function $F(x_1, \dots, x_n)$. Our aim is to compute an ideal J directly from I , which annihilates $- F(0, \dots, 0, x_k, \dots, x_n)$ in case of restriction or
- the integral of F with respect to x_1, \dots, x_m in case of integration. The corresponding procedures are `restrictionModule`, `restrictionIdeal`, `integralModule` and `integralIdeal`.
- characteristic varieties defined by ideals in Weyl algebras can be computed with `charVariety` and `charInfo`.
- `appelF1`, `appelF2` and `appelF4` return ideals in parametric Weyl algebras, which annihilate corresponding Appel hypergeometric functions.

References:

- (SST) Saito, Sturmfels, Takayama 'Groebner Deformations of Hypergeometric Differential Equations', Springer, 2000
 (OTW) Oaku, Takayama, Walther 'A Localization Algorithm for D-modules', Journal of Symbolic Computation, 2000
 (OT) Oaku, Takayama 'Algorithms for D-modules', Journal of Pure and Applied Algebra, 1998

Procedures: D-module See also: Section 7.5.2 [bfun_lib], page 370; Section 7.5.4 [dmod_lib], page 395; Section 7.5.7 [dmodvar_lib], page 446; Section D.6.13 [gmssing_lib], page 862.

7.5.5.1 annPoly

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp_lib], page 414).

Usage: `annPoly(f)`; f a poly

Return: ring (a Weyl algebra) containing an ideal 'LD'

Purpose: compute the complete annihilator ideal of f in the corresponding Weyl algebra

Assume: basering is commutative and over a field of characteristic 0

Note: Activate the output ring with the `setring` command.
 In the output ring, the ideal 'LD' (in Groebner basis) is the annihilator.

Display: If `printlevel=1`, progress debug messages will be printed,
 if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmodapp.lib";
ring r = 0, (x,y,z), dp;
poly f = x^2*z - y^3;
def A = annPoly(f);
setring A; // A is the 3rd Weyl algebra in 6 variables
LD; // the Groebner basis of annihilator
↳ LD[1]=Dz^2
↳ LD[2]=Dy*Dz
↳ LD[3]=Dx*Dy
↳ LD[4]=y*Dy+3*z*Dz-3
↳ LD[5]=x*Dx-2*z*Dz
↳ LD[6]=z*Dx*Dz-Dx
↳ LD[7]=Dy^3+3*Dx^2*Dz
↳ LD[8]=x*Dy^2+3*y*Dx*Dz
↳ LD[9]=x^2*Dy+3*y^2*Dz
↳ LD[10]=Dx^3
↳ LD[11]=3*y*Dx^2+z*Dy^2
↳ LD[12]=3*y^2*Dx+2*x*z*Dy
↳ LD[13]=y^3*Dz-x^2*z*Dz+x^2
gkdim(LD); // must be 3 = 6/2, since A/LD is holonomic module
↳ 3
NF(Dy^4, LD); // must be 0 since Dy^4 clearly annihilates f
↳ 0
poly f = imap(r,f);
```

```

NF(LD*f,std(ideal(Dx,Dy,Dz))); // must be zero if LD indeed annihilates f
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0
↳ _[4]=0
↳ _[5]=0
↳ _[6]=0
↳ _[7]=0
↳ _[8]=0
↳ _[9]=0
↳ _[10]=0
↳ _[11]=0
↳ _[12]=0
↳ _[13]=0

```

See also: Section 7.5.5.2 [annRat], page 416.

7.5.5.2 annRat

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp.lib], page 414).

Usage: `annRat(g,f)`; f, g polynomials

Return: ring (a Weyl algebra) containing an ideal 'LD'

Purpose: compute the annihilator of the rational function g/f in the corresponding Weyl algebra

Assume: basering is commutative and over a field of characteristic 0

Note: Activate the output ring with the `setring` command.
 In the output ring, the ideal 'LD' (in Groebner basis) is the annihilator of g/f .
 The algorithm uses the computation of $\text{Ann}(f^{-1})$ via D-modules, see (SST).

Display: If `printlevel=1`, progress debug messages will be printed,
 if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmodapp.lib";
ring r = 0,(x,y),dp;
poly g = 2*x*y; poly f = x^2 - y^3;
def B = annRat(g,f);
setring B;
LD;
↳ LD[1]=3*y^2*Dx^2*Dy+2*x*Dx*Dy^2+9*y*Dx^2+4*Dy^2
↳ LD[2]=3*y^3*Dx^2-10*x*y*Dx*Dy-8*y^2*Dy^2+10*x*Dx
↳ LD[3]=y^3*Dy^2-x^2*Dy^2-6*x*y*Dx+2*y^2*Dy+4*y
↳ LD[4]=3*x*Dx+2*y*Dy+1
↳ LD[5]=y^4*Dy-x^2*y*Dy+2*y^3+x^2
// Now, compare with the output of Macaulay2:
ideal tst = 3*x*Dx + 2*y*Dy + 1, y^3*Dy^2 - x^2*Dy^2 + 6*y^2*Dy + 6*y,
9*y^2*Dx^2*Dy-4*y*Dy^3+27*y*Dx^2+2*Dy^2, 9*y^3*Dx^2-4*y^2*Dy^2+10*y*Dy -10;
option(redSB); option(redTail);
LD = groebner(LD);
tst = groebner(tst);

```

```

print(matrix(NF(LD,tst))); print(matrix(NF(tst,LD)));
↳ 0,0,0,0,0
↳ 0,0,0,0,0
// So, these two answers are the same

```

See also: Section 7.5.5.1 [annPoly], page 415.

7.5.5.3 DLoc

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp.lib], page 414).

Usage: DLoc(I, f); I an ideal, f a poly

Return: list of ideal and list

Assume: the basering is a Weyl algebra

Purpose: compute the presentation of the localization of D/I w.r.t. f 's

Note: In the output list L,
- L[1] is an ideal (given as Groebner basis), the presentation of the localization,
- L[2] is a list containing roots with multiplicities of Bernstein polynomial of $(D/I)_f$.

Display: If `printlevel=1`, progress debug messages will be printed,
if `printlevel>=2`, all the debug messages will be printed.

Example:

```

LIB "dmodapp.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def R = Weyl(); setring R; // Weyl algebra in variables x,y,Dx,Dy
poly F = x2-y3;
ideal I = (y^3 - x^2)*Dx - 2*x, (y^3 - x^2)*Dy + 3*y^2; // I = Dx*F, Dy*F;
// I is not holonomic, since its dimension is not 4/2=2
gkdim(I);
↳ 3
list L = DLoc(I, x2-y3);
L[1]; // localized module (R/I)_f is isomorphic to R/LD0
↳ _[1]=3*x*Dx+2*y*Dy+12
↳ _[2]=3*y^2*Dx+2*x*Dy
↳ _[3]=y^3*Dy-x^2*Dy+6*y^2
L[2]; // description of b-function for localization
↳ [1]:
↳ _[1]=0
↳ _[2]=1/6
↳ _[3]=-1/6
↳ [2]:
↳ 1,1,1

```

7.5.5.4 SLoc

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp.lib], page 414).

Usage: SLoc(I, f); I an ideal, f a poly

Return: ring (basing extended by a new variable) containing an ideal 'LD'

Purpose: compute a generic presentation of the localization of D/I w.r.t. f^s

Assume: the basering D is a Weyl algebra over a field of characteristic 0

Note: Activate this ring with the `setring` command. In this ring, the ideal LD (given as Groebner basis) is the presentation of the localization.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmodapp.lib";
ring r = 0,(x,y,Dx,Dy),dp;
def R = Weyl(); // Weyl algebra on the variables x,y,Dx,Dy
setring R;
poly F = x2-y3;
ideal I = Dx*F, Dy*F;
// note, that I is not holonomic, since it's dimension is not 2
gkdim(I); // 3, while dim R = 4
 $\mapsto$  3
def W = SDLoc(I,F);
setring W; // = R[s], where s is a new variable
LD; // Groebner basis of s-parametric presentation
 $\mapsto$  LD[1]=3*x*Dx*s+2*y*Dy*s-6*s^2+6*s
 $\mapsto$  LD[2]=3*y^2*Dx*s+2*x*Dy*s
 $\mapsto$  LD[3]=y^3*Dy-x^2*Dy-3*y^2*s+3*y^2
 $\mapsto$  LD[4]=y^3*Dx-x^2*Dx+2*x*s-2*x
```

7.5.5.5 DLoc0

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

Usage: `DLoc0(I, f)`; I an ideal, f a poly

Return: ring (a Weyl algebra) containing an ideal 'LD0' and a list 'BS'

Purpose: compute the presentation of the localization of D/I w.r.t. f^s , where D is a Weyl Algebra, based on the output of procedure `SDLoc`

Assume: the basering is similar to the output ring of `SDLoc` procedure

Note: activate the output ring with the `setring` command. In this ring,
 - the ideal $LD0$ (given as Groebner basis) is the presentation of the localization,
 - the list BS contains roots and multiplicities of Bernstein polynomial of $(D/I).f$.

Display: If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

Example:

```
LIB "dmodapp.lib";
ring r = 0,(x,y,Dx,Dy),dp;
def R = Weyl(); setring R; // Weyl algebra in variables x,y,Dx,Dy
poly F = x2-y3;
ideal I = (y^3 - x^2)*Dx - 2*x, (y^3 - x^2)*Dy + 3*y^2; // I = Dx*F, Dy*F;
```

```

// moreover I is not holonomic, since its dimension is not 2 = 4/2
gkdim(I); // 3
↳ 3
def W = SLoc(I,F); setring W; // creates ideal LD in W = R[s]
def U = DLoc0(LD, x2-y3); setring U; // compute in R
LD0; // Groebner basis of the presentation of localization
↳ LD0[1]=3*x*Dx+2*y*Dy+12
↳ LD0[2]=3*y^2*Dx+2*x*Dy
↳ LD0[3]=y^3*Dy-x^2*Dy+6*y^2
BS; // description of b-function for localization
↳ [1]:
↳   _[1]=0
↳   _[2]=1/6
↳   _[3]=-1/6
↳ [2]:
↳   1,1,1

```

7.5.5.6 GBWeight

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

- Usage:** `GBWeight(I,u,v [s,t,w]);`
 I ideal, u,v intvecs, s,t optional ints, w an optional intvec
- Return:** ideal, Groebner basis of I w.r.t. the weights u and v
- Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all $1 \leq i \leq n$ the identity $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$ holds, i.e. the sequence of variables is given by $x(1), \dots, x(n), D(1), \dots, D(n)$, where $D(i)$ is the differential operator belonging to $x(i)$.
- Purpose:** computes a Groebner basis with respect to given weights
- Note:** The weights u and v are understood as weight vectors for $x(i)$ and $D(i)$, respectively. According to (SST), one computes the homogenization of a given ideal relative to (u,v), then one computes a Groebner basis and returns the dehomogenization of the result.
 If $s < 0$, `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.
 If $t < 0$, a matrix ordering is used for Groebner basis computations, otherwise, and by default, a block ordering is used.
 If w is given and consists of exactly $2 \cdot n$ strictly positive entries, w is used for constructing the weighted homogenized Weyl algebra, see Noro (2002). Otherwise, and by default, the homogenization weight $(1, \dots, 1)$ is used.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel` ≥ 2 , all the debug messages will be printed.
- Example:**
- ```

LIB "dmodapp.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def D2 = Weyl();
setring D2;

```

```

ideal I = 3*x^2*Dy+2*y*Dx, 2*x*Dx+3*y*Dy+6;
intvec u = -2,-3;
intvec v = -u;
GBWeight(I,u,v);
↳ _[1]=2*x*Dx+3*y*Dy+6
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
↳ _[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
ideal J = std(I);
GBWeight(J,u,v); // same as above
↳ _[1]=2*x*Dx+3*y*Dy+6
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
↳ _[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
u = 0,1;
GBWeight(I,u,v);
↳ _[1]=2*x*Dx+3*y*Dy+6
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=-x^3*Dx+y^2*Dx-3*x^2

```

### 7.5.5.7 initialMalgrange

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

- Usage:** `initialMalgrange(f,[a,b,v]);`  $f$  poly,  $a,b$  optional ints,  $v$  opt. intvec
- Return:** ring, Weyl algebra induced by basering, extended by two new vars  $t,Dt$
- Purpose:** computes the initial Malgrange ideal of a given polynomial w.r.t. the weight vector  $(-1,0,\dots,0,1,0,\dots,0)$  such that the weight of  $t$  is  $-1$  and the weight of  $Dt$  is  $1$ .
- Assume:** The basering is commutative and over a field of characteristic  $0$ .
- Note:** Activate the output ring with the `setring` command.  
 The returned ring contains the ideal 'inF', being the initial ideal of the Malgrange ideal of  $f$ .  
 Varnames of the basering should not include  $t$  and  $Dt$ .  
 If  $a < 0$ , `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.  
 If  $b < 0$ , a matrix ordering is used for Groebner basis computations, otherwise, and by default, a block ordering is used.  
 If a positive weight vector  $v$  is given, the weight  $(d,v[1],\dots,v[n],1,d+1-v[1],\dots,d+1-v[n])$  is used for homogenization computations, where  $d$  denotes the weighted degree of  $f$ .  
 Otherwise and by default,  $v$  is set to  $(1,\dots,1)$ . See Noro (2002).
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel`  $\geq 2$ , all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0,(x,y),dp;
poly f = x^2+y^3+x*y^2;
def D = initialMalgrange(f);
setring D;

```

```

inF;
↳ inF [1]=x*Dt
↳ inF [2]=2*x*y*Dx+3*y^2*Dx-y^2*Dy-2*x*Dy
↳ inF [3]=2*x^2*Dx+x*y*Dx+x*y*Dy+18*t*Dt+9*x*Dx-x*Dy+6*y*Dy+4*x+18
↳ inF [4]=18*t*Dt^2+6*y*Dt*Dy-y*Dt+27*Dt
↳ inF [5]=y^2*Dt
↳ inF [6]=2*t*y*Dt+2*x*y*Dx+2*y^2*Dx-6*t*Dt-3*x*Dx-x*Dy-2*y*Dy+2*y-6
↳ inF [7]=x*y^2+y^3+x^2
↳ inF [8]=2*y^3*Dx-2*y^3*Dy-3*y^2*Dx-2*x*y*Dy+y^2*Dy-4*y^2+36*t*Dt+18*x*Dx+1\
2*y*Dy+36
setring r;
intvec v = 3,2;
def D2 = initialMalgrange(f,1,1,v);
setring D2;
inF;
↳ inF [1]=x*Dt
↳ inF [2]=2*x*y*Dx+3*y^2*Dx-y^2*Dy-2*x*Dy
↳ inF [3]=4*x^2*Dx-3*y^2*Dx+2*x*y*Dy+y^2*Dy+36*t*Dt+18*x*Dx+12*y*Dy+8*x+36
↳ inF [4]=18*t*Dt^2+6*y*Dt*Dy-y*Dt+27*Dt
↳ inF [5]=y^2*Dt
↳ inF [6]=2*t*y*Dt-y^2*Dx+y^2*Dy-6*t*Dt-3*x*Dx+x*Dy-2*y*Dy+2*y-6
↳ inF [7]=x*y^2+y^3+x^2
↳ inF [8]=2*y^3*Dx-2*y^3*Dy-3*y^2*Dx-2*x*y*Dy+y^2*Dy-4*y^2+36*t*Dt+18*x*Dx+1\
2*y*Dy+36

```

### 7.5.5.8 initialIdealW

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

- Usage:** `initialIdealW(I,u,v [,s,t,w]);`  
 $I$  ideal,  $u,v$  intvecs,  $s,t$  optional ints,  $w$  an optional intvec
- Return:** ideal, GB of initial ideal of the input ideal wrt the weights  $u$  and  $v$
- Assume:** The basering is the  $n$ -th Weyl algebra in characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .
- Purpose:** computes the initial ideal with respect to given weights.
- Note:**  $u$  and  $v$  are understood as weight vectors for  $x(1..n)$  and  $D(1..n)$  respectively.  
If  $s < 0$ , `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.  
If  $t < 0$ , a matrix ordering is used for Groebner basis computations, otherwise, and by default, a block ordering is used.  
If  $w$  is given and consists of exactly  $2 \cdot n$  strictly positive entries,  $w$  is used as homogenization weight.  
Otherwise, and by default, the homogenization weight  $(1, \dots, 1)$  is used.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel`  $\geq 2$ , all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def D2 = Weyl();
setring D2;
ideal I = 3*x^2*Dy+2*y*Dx, 2*x*Dx+3*y*Dy+6;
intvec u = -2,-3;
intvec v = -u;
initialIdealW(I,u,v);
↳ _[1]=2*x*Dx+3*y*Dy+6
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
↳ _[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
ideal J = std(I);
initialIdealW(J,u,v); // same as above
↳ _[1]=2*x*Dx+3*y*Dy+6
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
↳ _[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
u = 0,1;
initialIdealW(I,u,v);
↳ _[1]=Dx
↳ _[2]=Dy

```

### 7.5.5.9 inForm

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `inForm(I,w)`; I ideal or poly, w intvec

**Return:** ideal, generated by initial forms of generators of I w.r.t. w, or poly, initial form of input poly w.r.t. w

**Purpose:** computes the initial form of an ideal or a poly w.r.t. the weight w

**Note:** The size of the weight vector must be equal to the number of variables of the basering.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def D = Weyl(); setring D;
poly F = 3*x^2*Dy+2*y*Dx;
poly G = 2*x*Dx+3*y*Dy+6;
ideal I = F,G;
intvec w1 = -1,-1,1,1;
intvec w2 = -1,-2,1,2;
intvec w3 = -2,-3,2,3;
inForm(I,w1);
↳ _[1]=2*y*Dx
↳ _[2]=2*x*Dx+3*y*Dy+6
inForm(I,w2);
↳ _[1]=3*x^2*Dy
↳ _[2]=2*x*Dx+3*y*Dy+6
inForm(I,w3);
↳ _[1]=3*x^2*Dy+2*y*Dx

```



```

↳ _[2]=2*x*Dx+3*y*Dy+6
inForm(F,w1);
↳ 2*y*Dx

```

### 7.5.5.10 restrictionIdeal

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

- Usage:** `restrictionIdeal(I,w,[eng,m,G]);`  
 $I$  ideal,  $w$  intvec,  $eng$  and  $m$  optional ints,  $G$  optional ideal
- Return:** ring (a Weyl algebra) containing an ideal 'resIdeal'
- Assume:** The basering is the  $n$ -th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) * \text{var}(i) = \text{var}(i) * \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .  
Further, assume that  $I$  is holonomic and that  $w$  is  $n$ -dimensional with non-negative entries.
- Purpose:** computes the restriction ideal of a holonomic ideal to the subspace defined by the variables corresponding to the non-zero entries of the given intvec
- Note:** The output ring is the Weyl algebra defined by the zero entries of  $w$ . It contains an ideal 'resIdeal' being the restriction ideal of  $I$  wrt  $w$ . If there are no zero entries, the input ring is returned. If  $eng < 0$ , `std` is used for Groebner basis computations, otherwise, and by default, `slingb` is used. The minimal integer root of the  $b$ -function of  $I$  wrt the weight  $(-w, w)$  can be specified via the optional argument  $m$ . The optional argument  $G$  is used for specifying a Groebner basis of  $I$  wrt the weight  $(-w, w)$ , that is, the initial form of  $G$  generates the initial ideal of  $I$  wrt the weight  $(-w, w)$ . Further note, that the assumptions on  $m$  and  $G$  (if given) are not checked.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (a,x,b,Da,Dx,Db), dp;
def D3 = Weyl();
setring D3;
ideal I = a*Db-Dx+2*Da,
x*Db-Da,
x*Da+a*Da+b*Db+1,
x*Dx-2*x*Da-a*Da,
b*Db^2+Dx*Da-Da^2+Db,
a*Dx*Da+2*x*Da^2+a*Da^2+b*Dx*Db+Dx+2*Da;
intvec w = 1,0,0;
def D2 = restrictionIdeal(I,w);
setring D2; D2;

```

```

⇒ // coefficients: QQ
⇒ // number of vars : 4
⇒ // block 1 : ordering C
⇒ // block 2 : ordering dp
⇒ // : names x b Dx Db
⇒ // noncommutative relations:
⇒ // Dxx=x*Dx+1
⇒ // Dbb=b*Db+1
resIdeal;
⇒ resIdeal[1]=2*x*Db-Dx
⇒ resIdeal[2]=x*Dx+2*b*Db+2
⇒ resIdeal[3]=4*b*Db^2+Dx^2+6*Db

```

### 7.5.5.11 restrictionModule

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

- Usage:** `restrictionModule(I,w,[,eng,m,G]);`  
 I ideal, w intvec, eng and m optional ints, G optional ideal
- Return:** ring (a Weyl algebra) containing a module 'resMod'
- Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .  
 Further, assume that I is holonomic and that w is n-dimensional with non-negative entries.
- Purpose:** computes the restriction module of a holonomic ideal to the subspace defined by the variables corresponding to the non-zero entries of the given intvec
- Note:** The output ring is the Weyl algebra defined by the zero entries of w. It contains a module 'resMod' being the restriction module of I wrt w. If there are no zero entries, the input ring is returned.  
 If `eng < 0`, `std` is used for Groebner basis computations, otherwise, and by default, `slingb` is used.  
 The minimal integer root of the b-function of I wrt the weight  $(-w, w)$  can be specified via the optional argument m.  
 The optional argument G is used for specifying a Groebner Basis of I wrt the weight  $(-w, w)$ , that is, the initial form of G generates the initial ideal of I wrt the weight  $(-w, w)$ .  
 Further note, that the assumptions on m and G (if given) are not checked.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel ≥ 2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (a, x, b, Da, Dx, Db), dp;
def D3 = Weyl();
setring D3;

```

```

ideal I = a*Db-Dx+2*Da, x*Db-Da, x*Da+a*Da+b*Db+1,
x*Dx-2*x*Da-a*Da, b*Db^2+Dx*Da-Da^2+Db,
a*Dx*Da+2*x*Da^2+a*Da^2+b*Dx*Db+Dx+2*Da;
intvec w = 1,0,0;
def rm = restrictionModule(I,w);
setring rm; rm;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering C
↳ // block 2 : ordering dp
↳ // : names x b Dx Db
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dbb=b*Db+1
print(resMod);
↳ 2*x*Db-Dx,x*Dx+2*b*Db+2,4*b*Db^2+Dx^2+6*Db

```

### 7.5.5.12 integralIdeal

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `integralIdeal(I,w,[,eng,m,G]);`

`I` ideal, `w` intvec, `eng` and `m` optional ints, `G` optional ideal

**Return:** ring (a Weyl algebra) containing an ideal 'intIdeal'

**Assume:** The basering is the  $n$ -th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

Further, assume that  $I$  is holonomic and that  $w$  is  $n$ -dimensional with non-negative entries.

**Purpose:** computes the integral ideal of a holonomic ideal w.r.t. the subspace defined by the variables corresponding to the non-zero entries of the given intvec.

**Note:** The output ring is the Weyl algebra defined by the zero entries of  $w$ . It contains ideal 'intIdeal' being the integral ideal of  $I$  w.r.t.  $w$ .

If there are no zero entries, the input ring is returned.

If `eng <> 0`, `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.

The minimal integer root of the  $b$ -function of  $I$  wrt the weight  $(-w,w)$  can be specified via the optional argument `m`.

The optional argument `G` is used for specifying a Groebner basis of  $I$  wrt the weight  $(-w,w)$ , that is, the initial form of  $G$  generates the initial ideal of  $I$  wrt the weight  $(-w,w)$ .

Further note, that the assumptions on `m` and `G` (if given) are not checked.

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel >= 2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,b,Dx,Db), dp;
def D2 = Weyl();
setring D2;
ideal I = x*Dx+2*b*Db+2, x^2*Dx+b*Dx+2*x;
intvec w = 1,0;
def D1 = integralIdeal(I,w);
setring D1; D1;
↪ // coefficients: QQ
↪ // number of vars : 2
↪ // block 1 : ordering C
↪ // block 2 : ordering dp
↪ // : names b Db
↪ // noncommutative relations:
↪ // Dbb=b*Db+1
intIdeal;
↪ intIdeal[1]=2*b*Db+1

```

### 7.5.5.13 integralModule

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

- Usage:** `integralModule(I,w,[,eng,m,G]);`  
 I ideal, w intvec, eng and m optional ints, G optional ideal
- Return:** ring (a Weyl algebra) containing a module 'intMod'
- Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .  
 Further, assume that I is holonomic and that w is n-dimensional with non-negative entries.
- Purpose:** computes the integral module of a holonomic ideal w.r.t. the subspace defined by the variables corresponding to the non-zero entries of the given intvec
- Note:** The output ring is the Weyl algebra defined by the zero entries of w. It contains a module 'intMod' being the integral module of I wrt w. If there are no zero entries, the input ring is returned. If `eng <> 0`, `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used. Let  $F(I)$  denote the Fourier transform of I w.r.t. w. The minimal integer root of the b-function of  $F(I)$  w.r.t. the weight  $(-w, w)$  can be specified via the optional argument m. The optional argument G is used for specifying a Groebner Basis of  $F(I)$  wrt the weight  $(-w, w)$ , that is, the initial form of G generates the initial ideal of  $F(I)$  w.r.t. the weight  $(-w, w)$ . Further note, that the assumptions on m and G (if given) are not checked.
- Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel >= 2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,b,Dx,Db), dp;
def D2 = Weyl();
setring D2;
ideal I = x*Dx+2*b*Db+2, x^2*Dx+b*Dx+2*x;
intvec w = 1,0;
def im = integralModule(I,w);
setring im; im;
↳ // coefficients: QQ
↳ // number of vars : 2
↳ // block 1 : ordering C
↳ // block 2 : ordering dp
↳ // : names b Db
↳ // noncommutative relations:
↳ // Dbb=b*Db+1
print(intMod);
↳ 2*b*Db+1,0,
↳ 0, b*Db

```

**7.5.5.14 deRhamCohom**

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `deRhamCohom(f[,w,eng,m]);` `f` poly, `w` optional intvec, `eng` and `m` optional ints

**Return:** ring (a Weyl Algebra) containing a list 'DR' of ideal and int

**Assume:** Basing is commutative and over a field of characteristic 0.

**Purpose:** computes a basis of the  $n$ -th de Rham cohomology group of the complement of the hypersurface defined by `f`, where  $n$  denotes the number of variables of the basering

**Note:** The output ring is the  $n$ -th Weyl algebra. It contains a list 'DR' with two entries (ideal `J` and int `m`) such that  $\{f^m \cdot J[i] : i=1..size(I)\}$  is a basis of the  $n$ -th de Rham cohomology group of the complement of the hypersurface defined by `f`.

If `w` is an intvec with exactly  $n$  strictly positive entries, `w` is used in the computation. Otherwise, and by default, `w` is set to  $(1, \dots, 1)$ .

If `eng`  $\neq 0$ , `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.

If `m` is given, it is assumed to be less than or equal to the minimal integer root of the Bernstein-Sato polynomial of `f`. This assumption is not checked. If not specified, `m` is set to the minimal integer root of the Bernstein-Sato polynomial of `f`.

**Theory:** (SST) pp. 232-235

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel`  $\geq 2$ , all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,y,z), dp;

```

```

poly f = x^3+y^3+z^3;
def A = deRhamCohom(f); // we see that the K-dim is 2
setring A;
DR;
↳ [1]:
↳ _[1]=-x^3*Dx*Dy*Dz
↳ _[2]=-x*y*z*Dx*Dy*Dz
↳ [2]:
↳ -2

```

See also: Section 7.5.5.15 [deRhamCohomIdeal], page 428.

### 7.5.5.15 deRhamCohomIdeal

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp.lib], page 414).

- Usage:** `deRhamCohomIdeal (I[,w,eng,k,G]);`  
 I ideal, w optional intvec, eng and k optional ints, G optional ideal
- Return:** ideal
- Assume:** The basering is the n-th Weyl algebra D over a field of characteristic zero and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .  
 Further, assume that I is of special kind, namely let  $f$  in  $K[x]$  and consider the module  $K[x, 1/f]f^m$ , where  $m$  is smaller than or equal to the minimal integer root of the Bernstein-Sato polynomial of  $f$ .  
 Since this module is known to be a holonomic D-module, it has a cyclic presentation  $D/I$ .
- Purpose:** computes a basis of the n-th de Rham cohomology group of the complement of the hypersurface defined by  $f$
- Note:** The elements of the basis are of the form  $f^m \cdot p$ , where  $p$  runs over the entries of the returned ideal.  
 If I does not satisfy the assumptions described above, the result might have no meaning. Note that I can be computed with `annfs`.  
 If w is an intvec with exactly n strictly positive entries, w is used in the computation. Otherwise, and by default, w is set to  $(1, \dots, 1)$ .  
 If  $\text{eng} < 0$ , `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.  
 Let  $F(I)$  denote the Fourier transform of I wrt w.  
 An integer smaller than or equal to the minimal integer root of the b-function of  $F(I)$  wrt the weight  $(-w, w)$  can be specified via the optional argument k.  
 The optional argument G is used for specifying a Groebner Basis of  $F(I)$  wrt the weight  $(-w, w)$ , that is, the initial form of G generates the initial ideal of  $F(I)$  wrt the weight  $(-w, w)$ .  
 Further note, that the assumptions on I, k and G (if given) are not checked.
- Theory:** (SST) pp. 232-235

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodapp.lib";
ring r = 0, (x,y,z), dp;
poly F = x^3+y^3+z^3;
bfctAnn(F); // Bernstein-Sato poly of F has minimal integer root -2
⇨ [1]:
⇨ _[1]==-1
⇨ _[2]==-4/3
⇨ _[3]==-5/3
⇨ _[4]==-2
⇨ [2]:
⇨ 2,1,1,1
def W = annRat(1,F^2); // so we compute the annihilator of 1/F^2
setring W; W; // Weyl algebra, contains LD = Ann(1/F^2)
⇨ // coefficients: QQ
⇨ // number of vars : 6
⇨ // block 1 : ordering dp
⇨ // : names x y z Dx Dy Dz
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // Dxx=x*Dx+1
⇨ // Dyy=y*Dy+1
⇨ // Dzz=z*Dz+1
LD; // K[x,y,z,1/F]F^(-2) is isomorphic to W/LD as W-module
⇨ LD[1]=x*Dx+y*Dy+z*Dz+6
⇨ LD[2]=z^2*Dy-y^2*Dz
⇨ LD[3]=z^2*Dx-x^2*Dz
⇨ LD[4]=y^2*Dx-x^2*Dy
⇨ LD[5]=x^3*Dz+y^3*Dz+z^3*Dz+6*z^2
⇨ LD[6]=x^3*Dy+y^3*Dy+y^2*z*Dz+6*y^2
deRhamCohomIdeal(LD); // we see that the K-dim is 2
⇨ _[1]=-x^3*Dx*Dy*Dz
⇨ _[2]=-x*y*z*Dx*Dy*Dz
```

See also: Section 7.5.5.14 [deRhamCohom], page 427.

### 7.5.5.16 charVariety

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp.lib], page 414).

**Usage:** `charVariety(I [,eng]);` I an ideal, eng an optional int

**Return:** ring (commutative) containing an ideal 'charVar'

**Purpose:** computes an ideal whose zero set is the characteristic variety of I in the sense of D-module theory

**Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) * \text{var}(i) = \text{var}(i) * \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Note:** The output ring is commutative. It contains an ideal 'charVar'.  
If `eng<>0`, `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodapp.lib";
ring r = 0, (x,y), Dp;
poly F = x3-y2;
printlevel = 0;
def A = annfs(F);
setring A; // Weyl algebra
LD; // the annihilator of F
⇨ LD[1]=2*x*Dx+3*y*Dy+6
⇨ LD[2]=3*x^2*Dy+2*y*Dx
⇨ LD[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
⇨ LD[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
def CA = charVariety(LD);
setring CA; CA; // commutative ring
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x y Dx Dy
⇨ // block 2 : ordering C
charVar;
⇨ charVar[1]=2*x*Dx+3*y*Dy
⇨ charVar[2]=3*x^2*Dy+2*y*Dx
⇨ charVar[3]=9*x*y*Dy^2-4*y*Dx^2
⇨ charVar[4]=27*y^2*Dy^3+8*y*Dx^3
dim(std(charVar)); // hence I is holonomic
⇨ 2
```

See also: Section 7.5.5.17 [charInfo], page 430.

### 7.5.5.17 charInfo

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp.lib], page 414).

**Usage:** `charInfo(I)`; I an ideal

**Return:** ring (commut.) containing ideals 'charVar', 'singLoc' and list 'primDec'

**Purpose:** computes characteristic variety of I (in the sense of D-module theory), its singular locus and primary decomposition

**Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Note:** In the output ring, which is commutative:  
- the ideal 'charVar' is the characteristic variety  $\text{char}(I)$ ,  
- the ideal 'SingLoc' is the singular locus of  $\text{char}(I)$ ,  
- the list 'primDec' is the primary decomposition of  $\text{char}(I)$ .



**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodapp.lib";
ring r = 0, (x,y), Dp;
poly F = x^3-y^2;
printlevel = 0;
def A = annfs(F);
setring A; // Weyl algebra
LD; // the annihilator of F
↳ LD[1]=2*x*Dx+3*y*Dy+6
↳ LD[2]=3*x^2*Dy+2*y*Dx
↳ LD[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
↳ LD[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
def CA = charInfo(LD);
setring CA; CA; // commutative ring
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
charVar; // characteristic variety
↳ charVar[1]=2*x*Dx+3*y*Dy
↳ charVar[2]=3*x^2*Dy+2*y*Dx
↳ charVar[3]=9*x*y*Dy^2-4*y*Dx^2
↳ charVar[4]=27*y^2*Dy^3+8*y*Dx^3
singLoc; // singular locus
↳ singLoc[1]=y*Dy
↳ singLoc[2]=y*Dx
↳ singLoc[3]=2*x*Dx-3*y*Dy
↳ singLoc[4]=9*x*Dy^2-2*Dx^2
↳ singLoc[5]=3*x^2*Dy-y*Dx
↳ singLoc[6]=Dx^3
↳ singLoc[7]=x^3-y^2
primDec; // primary decomposition
↳ [1]:
↳ [1]:
↳ _[1]=Dy
↳ _[2]=Dx
↳ [2]:
↳ _[1]=Dy
↳ _[2]=Dx
↳ [2]:
↳ [1]:
↳ _[1]=27*y*Dy^3+8*Dx^3
↳ _[2]=9*x*Dy^2-4*Dx^2
↳ _[3]=2*x*Dx+3*y*Dy
↳ _[4]=3*x^2*Dy+2*y*Dx
↳ _[5]=x^3-y^2
↳ [2]:
↳ _[1]=27*y*Dy^3+8*Dx^3
↳ _[2]=9*x*Dy^2-4*Dx^2
```

```

↳ _[3]=2*x*Dx+3*y*Dy
↳ _[4]=3*x^2*Dy+2*y*Dx
↳ _[5]=x^3-y^2
↳ [3] :
↳ [1] :
↳ _[1]=y
↳ _[2]=x
↳ [2] :
↳ _[1]=y
↳ _[2]=x

```

### 7.5.5.18 isFsat

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `isFsat(I, F)`; I an ideal, F a poly

**Return:** int, 1 if I is F-saturated and 0 otherwise

**Purpose:** checks whether the ideal I is F-saturated

**Note:** We check indeed that  $\text{Ker}(D \rightarrow F \rightarrow D/I)$  is 0, where D is the basering.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,y), dp;
poly G = x*(x-y)*y;
def A = annfs(G);
setring A;
poly F = x3-y2;
isFsat(LD,F);
↳ 1
ideal J = LD*F;
isFsat(J,F);
↳ 0

```

### 7.5.5.19 appelF1

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `appelF1()`;

**Return:** ring (a parametric Weyl algebra) containing an ideal 'IAappel1'

**Purpose:** defines the ideal in a parametric Weyl algebra,  
which annihilates Appel F1 hypergeometric function

**Note:** The output ring is a parametric Weyl algebra. It contains an ideal  
'IAappel1' annihilating Appel F1 hypergeometric function.  
See (SST) p. 48.

**Example:**

```

LIB "dmodapp.lib";
def A = appelF1();
setring A;
IAappel1;
↳ IAappel1[1]=-x^3*Dx^2+x^2*Dx^2-x^2*y*Dx*Dy+x*y*Dx*Dy+(-a-b-1)*x^2*Dx+(c)*x\

```

```

 *Dx+(-b)*x*y*Dy+(-a*b)*x
 ↪ IAppel1[2]=-x*y^2*Dx*Dy+x*y*Dx*Dy-y^3*Dy^2+y^2*Dy^2+(-d)*x*y*Dx+(-a-d-1)*\
 y^2*Dy+(c)*y*Dy+(-a*d)*y
 ↪ IAppel1[3]=x*Dx*Dy-y*Dx*Dy+(-d)*Dx+(b)*Dy

```

### 7.5.5.20 appelF2

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `appelF2();`

**Return:** ring (a parametric Weyl algebra) containing an ideal 'IAppel2'

**Purpose:** defines the ideal in a parametric Weyl algebra,  
which annihilates Appel F2 hypergeometric function

**Note:** The output ring is a parametric Weyl algebra. It contains an ideal  
'IAppel2' annihilating Appel F2 hypergeometric function.  
See (SST) p. 85.

**Example:**

```

LIB "dmodapp.lib";
def A = appelF2();
setring A;
IAppel2;
 ↪ IAppel2[1]=-x^3*Dx^2+x^2*Dx^2-x^2*y*Dx*Dy+(-a-b-1)*x^2*Dx+x*Dx+(-b)*x*y*D\
 y+(-a*b)*x
 ↪ IAppel2[2]=-x*y^2*Dx*Dy-y^3*Dy^2+y^2*Dy^2+(-c)*x*y*Dx+(-a-c-1)*y^2*Dy+y*D\
 y+(-a*c)*y

```

### 7.5.5.21 appelF4

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `appelF4();`

**Return:** ring (a parametric Weyl algebra) containing an ideal 'IAppel4'

**Purpose:** defines the ideal in a parametric Weyl algebra,  
which annihilates Appel F4 hypergeometric function

**Note:** The output ring is a parametric Weyl algebra. It contains an ideal  
'IAppel4' annihilating Appel F4 hypergeometric function.  
See (SST) p. 39.

**Example:**

```

LIB "dmodapp.lib";
def A = appelF4();
setring A;
IAppel4;
 ↪ IAppel4[1]=-x^2*Dx^2+x*Dx^2-2*x*y*Dx*Dy-y^2*Dy^2+(-a-b-1)*x*Dx+(c)*Dx+(-a\
 -b-1)*y*Dy+(-a*b)
 ↪ IAppel4[2]=-x^2*Dx^2-2*x*y*Dx*Dy-y^2*Dy^2+y*Dy^2+(-a-b-1)*x*Dx+(-a-b-1)*y\
 *Dy+(d)*Dy+(-a*b)

```

### 7.5.5.22 fourier

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `fourier(I,[v]);` I an ideal, v an optional intvec

**Return:** ideal

**Purpose:** computes the Fourier transform of an ideal in a Weyl algebra

**Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Note:** The Fourier automorphism is defined by mapping  $x(i)$  to  $-D(i)$  and  $D(i)$  to  $x(i)$ .

If v is an intvec with entries ranging from 1 to n, the Fourier transform of I restricted to the variables given by v is computed.

**Example:**

```
LIB "dmodapp.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def D2 = Weyl();
setring D2;
ideal I = x*Dx+2*y*Dy+2, x^2*Dx+y*Dx+2*x;
intvec v = 2;
fourier(I,v);
↪ _[1]=x*Dx-2*y*Dy
↪ _[2]=x^2*Dx-Dx*Dy+2*x
fourier(I);
↪ _[1]=-x*Dx-2*y*Dy-1
↪ _[2]=x*Dx^2-x*Dy
```

See also: Section 7.5.5.23 [`inverseFourier`], page 434.

### 7.5.5.23 inverseFourier

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `inverseFourier(I,[v]);` I an ideal, v an optional intvec

**Return:** ideal

**Purpose:** computes the inverse Fourier transform of an ideal in a Weyl algebra

**Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Note:** The Fourier automorphism is defined by mapping  $x(i)$  to  $-D(i)$  and  $D(i)$  to  $x(i)$ .

If v is an intvec with entries ranging from 1 to n, the inverse Fourier transform of I restricted to the variables given by v is computed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def D2 = Weyl();
setring D2;
ideal I = x*Dx+2*y*Dy+2, x^2*Dx+y*Dx+2*x;
intvec v = 2;
ideal FI = fourier(I);
inverseFourier(FI);
 \mapsto _[1]=x*Dx+2*y*Dy+2
 \mapsto _[2]=x^2*Dx+y*Dx+2*x

```

See also: Section 7.5.5.22 [fourier], page 434.

### 7.5.5.24 bFactor

Procedure from library `dmodapp.lib` (see Section 7.5.5 [dmodapp.lib], page 414).

**Usage:** `bFactor(f)`;  $f$  poly

**Return:** list of ideal and intvec and possibly a string

**Purpose:** tries to compute the roots of a univariate poly  $f$

**Note:** The output list consists of two or three entries:  
 roots of  $f$  as an ideal, their multiplicities as intvec, and,  
 if present, a third one being the product of all irreducible factors  
 of degree greater than one, given as string.  
 If  $f$  is the zero polynomial or if  $f$  has no roots in the ground field,  
 this is encoded as root 0 with multiplicity 0.

**Display:** If `printlevel=1`, progress debug messages will be printed,  
 if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0, (x,y), dp;
bFactor((x^2-1)^2);
 \mapsto [1]:
 \mapsto _[1]=1
 \mapsto _[2]=-1
 \mapsto [2]:
 \mapsto 2,2
bFactor((x^2+1)^2);
 \mapsto [1]:
 \mapsto _[1]=0
 \mapsto [2]:
 \mapsto 0
 \mapsto [3]:
 \mapsto x4+2x2+1
bFactor((y^2+1/2)*(y+9)*(y-7));
 \mapsto [1]:
 \mapsto _[1]=7
 \mapsto _[2]=-9
 \mapsto [2]:
 \mapsto 1,1
 \mapsto [3]:

```

```

↳ 2y2+1
bFactor(1);
↳ [1]:
↳ _[1]=0
↳ [2]:
↳ 0
↳ [3]:
↳ 1
bFactor(0);
↳ [1]:
↳ _[1]=0
↳ [2]:
↳ 0
↳ [3]:
↳ 0

```

### 7.5.5.25 intRoots

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `isInt(L)`; L a list

**Return:** list

**Purpose:** extracts integer roots from a list given in `bFactor` format

**Assume:** The input list must be given in the format of `bFactor`.

**Note:** Parameters are treated as integers.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0,x,dp;
list L = bFactor((x-4/3)*(x+3)^2*(x-5)^4); L;
↳ [1]:
↳ _[1]=5
↳ _[2]=4/3
↳ _[3]=-3
↳ [2]:
↳ 4,1,2
intRoots(L);
↳ [1]:
↳ _[1]=5
↳ _[2]=-3
↳ [2]:
↳ 4,2

```

See also: Section 7.5.5.24 [`bFactor`], page 435.

### 7.5.5.26 poly2list

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `poly2list(f)`; f a poly

**Return:** list of exponents and corresponding terms of f

**Purpose:** converts a poly to a list of pairs consisting of intvecs (1st entry) and polys (2nd entry), where the  $i$ -th pair contains the exponent of the  $i$ -th term of  $f$  and the  $i$ -th term (with coefficient) itself.

**Example:**

```
LIB "dmodapp.lib";
ring r = 0,x,dp;
poly F = x;
poly2list(F);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ x
ring r2 = 0,(x,y),dp;
poly F = x2y+5xy2;
poly2list(F);
↳ [1]:
↳ [1]:
↳ 2,1
↳ [2]:
↳ x2y
↳ [2]:
↳ [1]:
↳ 1,2
↳ [2]:
↳ 5xy2
poly2list(0);
↳ [1]:
↳ [1]:
↳ 0,0
↳ [2]:
↳ 0
```

### 7.5.5.27 fl2poly

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `fl2poly(L,s)`;  $L$  a list,  $s$  a string

**Return:** poly

**Purpose:** reconstruct a monic polynomial in one variable from its factorization

**Assume:**  $s$  is a string with the name of some variable and

$L$  is supposed to consist of two entries:

-  $L[1]$  of the type ideal with the roots of a polynomial

-  $L[2]$  of the type intvec with the multiplicities of corr. roots

**Example:**

```
LIB "dmodapp.lib";
ring r = 0,(x,y,z,s),Dp;
ideal I = -1,-4/3,0,-5/3,-2;
intvec mI = 2,1,2,1,1;
list BS = I,mI;
```

```

poly p = f12poly(BS,"s");
p;
↳ s7+7s6+173/9s5+233/9s4+154/9s3+40/9s2
factorize(p,2);
↳ [1]:
↳ _[1]=s+2
↳ _[2]=3s+4
↳ _[3]=3s+5
↳ _[4]=s
↳ _[5]=s+1
↳ [2]:
↳ 1,1,1,2,2

```

### 7.5.5.28 insertGenerator

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `insertGenerator(id,p[,k]);`  
`id` an ideal/module, `p` a poly/vector, `k` an optional int

**Return:** of the same type as `id`

**Purpose:** inserts `p` into `id` at `k`-th position and returns the enlarged object

**Note:** If `k` is given, `p` is inserted at position `k`, otherwise (and by default), `p` is inserted at the beginning (`k=1`).

**Example:**

```

LIB "dmodapp.lib";
ring r = 0,(x,y,z),dp;
ideal I = x^2,z^4;
insertGenerator(I,y^3);
↳ _[1]=y3
↳ _[2]=x2
↳ _[3]=z4
insertGenerator(I,y^3,2);
↳ _[1]=x2
↳ _[2]=y3
↳ _[3]=z4
module M = I*gen(3);
insertGenerator(M,[x^3,y^2,z],2);
↳ _[1]=x2*gen(3)
↳ _[2]=x3*gen(1)+y2*gen(2)+z*gen(3)
↳ _[3]=z4*gen(3)
insertGenerator(M,x+y+z,4);
↳ _[1]=x2*gen(3)
↳ _[2]=z4*gen(3)
↳ _[3]=0
↳ _[4]=x*gen(1)+y*gen(1)+z*gen(1)

```

See also: Section 7.5.5.29 [`deleteGenerator`], page 438.

### 7.5.5.29 deleteGenerator

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `deleteGenerator(id,k);` `id` an ideal/module, `k` an int



**Return:** of the same type as `id`

**Purpose:** deletes the  $k$ -th generator from the first argument and returns the altered object

**Example:**

```
LIB "dmodapp.lib";
ring r = 0, (x,y,z), dp;
ideal I = x^2, y^3, z^4;
deleteGenerator(I, 2);
↪ _[1]=x2
↪ _[2]=z4
module M = [x,y,z], [x2,y2,z2], [x3,y3,z3];
print(deleteGenerator(M, 2));
↪ x, x3,
↪ y, y3,
↪ z, z3
M = M[1];
deleteGenerator(M, 1);
↪ _[1]=0
```

See also: Section 7.5.5.28 [`insertGenerator`], page 438.

### 7.5.5.30 `isInt`

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `isInt(n)`;  $n$  a number

**Return:** int, 1 if  $n$  is an integer or 0 otherwise

**Purpose:** check whether given object of type number is actually an int

**Note:** Parameters are treated as integers.

**Example:**

```
LIB "dmodapp.lib";
ring r = 0, x, dp;
number n = 4/3;
isInt(n);
↪ 0
n = 11;
isInt(n);
↪ 1
```

### 7.5.5.31 `sortIntvec`

Procedure from library `dmodapp.lib` (see Section 7.5.5 [`dmodapp.lib`], page 414).

**Usage:** `sortIntvec(v)`;  $v$  an intvec

**Return:** list of two intvecs

**Purpose:** sorts an intvec

**Note:** In the output list  $L$ , the first entry consists of the entries of  $v$  satisfying  $L[1][i] \geq L[1][i+1]$ . The second entry is a permutation such that  $v[L[2]] = L[1]$ .

Unlike in the procedure `sort`, zeros are not dismissed.

**Example:**

```

LIB "dmodapp.lib";
ring r = 0,x,dp;
intvec v = -1,0,1,-2,0,2;
list L = sortIntvec(v); L;
⇒ [1]:
⇒ 2,1,0,0,-1,-2
⇒ [2]:
⇒ 6,3,2,5,1,4
v[L[2]];
⇒ 2 1 0 0 -1 -2
v = -3,0;
sortIntvec(v);
⇒ [1]:
⇒ 0,-3
⇒ [2]:
⇒ 2,1
v = 0,-3;
sortIntvec(v);
⇒ [1]:
⇒ 0,-3
⇒ [2]:
⇒ 1,2

```

See also: [sort], page 791.

**7.5.6 dmodideal.lib**

**Library:** dmodideal.lib

**Purpose:** Algorithms for Bernstein-Sato ideals of morphisms

**Authors:** Robert Loew, robert.loew at rwth-aachen.de  
 Viktor Levandovskyy, levandov at math.rwth-aachen.de Jorge Martin Morales, jorge at unizar.es

**Overview:** Let  $K$  be a field of characteristic 0. Given a polynomial ring  $R = K[x_1, \dots, x_n]$  and a map, given by polynomials  $F_1, \dots, F_r$  from  $R$ , one is interested in the  $R[1/(F_1 \cdots F_r)]$ -module of rank one, generated by the symbol  $F^s = F_1^{s_1} \cdots F_r^{s_r}$  for symbolic discrete variables  $s_1, \dots, s_r$ . This module  $R[1/(F_1 \cdots F_r)] F^s$  has a structure of a  $D(R)[s_1, \dots, s_r]$ -module, where  $D(R)$  is an  $n$ -th Weyl algebra  $K\langle x_1, \dots, x_n, d_1, \dots, d_n \mid d_j x_j = x_j d_j + 1 \rangle$  and  $D(R)[s] := D(R)$  tensored with  $K[s] := K[s_1, \dots, s_r]$  over  $K$ . We often write just  $D$  for  $D(R)$  and  $D[s]$  for  $D(R)[s]$ .

One is interested in the computation of the following data:

- $\text{Ann}_{D[s]} F^s$ , the annihilator of  $F^s$  in  $D[s]$ ; see `annihilatorMultiFs`
- $\text{Ann}^{\{1\}}_{D[s]} F^s$ , the logarithmic annihilator of  $F^s$  in  $D[s]$ ; see `annfsLogIdeal`
- several kinds of global Bernstein-Sato ideals in  $K[s]$ , cf. (CU) and (Bud12); see `BernsteinSatoIdeal` and `BSidealFromAnn`
- $\text{Ann}_{D} F^\alpha$  for  $\alpha$  from  $K^r$ , the annihilator of  $F^\alpha$  in  $D$ ; see `annfalphaI`
- sub- and over-ideals, bounding the Bernstein-Sato ideal; see `BFBoundsBudur`

**References:**

(BM) the  $\text{Ann} F^s$  algorithm by Briancon and Maisonobe (Remarques sur l'ideal de Bernstein associe a des polynomes, preprint, 2002)

(LM08) V. Levandovskyy and J. Martin-Morales, ISSAC 2008

(CU) Castro and Ucha, On the computation of Bernstein-Sato ideals, JSC 2005

(SST) Saito, Sturmfels, Takayama 'Groebner Deformations of Hypergeometric Differential Equations', Springer, 2000

(Bud12) N. Budur, Bernstein-Sato ideals and local systems, Annales de l'Institut Fourier, Volume 65 (2015) no. 2

(OT99) T. Oaku and N. Takayama, An algorithm for de Rham cohomology groups of the complement of an affine variety via D-module computation, Journal of Pure and Applied Algebra, 1999

**Procedures:** See also: Section 7.5.2 [bfun.lib], page 370; Section 7.5.4 [dmod.lib], page 395; Section 7.5.5 [dmodapp.lib], page 414; Section 7.5.14 [dmodloc.lib], page 519; Section D.6.13 [gmssing.lib], page 862.

### 7.5.6.1 annfsLogIdeal

Procedure from library `dmodideal.lib` (see Section 7.5.6 [dmodideal.lib], page 440).

**Usage:** `annfsLogIdeal(F)`;  $F$  an ideal

**Return:** ring

**Purpose:** compute the logarithmic annihilator of  $F[1]^s(1)*\dots*F[P]^s(P)$

**Assume:** basering is a commutative polynomial ring in characteristic 0

**Note:** activate the output ring with the `setring` command. In this ring, `annfsLog` is the logarithmic annihilator of  $F^s$  (no Groebner basis). If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodideal.lib";
ring R = 0, (x,y), dp;
ideal F = x^3+y^4+x*y^3, x;
def S1 = annfsLogIdeal(F);
setring S1;
annfsLog;
↳ annfsLog[1]=-9*s(1)*x-12*s(1)*y-3*s(2)*x-4*s(2)*y+3*x^2*Dx+4*x*y*Dx+2*x*y\
 *Dy+3*y^2*Dy
↳ annfsLog[2]=-36*s(1)*y^2-192*s(1)*y-9*s(2)*y^2-64*s(2)*y+9*x*y^2*Dx+9*y^3\
 *Dy+64*x*y*Dx+3*x^2*Dy-4*x*y*Dy+48*y^2*Dy
setring R; // now compare with the full annihilator
def S = annihilatorMultiFs(F);
setring S;
annFs;
↳ annFs[1]=-3*x^2*Dx-4*x*y*Dx-2*x*y*Dy-3*y^2*Dy+9*x*s(1)+12*y*s(1)+3*x*s(2)\
 +4*y*s(2)
↳ annFs[2]=-9*x*y^2*Dx-9*y^3*Dy-64*x*y*Dx-3*x^2*Dy+4*x*y*Dy-48*y^2*Dy+36*y^3\
 *s(1)+9*y^2*s(2)+192*y*s(1)+64*y*s(2)
↳ annFs[3]=-3*x^2*y^2*Dx-4*x*y^3*Dx+x*y^3*Dy+3*x^3*Dy+3*x*y^2*s(2)+4*y^3*s(\
 2)
↳ annFs[4]=-144*x^2*y*Dx^2-84*x*y^2*Dx^2-141*x*y^2*Dx*Dy-81*y^3*Dy^2-768*x^2\
 *Dx^2-256*x*y*Dx^2+84*x^2*Dx*Dy-1008*x*y*Dx*Dy-192*y^2*Dx*Dy-51*x^2*Dy^2\
 +64*x*y*Dy^2-336*y^2*Dy^2+36*x*y*Dx*s(2)+84*y^2*Dx*s(2)-27*y^2*Dy*s(2)-70\
 *2*x*y*Dx-84*y^2*Dx-555*y^2*Dy+768*y*Dx*s(1)-384*y*Dy*s(1)+1296*y*s(1)^2+2\
 *56*y*Dx*s(2)-128*y*Dy*s(2)+756*y*s(1)*s(2)+108*y*s(2)^2-3712*x*Dx-256*y*D\
```

```

x+244*x*Dy-2528*y*Dy+1980*y*s(1)+6912*s(1)^2+558*y*s(2)+4608*s(1)*s(2)+76\
8*s(2)^2+8832*s(1)+2944*s(2)
lead(groebner(imap(S1,annfsLog)));
↳ _[1]=3*x^2*Dx
↳ _[2]=9*x*y^2*Dx
↳ _[3]=x*y^3*Dy
↳ _[4]=27*y^4*Dx*Dy
lead(groebner(annFs)); // and we see the difference
↳ _[1]=3*x^2*Dx
↳ _[2]=9*x*y^2*Dx
↳ _[3]=3*y^3*Dx*Dy
↳ _[4]=x*y^3*Dy

```

### 7.5.6.2 annihilatorMultiFs

Procedure from library `dmodideal.lib` (see Section 7.5.6 [`dmodideal.lib`], page 440).

**Usage:** `annihilatorMultiFs(F [,eng,us,ord]);` F an ideal, eng, us, ord optional ints

**Return:** ring

**Purpose:** compute  $\text{Ann}(F[1]^s(1)*\dots*F[P]^s(P))$   
with the multivariate algorithm by Briancon and Maisonobe.

**Assume:** basering is a commutative polynomial ring in characteristic 0

**Note:** activate the output ring with the `setring` command. In this ring, the ideal `annFs` is the annihilator of  $F[1]^s(1)*\dots*F[P]^s(p)$ . If `eng <> 0`, `std` is used for Groebner basis computations, otherwise, and by default `slimgb` is used. If `us <> 0`, then syzygies-driven method is used additionally. If specified, `ord` describes the desired order from the following choices: 0 - 'dp'

1 - elimination order for x, 'dp' in the parts

2 - elimination order for s, 'dp' in the parts

3 - elimination order for x and s, 'dp' in the parts

4 - elimination order for x and D, 'dp' in the parts

(used for the further work in the Bernstein-Sato ideal) If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodideal.lib";
ring R = 0, (x,y), dp;
ideal F = x^2-y,y;
def S = annihilatorMultiFs(F,0,0,0);
setring S;
annFs;
↳ annFs[1]=-2*x*y*Dy-y*Dx+2*x*s(2)
↳ annFs[2]=-x*Dx-2*y*Dy+2*s(1)+2*s(2)
groebner(annFs);
↳ _[1]=x*Dx+2*y*Dy-2*s(1)-2*s(2)
↳ _[2]=2*x*y*Dy+y*Dx-2*x*s(2)
↳ _[3]=4*y^2*Dy^2-y*Dx^2-4*y*Dy*s(1)-8*y*Dy*s(2)+2*y*Dy+4*s(1)*s(2)+4*s(2)^2+2*s(2)

```

### 7.5.6.3 BSidealFromAnn

Procedure from library `dmodideal.lib` (see Section 7.5.6 [`dmodideal.lib`], page 440).

- Usage:** BSidealFromAnn(F, @R [,eng,met]); F an ideal, @R a ring, eng, met optional ints
- Return:** ring
- Purpose:** compute several kinds of Bernstein-Sato ideals, associated to  $f = F[1]^{*..*}F[P]$ , with the multivariate algorithm by Briancon and Maisonobe from  $\text{ann}(F^s)$  as input.
- Assume:** basering is a commutative polynomial ring in characteristic 0 @R is a ring as returned from annihilatorMultiFs.
- Note:** activate the output ring with the `setring` command. In this ring, the ideal BS is a Bernstein-Sato ideal of a polynomial  $f = F[1]^{*..*}F[P]$ . If  $\text{eng} < 0$ , `std` is used for Groebner basis computations, otherwise, and by default `slimgb` is used. If `met` is of type `int`:  
 if  $\text{met} < 0$ , the B-Sigma ideal (cf. (CU)) is computed.  
 If  $0 < \text{met} < P$ , then the ideal `B.met` (cf. (CU)) is computed. If `met` is an `intvec` or a list of `intvecs`, Budurs generalized Bernstein-Sato ideal associated to `met` is computed. Otherwise, and by default, the ideal `B` (cf. (CU)) is computed. If `met` is of type `intvec`: Budurs generalized Bernstein-Sato ideal `B^met.F` is computed. If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodideal.lib";
ring R = 0,(x,y),dp;
ideal F = x+y,x-y,x;
def @R = annihilatorMultiFs(F, 0, 0, 4);
// first we compute the ideal B
def @R2 = BSidealFromAnn(F, @R, 0, 0);
setring @R2;
BS;
↪ BS[1]=s(1)^4*s(2)*s(3)+s(1)^4*s(2)+s(1)^4*s(3)+s(1)^4+3*s(1)^3*s(2)^2*s(3)\
)+3*s(1)^3*s(2)^2+3*s(1)^3*s(2)*s(3)^2+16*s(1)^3*s(2)*s(3)+13*s(1)^3*s(2)\
+3*s(1)^3*s(3)^2+13*s(1)^3*s(3)+10*s(1)^3+3*s(1)^2*s(2)^3*s(3)+3*s(1)^2*s\
(2)^3+6*s(1)^2*s(2)^2*s(3)^2+30*s(1)^2*s(2)^2*s(3)+24*s(1)^2*s(2)^2+3*s(1)\
)^2*s(2)*s(3)^3+30*s(1)^2*s(2)*s(3)^2+83*s(1)^2*s(2)*s(3)+56*s(1)^2*s(2)+\
3*s(1)^2*s(3)^3+24*s(1)^2*s(3)^2+56*s(1)^2*s(3)+35*s(1)^2+s(1)*s(2)^4*s(3)\
)+s(1)*s(2)^4+3*s(1)*s(2)^3*s(3)^2+16*s(1)*s(2)^3*s(3)+13*s(1)*s(2)^3+3*s\
(1)*s(2)^2*s(3)^3+30*s(1)*s(2)^2*s(3)^2+83*s(1)*s(2)^2*s(3)+56*s(1)*s(2)^\
2+s(1)*s(2)*s(3)^4+16*s(1)*s(2)*s(3)^3+83*s(1)*s(2)*s(3)^2+162*s(1)*s(2)*\
s(3)+94*s(1)*s(2)+s(1)*s(3)^4+13*s(1)*s(3)^3+56*s(1)*s(3)^2+94*s(1)*s(3)+\
50*s(1)+s(2)^4*s(3)+s(2)^4+3*s(2)^3*s(3)^2+13*s(2)^3*s(3)+10*s(2)^3+3*s(2)\
)^2*s(3)^3+24*s(2)^2*s(3)^2+56*s(2)^2*s(3)+35*s(2)^2+s(2)*s(3)^4+13*s(2)*\
s(3)^3+56*s(2)*s(3)^2+94*s(2)*s(3)+50*s(2)+s(3)^4+10*s(3)^3+35*s(3)^2+50*\
s(3)+24
setring R;
// secondly we compute the ideal B_1
@R2 = BSidealFromAnn(F, @R, 0, 1);
setring @R2;
BS;
↪ BS[1]=s(1)^2+s(1)*s(2)+s(1)*s(3)+3*s(1)+s(2)+s(3)+2
```

**7.5.6.4 BernsteinSatoIdeal**

Procedure from library `dmodideal.lib` (see Section 7.5.6 [`dmodideal.lib`], page 440).

- Usage:** BernsteinSatoIdeal(F [,eng,met,us]); F an ideal, eng, us optional ints, met optional int or intvec
- Return:** ring
- Purpose:** compute two kinds of Bernstein-Sato ideals, associated to  $f = F[1]^*..*F[P]$ , with the multivariate algorithm by Briancon and Maisonobe.
- Assume:** basering is a commutative polynomial ring in characteristic 0
- Note:** activate the output ring with the `setring` command. In this ring,  
 - the ideal LD is the annihilator of  $F[1]^s_1*..*F[P]^s_p$ ,  
 - the list or ideal BS is a Bernstein-Sato ideal of a polynomial  $f = F[1]^*..*F[P]$ . If `eng <> 0`, `std` is used for Groebner basis computations, otherwise, and by default `slimgb` is used. If `met < 0`, the B-Sigma ideal (cf. Castro and Ucha, 'On the computation of Bernstein-Sato ideals', 2005) is computed. If  $0 < \text{met} < P$ , then the ideal B.P (cf. the paper) is computed. If `met` is an intvec, Budurs generalized Bernstein-Sato ideal associated to `met` is computed.  
 Otherwise, and by default, the ideal B (cf. the paper) is computed. If `us <> 0`, then syzygies-driven method is used.  
 If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodideal.lib";
ring R = 0,(x,y),dp;
ideal F = x^2-y,y;
// first we compute the ideal B:
def S = BernsteinSatoIdeal(F);
setring S;
BS;
↳ BS[1]=4*s(1)^3*s(2)+4*s(1)^3+8*s(1)^2*s(2)^2+28*s(1)^2*s(2)+20*s(1)^2+4*s(1)*s(2)^3+28*s(1)*s(2)^2+55*s(1)*s(2)+31*s(1)+4*s(2)^3+20*s(2)^2+31*s(2)+15
// secondly we compute the ideal B_1:
setring R;
def S = BernsteinSatoIdeal(F,0,1);
↳ // ** redefining S (def S = BernsteinSatoIdeal(F,0,1);) ./examples/Bernst\
 einSatoIdeal.sing:10
setring S;
BS;
↳ BS[1]=2*s(1)^2+2*s(1)*s(2)+5*s(1)+2*s(2)+3
// thirdly we compute the ideal B_sigma:
setring R;
def S = BernsteinSatoIdeal(F,0,-1);
↳ // ** redefining S (def S = BernsteinSatoIdeal(F,0,-1);) ./examples/Berns\
 teinSatoIdeal.sing:15
setring S;
BS;
↳ BS[1]=2*s(1)*s(2)+2*s(1)+2*s(2)^2+5*s(2)+3
↳ BS[2]=2*s(1)^2+3*s(1)-2*s(2)^2-3*s(2)
```

**7.5.6.5 BFBoundsBudur**

Procedure from library `dmodideal.lib` (see Section 7.5.6 [`dmodideal.lib`], page 440).

- Usage:** BFBoundsBudur(F,m); F an ideal, m an intvec
- Return:** ring
- Assume:** basering is a commutative polynomial ring in characteristic 0
- Purpose:** determine upper and lower bounds of the Bernstein-Sato ideal associated to m with the method of (Bud12)
- Note:** The returned ring contains lists Bj, containing the Bernstein-Sato ideals associated to  $e_j$ , shiftedIdeals, containing the shifted ideals from (Bud12) 4.7, and ideals upperBound, lowerBound which give upper bound and lower bound for the Bernstein-Sato-Ideal associated to m respectively.

**Example:**

```
LIB "dmodideal.lib";
ring r = 0, (x,y,z), dp;
setring r;
ideal F = x*z, 2*x^2*y^2*z+x^4+y^4;
def A = BFBoundsBudur(F, intvec(1,1));
setring A;
lead(upperBound);
↪ _[1]=2*s(1)^8*s(2)^2
↪ _[2]=s(1)^9*s(2)
lead(lowerBound);
↪ _[1]=s(1)^11*s(2)
↪ _[2]=2*s(1)^10*s(2)^2
↪ _[3]=2*s(1)^10*s(2)^2
↪ _[4]=4*s(1)^9*s(2)^3
```

**7.5.6.6 annfalphi**

Procedure from library `dmodideal.lib` (see Section 7.5.6 [dmodideal.lib], page 440).

- Usage:** annfalphi(f,alpha); f,alpha ideals
- Return:** ring
- Assume:** basering is a commutative polynomial ring in characteristic 0
- Purpose:** determine annihilator of  $f^\alpha$  with the method of (OT99)
- Note:** The returned ring contains the annihilator of  $f^\alpha$  over D as annfalphi. alpha should contain the desired rational exponents.  
The procedure may also be applied to the univariate case, i.e. for  $r=1$ .

**Example:**

```
LIB "dmodideal.lib";
ring R = 0, (x,y,z), dp;
ideal f = x,y,z;
ideal alpha = 1/4, 2/3, 1;
def A = annfalphi(f,alpha);
setring A;
annfalphi;
↪ annfalphi[1]=Dz^2
↪ annfalphi[2]=z*Dz-1
↪ annfalphi[3]=3*y*Dy-2
↪ annfalphi[4]=4*x*Dx-1
```

### 7.5.6.7 extractS

Procedure from library `dmodideal.lib` (see Section 7.5.6 [`dmodideal.lib`], page 440).

**Usage:** `extractS(I,r)`; I ideal, r int

**Return:** ring

**Assume:** I is an ideal in the first r variables of the basering and these r variables generate a commutative subring

**Purpose:** give the ideal generated by I in the commutative subring generated by the first r variables, ordering dp

**Note:** The returned ring contains I.

**Example:**

```
LIB "dmodideal.lib";
ring R = 0,(x,y),dp;
ideal f = x^2-y^2,y;
def S = BernsteinSatoIdeal(f);
setring S;
BS;
⇒ BS[1]=8*s(1)^4*s(2)+8*s(1)^4+12*s(1)^3*s(2)^2+56*s(1)^3*s(2)+44*s(1)^3+6*s(1)^2*s(2)^3+54*s(1)^2*s(2)^2+136*s(1)^2*s(2)+88*s(1)^2+s(1)*s(2)^4+16*s(1)*s(2)^3+77*s(1)*s(2)^2+138*s(1)*s(2)+76*s(1)+s(2)^4+10*s(2)^3+35*s(2)^2+50*s(2)+24
def T = extractS(BS,2);
setring T;
I;
⇒ I[1]=8*s(1)^4*s(2)+12*s(1)^3*s(2)^2+6*s(1)^2*s(2)^3+s(1)*s(2)^4+8*s(1)^4+56*s(1)^3*s(2)+54*s(1)^2*s(2)^2+16*s(1)*s(2)^3+s(2)^4+44*s(1)^3+136*s(1)^2*s(2)+77*s(1)*s(2)^2+10*s(2)^3+88*s(1)^2+138*s(1)*s(2)+35*s(2)^2+76*s(1)+50*s(2)+24
factorize(I[1]);
⇒ [1]:
⇒ _[1]=1
⇒ _[2]=s(1)+1
⇒ _[3]=s(2)+1
⇒ _[4]=2*s(1)+s(2)+2
⇒ _[5]=2*s(1)+s(2)+3
⇒ _[6]=2*s(1)+s(2)+4
⇒ [2]:
⇒ 1,1,1,1,1,1
```

### 7.5.7 dmodvar\_lib

**Library:** `dmodvar.lib`

**Purpose:** Algebraic D-modules for varieties

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Support: DFG Graduiertenkolleg 1632 'Experimentelle und konstruktive Algebra'



**Overview:** Let  $K$  be a field of characteristic 0. Given a polynomial ring  $R = K[x_1, \dots, x_n]$  and polynomials  $f_1, \dots, f_r$  in  $R$ , define  $F = f_1 \cdots f_r$  and  $F^s = f_1^{s_1} \cdots f_r^{s_r}$  for symbolic discrete (that is shiftable) variables  $s_1, \dots, s_r$ . The module  $R[1/F]^*F^s$  has the structure of a  $D\langle S \rangle$ -module, where  $D\langle S \rangle = D(R)$  tensored with  $S$  over  $K$ , where

- $D(R)$  is the  $n$ -th Weyl algebra  $K\langle x_1, \dots, x_n, d_1, \dots, d_n \mid d_j x_j = x_j d_j + 1 \rangle$
- $S$  is the universal enveloping algebra of  $\mathfrak{gl}_r$ , generated by  $s_i = s_{\{i\}}$ .

One is interested in the following data:

- the left ideal  $\text{Ann } F^s$  in  $D\langle S \rangle$ , usually denoted by  $LD$  in the output
- global Bernstein polynomial in one variable  $s = s_1 + \dots + s_r$ , denoted by  $bs$ ,
- its minimal integer root  $s_0$ , the list of all roots of  $bs$ , which are known to be negative rational numbers, with their multiplicities, which is denoted by  $BS$
- an  $r$ -tuple of operators in  $D\langle S \rangle$ , denoted by  $PS$ , such that the functional equality  $\sum_{k=1}^r P_k * f_k * F^s = bs * F^s$  holds in  $R[1/F]^*F^s$ .

**References:**

- (BMS06) Budur, Mustata, Saito: Bernstein-Sato polynomials of arbitrary varieties (2006).
- (ALM09) Andres, Levandovskyy, Martin-Morales: Principal Intersection and Bernstein-Sato Polynomial of an Affine Variety (2009).

**Procedures:** See also: Section 7.5.2 [`bfun.lib`], page 370; Section 7.5.4 [`dmod.lib`], page 395; Section 7.5.5 [`dmodapp.lib`], page 414; Section D.6.13 [`gmssing.lib`], page 862.

### 7.5.7.1 `bfcTVarIn`

Procedure from library `dmodvar.lib` (see Section 7.5.7 [`dmodvar.lib`], page 446).

**Usage:** `bfcTVarIn(I [,a,b,c]);`  $I$  an ideal,  $a, b, c$  optional ints

**Return:** list of ideal and `intvec`

**Purpose:** computes the roots of the Bernstein-Sato polynomial and their multiplicities for an affine algebraic variety defined by  $I$ .

**Assume:** The basering is commutative and over a field of characteristic 0. Varnames of the basering do not include  $t(1), \dots, t(r)$  and  $Dt(1), \dots, Dt(r)$ , where  $r$  is the number of entries of the input ideal.

**Note:** In the output list, say  $L$ ,

- $L[1]$  of type ideal contains all the rational roots of a b-function,
- $L[2]$  of type `intvec` contains the multiplicities of above roots,
- optional  $L[3]$  of type string is the part of b-function without rational roots.

Note, that a b-function of degree 0 is encoded via  $L[1][1]=0$ ,  $L[2]=0$  and  $L[3]$  is 1 (for nonzero constant) or 0 (for zero b-function).

If  $a < 0$ , the ideal is used as given. Otherwise, and by default, a heuristically better suited generating set is used to reduce computation time.

If  $b < 0$ , `std` is used for GB computations in characteristic 0, otherwise, and by default, `slingb` is used.

If  $c < 0$ , a matrix ordering is used for GB computations, otherwise, and by default, a block ordering is used.

Further note, that in this proc, the initial ideal of the multivariate Malgrange ideal defined by  $I$  is computed and then a system of linear equations is solved by linear reductions following the ideas by Noro.

The result is shifted by  $1 - \text{codim}(\text{Var}(F))$  following (BMS06).

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodvar.lib";
ring R = 0, (x,y,z), dp;
ideal F = x^2+y^3, z;
list L = bfctVarIn(F);
L;
⇒ [1]:
⇒ _[1]=-5/6
⇒ _[2]=-1
⇒ _[3]=-7/6
⇒ [2]:
⇒ 1,1,1
```

### 7.5.7.2 bfctVarAnn

Procedure from library `dmodvar.lib` (see Section 7.5.7 [`dmodvar.lib`], page 446).

**Usage:** `bfctVarAnn(F[,gid,eng])`;  $F$  an ideal, `gid,eng` optional ints

**Return:** list of an ideal and an intvec

**Purpose:** computes the roots of the Bernstein-Sato polynomial and their multiplicities for an affine algebraic variety defined by  $F = F[1], \dots, F[r]$ .

**Assume:** The basering is commutative and over a field in char 0.

**Note:** In the output list, the ideal contains all the roots and the intvec their multiplicities. If `gid<>0`, the ideal is used as given. Otherwise, and by default, a heuristically better suited generating set is used. If `eng<>0`, `std` is used for GB computations, otherwise, and by default, `slingb` is used. Computational remark: The time of computation can be very different depending on the chosen generators of  $F$ , although the result is always the same. Further note that in this proc, the annihilator of  $f^s$  in  $D[s]$  is computed and then a system of linear equations is solved by linear reductions in order to find the minimal polynomial of  $S = s(1)(1) + \dots + s(P)(P)$ . The result is shifted by  $1 - \text{codim}(\text{Var}(F))$  following (BMS06).

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodvar.lib";
ring R = 0, (x,y,z), Dp;
ideal F = x^2+y^3, z;
bfctVarAnn(F);
⇒ [1]:
⇒ _[1]=-5/6
⇒ _[2]=-1
⇒ _[3]=-7/6
⇒ [2]:
⇒ 1,1,1
```

### 7.5.7.3 SannfsVar

Procedure from library `dmodvar.lib` (see Section 7.5.7 [`dmodvar.lib`], page 446).

**Usage:** `SannfsVar(F [,ORD,eng]);`  $F$  an ideal,  $ORD$  an optional string,  $eng$  an optional int

**Return:** ring (Weyl algebra tensored with  $U(\mathfrak{gl}_P)$ ), containing an ideal  $LD$

**Purpose:** compute the  $D\langle S \rangle$ -module structure of  $D\langle S \rangle * f^s$  where  $f = F[1]^* \dots * F[P]$  and  $D\langle S \rangle$  is the Weyl algebra  $D$  tensored with  $K\langle S \rangle = U(\mathfrak{gl}_P)$ , according to the generalized algorithm by Briancon and Maisonobe for affine varieties

**Assume:** The basering is commutative and over a field of characteristic 0.

**Note:** Activate the output ring  $D\langle S \rangle$  with the `setring` command. In the ring  $D\langle S \rangle$ , the ideal  $LD$  is the needed  $D\langle S \rangle$ -module structure.

The value of  $ORD$  must be an elimination ordering in  $D\langle Dt, S \rangle$  for  $Dt$  written in the string form, otherwise the result may have no meaning. By default  $ORD = '(a(1..(P)..1), a(1..(P+P^2)..1), dp)'$ .

If  $eng > 0$ , `std` is used for Groebner basis computations, otherwise, and by default `slimgb` is used.

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodvar.lib";
ring R = 0, (x,y), Dp;
ideal F = x^3, y^5;
//ORD = "(a(1,1), a(1,1,1,1,1,1), dp)";
//eng = 0;
def A = SannfsVar(F);
setring A;
A;
↳ // coefficients: QQ
↳ // number of vars : 8
↳ // block 1 : ordering a
↳ // : names s(1)(1) s(1)(2) s(2)(1) s(2)(2)
↳ // : weights 1 1 1 1
↳ // block 2 : ordering dp
↳ // : names s(1)(1) s(1)(2) s(2)(1) s(2)(2) x y Dx Dy
↳ // block 3 : ordering C
↳ // noncommutative relations:
↳ // s(1)(2)s(1)(1)=s(1)(1)*s(1)(2)-s(1)(2)
↳ // s(2)(1)s(1)(1)=s(1)(1)*s(2)(1)+s(2)(1)
↳ // s(2)(1)s(1)(2)=s(1)(2)*s(2)(1)-s(1)(1)+s(2)(2)
↳ // s(2)(2)s(1)(2)=s(1)(2)*s(2)(2)-s(1)(2)
↳ // s(2)(2)s(2)(1)=s(2)(1)*s(2)(2)+s(2)(1)
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
LD;
↳ LD[1]=5*s(2)(2)-y*Dy
↳ LD[2]=3*s(1)(1)-x*Dx
↳ LD[3]=15*s(1)(2)*s(2)(1)-x*y*Dx*Dy-5*x*Dx
↳ LD[4]=5*s(2)(1)*y^4-x^3*Dy
↳ LD[5]=3*s(1)(2)*x^2-y^5*Dx
```

### 7.5.7.4 makeMalgrange

Procedure from library `dmodvar.lib` (see Section 7.5.7 [`dmodvar.lib`], page 446).

**Usage:** `makeMalgrange(F [,ORD]);` F an ideal, ORD an optional string

**Return:** ring (Weyl algebra) containing an ideal IF

**Purpose:** create the ideal by Malgrange associated with  $F = F[1], \dots, F[P]$ .

**Note:** Activate the output ring with the `setring` command. In this ring, the ideal IF is the ideal by Malgrange corresponding to F.

The value of ORD must be an arbitrary ordering in  $K\langle t, x, Dt, Dx \rangle$  written in the string form. By default `ORD = 'dp'`.

**Display:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodvar.lib";
ring R = 0, (x,y,z), Dp;
ideal I = x^2+y^3, z;
def W = makeMalgrange(I);
setring W;
W;
↳ // coefficients: QQ
↳ // number of vars : 10
↳ // block 1 : ordering dp
↳ // : names t(1) t(2) x y z Dt(1) Dt(2) Dx Dy Dz
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dt(1)t(1)=t(1)*Dt(1)+1
↳ // Dt(2)t(2)=t(2)*Dt(2)+1
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
↳ // Dzz=z*Dz+1
IF;
↳ IF[1]=-y^3-x^2+t(1)
↳ IF[2]=t(2)-z
↳ IF[3]=2*x*Dt(1)+Dx
↳ IF[4]=3*y^2*Dt(1)+Dy
↳ IF[5]=Dt(2)+Dz
```

### 7.5.8 involut.lib

**Library:** `involut.lib`

**Purpose:** Computations and operations with involutions

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**Overview:** Involution is an anti-automorphism of a non-commutative K-algebra with the property that applied an involution twice, one gets an identity. Involution is linear with respect to the ground field. In this library we compute linear involutions, distinguishing the case of a diagonal matrix (such involutions are called homothetic) and a general one. Also, linear automorphisms of different order can be computed.

**Support:** Forschungsschwerpunkt 'Mathematik und Praxis' (Project of Dr. E. Zerz and V. Levandovskyy), Uni Kaiserslautern

**Remark:** This library provides algebraic tools for computations and operations with algebraic involutions and linear automorphisms of non-commutative algebras

**Procedures:**

### 7.5.8.1 findInvo

Procedure from library `involut.lib` (see Section 7.5.8 [`involut.lib`], page 450).

**Usage:** `findInvo();`

**Return:** a ring containing a list `L` of pairs, where  
`L[i][1]` = ideal; a Groebner Basis of an  $i$ -th associated prime,  
`L[i][2]` = matrix, defining a linear map, with entries, reduced with respect to `L[i][1]`

**Purpose:** computed the ideal of linear involutions of the basering

**Assume:** the relations on the algebra are of the form  $YX = XY + D$ , that is the current ring is a  $G$ -algebra of Lie type.

**Note:** for convenience, the full ideal of relations `idJ` and the initial matrix with indeterminates `matD` are exported in the output ring

**Example:**

```
LIB "involut.lib";
def a = makeWeyl(1);
setring a; // this algebra is a first Weyl algebra
a;
⇒ // coefficients: QQ
⇒ // number of vars : 2
⇒ // block 1 : ordering dp
⇒ // : names x D
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Dx=xD+1
def X = findInvo();
setring X; // ring with new variables, corr. to unknown coefficients
X;
⇒ // coefficients: QQ
⇒ // number of vars : 4
⇒ // block 1 : ordering dp
⇒ // : names a11 a12 a21 a22
⇒ // block 2 : ordering C
L;
⇒ [1]:
⇒ [1]:
⇒ _[1]=a11+a22
⇒ _[2]=a12*a21+a22^2-1
⇒ [2]:
⇒ _[1,1]=-a22
⇒ _[1,2]=a12
⇒ _[2,1]=a21
⇒ _[2,2]=a22
```

```

// look at the matrix in the new variables, defining the linear involution
print(L[1][2]);
↳ -a22, a12,
↳ a21, a22
L[1][1]; // where new variables obey these relations
↳ _[1]=a11+a22
↳ _[2]=a12*a21+a22^2-1
idJ;
↳ idJ[1]=-a12*a21+a11*a22+1
↳ idJ[2]=a11^2+a12*a21-1
↳ idJ[3]=a11*a12+a12*a22
↳ idJ[4]=a11*a21+a21*a22
↳ idJ[5]=a12*a21+a22^2-1

```

See also: Section 7.5.8.2 [findInvoDiag], page 452; Section 7.5.8.5 [involution], page 455.

### 7.5.8.2 findInvoDiag

Procedure from library `involut.lib` (see Section 7.5.8 [involut.lib], page 450).

**Usage:** `findInvoDiag();`

**Return:** a ring together with a list of pairs  $L$ , where  
 $L[i][1]$  = ideal; a Groebner Basis of an  $i$ -th associated prime,  
 $L[i][2]$  = matrix, defining a linear map, with entries, reduced with respect to  $L[i][1]$

**Purpose:** compute homothetic (diagonal) involutions of the basering

**Assume:** the relations on the algebra are of the form  $YX = XY + D$ , that is the current ring is a  $G$ -algebra of Lie type.

**Note:** for convenience, the full ideal of relations `idJ` and the initial matrix with indeterminates `matD` are exported in the output ring

**Example:**

```

LIB "involut.lib";
def a = makeWeyl(1);
setring a; // this algebra is a first Weyl algebra
a;
↳ // coefficients: QQ
↳ // number of vars : 2
↳ // block 1 : ordering dp
↳ // : names x D
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dx=xD+1
def X = findInvoDiag();
setring X; // ring with new variables, corresponding to unknown coefficients
X;
↳ // coefficients: QQ
↳ // number of vars : 2
↳ // block 1 : ordering dp
↳ // : names a11 a22
↳ // block 2 : ordering C
// print matrices, defining linear involutions
print(L[1][2]); // a first matrix: we see it is constant

```

```

↳ -1,0,
↳ 0, 1
print(L[2][2]); // and a second possible matrix; it is constant too
↳ 1,0,
↳ 0,-1
L; // let us take a look on the whole list
↳ [1]:
↳ [1]:
↳ _[1]=a22-1
↳ _[2]=a11+1
↳ [2]:
↳ _[1,1]=-1
↳ _[1,2]=0
↳ _[2,1]=0
↳ _[2,2]=1
↳ [2]:
↳ [1]:
↳ _[1]=a22+1
↳ _[2]=a11-1
↳ [2]:
↳ _[1,1]=1
↳ _[1,2]=0
↳ _[2,1]=0
↳ _[2,2]=-1
idJ;
↳ idJ[1]=a11*a22+1
↳ idJ[2]=a11^2-1
↳ idJ[3]=a22^2-1

```

See also: Section 7.5.8.1 [findInvo], page 451; Section 7.5.8.5 [involution], page 455.

### 7.5.8.3 findAuto

Procedure from library `involut.lib` (see Section 7.5.8 [involut.lib], page 450).

**Usage:** `findAuto(n)`;  $n$  an integer

**Return:** a ring together with a list of pairs  $L$ , where  
 $L[i][1]$  = ideal; a Groebner Basis of an  $i$ -th associated prime,  
 $L[i][2]$  = matrix, defining a linear map, with entries, reduced with respect to  $L[i][1]$

**Purpose:** compute the ideal of linear automorphisms of the basering,  
given by a matrix,  $n$ -th power of which gives identity (i.e. unipotent matrix)

**Assume:** the relations on the algebra are of the form  $YX = XY + D$ , that is the current ring is  
a  $G$ -algebra of Lie type.

**Note:** if  $n=0$ , a matrix, defining an automorphism is not assumed to be unipotent  
but just non-degenerate. A nonzero parameter  $\mathcal{O}p$  is introduced as the value of  
the determinant of the matrix above.  
For convenience, the full ideal of relations `idJ` and the initial matrix with indetermi-  
nates  
`matD` are mutually exported in the output ring

**Example:**

```

LIB "involut.lib";
def a = makeWeyl(1);
setring a; // this algebra is a first Weyl algebra
a;
↳ // coefficients: QQ
↳ // number of vars : 2
↳ // block 1 : ordering dp
↳ // : names x D
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dx=xD+1
def X = findAuto(2); // in contrast to findInvo look for automorphisms
setring X; // ring with new variables - unknown coefficients
X;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names a11 a12 a21 a22
↳ // block 2 : ordering C
size(L); // we have (size(L)) families in the answer
↳ 2
// look at matrices, defining linear automorphisms:
print(L[1][2]); // a first one: we see it is the identity
↳ 1,0,
↳ 0,1
print(L[2][2]); // and a second possible matrix; it is diagonal
↳ -1,0,
↳ 0, -1
// L; // we can take a look on the whole list, too
idJ;
↳ idJ[1]=-a12*a21+a11*a22-1
↳ idJ[2]=a11^2+a12*a21-1
↳ idJ[3]=a11*a12+a12*a22
↳ idJ[4]=a11*a21+a21*a22
↳ idJ[5]=a12*a21+a22^2-1
kill X; kill a;
//----- find all the linear automorphisms -----
//----- use the call findAuto(0) -----
ring R = 0,(x,s),dp;
def r = nc_algebra(1,s); setring r; // the shift algebra
s*x; // the only relation in the algebra is:
↳ xs+s
def Y = findAuto(0);
setring Y;
size(L); // here, we have 1 parametrized family
↳ 1
print(L[1][2]); // here, @p is a nonzero parameter
↳ 1,a12,
↳ 0,(@p)
det(L[1][2]-@p); // check whether determinante is zero
↳ 0

```

See also: Section 7.5.8.1 [findInvo], page 451.



### 7.5.8.4 ncdetection

Procedure from library `involut.lib` (see Section 7.5.8 [`involut.lib`], page 450).

**Usage:** `ncdetection()`;

**Return:** ideal, representing an involution map

**Purpose:** compute classical involutions (i.e. acting rather on operators than on variables) for some particular noncommutative algebras

**Assume:** the procedure is aimed at non-commutative algebras with differential, shift or advance operators arising in Control Theory. It has to be executed in a ring.

**Example:**

```
LIB "involut.lib";
ring R = 0, (x,y,z,D(1..3)), dp;
matrix D[6][6];
D[1,4]=1; D[2,5]=1; D[3,6]=1;
def r = nc_algebra(1,D); setring r;
ncdetection();
↳ _[1]=x
↳ _[2]=y
↳ _[3]=z
↳ _[4]=-D(1)
↳ _[5]=-D(2)
↳ _[6]=-D(3)
kill r, R;
//-----
ring R=0, (x,S), dp;
def r = nc_algebra(1,-S); setring r;
ncdetection();
↳ _[1]=-x
↳ _[2]=S
kill r, R;
//-----
ring R=0, (x,D(1),S), dp;
matrix D[3][3];
D[1,2]=1; D[1,3]=-S;
def r = nc_algebra(1,D); setring r;
ncdetection();
↳ _[1]=-x
↳ _[2]=D(1)
↳ _[3]=S
```

### 7.5.8.5 involution

Procedure from library `involut.lib` (see Section 7.5.8 [`involut.lib`], page 450).

**Usage:** `involution(m, theta)`; `m` is a poly/vector/ideal/matrix/module, `theta` is a map

**Return:** object of the same type as `m`

**Purpose:** applies the involution, presented by `theta` to the object `m`

**Theory:** for an involution `theta` and two polynomials `a, b` from the algebra,  $\text{theta}(ab) = \text{theta}(b) \text{theta}(a)$ ; `theta` is linear with respect to the ground field

**Note:** This is generalized "theta(m)" for data types unsupported by "map".

**Example:**

```
LIB "involut.lib";
ring R = 0,(x,d),dp;
def r = nc_algebra(1,1); setring r; // Weyl-Algebra
map F = r,x,-d;
F(F); // should be maxideal(1) for an involution
↪ _[1]=x
↪ _[2]=d
poly f = x*d^2+d;
poly If = involution(f,F);
f-If;
↪ 0
poly g = x^2*d+2*x*d+3*x+7*d;
poly tg = -d*x^2-2*d*x+3*x-7*d;
poly Ig = involution(g,F);
tg-Ig;
↪ 0
ideal I = f,g;
ideal II = involution(I,F);
II;
↪ II[1]=xd2+d
↪ II[2]=-x2d-2xd+x-7d-2
matrix(I) - involution(II,F);
↪ _[1,1]=0
↪ _[1,2]=0
module M = [f,g,0],[g,0,x^2*d];
module IM = involution(M,F);
print(IM);
↪ xd2+d, -x2d-2xd+x-7d-2,
↪ -x2d-2xd+x-7d-2,0,
↪ 0, -x2d-2x
print(matrix(M) - involution(IM,F));
↪ 0,0,
↪ 0,0,
↪ 0,0
```

### 7.5.8.6 isInvolution

Procedure from library `involut.lib` (see Section 7.5.8 [`involut.lib`], page 450).

**Usage:** `isInvolution(F)`; F is a map from current ring to itself

**Return:** integer, 1 if F determines an involution and 0 otherwise

**Theory:** involution is an antiautomorphism of order 2

**Assume:** F is a map from current ring to itself

**Example:**

```
LIB "involut.lib";
def A = makeUsl(2); setring A;
map I = A,-e,-f,-h; //correct antiauto involution
isInvolution(I);
↪ 1
```

```

map J = A,3*e,1/3*f,-h; // antiauto but not involution
isInvolution(J);
↳ 0
map K = A,f,e,-h; // not antiauto
isInvolution(K);
↳ 0

```

See also: Section 7.5.8.1 [findInvo], page 451; Section 7.5.8.5 [involution], page 455; Section 7.5.8.7 [isAntiEndo], page 457.

### 7.5.8.7 isAntiEndo

Procedure from library `involut.lib` (see Section 7.5.8 [involut.lib], page 450).

**Usage:** `isAntiEndo(F)`;  $F$  is a map from current ring to itself

**Return:** integer, 1 if  $F$  determines an antiendomorphism of current ring and 0 otherwise

**Assume:**  $F$  is a map from current ring to itself

**Example:**

```

LIB "involut.lib";
def A = makeUs1(2); setring A;
map I = A,-e,-f,-h; //correct antiauto involution
isAntiEndo(I);
↳ 1
map J = A,3*e,1/3*f,-h; // antiauto but not involution
isAntiEndo(J);
↳ 1
map K = A,f,e,-h; // not antiendo
isAntiEndo(K);
↳ 0

```

See also: Section 7.5.8.1 [findInvo], page 451; Section 7.5.8.5 [involution], page 455; Section 7.5.8.6 [isInvolution], page 456.

### 7.5.9 gkdim\_lib

**Library:** `gkdim.lib`

**Purpose:** Procedures for calculating the Gelfand-Kirillov dimension

**Authors:** Lobillo, F.J., [jlobillo@ugr.es](mailto:jlobillo@ugr.es)  
 Rabelo, C., [crabelo@ugr.es](mailto:crabelo@ugr.es)

Support: 'Metodos algebraicos y efectivos en grupos cuanticos', BFM2001-3141, MCYT, Jose Gomez-Torrecillas (Main researcher).

**Note:** The built-in command `dim`, executed for a module in `@plural`, computes the Gelfand-Kirillov dimension.

**Procedures:**

#### 7.5.9.1 GKdim

Procedure from library `gkdim.lib` (see Section 7.5.9 [gkdim.lib], page 457).

**Usage:** `GKdim(L)`;  $L$  is a left ideal/module/matrix

**Return:** int

**Purpose:** compute the Gelfand-Kirillov dimension of the factor-module, whose presentation is given by  $L$ , e.g.  $R^r/L$

**Note:** if the factor-module is zero, -1 is returned

**Example:**

```
LIB "gkdim.lib";
ring R = 0,(x,y,z),Dp;
matrix C[3][3]=0,1,1,0,0,-1,0,0,0;
matrix D[3][3]=0,0,0,0,0,x;
def r = nc_algebra(C,D); setring r;
r;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ // block 1 : ordering Dp
↳ // : names x y z
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // zy=-yz+x
ideal I=x;
GKdim(I);
↳ 2
ideal J=x^2,y;
GKdim(J);
↳ 1
module M=[x^2,y,1],[x,y^2,0];
GKdim(M);
↳ 3
ideal A = x,y,z;
GKdim(A);
↳ 0
ideal B = 1;
GKdim(B);
↳ -1
GKdim(ideal(0)) == nvars(basing); // should be true, i.e., evaluated to 1
↳ 1
```

### 7.5.10 ncalg\_lib

**Library:** ncalg.lib

**Purpose:** Definitions of important G- and GR-algebras

**Authors:** Viktor Levandovskyy, levandov@mathematik.uni-kl.de,  
Oleksandr Motsak, U@D, where U={motsak}, D={mathematik.uni-kl.de}

**Conventions:**

This library provides pre-defined important noncommutative algebras. For universal enveloping algebras of finite dimensional Lie algebras  $sl_n$ ,  $gl_n$ ,  $g_2$  etc. there are functions `makeUs1`, `makeUg1`, `makeUg2` etc. For quantized enveloping algebras  $U_q(sl_2)$  and  $U_q(sl_3)$ , there are functions `makeQs12`, `makeQs13`) and for non-standard quantum deformation of  $so_3$ , there is the function `makeQso3`.

For bigger algebras we suppress the output of the (lengthy) list of non-commutative relations and provide only the number of these relations instead.

**Procedures:**

### 7.5.10.1 makeUsl2

Procedure from library `ncalg.lib` (see Section 7.5.10 [`ncalg.lib`], page 458).

**Usage:** `makeUsl2([p])`, `p` an optional integer (field characteristic)

**Return:** ring

**Purpose:** set up the  $U(\mathfrak{sl}_2)$  in the variables  $e, f, h$  over the field of char  $p$

**Note:** activate this ring with the `setring` command

**Example:**

```
LIB "ncalg.lib";
def a=makeUsl2();
setring a;
a;
⇒ // coefficients: QQ
⇒ // number of vars : 3
⇒ // block 1 : ordering dp
⇒ // : names e f h
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // fe=ef-h
⇒ // he=eh+2e
⇒ // hf=fh-2f
```

See also: Section 7.5.10.17 [`makeUg2`], page 469; Section 7.5.10.3 [`makeUgl`], page 460; Section 7.5.10.2 [`makeUsl`], page 459.

### 7.5.10.2 makeUsl

Procedure from library `ncalg.lib` (see Section 7.5.10 [`ncalg.lib`], page 458).

**Usage:** `makeUsl(n,[p])`; `n` an integer,  $n > 1$ ; `p` an optional integer (field characteristic)

**Return:** ring

**Purpose:** set up the  $U(\mathfrak{sl}_n)$  in the variables  $(x(i), y(i), h(i) \mid i=1..n+1)$  over the field of char  $p$

**Note:** activate this ring with the `setring` command

This presentation of  $U(\mathfrak{sl}_n)$  is the standard one, i.e. positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$  and the Cartan elements are denoted by  $h(i)$ .

The variables are ordered as  $x(1), \dots, x(n), y(1), \dots, y(n), h(1), \dots, h(n)$ .

**Example:**

```
LIB "ncalg.lib";
def a=makeUsl(3);
setring a;
a;
⇒ // coefficients: QQ
⇒ // number of vars : 8
⇒ // block 1 : ordering dp
```

```

⇒ // : names x(1) x(2) x(3) y(1) y(2) y(3) h(1) h(2)
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // x(2)x(1)=x(1)*x(2)+x(3)
⇒ // y(1)x(1)=x(1)*y(1)-h(1)
⇒ // y(3)x(1)=x(1)*y(3)-y(2)
⇒ // h(1)x(1)=x(1)*h(1)+2*x(1)
⇒ // h(2)x(1)=x(1)*h(2)-x(1)
⇒ // y(2)x(2)=x(2)*y(2)-h(2)
⇒ // y(3)x(2)=x(2)*y(3)+y(1)
⇒ // h(1)x(2)=x(2)*h(1)-x(2)
⇒ // h(2)x(2)=x(2)*h(2)+2*x(2)
⇒ // y(1)x(3)=x(3)*y(1)-x(2)
⇒ // y(2)x(3)=x(3)*y(2)+x(1)
⇒ // y(3)x(3)=x(3)*y(3)-h(1)-h(2)
⇒ // h(1)x(3)=x(3)*h(1)+x(3)
⇒ // h(2)x(3)=x(3)*h(2)+x(3)
⇒ // y(2)y(1)=y(1)*y(2)-y(3)
⇒ // h(1)y(1)=y(1)*h(1)-2*y(1)
⇒ // h(2)y(1)=y(1)*h(2)+y(1)
⇒ // h(1)y(2)=y(2)*h(1)+y(2)
⇒ // h(2)y(2)=y(2)*h(2)-2*y(2)
⇒ // h(1)y(3)=y(3)*h(1)-y(3)
⇒ // h(2)y(3)=y(3)*h(2)-y(3)

```

See also: Section 7.5.10.24 [makeQsl3], page 473; Section 7.5.10.22 [makeQso3], page 472; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.3 [makeUgl], page 460; Section 7.5.10.1 [makeUsl2], page 459.

### 7.5.10.3 makeUgl

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg.lib], page 458).

**Usage:** `makeUgl(n,[p]);`  $n$  an int,  $n > 1$ ;  $p$  an optional int (field characteristic)

**Return:** ring

**Purpose:** set up the  $U(\mathfrak{gl}_n)$  in the  $(e_{ij} \ (1 < i, j < n))$  presentation (where  $e_{ij}$  corresponds to a matrix with 1 at  $i, j$  only) over the field of char  $p$

**Note:** activate this ring with the `setring` command  
the variables are ordered as  $e_{12}, e_{13}, \dots, e_{1n}, e_{21}, \dots, e_{nn}$ .

**Example:**

```

LIB "ncalg.lib";
def a=makeUgl(3);
setring a; a;
⇒ // coefficients: QQ
⇒ // number of vars : 9
⇒ // block 1 : ordering dp
⇒ // : names e_1_1 e_1_2 e_1_3 e_2_1 e_2_2 e_2_3 e_3_1 \
 e_3_2 e_3_3
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // e_1_2e_1_1=e_1_1*e_1_2-e_1_2
⇒ // e_1_3e_1_1=e_1_1*e_1_3-e_1_3

```

```

⇒ // e_2_1e_1_1=e_1_1*e_2_1+e_2_1
⇒ // e_3_1e_1_1=e_1_1*e_3_1+e_3_1
⇒ // e_2_1e_1_2=e_1_2*e_2_1-e_1_1+e_2_2
⇒ // e_2_2e_1_2=e_1_2*e_2_2-e_1_2
⇒ // e_2_3e_1_2=e_1_2*e_2_3-e_1_3
⇒ // e_3_1e_1_2=e_1_2*e_3_1+e_3_2
⇒ // e_2_1e_1_3=e_1_3*e_2_1+e_2_3
⇒ // e_3_1e_1_3=e_1_3*e_3_1-e_1_1+e_3_3
⇒ // e_3_2e_1_3=e_1_3*e_3_2-e_1_2
⇒ // e_3_3e_1_3=e_1_3*e_3_3-e_1_3
⇒ // e_2_2e_2_1=e_2_1*e_2_2+e_2_1
⇒ // e_3_2e_2_1=e_2_1*e_3_2+e_3_1
⇒ // e_2_3e_2_2=e_2_2*e_2_3-e_2_3
⇒ // e_3_2e_2_2=e_2_2*e_3_2+e_3_2
⇒ // e_3_1e_2_3=e_2_3*e_3_1-e_2_1
⇒ // e_3_2e_2_3=e_2_3*e_3_2-e_2_2+e_3_3
⇒ // e_3_3e_2_3=e_2_3*e_3_3-e_2_3
⇒ // e_3_3e_3_1=e_3_1*e_3_3+e_3_1
⇒ // e_3_3e_3_2=e_3_2*e_3_3+e_3_2

```

See also: Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459.

#### 7.5.10.4 makeUso5

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeUso5([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_5)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_5)$  is derived from the Chevalley representation of  $\mathfrak{so}_5$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```

LIB "ncalg.lib";
def ncAlgebra = makeUso5();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 10
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) Y(1) Y(2) Y(3) Y(4) H(\
1) H(2)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 28 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.12 [makeUsp1], page 465.

#### 7.5.10.5 makeUso6

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeUso6([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_6)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_6)$  is derived from the Chevalley representation of  $\mathfrak{so}_6$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUso6();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 15
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) Y(1) Y(2) Y(\
 3) Y(4) Y(5) Y(6) H(1) H(2) H(3)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 60 noncommutative relations
```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.6 makeUso7

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg.lib], page 458).

**Usage:** `makeUso7([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_7)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_7)$  is derived from the Chevalley representation of  $\mathfrak{so}_7$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUso7();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 21
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
 9) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) H(1) H(2) H(3)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 107 noncommutative relations
```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.



### 7.5.10.7 makeUso8

Procedure from library `ncalg.lib` (see Section 7.5.10 [`ncalg.lib`], page 458).

**Usage:** `makeUso8([p]);` `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_8)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_8)$  is derived from the Chevalley representation of  $\mathfrak{so}_8$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUso8();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 28
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) Y(10) Y\
(11) Y(12) H(1) H(2) H(3) H(4)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 180 noncommutative relations
```

See also: Section 7.5.10.19 [`makeUe6`], page 470; Section 7.5.10.20 [`makeUe7`], page 470; Section 7.5.10.21 [`makeUe8`], page 471; Section 7.5.10.18 [`makeUf4`], page 469; Section 7.5.10.17 [`makeUg2`], page 469; Section 7.5.10.2 [`makeUsl`], page 459; Section 7.5.10.4 [`makeUso5`], page 461; Section 7.5.10.12 [`makeUsp1`], page 465.

### 7.5.10.8 makeUso9

Procedure from library `ncalg.lib` (see Section 7.5.10 [`ncalg.lib`], page 458).

**Usage:** `makeUso9([p]);` `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_9)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_9)$  is derived from the Chevalley representation of  $\mathfrak{so}_9$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUso9();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 36
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6\
) Y(7) Y(8) Y(9) Y(10) Y(11) Y(12) Y(13) Y(14) Y(15) Y(16) H(1) H(2) H(3)\
H(4)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
```

```

setring ncAlgebra;
// ... 264 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.9 makeUso10

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeUso10([p]);` `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_{-10})$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_{-10})$  is derived from the Chevalley representation of  $\mathfrak{so}_{-10}$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```

LIB "ncalg.lib";
def ncAlgebra = makeUso10();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 45
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) Y(1)\
Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) Y(10) Y(11) Y(12) Y(13) Y(14) Y(\
15) Y(16) Y(17) Y(18) Y(19) Y(20) H(1) H(2) H(3) H(4) H(5)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 390 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.10 makeUso11

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeUso11([p]);` `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_{-11})$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_{-11})$  is derived from the Chevalley representation of  $\mathfrak{so}_{-11}$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```

LIB "ncalg.lib";
def ncAlgebra = makeUso11();
ncAlgebra;

```

```

⇒ // coefficients: QQ
⇒ // number of vars : 55
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) X(21\
) X(22) X(23) X(24) X(25) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) Y(\
10) Y(11) Y(12) Y(13) Y(14) Y(15) Y(16) Y(17) Y(18) Y(19) Y(20) Y(21) Y(2\
2) Y(23) Y(24) Y(25) H(1) H(2) H(3) H(4) H(5)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 523 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.11 makeUso12

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg\_lib], page 458).

**Usage:** `makeUso12([p]);` `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{so}_{-12})$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{so}_{-12})$  is derived from the Chevalley representation of  $\mathfrak{so}_{-12}$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```

LIB "ncalg.lib";
def ncAlgebra = makeUso12();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 66
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) X(21\
) X(22) X(23) X(24) X(25) X(26) X(27) X(28) X(29) X(30) Y(1) Y(2) Y(3) Y(\
4) Y(5) Y(6) Y(7) Y(8) Y(9) Y(10) Y(11) Y(12) Y(13) Y(14) Y(15) Y(16) Y(1\
7) Y(18) Y(19) Y(20) Y(21) Y(22) Y(23) Y(24) Y(25) Y(26) Y(27) Y(28) Y(29\
) Y(30) H(1) H(2) H(3) H(4) H(5) H(6)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 714 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.12 makeUsp1

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg\_lib], page 458).

**Usage:** `makeUsp1([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{sp}_1)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{sp}_1)$  is derived from the Chevalley representation of  $\mathfrak{sp}_1$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUsp1();
setring ncAlgebra;
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 3
⇒ // block 1 : ordering dp
⇒ // : names X(1) Y(1) H(1)
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Y(1)X(1)=X(1)*Y(1)-H(1)
⇒ // H(1)X(1)=X(1)*H(1)+2*X(1)
⇒ // H(1)Y(1)=Y(1)*H(1)-2*Y(1)
```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461.

### 7.5.10.13 makeUsp2

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeUsp2([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{sp}_2)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{sp}_2)$  is derived from the Chevalley representation of  $\mathfrak{sp}_2$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUsp2();
setring ncAlgebra;
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 10
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) Y(1) Y(2) Y(3) Y(4) H(\
 1) H(2)
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // X(2)X(1)=X(1)*X(2)+X(3)
⇒ // X(3)X(1)=X(1)*X(3)+2*X(4)
⇒ // Y(1)X(1)=X(1)*Y(1)-H(1)
⇒ // Y(3)X(1)=X(1)*Y(3)-2*Y(2)
⇒ // Y(4)X(1)=X(1)*Y(4)-Y(3)
```

```

⇒ // H(1)X(1)=X(1)*H(1)+2*X(1)
⇒ // H(2)X(1)=X(1)*H(2)-X(1)
⇒ // Y(2)X(2)=X(2)*Y(2)-H(2)
⇒ // Y(3)X(2)=X(2)*Y(3)+Y(1)
⇒ // H(1)X(2)=X(2)*H(1)-2*X(2)
⇒ // H(2)X(2)=X(2)*H(2)+2*X(2)
⇒ // Y(1)X(3)=X(3)*Y(1)-2*X(2)
⇒ // Y(2)X(3)=X(3)*Y(2)+X(1)
⇒ // Y(3)X(3)=X(3)*Y(3)-H(1)-2*H(2)
⇒ // Y(4)X(3)=X(3)*Y(4)+Y(1)
⇒ // H(2)X(3)=X(3)*H(2)+X(3)
⇒ // Y(1)X(4)=X(4)*Y(1)-X(3)
⇒ // Y(3)X(4)=X(4)*Y(3)+X(1)
⇒ // Y(4)X(4)=X(4)*Y(4)-H(1)-H(2)
⇒ // H(1)X(4)=X(4)*H(1)+2*X(4)
⇒ // Y(2)Y(1)=Y(1)*Y(2)-Y(3)
⇒ // Y(3)Y(1)=Y(1)*Y(3)-2*Y(4)
⇒ // H(1)Y(1)=Y(1)*H(1)-2*Y(1)
⇒ // H(2)Y(1)=Y(1)*H(2)+Y(1)
⇒ // H(1)Y(2)=Y(2)*H(1)+2*Y(2)
⇒ // H(2)Y(2)=Y(2)*H(2)-2*Y(2)
⇒ // H(2)Y(3)=Y(3)*H(2)-Y(3)
⇒ // H(1)Y(4)=Y(4)*H(1)-2*Y(4)

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.14 makeUsp3

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg.lib], page 458).

**Usage:** `makeUsp3([p]);` `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{sp}_3)$

**Note:** You have to activate this ring with the `'setring'` command. The presentation of  $U(\mathfrak{sp}_3)$  is derived from the Chevalley representation of  $\mathfrak{sp}_3$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```

LIB "ncalg.lib";
def ncAlgebra = makeUsp3();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 21
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) H(1) H(2) H(3)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 107 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17

[makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.15 makeUsp4

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg.lib], page 458).

**Usage:** `makeUsp4([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{sp}_4)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{sp}_4)$  is derived from the Chevalley representation of  $\mathfrak{sp}_4$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUsp4();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 36
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6\
) Y(7) Y(8) Y(9) Y(10) Y(11) Y(12) Y(13) Y(14) Y(15) Y(16) H(1) H(2) H(3)\
H(4)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 264 noncommutative relations
```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.16 makeUsp5

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg.lib], page 458).

**Usage:** `makeUsp5([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{sp}_5)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{sp}_5)$  is derived from the Chevalley representation of  $\mathfrak{sp}_5$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUsp5();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 55
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) X(21\
```

```

) X(22) X(23) X(24) X(25) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) Y(\
10) Y(11) Y(12) Y(13) Y(14) Y(15) Y(16) Y(17) Y(18) Y(19) Y(20) Y(21) Y(2\
2) Y(23) Y(24) Y(25) H(1) H(2) H(3) H(4) H(5)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 523 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.17 makeUg2

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg.lib], page 458).

**Usage:** `makeUg2([p])`, `p` an optional int (field characteristic)

**Return:** ring

**Purpose:** set up the  $U(\mathfrak{g}_2)$  in variables  $(x(i), y(i), Ha, Hb)$  for  $i=1..6$  over the field of char `p`

**Note:** activate this ring with the `setring` command  
the variables are ordered as  $x(1), \dots, x(6), y(1), \dots, y(6), Ha, Hb$ .

**Example:**

```

LIB "ncalg.lib";
def a = makeUg2();
a;
⇒ // coefficients: QQ
⇒ // number of vars : 14
⇒ // block 1 : ordering dp
⇒ // : names x(1) x(2) x(3) x(4) x(5) x(6) y(1) y(2) y(\
3) y(4) y(5) y(6) Ha Hb
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring a;
// ... 56 noncommutative relations

```

See also: Section 7.5.10.3 [makeUgl], page 460; Section 7.5.10.2 [makeUsl], page 459.

### 7.5.10.18 makeUf4

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg.lib], page 458).

**Usage:** `makeUf4([p])`; `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(\mathfrak{f}_4)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(\mathfrak{f}_4)$  is derived from the Chevalley representation of  $\mathfrak{f}_4$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```

LIB "ncalg.lib";
def ncAlgebra = makeUf4();
ncAlgebra;

```

```

⇒ // coefficients: QQ
⇒ // number of vars : 52
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) X(21\
) X(22) X(23) X(24) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) Y(10) Y(\
11) Y(12) Y(13) Y(14) Y(15) Y(16) Y(17) Y(18) Y(19) Y(20) Y(21) Y(22) Y(2\
3) Y(24) H(1) H(2) H(3) H(4)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 552 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.19 makeUe6

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg\_lib], page 458).

**Usage:** `makeUe6([p]);` `p` an optional integer (field characteristic)

**Return:** a ring, describing  $U(e_6)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(e_6)$  is derived from the Chevalley representation of  $e_6$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```

LIB "ncalg.lib";
def ncAlgebra = makeUe6();
ncAlgebra;
⇒ // coefficients: QQ
⇒ // number of vars : 78
⇒ // block 1 : ordering dp
⇒ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) X(21\
) X(22) X(23) X(24) X(25) X(26) X(27) X(28) X(29) X(30) X(31) X(32) X(33)\
X(34) X(35) X(36) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) Y(9) Y(10) Y(1\
1) Y(12) Y(13) Y(14) Y(15) Y(16) Y(17) Y(18) Y(19) Y(20) Y(21) Y(22) Y(23\
) Y(24) Y(25) Y(26) Y(27) Y(28) Y(29) Y(30) Y(31) Y(32) Y(33) Y(34) Y(35)\
Y(36) H(1) H(2) H(3) H(4) H(5) H(6)
⇒ // block 2 : ordering C
⇒ // noncommutative relations: ...
setring ncAlgebra;
// ... 1008 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.20 makeUe7

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg\_lib], page 458).



**Usage:** `makeUe7([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(e_7)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(e_7)$  is derived from the Chevalley representation of  $e_7$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUe7();
ncAlgebra;
↳ // coefficients: QQ
↳ // number of vars : 133
↳ // block 1 : ordering dp
↳ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) X(21\
) X(22) X(23) X(24) X(25) X(26) X(27) X(28) X(29) X(30) X(31) X(32) X(33)\
X(34) X(35) X(36) X(37) X(38) X(39) X(40) X(41) X(42) X(43) X(44) X(45) \
X(46) X(47) X(48) X(49) X(50) X(51) X(52) X(53) X(54) X(55) X(56) X(57) X\
(58) X(59) X(60) X(61) X(62) X(63) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8\
) Y(9) Y(10) Y(11) Y(12) Y(13) Y(14) Y(15) Y(16) Y(17) Y(18) Y(19) Y(20) \
Y(21) Y(22) Y(23) Y(24) Y(25) Y(26) Y(27) Y(28) Y(29) Y(30) Y(31) Y(32) Y\
(33) Y(34) Y(35) Y(36) Y(37) Y(38) Y(39) Y(40) Y(41) Y(42) Y(43) Y(44) Y(\
45) Y(46) Y(47) Y(48) Y(49) Y(50) Y(51) Y(52) Y(53) Y(54) Y(55) Y(56) Y(5\
7) Y(58) Y(59) Y(60) Y(61) Y(62) Y(63) H(1) H(2) H(3) H(4) H(5) H(6) H(7)
↳ // block 2 : ordering C
↳ // noncommutative relations: ...
setring ncAlgebra;
// ... 2541 noncommutative relations
```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.21 makeUe8

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeUe8([p]);`  $p$  an optional integer (field characteristic)

**Return:** a ring, describing  $U(e_8)$

**Note:** You have to activate this ring with the 'setring' command. The presentation of  $U(e_8)$  is derived from the Chevalley representation of  $e_8$ , positive resp. negative roots are denoted by  $x(i)$  resp.  $y(i)$ ; Cartan elements are denoted by  $h(i)$ .

**Example:**

```
LIB "ncalg.lib";
def ncAlgebra = makeUe8();
ncAlgebra;
↳ // coefficients: QQ
↳ // number of vars : 248
↳ // block 1 : ordering dp
↳ // : names X(1) X(2) X(3) X(4) X(5) X(6) X(7) X(8) X(\
```

```

9) X(10) X(11) X(12) X(13) X(14) X(15) X(16) X(17) X(18) X(19) X(20) X(21\
) X(22) X(23) X(24) X(25) X(26) X(27) X(28) X(29) X(30) X(31) X(32) X(33)\
 X(34) X(35) X(36) X(37) X(38) X(39) X(40) X(41) X(42) X(43) X(44) X(45) \
X(46) X(47) X(48) X(49) X(50) X(51) X(52) X(53) X(54) X(55) X(56) X(57) X\
(58) X(59) X(60) X(61) X(62) X(63) X(64) X(65) X(66) X(67) X(68) X(69) X(\
70) X(71) X(72) X(73) X(74) X(75) X(76) X(77) X(78) X(79) X(80) X(81) X(8\
2) X(83) X(84) X(85) X(86) X(87) X(88) X(89) X(90) X(91) X(92) X(93) X(94\
) X(95) X(96) X(97) X(98) X(99) X(100) X(101) X(102) X(103) X(104) X(105)\
 X(106) X(107) X(108) X(109) X(110) X(111) X(112) X(113) X(114) X(115) X(\
116) X(117) X(118) X(119) X(120) Y(1) Y(2) Y(3) Y(4) Y(5) Y(6) Y(7) Y(8) \
Y(9) Y(10) Y(11) Y(12) Y(13) Y(14) Y(15) Y(16) Y(17) Y(18) Y(19) Y(20) Y(\
21) Y(22) Y(23) Y(24) Y(25) Y(26) Y(27) Y(28) Y(29) Y(30) Y(31) Y(32) Y(3\
3) Y(34) Y(35) Y(36) Y(37) Y(38) Y(39) Y(40) Y(41) Y(42) Y(43) Y(44) Y(45\
) Y(46) Y(47) Y(48) Y(49) Y(50) Y(51) Y(52) Y(53) Y(54) Y(55) Y(56) Y(57)\
 Y(58) Y(59) Y(60) Y(61) Y(62) Y(63) Y(64) Y(65) Y(66) Y(67) Y(68) Y(69) \
Y(70) Y(71) Y(72) Y(73) Y(74) Y(75) Y(76) Y(77) Y(78) Y(79) Y(80) Y(81) Y\
(82) Y(83) Y(84) Y(85) Y(86) Y(87) Y(88) Y(89) Y(90) Y(91) Y(92) Y(93) Y(\
94) Y(95) Y(96) Y(97) Y(98) Y(99) Y(100) Y(101) Y(102) Y(103) Y(104) Y(10\
5) Y(106) Y(107) Y(108) Y(109) Y(110) Y(111) Y(112) Y(113) Y(114) Y(115) \
Y(116) Y(117) Y(118) Y(119) Y(120) H(1) H(2) H(3) H(4) H(5) H(6) H(7) H(8\
)
↳ // block 2 : ordering C
↳ // noncommutative relations: ...
setring ncAlgebra;
// ... 7752 noncommutative relations

```

See also: Section 7.5.10.19 [makeUe6], page 470; Section 7.5.10.20 [makeUe7], page 470; Section 7.5.10.21 [makeUe8], page 471; Section 7.5.10.18 [makeUf4], page 469; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.2 [makeUsl], page 459; Section 7.5.10.4 [makeUso5], page 461; Section 7.5.10.12 [makeUsp1], page 465.

### 7.5.10.22 makeQso3

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeQso3([n])`, `n` an optional int

**Purpose:** set up the  $U_q(\mathfrak{so}_3)$  in the presentation of Klimyk; if `n` is specified, the quantum parameter  $Q$  will be specialized at the  $(2n)$ -th root of unity

**Return:** ring

**Note:** activate this ring with the `setring` command

**Example:**

```

LIB "ncalg.lib";
def K = makeQso3(3);
setring K;
K;
↳ // coefficients: QQ[Q]/(Q2-Q+1)
↳ // number of vars : 3
↳ // block 1 : ordering dp
↳ // : names x y z
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // yx=(Q-1)*xy+(-Q)*z

```

```

↳ // zx=(-Q)*xz+(-Q+1)*y
↳ // zy=(Q-1)*yz+(-Q)*x

```

See also: Section 7.5.10.25 [Qso3Casimir], page 475; Section 7.5.10.23 [makeQsl2], page 473; Section 7.5.10.24 [makeQsl3], page 473; Section 7.5.10.17 [makeUg2], page 469; Section 7.5.10.3 [makeUgl], page 460; Section 7.5.10.2 [makeUsl], page 459.

### 7.5.10.23 makeQsl2

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeQsl2([n])`, `n` an optional int

**Return:** ring

**Purpose:** define the  $U_q(\mathfrak{sl}_2)$  as a factor-ring of a ring  $V_q(\mathfrak{sl}_2)$  modulo the ideal `Qideal`

**Note:** the output consists of a ring, presenting  $V_q(\mathfrak{sl}_2)$  together with the ideal called `Qideal` in this ring  
 activate this ring with the `setring` command  
 in order to create the  $U_q(\mathfrak{sl}_2)$  from the output, execute the command like `qring Usl2q = Qideal;`  
 If `n` is specified, the quantum parameter `q` will be specialized at the `n`-th root of unity

**Example:**

```

LIB "ncalg.lib";
def A = makeQsl2(3);
setring A;
Qideal;
↳ Qideal[1]=Ke*Kf-1
qring Usl2q = Qideal;
Usl2q;
↳ // coefficients: QQ[q]/(q^2+q+1)
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names E F Ke Kf
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // FE=E*F+(2/3*q+1/3)*Ke+(-2/3*q-1/3)*Kf
↳ // KeE=(-q-1)*E*Ke
↳ // KfE=(q)*E*Kf
↳ // KeF=(q)*F*Ke
↳ // KfF=(-q-1)*F*Kf
↳ // quotient ring from ideal
↳ _[1]=Ke*Kf-1

```

See also: Section 7.5.10.24 [makeQsl3], page 473; Section 7.5.10.22 [makeQso3], page 472; Section 7.5.10.2 [makeUsl], page 459.

### 7.5.10.24 makeQsl3

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `makeQsl3([n])`, `n` an optional int

**Return:** ring

**Purpose:** define the  $U_q(\mathfrak{sl}_3)$  as a factor-ring of a ring  $V_q(\mathfrak{sl}_3)$  modulo the ideal `Qideal`

**Note:** the output consists of a ring, presenting  $V_q(\mathfrak{sl}_3)$  together with the ideal called `Qideal` in this ring  
 activate this ring with the `setring` command  
 in order to create the  $U_q(\mathfrak{sl}_3)$  from the output, execute the command like `qring`  
`Usl3q = Qideal;`  
 If `n` is specified, the quantum parameter `q` will be specialized at the `n`-th root of unity

**Example:**

```
LIB "ncalg.lib";
def B = makeQsl3(5);
setring B;
qring Usl3q = Qideal;
Usl3q;
⇒ // coefficients: QQ[q]/(q^4+q^3+q^2+q+1)
⇒ // number of vars : 10
⇒ // block 1 : ordering wp
⇒ // : names f12 f13 f23 k1 k2 l1 l2 e12 e13 e23
⇒ // : weights 2 3 2 1 1 1 1 2 3 2
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // f13f12=(q^3)*f12*f13
⇒ // f23f12=(q^2)*f12*f23+(-q)*f13
⇒ // k1f12=(q^3)*f12*k1
⇒ // k2f12=(q)*f12*k2
⇒ // l1f12=(q^2)*f12*l1
⇒ // l2f12=(-q^3-q^2-q-1)*f12*l2
⇒ // e12f12=f12*e12+(1/5*q^3-3/5*q^2-2/5*q-1/5)*k1^2+(-1/5*q^3+3/5*q^2+2\
/5*q+1/5)*l1^2
⇒ // e13f12=f12*e13+(q^3+q^2+q+1)*l1^2*e23
⇒ // f23f13=(q^3)*f13*f23
⇒ // k1f13=(-q^3-q^2-q-1)*f13*k1
⇒ // k2f13=(-q^3-q^2-q-1)*f13*k2
⇒ // l1f13=(q)*f13*l1
⇒ // l2f13=(q)*f13*l2
⇒ // e12f13=f13*e12+(q)*f23*k1^2
⇒ // e13f13=f13*e13+(-1/5*q^3+3/5*q^2+2/5*q+1/5)*k1^2*k2^2+(1/5*q^3-3/5*\
q^2-2/5*q-1/5)*l1^2*l2^2
⇒ // e23f13=f13*e23+(q^3+q^2+q+1)*f12*l2^2
⇒ // k1f23=(q)*f23*k1
⇒ // k2f23=(q^3)*f23*k2
⇒ // l1f23=(-q^3-q^2-q-1)*f23*l1
⇒ // l2f23=(q^2)*f23*l2
⇒ // e13f23=f23*e13+(q)*k2^2*e12
⇒ // e23f23=f23*e23+(1/5*q^3-3/5*q^2-2/5*q-1/5)*k2^2+(-1/5*q^3+3/5*q^2+2\
/5*q+1/5)*l2^2
⇒ // e12k1=(q^3)*k1*e12
⇒ // e13k1=(-q^3-q^2-q-1)*k1*e13
⇒ // e23k1=(q)*k1*e23
⇒ // e12k2=(q)*k2*e12
⇒ // e13k2=(-q^3-q^2-q-1)*k2*e13
⇒ // e23k2=(q^3)*k2*e23
⇒ // e12l1=(q^2)*l1*e12
⇒ // e13l1=(q)*l1*e13
```

```

⇒ // e23l1=(-q^3-q^2-q-1)*l1*e23
⇒ // e12l2=(-q^3-q^2-q-1)*l2*e12
⇒ // e13l2=(q)*l2*e13
⇒ // e23l2=(q^2)*l2*e23
⇒ // e13e12=(q^3)*e12*e13
⇒ // e23e12=(q^2)*e12*e23+(-q)*e13
⇒ // e23e13=(q^3)*e13*e23
⇒ // quotient ring from ideal
⇒ _[1]=k2*l2-1
⇒ _[2]=k1*l1-1

```

See also: Section 7.5.10.23 [makeQsl2], page 473; Section 7.5.10.22 [makeQso3], page 472; Section 7.5.10.2 [makeUsl], page 459.

### 7.5.10.25 Qso3Casimir

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `Qso3Casimir(n [,m])`,  $n$  an integer,  $m$  an optional integer

**Return:** list (of polynomials)

**Purpose:** compute the Casimir (central) elements of  $U_q(\mathfrak{so}_3)$  for the quantum parameter specialized at the  $n$ -th root of unity; if  $m \neq 0$  is given, polynomials will be normalized

**Assume:** the basering must be  $U_q(\mathfrak{so}_3)$

**Example:**

```

LIB "ncalg.lib";
def R = makeQso3(5);
setring R;
list C = Qso3Casimir(5);
C;
⇒ [1]:
⇒ 1/5*x5+(1/5Q3-1/5Q2+2/5)*x3+(1/5Q3-1/5Q2+1/5)*x
⇒ [2]:
⇒ 1/5*y5+(1/5Q3-1/5Q2+2/5)*y3+(1/5Q3-1/5Q2+1/5)*y
⇒ [3]:
⇒ 1/5*z5+(1/5Q3-1/5Q2+2/5)*z3+(1/5Q3-1/5Q2+1/5)*z
list Cnorm = Qso3Casimir(5,1);
Cnorm;
⇒ [1]:
⇒ x5+(Q3-Q2+2)*x3+(Q3-Q2+1)*x
⇒ [2]:
⇒ y5+(Q3-Q2+2)*y3+(Q3-Q2+1)*y
⇒ [3]:
⇒ z5+(Q3-Q2+2)*z3+(Q3-Q2+1)*z

```

See also: Section 7.5.10.22 [makeQso3], page 472.

### 7.5.10.26 GKZsystem

Procedure from library `ncalg.lib` (see Section 7.5.10 [ncalg-lib], page 458).

**Usage:** `GKZsystem(A, sord, alg, [,v])`;  $A$  intmat, `sord`, `alg` string,  $v$  intvec

**Return:** ring

- Purpose:** define a ring (Weyl algebra) and create a Gelfand-Kapranov-Zelevinsky (GKZ) system of equations in a ring from the following data:  
 A is an intmat, defining the system,  
 sord is a string with desired term ordering,  
 alg is a string, saying which algorithm to use (exactly like in toric\_lib),  
 v is an optional intvec.  
 In addition, the ideal called GKZid containing actual equations is calculated and exported to the ring.
- Note:** activate the output ring with the `setring` command. This procedure is elaborated by Oleksandr Iena
- Assume:** This procedure uses `toric_lib` and therefore inherits its input requirements:  
 possible values for input variable `alg` are: "ect", "pt", "blr", "hs", "du".  
 As for the term ordering, it should be a string `sord` in Singular format like "lp", "dp", etc.  
 Please consult the `toric_lib` for allowed orderings and more details.

**Example:**

```
LIB "ncalg.lib";
// example 3.1.4 from the [SST] without the vector w
intmat A[2][4]=3,2,1,0,0,1,2,3;
print(A);
↳ 3 2 1 0
↳ 0 1 2 3
def D1 = GKZsystem(A,"lp","ect");
setring D1;
D1;
↳ // coefficients: QQ(b(1), b(2))
↳ // number of vars : 8
↳ // block 1 : ordering a
↳ // : names x(1) x(2) x(3) x(4)
↳ // : weights 0 0 0 0
↳ // block 2 : ordering lp
↳ // : names x(1) x(2) x(3) x(4) d(1) d(2) d(3) d(4)
↳ // block 3 : ordering C
↳ // noncommutative relations:
↳ // d(1)x(1)=x(1)*d(1)+1
↳ // d(2)x(2)=x(2)*d(2)+1
↳ // d(3)x(3)=x(3)*d(3)+1
↳ // d(4)x(4)=x(4)*d(4)+1
print(GKZid);
↳ 3*x(1)*d(1)+2*x(2)*d(2)+x(3)*d(3)+(-b(1)),
↳ x(2)*d(2)+2*x(3)*d(3)+3*x(4)*d(4)+(-b(2)),
↳ d(2)*d(4)-d(3)^2,
↳ d(1)*d(4)-d(2)*d(3),
↳ d(1)*d(3)-d(2)^2
// now, consider A with the vector w=1,1,1,1
intvec v=1,1,1,1;
def D2 = GKZsystem(A,"lp","blr",v);
setring D2;
print(GKZid);
↳ 3*x(1)*d(1)+2*x(2)*d(2)+x(3)*d(3)+(-b(1)),
↳ x(2)*d(2)+2*x(3)*d(3)+3*x(4)*d(4)+(-b(2)),
```

$\mapsto d(2)*d(4)-d(3)^2,$   
 $\mapsto d(1)*d(4)-d(2)*d(3),$   
 $\mapsto d(1)*d(3)-d(2)^2$

See also: Section D.4.35 [toric\_lib], page 834.

### 7.5.11 ncdecomp\_lib

**Library:** ncdecomp.lib

**Purpose:** Decomposition of a module into its central characters

**Authors:** Viktor Levandovskyy, levandov@mathematik.uni-kl.de.

**Overview:**

This library presents algorithms for the central character decomposition of a module, i.e. a decomposition into generalized weight modules with respect to the center. Based on ideas of O. Khomenko and V. Levandovskyy (see the article [L2] in the References for details).

**Procedures:**

#### 7.5.11.1 CentralQuot

Procedure from library ncdecomp.lib (see Section 7.5.11 [ncdecomp\_lib], page 477).

**Usage:** CentralQuot(M, G), M a module, G an ideal

**Assume:** G is an ideal in the center of the base ring

**Return:** module

**Purpose:** compute the central quotient M:G

**Theory:** for an ideal G of the center of an algebra and a submodule M of  $A^n$ , the central quotient of M by G is defined to be  $M:G := \{ v \text{ in } A^n \mid z*v \text{ in } M, \text{ for all } z \text{ in } G \}$ .

**Note:** the output module is not necessarily given in a Groebner basis

**Example:**

```

LIB "ncdecomp.lib";
option(returnSB);
def a = makeUs12();
setring a;
ideal I = e3,f3,h3-4*h;
I = std(I);
poly C=4*e*f+h^2-2*h; // C in Z(U(sl2)), the central element
ideal G = (C-8)*(C-24); // G normal factor in Z(U(sl2)) as an ideal in the center
ideal R = CentralQuot(I,G); // same as I:G
R;
 $\mapsto R[1]=h$
 $\mapsto R[2]=f$
 $\mapsto R[3]=e$

```

See also: Section 7.5.11.3 [CenCharDec], page 478; Section 7.5.11.2 [CentralSaturation], page 478.

### 7.5.11.2 CentralSaturation

Procedure from library `ncdecomp.lib` (see Section 7.5.11 [`ncdecomp.lib`], page 477).

**Usage:** `CentralSaturation(M, T)`, for a module  $M$  and an ideal  $T$

**Assume:**  $T$  is an ideal in the center of the base ring

**Return:** module

**Purpose:** compute the central saturation of  $M$  by  $T$ , that is  $M:T^{\infty}$ , by repetitive application of `CentralQuot`

**Note:** the output module is not necessarily a Groebner basis

**Example:**

```
LIB "ncdecomp.lib";
option(returnSB);
def a = makeUs12();
setring a;
ideal I = e3,f3,h3-4*h;
I = std(I);
poly C=4*e*f+h^2-2*h;
ideal G = C*(C-8);
ideal R = CentralSaturation(I,G);
R=std(R);
vdim(R);
↪ 5
R;
↪ R[1]=h
↪ R[2]=ef-6
↪ R[3]=f3
↪ R[4]=e3
```

See also: Section 7.5.11.3 [`CenCharDec`], page 478; Section 7.5.11.1 [`CentralQuot`], page 477.

### 7.5.11.3 CenCharDec

Procedure from library `ncdecomp.lib` (see Section 7.5.11 [`ncdecomp.lib`], page 477).

**Usage:** `CenCharDec(I, C)`;  $I$  a module,  $C$  an ideal

**Assume:**  $C$  consists of generators of the center of the base ring

**Return:** a list  $L$ , where each entry consists of three records (if a finite decomposition exists)  
 $L[*][1]$  ('ideal' type), the central character as a maximal ideal in the center,  
 $L[*][2]$  ('module' type), the Groebner basis of the weight module, corresponding to the character in  $L[*][1]$ ,  
 $L[*][3]$  ('int' type) is the vector space dimension of the weight module (-1 in case of infinite dimension);

**Purpose:** compute a finite decomposition of  $C$  into central characters or determine that there is no finite decomposition

**Note:** actual decomposition is the sum of  $L[i][2]$  above;  
 some modules have no finite decomposition (in such case one gets warning message)  
 The function `central` in `central.lib` may be used to obtain  $C$ , when needed.

**Example:**



```

LIB "ncdecomp.lib";
printlevel=0;
option(returnSB);
def a = makeUs12(); // U(sl_2) in characteristic 0
setring a;
ideal I = e3,f3,h3-4*h;
I = twostd(I); // two-sided ideal generated by I
vdim(I); // it is finite-dimensional
⇒ 10
ideal Cn = 4*e*f+h^2-2*h; // the only central element
list T = CenCharDec(I,Cn);
T;
⇒ [1]:
⇒ [1]:
⇒ _[1]=4ef+h2-2h-8
⇒ [2]:
⇒ _[1]=h
⇒ _[2]=f
⇒ _[3]=e
⇒ [3]:
⇒ 1
⇒ [2]:
⇒ [1]:
⇒ _[1]=4ef+h2-2h
⇒ [2]:
⇒ _[1]=4ef+h2-2h-8
⇒ _[2]=h3-4h
⇒ _[3]=fh2-2fh
⇒ _[4]=eh2+2eh
⇒ _[5]=f2h-2f2
⇒ _[6]=e2h+2e2
⇒ _[7]=f3
⇒ _[8]=e3
⇒ [3]:
⇒ 9
// consider another example
ideal J = e*f*h;
CenCharDec(J,Cn);
⇒ There is no finite decomposition
⇒ 0

```

See also: Section 7.5.11.1 [CentralQuot], page 477; Section 7.5.11.2 [CentralSaturation], page 478.

#### 7.5.11.4 IntersectWithSub

Procedure from library `ncdecomp.lib` (see Section 7.5.11 [ncdecomp.lib], page 477).

**Usage:** `IntersectWithSub(M,Z)`, `M` an ideal, `Z` an ideal

**Assume:** `Z` consists of pairwise commutative elements

**Return:** ideal of two-sided generators, not a Groebner basis

**Purpose:** computes the intersection of `M` with the subalgebra, generated by `Z`

**Note:** usually `Z` consists of generators of the center  
The function `central` from `central.lib` may be used to obtain the center `Z`, if needed.

**Example:**

```

LIB "ncdecomp.lib";
ring R=(0,a),(e,f,h),Dp;
matrix @d[3][3];
@d[1,2]=-h; @d[1,3]=2e; @d[2,3]=-2f;
def r = nc_algebra(1,@d); setring r; // parametric U(sl_2)
ideal I = e,h-a;
ideal C;
C[1] = h^2-2*h+4*e*f; // the center of U(sl_2)
ideal X = IntersectWithSub(I,C);
X;
↪ X[1]=4*ef+h^2-2*h+(-a^2-2a)
ideal G = e*f, h; // the biggest comm. subalgebra of U(sl_2)
ideal Y = IntersectWithSub(I,G);
Y;
↪ Y[1]=h+(-a)
↪ Y[2]=ef+(-a)

```

**7.5.12 ncfactor\_lib****Library:** ncfactor.lib**Purpose:** Tools for factorization in some noncommutative algebras**Authors:** Albert Heinle, aheinle at uwaterloo.ca  
Viktor Levandovskyy, levandov at math.rwth-aachen.de

**Overview:** In this library, new methods for factorization on polynomials are implemented for several types of algebras, namely

- finitely presented (and also free) associative algebras (Letterplace subsystem)
- G-algebras (Plural subsystem), including (q)-Weyl and (q)-shift algebras in  $2n$  variables

The determination of the best algorithm available for users input is done automatically in the procedure ncfactor().

More detailed description of the algorithms and related publications can be found at [@url{https://cs.uwaterloo.ca/~aheinle/}](https://cs.uwaterloo.ca/~aheinle/).

**Procedures:****7.5.12.1 ncfactor**

Procedure from library ncfactor.lib (see Section 7.5.12 [ncfactor\_lib], page 480).

**Usage:** ncfactor(h); h is a polynomial in a non-commutative polynomial algebra over a field k.**Return:** list(list)**Purpose:** Compute all factorizations of h.**Theory:** Implements an ansatz-driven factorization method as outlined by Bell, Heinle and Levandovskyy in "On Noncommutative Finite Factorization Domains".

**Assume:**

- k is a ring, such that factorize can factor any univariate and multivariate commutative polynomial over k.
- There exists at least one variable in the ring.

**Note:** - works for both PLURAL and LETTERPLACE subsystems  
 - Every entry of the output list is a list with factors for one possible factorization. The first factor is always a constant (1, if no nontrivial constant could be excluded).

**Example:**

```
LIB "ncfactor.lib";
// first, an example with PLURAL
def R = makeUsl2();
setring(R);
poly p = e^3*f+e^2*f^2-e^3+e^2*f+2*e*f^2-3*e^2*h-2*e*f*h-8*e^2
+e*f+f^2-4*e*h-2*f*h-7*e+f-h;
ncfactor(p);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ e+1
↳ [3]:
↳ ef-e+f-2h-3
↳ [4]:
↳ e+f
↳ [2]:
↳ [1]:
↳ 1
↳ [2]:
↳ e2f+ef2-e2+f2-2eh-3e-f-2h
↳ [3]:
↳ e+1
kill R;
// an example with LETTERPLACE
LIB "freegb.lib";
↳ // ** redefining tstfreegb (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
0
↳ // ** redefining tstfreegb (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
0
↳ // ** redefining setLetterplaceAttributes (LIB "freegb.lib"); ./examples/\
ncfactor.sing:10
↳ // ** redefining setLetterplaceAttributes (LIB "freegb.lib"); ./examples/\
ncfactor.sing:10
↳ // ** redefining lst2str (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining mod2str (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining vct2str (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining isVar (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining letplaceGBasis (LIB "freegb.lib"); ./examples/ncfactor.s\
ing:10
↳ // ** redefining letplaceGBasis (LIB "freegb.lib"); ./examples/ncfactor.s\
ing:10
↳ // ** redefining lieBracket (LIB "freegb.lib"); ./examples/ncfactor.sing:\
10
↳ // ** redefining lieBracket (LIB "freegb.lib"); ./examples/ncfactor.sing:\
10
↳ // ** redefining lpPrint (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining lpPrint (LIB "freegb.lib"); ./examples/ncfactor.sing:10
```

```

⇒ // ** redefining freeGBasis (LIB "freegb.lib"); ./examples/ncfactor.sing:\
 10
⇒ // ** redefining freeGBasis (LIB "freegb.lib"); ./examples/ncfactor.sing:\
 10
⇒ // ** redefining crs (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining polylen (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpDegBound (LIB "freegb.lib"); ./examples/ncfactor.sing:\
 10
⇒ // ** redefining lpDegBound (LIB "freegb.lib"); ./examples/ncfactor.sing:\
 10
⇒ // ** redefining lpVarBlockSize (LIB "freegb.lib"); ./examples/ncfactor.s\
 ing:10
⇒ // ** redefining lpVarBlockSize (LIB "freegb.lib"); ./examples/ncfactor.s\
 ing:10
⇒ // ** redefining isFreeAlgebra (LIB "freegb.lib"); ./examples/ncfactor.si\
 ng:10
⇒ // ** redefining isFreeAlgebra (LIB "freegb.lib"); ./examples/ncfactor.si\
 ng:10
⇒ // ** redefining lpNcgenCount (LIB "freegb.lib"); ./examples/ncfactor.sin\
 g:10
⇒ // ** redefining lpNcgenCount (LIB "freegb.lib"); ./examples/ncfactor.sin\
 g:10
⇒ // ** redefining makeLetterplaceRing (LIB "freegb.lib"); ./examples/ncfac\
 tor.sing:10
⇒ // ** redefining makeLetterplaceRing (LIB "freegb.lib"); ./examples/ncfac\
 tor.sing:10
⇒ // ** redefining makeLetterplaceRing1 (LIB "freegb.lib"); ./examples/ncfa\
 ctor.sing:10
⇒ // ** redefining makeLetterplaceRing2 (LIB "freegb.lib"); ./examples/ncfa\
 ctor.sing:10
⇒ // ** redefining makeLetterplaceRing4 (LIB "freegb.lib"); ./examples/ncfa\
 ctor.sing:10
⇒ // ** redefining makeLetterplaceRing3 (LIB "freegb.lib"); ./examples/ncfa\
 ctor.sing:10
⇒ // ** redefining freegbold (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
 0
⇒ // ** redefining stringpoly2lplace (LIB "freegb.lib"); ./examples/ncfacto\
 r.sing:10
⇒ // ** redefining addplaces (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
 0
⇒ // ** redefining sent2lplace (LIB "freegb.lib"); ./examples/ncfactor.sing\
 :10
⇒ // ** redefining testnumber (LIB "freegb.lib"); ./examples/ncfactor.sing:\
 10
⇒ // ** redefining str2lplace (LIB "freegb.lib"); ./examples/ncfactor.sing:\
 10
⇒ // ** redefining strpower2rep (LIB "freegb.lib"); ./examples/ncfactor.sin\
 g:10
⇒ // ** redefining shiftPoly (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
 0
⇒ // ** redefining lastBlock (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
 0
⇒ // ** redefining test_shift (LIB "freegb.lib"); ./examples/ncfactor.sing:\

```

```

10
⇒ // ** redefining lp2lstr (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lp2lstr (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining strList2poly (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining file2lplace (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpPower (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpNF (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpNF (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpDivision (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpDivision (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpGBPres2Poly (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpGBPres2Poly (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining getExpVecs (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining delSupZero (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining delSupZeroList (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining makeDVec (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining makeDVecL (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining makeDVecI (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining dShiftDiv (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpNormalForm1 (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpNormalForm2 (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining isOrderingShiftInvariant (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining isOrderingShiftInvariant (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpMonomialsWithHoles (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining getlpCoeffs (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining lpReduce (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining entryViolation (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining checkAssumptionsLPIV (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining checkAssumptions (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining checkLPRing (LIB "freegb.lib"); ./examples/ncfactor.sing:10
⇒ // ** redefining checkAssumptionIdeal (LIB "freegb.lib"); ./examples/ncfactor.sing:10

```

```

ctor.sing:10
↳ // ** redefining checkAssumptionPoly (LIB "freegb.lib"); ./examples/ncfac\
tor.sing:10
↳ // ** redefining isContainedInVp (LIB "freegb.lib"); ./examples/ncfactor.\
sing:10
↳ // ** redefining extractLinearPart (LIB "freegb.lib"); ./examples/ncfacto\
r.sing:10
↳ // ** redefining isLinearVector (LIB "freegb.lib"); ./examples/ncfactor.s\
ing:10
↳ // ** redefining lpAssumeViolation (LIB "freegb.lib"); ./examples/ncfacto\
r.sing:10
↳ // ** redefining skip0 (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining ivL2lpI (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining ivL2lpI (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining iv2lp (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining iv2lp (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining iv2lpList (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
0
↳ // ** redefining iv2lpList (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
0
↳ // ** redefining iv2lpMat (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining iv2lpMat (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining lpId2ivLi (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
0
↳ // ** redefining lpId2ivLi (LIB "freegb.lib"); ./examples/ncfactor.sing:1\
0
↳ // ** redefining lp2iv (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining lp2iv (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining lp2ivId (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining lp2ivId (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining testLift (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining testLift (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining testSyz (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining testSyz (LIB "freegb.lib"); ./examples/ncfactor.sing:10
↳ // ** redefining mod_init (LIB "freegb.lib"); ./examples/ncfactor.sing:10
ring r = 0, (x,y), Dp;
def R = freeAlgebra(r,5); setring(R);
poly p = x*y*x - x;
ncfactor(p);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ x*y-1
↳ [3]:
↳ x
↳ [2]:
↳ [1]:
↳ 1
↳ [2]:
↳ x
↳ [3]:
↳ y*x-1

```

See also: Section 7.5.12.5 [facSubWeyl], page 487; Section 7.5.12.2 [facWeyl], page 485; Section 7.5.12.4 [testNCfac], page 486.

### 7.5.12.2 facWeyl

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Usage:** `facWeyl(h)`;  $h$  a polynomial in the  $n$ th Weyl algebra

**Return:** list

**Purpose:** compute all factorizations of a polynomial in the first Weyl algebra

**Theory:** Implements the new algorithm by A. Heinle and V. Levandovskyy, see the thesis of A. Heinle

**Assume:** `basering` is the  $n$ th Weyl algebra, where  $n$  in  $\mathbb{N}$ .

**Note:** Every entry of the output list is a list with factors for one possible factorization. The first factor is always a constant (1, if no nontrivial constant could be excluded).

**Example:**

```
LIB "ncfactor.lib";
ring R = 0, (x1,x2,d1,d2), dp;
matrix C[4][4] = 1,1,1,1,
1,1,1,1,
1,1,1,1,
1,1,1,1;
matrix D[4][4] = 0,0,1,0,
0,0,0,1,
-1,0,0,0,
0,-1,0,0;
def r = nc_algebra(C,D);
setring(r);
poly h = (d1+1)^2*(d1 + x1*d2);
facWeyl(h);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ d1+1
↳ [3]:
↳ d1+1
↳ [4]:
↳ x1*d2+d1
↳ [2]:
↳ [1]:
↳ 1
↳ [2]:
↳ x1*d1*d2+d1^2+x1*d2+d1+2*d2
↳ [3]:
↳ d1+1
```

See also: Section 7.5.12.7 [facFirstShift], page 489; Section 7.5.12.3 [facFirstWeyl], page 486; Section 7.5.12.5 [facSubWeyl], page 487; Section 7.5.12.4 [testNCfac], page 486.

### 7.5.12.3 facFirstWeyl

Procedure from library `ncfactor.lib` (see Section 7.5.12 [`ncfactor.lib`], page 480).

**Usage:** `facFirstWeyl(h)`;  $h$  a polynomial in the first Weyl algebra

**Return:** list

**Purpose:** compute all factorizations of a polynomial in the first Weyl algebra

**Theory:** This function is a wrapper for `facWeyl`. It exists to make this library downward-compatible with older versions.

**Assume:** `basing` is the first Weyl algebra

**Note:** Every entry of the output list is a list with factors for one possible factorization. The first factor is always a constant (1, if no nontrivial constant could be excluded).

**Example:**

```
LIB "ncfactor.lib";
ring R = 0,(x,y),dp;
def r = nc_algebra(1,1);
setring(r);
poly h = (x^2*y^2+x)*(x+1);
facFirstWeyl(h);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ x
↳ [3]:
↳ xy2+1
↳ [4]:
↳ x+1
```

See also: Section 7.5.12.6 [`facShift`], page 488; Section 7.5.12.5 [`facSubWeyl`], page 487; Section 7.5.12.4 [`testNCfac`], page 486.

### 7.5.12.4 testNCfac

Procedure from library `ncfactor.lib` (see Section 7.5.12 [`ncfactor.lib`], page 480).

**Usage:** `testNCfac(l[,p,b])`;  $l$  is a list,  $p$  is an optional poly,  $b$  is 1 or 0

**Return:** Case 1: No optional argument. In this case the output is 1, if the entries in the given list represent the same polynomial or 0 otherwise.

Case 2: One optional argument  $p$  is given. In this case it returns 1, if all the entries in  $l$  are factorizations of  $p$ , otherwise 0. Case 3: Second optional  $b$  is given. In this case a list is returned containing the difference between the product of each entry in  $l$  and  $p$ .

**Assume:** `basing` is the first Weyl algebra, the entries of  $l$  are polynomials

**Purpose:** Checks whether a list of factorizations contains factorizations of the same element in the first Weyl algebra

**Theory:** `testNCfac` multiplies out each factorization and checks whether each factorization was a factorization of the same element.

- if there is only a list given, the output will be 0, if it does not contain factorizations of the same element. Otherwise the output will be 1.



- if there is a polynomial in the second argument, then the procedure checks whether the given list contains factorizations of this polynomial. If it does, then the output depends on the third argument. If it is not given, the procedure will check whether the factorizations in the list  $l$  are associated to this polynomial and return either 1 or 0, respectively. If the third argument is given, the output will be a list with the length of the given one and in each entry is the product of one entry in  $l$  subtracted by the polynomial.

**Example:**

```
LIB "ncfactor.lib";
ring r = 0,(x,y),dp;
def R = nc_algebra(1,1);
setring R;
poly h = (x^2*y^2+1)*(x^2);
def t1 = facFirstWeyl(h);
//first a correct list
testNCfac(t1);
⇒ 1
//now a correct list with the factorized polynomial
testNCfac(t1,h);
⇒ 1
//now we put in an incorrect list without a polynomial
t1[3][3] = y;
testNCfac(t1);
⇒ 0
// take h as additional input
testNCfac(t1,h);
⇒ 0
// take h as additional input and output list of differences
testNCfac(t1,h,1);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ 0
⇒ [3]:
⇒ -x4y2+x3y3-4x3y+3x2y2-3x2+xy+1
```

See also: Section 7.5.12.7 [facFirstShift], page 489; Section 7.5.12.3 [facFirstWeyl], page 486; Section 7.5.12.5 [facSubWeyl], page 487.

**7.5.12.5 facSubWeyl**

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Usage:** `facSubWeyl(h,[x_1,...,x_n,d_1,...,d_n])`;  $h$  is a polynomial,  $x_i$ ,  $d_i$  are variables in the basering for  $i$  in  $\{1, \dots, n\}$

**Return:** `list(list)`

**Assume:**  $x_i$ ,  $d_i$  are variables of a basering with  $d_i * x_i = x_i * d_{i+1}$  for  $i$  in  $\{1, \dots, n\}$ .

That is, they generate the copy of the first Weyl algebra in a basering.

Moreover,  $h$  is a polynomial in the  $x_i, d_i$  only.

If the list of variables is omitted, this function will try to figure out itself if  $h$  is in a subalgebra that resembles the Weyl algebra.

This function produces an error if the conditions on the variables do not line up or if the

variables contained in  $h$  do not belong to a subalgebra of the basering that resembles the Weyl algebra.

**Purpose:** compute factorizations of the polynomial, depending on  $x_i$  and  $d_i$ .

**Example:**

```
LIB "ncfactor.lib";
ring r = 0, (x,y,z), dp;
matrix D[3][3]; D[1,3]=-1;
def R = nc_algebra(1,D); // x,z generate Weyl subalgebra
setring R;
poly h = (x^2*z^2+x)*x;
list fact1 = facSubWeyl(h,x,z);
// compare with facFirstWeyl:
ring s = 0, (z,x), dp;
def S = nc_algebra(1,1); setring S;
poly h = (x^2*z^2+x)*x;
list fact2 = facFirstWeyl(h);
map F = R,x,0,z;
list fact1 = F(fact1); // it is identical to list fact2
testNCfac(fact1); // check the correctness again
↳ 1
```

See also: Section 7.5.12.7 [facFirstShift], page 489; Section 7.5.12.3 [facFirstWeyl], page 486; Section 7.5.12.4 [testNCfac], page 486.

### 7.5.12.6 facShift

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Usage:** `facShift(h)`;  $h$  a polynomial in the  $n$ 'th shift algebra

**Return:** list

**Purpose:** compute all factorizations of a polynomial in the  $n$ th shift algebra

**Theory:** Currently, we do not have a specialized algorithm for the shift algebra in this library that takes advantage of the graded structure, hence this function is mapping to the general factorization algorithm for  $G$ -Algebras

**Note:** Every entry of the output list is a list with factors for one possible factorization.

**Example:**

```
LIB "ncfactor.lib";
ring R = 0, (x1,x2,s1,s2), dp;
matrix C[4][4] = 1,1,1,1,
1,1,1,1,
1,1,1,1,
1,1,1,1;
matrix D[4][4] = 0,0,s1,0,
0,0,0,s2,
-s1,0,0,0,
0,-s2,0,0;
def r = nc_algebra(C,D);
setring(r);
poly h = x1*(x1+1)*s1^2-2*x1*(x1+100)*s1+(x1+99)*(x1+100);
facShift(h);
```

```

↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ x1*s1-x1+s1-100
↳ [3]:
↳ x1*s1-x1-s1-99
↳ [2]:
↳ [1]:
↳ 1
↳ [2]:
↳ x1*s1-x1-100
↳ [3]:
↳ x1*s1-x1-99
↳ [3]:
↳ [1]:
↳ 1
↳ [2]:
↳ x1*s1-x1-99
↳ [3]:
↳ x1*s1-x1-100

```

See also: Section 7.5.12.3 [facFirstWeyl], page 486; Section 7.5.12.5 [facSubWeyl], page 487; Section 7.5.12.4 [testNCfac], page 486.

### 7.5.12.7 facFirstShift

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Usage:** `facFirstShift(h)`;  $h$  a polynomial in the first shift algebra

**Return:** list

**Purpose:** compute all factorizations of a polynomial in the first shift algebra

**Theory:** This function is a wrapper for `facShift`. It exists to make this library downward-compatible with older versions.

**Assume:** `basing` is the first shift algebra

**Note:** Every entry of the output list is a list with factors for one possible factorization.

**Example:**

```

LIB "ncfactor.lib";
ring R = 0,(x,s),dp;
def r = nc_algebra(1,s);
setring(r);
poly h = (s^2*x+x)*s;
facFirstShift(h);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ s
↳ [3]:
↳ s2+1
↳ [4]:

```

```

↳ x-1
↳ [2] :
↳ [1] :
↳ 1
↳ [2] :
↳ s2+1
↳ [3] :
↳ s
↳ [4] :
↳ x-1
↳ [3] :
↳ [1] :
↳ 1
↳ [2] :
↳ s2+1
↳ [3] :
↳ x
↳ [4] :
↳ s

```

See also: Section 7.5.12.3 [facFirstWeyl], page 486; Section 7.5.12.5 [facSubWeyl], page 487; Section 7.5.12.4 [testNCfac], page 486.

### 7.5.12.8 homogfacNthWeyl

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Usage:** `homogfacNthWeyl(h)`; `h` is a homogeneous polynomial in the `n`th Weyl algebra with respect to the `-1,1`-grading

**Return:** list

**Purpose:** Computes a factorization of a homogeneous polynomial `h` with respect to the `ZZ`-grading on the `n`-th Weyl algebra.

**Theory:** `homogfacFirstWeyl` returns a list with a factorization of the given, `[-1,1]`-homogeneous polynomial. For every `i` in `1..n`: If the degree of the polynomial in `[d.i,x.i]` is `k` with `k` positive, the last `k` entries in the output list are the second variable. If `k` is positive, the last `k` entries will be `x.i`. The other entries will be irreducible polynomials of degree zero or 1 resp. `-1`. resp. other variables

**General assumptions:**

- The basering is the `n`th Weyl algebra and has the form, that the first `n` variables represent `x1, ..., xn`, and the second `n` variables do represent the `d1, ..., dn`.

### 7.5.12.9 homogfacNthQWeyl

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Usage:** `homogfacNthQWeyl(h)`; `h` is a homogeneous polynomial in the `n`'th `q`-Weyl algebra with respect to the weight vector `@ [-1,...,-1,1,...,1]`.  
`@ \_ \_ / \_ \_ / @ \ / \ / @ n/2 n/2`

**Return:** list

**Purpose:** Computes a factorization of a homogeneous polynomial `h` in the `n`'th `q`-Weyl algebra

**Theory:** `homogfacNthQWeyl` returns a list with a factorization of the given,  $[-1,1]$ -homogeneous polynomial. For every  $i$  in  $1..n$ : If the degree of the polynomial in  $[d_i, x_i]$  is  $k$  with  $k$  positive, the last entries in the output list are the second variable. If  $k$  is positive, the last  $k$  entries will be  $x_i$ . The other entries will be irreducible polynomials of degree zero or 1 resp. -1. resp. other variables

**General assumptions:**

- The basering is the  $n$ th Weyl algebra and has the form, that the first  $n$  variables represent  $x_1, \dots, x_n$ , and the second  $n$  variables do represent the  $d_1, \dots, d_n$ .
- We have  $n$  parameters  $q_1, \dots, q_n$  given.

**Example:**

```
LIB "ncfactor.lib";
ring R = (0,q1,q2,q3),(x1,x2,x3,d1,d2,d3),dp;
matrix C[6][6] = 1,1,1,q1,1,1,
1,1,1,1,q2,1,
1,1,1,1,1,q3,
1,1,1,1,1,1,
1,1,1,1,1,1,
1,1,1,1,1,1;
matrix D[6][6] = 0,0,0,1,0,0,
0,0,0,0,1,0,
0,0,0,0,0,1,
-1,0,0,0,0,0,
0,-1,0,0,0,0,
0,0,-1,0,0,0;
def r = nc_algebra(C,D);
setring(r);
poly h =x1*x2^2*x3^3*d1*d2^2+x2*x3^3*d2;
homogfacNthQWeyl(h);
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ d2
↳ [4]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ x3
```

See also: Section 7.5.12.10 [`homogfacFirstQWeyl`], page 491; Section 7.5.12.12 [`homogfacFirstQWeyl_all`], page 510; Section 7.5.12.11 [`homogfacNthQWeyl_all`], page 492.

### 7.5.12.10 `homogfacFirstQWeyl`

Procedure from library `ncfactor.lib` (see Section 7.5.12 [`ncfactor.lib`], page 480).

**Usage:** `homogfacFirstQWeyl(h)`;  $h$  is a homogeneous polynomial in the first  $q$ -Weyl algebra with respect to the weight vector  $[-1,1]$

**Return:** list

**Purpose:** Computes a factorization of a homogeneous polynomial  $h$  with respect to the weight vector  $[-1,1]$  in the first  $q$ -Weyl algebra

**Theory:** This function is a wrapper for `homogfacNthQWeyl`. It exists to make this library downward-compatible with older versions.

**Example:**

```
LIB "ncfactor.lib";
ring R = (0,q),(x,d),dp;
def r = nc_algebra (q,1);
setring(r);
poly h = q^25*x^10*d^10+q^16*(q^4+q^3+q^2+q+1)^2*x^9*d^9+
q^9*(q^13+3*q^12+7*q^11+13*q^10+20*q^9+26*q^8+30*q^7+
31*q^6+26*q^5+20*q^4+13*q^3+7*q^2+3*q+1)*x^8*d^8+
q^4*(q^9+2*q^8+4*q^7+6*q^6+7*q^5+8*q^4+6*q^3+
4*q^2+2*q+1)*(q^4+q^3+q^2+q+1)*(q^2+q+1)*x^7*d^7+
q*(q^2+q+1)*(q^5+2*q^4+2*q^3+3*q^2+2*q+1)*(q^4+q^3+q^2+q+1)*(q^2+1)*(q+1)*x^6*d^6+
(q^10+5*q^9+12*q^8+21*q^7+29*q^6+33*q^5+31*q^4+24*q^3+15*q^2+7*q+12)*x^5*d^5+
6*x^3*d^3+24;
homogfacFirstQWeyl(h);
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ x5d5+6
⇒ [3]:
⇒ x5d5+x3d3+4
```

See also: Section 7.5.12.12 [`homogfacFirstQWeyl.all`], page 510.

### 7.5.12.11 `homogfacNthQWeyl.all`

Procedure from library `ncfactor.lib` (see Section 7.5.12 [`ncfactor.lib`], page 480).

**Usage:** `homogfacNthQWeyl.all(h)`;  $h$  is a homogeneous polynomial in the  $n$ 'th  $q$ -Weyl algebra with respect to the weight vector  
 $@ [-1, \dots, -1, 1, \dots, 1]$ .  
 $@ \setminus \dots \setminus / \setminus \dots \setminus / @ \setminus / \setminus / @ n/2 \ n/2$

**Return:** list

**Purpose:** Computes all factorizations of a homogeneous polynomial  $h$  in the  $n$ 'th  $q$ -Weyl algebra

**Theory:** `homogfacNthQWeyl` returns a list with lists representing each a factorization of the given,  $[-1, \dots, -1, 1, \dots, 1]$ -homogeneous polynomial.

**General assumptions:**

- The basering is the  $n$ th Weyl algebra and has the form, that the first  $n$  variables represent  $x_1, \dots, x_n$ , and the second  $n$  variables do represent the  $d_1, \dots, d_n$ .
- We have  $n$  parameters  $q_1, \dots, q_n$  given.

**Example:**

```
LIB "ncfactor.lib";
ring R = (0,q1,q2,q3),(x1,x2,x3,d1,d2,d3),dp;
matrix C[6][6] = 1,1,1,q1,1,1,
```

```

1,1,1,1,q2,1,
1,1,1,1,1,q3,
1,1,1,1,1,1,
1,1,1,1,1,1,
1,1,1,1,1,1;
matrix D[6][6] = 0,0,0,1,0,0,
0,0,0,0,1,0,
0,0,0,0,0,1,
-1,0,0,0,0,0,
0,-1,0,0,0,0,
0,0,-1,0,0,0;
def r = nc_algebra(C,D);
setring(r);
poly h =x1*x2^2*x3^3*d1*d2^2+x2*x3^3*d2;
homogfacNthQWeyl_all(h);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x1*x2*d1*d2+1
↳ [4]:
↳ d2
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [2]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x1*x2*d1*d2+1
↳ [4]:
↳ x3
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [3]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x1*x2*d1*d2+1
↳ [4]:

```

```

↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [4]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x1*x2*d1*d2+1
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [5]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x1*x2*d1*d2+1
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [6]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x1*x2*d1*d2+1
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [7]:

```



```

↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x1*x2*d1*d2+1
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [8]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x1*x2*d1*d2+1
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [9]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x1*x2*d1*d2+1
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [10]:
↳ [1]:
↳ 1
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x3

```

```

↳ [5]:
↳ x3
↳ [6]:
↳ x1*x2*d1*d2+1
↳ [7]:
↳ d2
↳ [11]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x1*x2*d1*d2+1
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [12]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x1*x2*d1*d2+1
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [13]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x1*x2*d1*d2+1
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [14]:
↳ [1]:

```

```

↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ x1*x2*d1*d2+1
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [15]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ x1*x2*d1*d2+1
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [16]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ x1*x2*d1*d2+1
↳ [7]:
↳ d2
↳ [17]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:

```

```

↳ x1*x2*d1*d2+1
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [18]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:
↳ x1*x2*d1*d2+1
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [19]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:
↳ x3
↳ [6]:
↳ x1*x2*d1*d2+1
↳ [7]:
↳ d2
↳ [20]:
↳ [1]:
↳ 1
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x2
↳ [6]:
↳ x1*x2*d1*d2+1
↳ [7]:
↳ d2
↳ [21]:
↳ [1]:
↳ 1/(q2)

```

```

↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x2
↳ [4]:
↳ d2
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [22]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [23]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [24]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ x3

```

```

↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [25]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [26]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [27]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [28]:
↳ [1]:
↳ 1/(q2)
↳ [2]:

```

```

↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x2
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [29]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x2
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [30]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ x2
↳ [7]:
↳ d2
↳ [31]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ d2
↳ [4]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [5]:
↳ x3
↳ [6]:

```

```

↳ x3
↳ [7]:
↳ x3
↳ [32]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ d2
↳ [4]:
↳ x3
↳ [5]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [33]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ d2
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [7]:
↳ x3
↳ [34]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ d2
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [35]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2

```



```

↳ [3]:
↳ x3
↳ [4]:
↳ d2
↳ [5]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [36]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ d2
↳ [5]:
↳ x3
↳ [6]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [7]:
↳ x3
↳ [37]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ d2
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [38]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ d2
↳ [6]:
↳ x1*x2*d1*d2-x1*d1+(q2)

```

```

↳ [7]:
↳ x3
↳ [39]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [40]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x2
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [41]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [4]:
↳ x2
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x3
↳ [42]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:

```

```

↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [4]:
↳ x2
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [43]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [4]:
↳ x2
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ d2
↳ [44]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [4]:
↳ x3
↳ [5]:
↳ x2
↳ [6]:
↳ d2
↳ [7]:
↳ x3
↳ [45]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [4]:
↳ x3
↳ [5]:
↳ x2
↳ [6]:
↳ x3
↳ [7]:

```

```

↳ d2
↳ [46] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [4] :
↳ x3
↳ [5] :
↳ x3
↳ [6] :
↳ x2
↳ [7] :
↳ d2
↳ [47] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x2
↳ [4] :
↳ d2
↳ [5] :
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [6] :
↳ x3
↳ [7] :
↳ x3
↳ [48] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x2
↳ [4] :
↳ d2
↳ [5] :
↳ x3
↳ [6] :
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [7] :
↳ x3
↳ [49] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x2

```

```

↳ [4]:
↳ d2
↳ [5]:
↳ x3
↳ [6]:
↳ x3
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [50]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ d2
↳ [6]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [7]:
↳ x3
↳ [51]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [52]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x2
↳ [4]:
↳ x3
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)

```

```

↳ [53] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x3
↳ [4] :
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [5] :
↳ x2
↳ [6] :
↳ d2
↳ [7] :
↳ x3
↳ [54] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x3
↳ [4] :
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [5] :
↳ x2
↳ [6] :
↳ x3
↳ [7] :
↳ d2
↳ [55] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x3
↳ [4] :
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [5] :
↳ x3
↳ [6] :
↳ x2
↳ [7] :
↳ d2
↳ [56] :
↳ [1] :
↳ 1/(q2)
↳ [2] :
↳ x3
↳ [3] :
↳ x3
↳ [4] :

```

```

↳ x2
↳ [5]:
↳ d2
↳ [6]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [7]:
↳ x3
↳ [57]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:
↳ d2
↳ [6]:
↳ x3
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [58]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x2
↳ [5]:
↳ x3
↳ [6]:
↳ d2
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [59]:
↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x1*x2*d1*d2-x1*d1+(q2)
↳ [6]:
↳ x2
↳ [7]:
↳ d2
↳ [60]:

```

```

↳ [1]:
↳ 1/(q2)
↳ [2]:
↳ x3
↳ [3]:
↳ x3
↳ [4]:
↳ x3
↳ [5]:
↳ x2
↳ [6]:
↳ d2
↳ [7]:
↳ x1*x2*d1*d2-x1*d1+(q2)

```

See also: Section 7.5.12.10 [homogfacFirstQWeyl], page 491; Section 7.5.12.12 [homogfacFirstQWeyl\_all], page 510; Section 7.5.12.8 [homogfacNthWeyl], page 490.

### 7.5.12.12 homogfacFirstQWeyl\_all

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Usage:** `homogfacFirstQWeyl_all(h)`;  $h$  is a homogeneous polynomial in the first  $q$ -Weyl algebra with respect to the weight vector  $[-1,1]$

**Return:** list

**Purpose:** Computes all factorizations of a homogeneous polynomial  $h$  with respect to the weight vector  $[-1,1]$  in the first  $q$ -Weyl algebra

**Theory:** This function is a wrapper for `homogFacNthQWeyl_all`. It exists to make this library downward-compatible with older versions.

**Example:**

```

LIB "ncfactor.lib";
ring R = (0,q),(x,d),dp;
def r = nc_algebra (q,1);
setring(r);
poly h = q^25*x^10*d^10+q^16*(q^4+q^3+q^2+q+1)^2*x^9*d^9+
q^9*(q^13+3*q^12+7*q^11+13*q^10+20*q^9+26*q^8+30*q^7+
31*q^6+26*q^5+20*q^4+13*q^3+7*q^2+3*q+1)*x^8*d^8+
q^4*(q^9+2*q^8+4*q^7+6*q^6+7*q^5+8*q^4+6*q^3+
4*q^2+2q+1)*(q^4+q^3+q^2+q+1)*(q^2+q+1)*x^7*d^7+
q*(q^2+q+1)*(q^5+2*q^4+2*q^3+3*q^2+2*q+1)*(q^4+q^3+q^2+q+1)*(q^2+1)*(q+1)*x^6*d^6+
(q^10+5*q^9+12*q^8+21*q^7+29*q^6+33*q^5+31*q^4+24*q^3+15*q^2+7*q+12)*x^5*d^5+
6*x^3*d^3+24;
homogfacFirstQWeyl_all(h);
↳ [1]:
↳ [1]:
↳ 1
↳ [2]:
↳ x5d5+6
↳ [3]:
↳ x5d5+x3d3+4
↳ [2]:
↳ [1]:

```



```

↳ 1
↳ [2]:
↳ x5d5+x3d3+4
↳ [3]:
↳ x5d5+6

```

See also: Section 7.5.12.10 [homogfacFirstQWeyl], page 491.

### 7.5.12.13 tst\_ncfactor

Procedure from library `ncfactor.lib` (see Section 7.5.12 [ncfactor.lib], page 480).

**Example:**

```

LIB "ncfactor.lib";
tst_ncfactor();
↳ Testing delete_duplicates_noteval.
↳ Successful.
↳ Testing delete_duplicates_noteval_and_sort
↳ Successful.
↳ Testing combinekfinlf
↳ Successful.
↳ Testing permpp
↳ Successful.
↳ Testing binarySearch
↳ Successful.
↳ Testing getAllDivisorsFromFactList
↳ Successful.
↳ Testing triangNum
↳ Successful.
↳ Testing fromListToIntvec
↳ Successful.
↳ Testing fromIntvecToList
↳ Successful.
↳ Testing produceHomogListForProduct
↳ Successful.
↳ Testing possibleHomogPartsInBetween
↳ Successful.
↳ Testing nextSmallerEntry
↳ Successful.
↳ Testing possibleHomogPartsInBetweenNonRecursive
↳ Successful.
↳ Testing increment_intvec
↳ Successful.
↳ Testing isListEqual
↳ Successful.
↳ Testing ncfactor_isWeyl
↳ Successful.
↳ Testing ncfactor_isQWeyl
↳ Successful.
↳ Testing checkIfProperNthWeyl
↳ Successful.
↳ Testing checkIfProperNthQWeyl
↳ Successful.
↳ Testing isInCommutativeSubRing

```

```
⇒ Successful.
⇒ Testing factor_commutative
⇒ Successful.
⇒ Testing factorizeInt
⇒ Successful.
⇒ Testing getAllCoeffTuplesComb
⇒ Successful.
⇒ Testing normalizeFactors
⇒ Successful.
⇒ Testing divides
⇒ Successful.
⇒ Testing multiplyFactIntOutput
⇒ Successful.
⇒ Testing testNCfac
⇒ Successful.
⇒ Testing monsSmallerThan
⇒ Successful.
⇒ Testing getMaxDegreeVec
⇒ Successful.
⇒ Testing isFactorizationSmaller
⇒ Successful.
⇒ Testing sortedInsert
⇒ Successful.
⇒ Testing isFactorizationEqual
⇒ Successful.
⇒ Testing sortFactorizations
⇒ Successful.
⇒ Testing homogfacNthWeyl
⇒ Successful.
⇒ Testing homogfacNthWeyl_all
⇒ Successful.
⇒ Testing facWeyl
⇒ Successful.
⇒ Testing checkForHomogInhomogInterchangabilityNthWeyl
⇒ Successful.
⇒ Testing sfacwa2NthWeyl
⇒ Successful.
⇒ Testing sfacwa2NthWeyl
⇒ Successful.
⇒ Testing determineRestOfHomogPartsNthWeyl
⇒ Successful.
⇒ Testing gammaForThetaNthWeyl
⇒ Successful.
⇒ Testing extractHomogeneousDivisorsNthWeyl
⇒ Successful.
⇒ Testing extractHomogeneousDivisorsLeftNthWeyl
⇒ Successful.
⇒ Testing extractHomogeneousDivisorsRightNthWeyl
⇒ Successful.
⇒ Testing computeCombinationsMinMaxHomogNthWeyl
⇒ Successful.
⇒ Testing getAllCombOfHomogFact
⇒ Successful.
```

```

↳ Testing getPossibilitiesForRightSides
↳ Successful.
↳ Testing homogDistributionNthWeyl
↳ Successful.
↳ Testing extractLeadingTermOfNthWeylPoly
↳ Successful.
↳ Testing degreeOfNthWeylPoly
↳ Successful.
↳ Testing degreeOfNthWeylPolyInverted
↳ Successful.
↳ Testing homogwithorderNthWeyl
↳ Successful.
↳ Testing isInWeylSubAlg
↳ Successful.
↳ Testing facSubWeyl
↳ Successful.
↳ Testing facShift
↳ Successful.
↳ Testing homogfacNthQWeyl
↳ Test 4 for homogfacNthQWeyl failed.
↳ Expected:n
↳ [1]:
↳ 1/2
↳ [2]:
↳ (2*q4^6)*x4^4*d4^4+(2*q4^5+4*q4^4+6*q4^3)*x4^3*d4^3+(2*q4^3+6*q4^2+6*q\
4)*x4^2*d4^2+3*x2*d2+2*x4*d4+10
↳ [3]:
↳ (q3)*x3^2*d3^2+(q4)*x4^2*d4^2+x3*d3+x4*d4+4
↳ obtained:n
↳ [1]:
↳ 1/2
↳ [2]:
↳ (q3)*x3^2*d3^2+(q4)*x4^2*d4^2+x3*d3+x4*d4+4
↳ [3]:
↳ (2*q4^6)*x4^4*d4^4+(2*q4^5+4*q4^4+6*q4^3)*x4^3*d4^3+(2*q4^3+6*q4^2+6*q\
4)*x4^2*d4^2+3*x2*d2+2*x4*d4+10
↳ Test 5 for homogfacNthQWeyl failed.
↳ Expected:n
↳ [1]:
↳ 1/2
↳ [2]:
↳ (2*q4^6)*x4^4*d4^4+(2*q4^5+4*q4^4+6*q4^3)*x4^3*d4^3+(2*q4^3+6*q4^2+6*q\
4)*x4^2*d4^2+3*x2*d2+2*x4*d4+10
↳ [3]:
↳ (q3)*x3^2*d3^2+(q4)*x4^2*d4^2+x3*d3+x4*d4+4
↳ [4]:
↳ d1
↳ [5]:
↳ x2
↳ [6]:
↳ d3
↳ [7]:
↳ x4

```

```

↳ obtained:n
↳ [1]:
↳ 1/2
↳ [2]:
↳ (q3)*x3^2*d3^2+(q4)*x4^2*d4^2+x3*d3+x4*d4+4
↳ [3]:
↳ (2*q4^6)*x4^4*d4^4+(2*q4^5+4*q4^4+6*q4^3)*x4^3*d4^3+(2*q4^3+6*q4^2+6*q\
4)*x4^2*d4^2+3*x2*d2+2*x4*d4+10
↳ [4]:
↳ d1
↳ [5]:
↳ x2
↳ [6]:
↳ d3
↳ [7]:
↳ x4
↳ ? homogfacNthQWeyl is not working properly.
↳ ? leaving ncfactor.lib::tst_ncfactor (0)

```

### 7.5.13 nchilbert.lib

**Library:** nchilbert.lib

**Purpose:** Hilbert series, polynomial and multiplicity for G-Algebras (Plural)

**Authors:** Andre Ranft, andre.ranft at rwth-aachen.de  
Viktor Levandovskyy, levandov at rwth-aachen.de

**Overview:** The theory is found in the book by Bueso, Gomez-Torrecillas, and Verschoren Algorithmic Methods in Non-Commutative Algebra. Applications to Quantum Groups. and in the bachelor thesis by Andre Ranft, Hilbert polynomials of modules over noncommutative G-algebras, RWTH Aachen, 2014.

**Procedures:** See also: Section 5.1.56 [hilb], page 193; Section D.15.13 [multigrading.lib], page 934; Section D.7.3 [rinvar.lib], page 873.

#### 7.5.13.1 ncHilb

Procedure from library nchilbert.lib (see Section 7.5.13 [nchilbert.lib], page 514).

**Usage:** ncHilb(M,j); M a module, j an int

**Return:** intvec

**Assume:** - M is given via a Groebner basis; j is 1 or 2;  
- the weights of all the ring variables are positive

**Note:** - computes the first (if j=1) or second (j=2) Hilbert series of I as intvec - the procedure works analogously to the commutative procedure hilb - If the returned vector has the form  $v=(v_0, v_1, \dots, v_d, 0)$ , then the Hilbert series is  $v_0 + v_1*t + \dots + v_d*t^d$

**Example:**

```

LIB "nchilbert.lib";
def A = makeUs12(); setring A;
ideal I = e,h-1; I = std(I);
ncHilb(I,1); // first Hilbert series of A/I

```

```

↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
↳ 1,-2,1,0
ncHilb(I,2); // second Hilbert series of A/I
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
↳ 1,0
ideal J = I, f^2; J = std(J);
ncHilb(J,2);
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
↳ 1,1,0
// now with weights 1,2,3
ring r = 0,(e,f,h),wp(1,2,3);
matrix D[3][3]; D[1,2]=-h; D[1,3]=2*e;D[2,3]=-2*f;
def R = nc_algebra(1,D); setring R;
ideal I = imap(A,I); I = std(I);
ncHilb(I,1); // first weighted Hilbert series of R/I
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
↳ 1,-1,0,-1,1,0
ncHilb(I,2); // second weighted Hilbert series of R/I
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
↳ 1,1,1,0
matrix M[2][5] =
e,h-1,f^2, 0,0,
0,0,0, e,h+1;
module G = std(M);
print(G);
↳ e,0,h-1,0, f2,
↳ 0,e,0, h+1,0
ncHilb(G,1); // first weighted Hilbert series of R^2/G
↳ 2,-2,0,-2,1,1,0,1,-1,0
ncHilb(G,2); // second weighted Hilbert series of R^2/G
↳ 2,2,2,0,-1,-1,-1,0

```

See also: Section 5.1.56 [hilb], page 193.

### 7.5.13.2 ncHilbertSeries

Procedure from library `nchilbert.lib` (see Section 7.5.13 [nchilbert.lib], page 514).

**Usage:** `ncHilbertSeries(M,j)`;  $M$  is a module,  $j$  is an int

**Return:** ring

**Purpose:** computes the first (if  $j=1$ ) and second ( $j=2$ ) Hilbert Series of  $A^m/M$

**Assume:** -  $M$  is given via a Groebner basis;  $j$  is 1 or 2;  
- the weights of all the ring variables are positive

**Note:** - the procedure returns an univariate ring and a polynomial called `ncHS` in it.

**Example:**

```
LIB "nchilbert.lib";
```

```

def A = makeUs12(); setring A;
ideal I = e,h-1; I = std(I);
def r = ncHilbertSeries(I,1); setring r;
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
ncHS;
↳ t2-2t+1
setring A; kill r;
def s= ncHilbertSeries(I,2); setring s;
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
ncHS;
↳ 1
// now consider admissible weights 1,2,3
ring r = 0,(e,f,h),wp(1,2,3);
matrix D[3][3]; D[1,2]=-h; D[1,3]=2*e;D[2,3]=-2*f;
def R = nc_algebra(1,D); setring R;
matrix M[2][5] =
e,h-1,f^2, 0,0,
0,0,0, e,h+1;
module G = std(M);
print(G);
↳ e,0,h-1,0, f2,
↳ 0,e,0, h+1,0
def r= ncHilbertSeries(G,1); setring r;
↳ // ** redefining r (def r= ncHilbertSeries(G,1); setring r;) ./examples/\
ncHilbertSeries.sing:18
ncHS; // first weighted Hilbert series of R^2/G
↳ -t8+t7+t5+t4-2t3-2t+2
setring R; kill r;
def s=ncHilbertSeries(G,2); setring s;
↳ // ** redefining s (def s=ncHilbertSeries(G,2); setring s;) ./examples/nc\
HilbertSeries.sing:21
ncHS;// second weighted Hilbert series of R^2/G
↳ -t6-t5-t4+2t2+2t+2

```

### 7.5.13.3 ncHilbertPolynomial

Procedure from library `nchilbert.lib` (see Section 7.5.13 [`nchilbert.lib`], page 514).

**Usage:** `ncHilbertPolynomial(M)`;  $M$  is a module

**Return:** ring

**Purpose:** computes the Hilbert polynomial of  $A^m/M$

**Assume:** -  $M$  is given via a Groebner basis  
- the weights of all the ring variables are positive

**Note:** - the procedure returns an univariate ring and a polynomial called `ncHP` in it.

**Example:**

```

LIB "nchilbert.lib";
def A = makeUs12(); setring A;
ideal I = h^4,e*f*h^3,e^2*f^2*h^2+2*e*f*h^2; I = std(I);
dim(I); // 2

```

```

↳ 2
def r = ncHilbertPolynomial(I); setring r;
ncHP; // 2t+7
↳ 2t+7
kill r;
// now consider admissible weights 1,2,3
ring r = 0, (e,f,h), wp(1,2,3);
matrix D[3][3]; D[1,2]=-h; D[1,3]=2*e;D[2,3]=-2*f;
def R = nc_algebra(1,D); setring R;
ideal I = imap(A,I);
I = std(I);
dim(I); // 2
↳ 2
def r = ncHilbertPolynomial(I); setring r;
↳ // ** redefining r (def r = ncHilbertPolynomial(I); setring r;) ./example\
 s/ncHilbertPolynomial.sing:15
ncHP; // 6t+18
↳ 6t+18

```

See also: [hilbPoly], page 799.

#### 7.5.13.4 ncHilbertMultiplicity

Procedure from library `nchilbert.lib` (see Section 7.5.13 [nchilbert.lib], page 514).

**Usage:** `ncHilbertMultiplicity(M)`; M is a module

**Return:** `int`

**Purpose:** compute the (Hilbert) multiplicity of the module `coker(M)`

**Assume:** - M is given via a Groebner basis  
- the weights of all the ring variables are positive

**Note:** the multiplicity depends on the selected weights of variables

**Example:**

```

LIB "nchilbert.lib";
def A = makeUs12(); setring A;
ideal I = e,h-1; I = std(I);
ncHilbertMultiplicity(I); // multiplicity of A/I
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
↳ 1
ideal J = I, f^2; J = std(J);
ncHilbertMultiplicity(J);
↳ Warning: the input generators are not a Groebner basis
↳ The result might have no meaning
↳ 2
// now the same algebra with weights 1,2,3
ring r = 0, (e,f,h), wp(1,2,3);
matrix D[3][3]; D[1,2]=-h; D[1,3]=2*e;D[2,3]=-2*f;
def R = nc_algebra(1,D); setring R;
ideal I = imap(A,I); I = std(I);
ncHilbertMultiplicity(I);
↳ Warning: the input generators are not a Groebner basis

```

```

↳ The result might have no meaning
↳ 3
matrix M[2][5] =
e,h-1,f^2, 0,0,
0,0,0, e,h+1;
module G = std(M);
print(G);
↳ e,0,h-1,0, f2,
↳ 0,e,0, h+1,0
nchilbertMultiplicity(G);
↳ 3

```

### 7.5.13.5 GKExp

Procedure from library `nchilbert.lib` (see Section 7.5.13 [`nchilbert.lib`], page 514).

**Usage:** GKExp(M); M a module

**Return:** int

**Purpose:** computes the Gelfand-Kirillov-Dimension of `coker(M)` via `Exp(M)`

**Assume:** basering is G-Algebra

**Note:** for zero module -1 is returned

**Example:**

```

LIB "nchilbert.lib";
def A = makeUs12(); setring A;
ideal I = e,h-1; I = std(I);
GKExp(I); // computes GKdim(A/I), should be 1
↳ Warning: the input generators are not a Groebner basis
↳ Proceed with Groebner basis computation
↳ 1
ideal J = I, f^2; J = std(J);
GKExp(J); // should be 0
↳ Warning: the input generators are not a Groebner basis
↳ Proceed with Groebner basis computation
↳ 0
matrix M[2][4] =
e,h-1,0,0,
0,0,e,h+1;
module G = std(M);
print(G);
↳ h-1,0, e,0,
↳ 0, h+1,0,e
GKExp(G);
↳ 1

```

See also: Section 7.5.9.1 [`GKdim`], page 457; Section 7.3.3 [`dim (plural)`], page 331.

### 7.5.13.6 mondlim

Procedure from library `nchilbert.lib` (see Section 7.5.13 [`nchilbert.lib`], page 514).

**Usage:** mondlim(B,i); B is list of elements of  $N_0^i$ ,

**Return:** int



**Purpose:** computes the dimension of the monoid ideal generated by B

**Example:**

```
LIB "nchilbert.lib";
ring A = 0,(x,y,z),dp;
setring A;
list I = [1,0,1],[0,1,1]; // belongs to the ideal <xz,yz>
mondim(I,3);
↳ 1
mondim(I,5); // treat generators of I as extended in N_0^5
↳ 3
```

### 7.5.14 dmodloc.lib

Status: experimental

**Library:** dmodloc.lib

**Purpose:** Localization of algebraic D-modules and applications

**Author:** Daniel Andres, daniel.andres@math.rwth-aachen.de

Support: DFG Graduiertenkolleg 1632 ‘Experimentelle und konstruktive Algebra’

**Overview:** Let  $I$  be a left ideal in the  $n$ -th polynomial Weyl algebra  $D=K[x]\langle d \rangle$  and let  $f$  be a polynomial in  $K[x]$ .

If  $D/I$  is a holonomic module over  $D$ , it is known that the localization of  $D/I$  at  $f$  is also holonomic. The procedure `Dlocalization` computes an ideal  $J$  in  $D$  such that this localization is isomorphic to  $D/J$  as  $D$ -modules.

If one regards  $I$  as an ideal in the rational Weyl algebra as above,  $K(x)\langle d \rangle * I$ , and intersects with  $K[x]\langle d \rangle$ , the result is called the Weyl closure of  $I$ . The procedures `WeylClosure` (if  $I$  has finite holonomic rank) and `WeylClosure1` (if  $I$  is in the first Weyl algebra) can be used for computations.

As an application of the Weyl closure, the procedure `annRatSyz` computes a holonomic part of the annihilator of a rational function by computing certain syzygies. The full annihilator can be obtained by taking the Weyl closure of the result.

If one regards the left ideal  $I$  as system of linear PDEs, one can find its polynomial solutions with `polSol` (if  $I$  is holonomic) or `polSolFiniteRank` (if  $I$  is of finite holonomic rank). Rational solutions can be obtained with `ratSol`.

The procedure `bfctBound` computes a possible multiple of the  $b$ -function for  $f^s * u$  at a generic root of  $f$ . Here,  $u$  stands for  $[1]$  in  $D/I$ .

This library also offers the procedures `holonomicRank` and `DsingularLocus` to compute the holonomic rank and the singular locus of the  $D$ -module  $D/I$ .

**References:**

(OT) T. Oaku, N. Takayama: ‘Algorithms for D-modules’, Journal of Pure and Applied Algebra, 1998.

(OTT) T. Oaku, N. Takayama, H. Tsai: ‘Polynomial and rational solutions of holonomic systems’, Journal of Pure and Applied Algebra, 2001.

(OTW) T. Oaku, N. Takayama, U. Walther: ‘A Localization Algorithm for D-modules’, Journal of Symbolic Computation, 2000.

(Tsa) H. Tsai: ‘Algorithms for algebraic analysis’, PhD thesis, 2000.

**Procedures:** See also: Section 7.5.2 [`bfun.lib`], page 370; Section 7.5.4 [`dmod.lib`], page 395; Section 7.5.5 [`dmodapp.lib`], page 414; Section 7.5.7 [`dmodvar.lib`], page 446; Section D.6.13 [`gmssing.lib`], page 862.

### 7.5.14.1 Dlocalization

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `Dlocalization(I,f[,k,e]);` I ideal, f poly, k,e optional ints

**Assume:** The basering is the n-th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity

$\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

Further, assume that f does not contain any  $D(i)$  and that I is holonomic on  $K^n \setminus V(f)$ .

**Return:** ideal or list, computes an ideal J such that  $D/J$  is isomorphic to  $D/I$  localized at f as D-modules.

If  $k > 0$ , a list consisting of J and an integer m is returned, such that  $f^m$  represents the natural map from  $D/I$  to  $D/J$ . Otherwise (and by default), only the ideal J is returned.

**Remarks:** It is known that a localization at f of a holonomic D-module is again a holonomic D-module.

Reference: (OTW)

**Note:** If  $e > 0$ , `std` is used for Groebner basis computations, otherwise (and by default) `slimgb` is used.

If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodloc.lib";
// (OTW), Example 8
ring r = 0, (x,y,z,Dx,Dy,Dz), dp;
def W = Weyl();
setring W;
poly f = x^3-y^2*z^2;
ideal I = f^2*Dx+3*x^2, f^2*Dy-2*y*z^2, f^2*Dz-2*y^2*z;
// I annihilates exp(1/f)
ideal J = Dlocalization(I,f);
J;
↳ J[1]=y*Dy-z*Dz
↳ J[2]=2*y*z^2*Dx+3*x^2*Dy
↳ J[3]=2*y^2*z*Dx+3*x^2*Dz
↳ J[4]=2*z^3*Dx*Dz+3*x^2*Dy^2+2*z^2*Dx
↳ J[5]=3*y^2*z^3*Dz-2*x^4*Dx-6*x^3*z*Dz+12*y^2*z^2-12*x^3-6
↳ J[6]=4*x^4*Dx^2+12*x^3*z*Dx*Dz+9*x^2*z^2*Dz^2+40*x^3*Dx+63*x^2*z*Dz+72*x^2+12*Dx
↳ J[7]=3*y*z^4*Dz^2-2*x^4*Dx*Dy-6*x^3*z*Dy*Dz+21*y*z^3*Dz-12*x^3*Dy+24*y*z^2-6*Dy
↳ J[8]=3*z^5*Dz^3-2*x^4*Dx*Dy^2-6*x^3*z*Dy^2*Dz+30*z^4*Dz^2-12*x^3*Dy^2+66*z^3*Dz+24*z^2-6*Dy^2
Dlocalization(I,f,1); // The natural map D/I -> D/J is given by 1/f^2
↳ [1]:
↳ _[1]=y*Dy-z*Dz
↳ _[2]=2*y*z^2*Dx+3*x^2*Dy
↳ _[3]=2*y^2*z*Dx+3*x^2*Dz
↳ _[4]=2*z^3*Dx*Dz+3*x^2*Dy^2+2*z^2*Dx
↳ _[5]=3*y^2*z^3*Dz-2*x^4*Dx-6*x^3*z*Dz+12*y^2*z^2-12*x^3-6
```

```

↳ _[6]=4*x^4*Dx^2+12*x^3*z*Dx*Dz+9*x^2*z^2*Dz^2+40*x^3*Dx+63*x^2*z*Dz+72\
*x^2+12*Dx
↳ _[7]=3*y*z^4*Dz^2-2*x^4*Dx*Dy-6*x^3*z*Dy*Dz+21*y*z^3*Dz-12*x^3*Dy+24*y\
*z^2-6*Dy
↳ _[8]=3*z^5*Dz^3-2*x^4*Dx*Dy^2-6*x^3*z*Dy^2*Dz+30*z^4*Dz^2-12*x^3*Dy^2+\
66*z^3*Dz+24*z^2-6*Dy^2
↳ [2]:
↳ 2

```

See also: Section 7.5.5.3 [DLoc], page 417; Section 7.5.5.5 [DLoc0], page 418; Section 7.5.5.4 [SDLoc], page 417.

### 7.5.14.2 WeylClosure

Procedure from library `dmodloc.lib` (see Section 7.5.14 [dmodloc.lib], page 519).

**Usage:** WeylClosure(I); I an ideal

**Assume:** The basering is the  $n$ -th Weyl algebra  $W$  over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ . Moreover, assume that the holonomic rank of  $W/I$  is finite.

**Return:** ideal, the Weyl closure of  $I$

**Remarks:** The Weyl closure of a left ideal  $I$  in the Weyl algebra  $W$  is defined to be the intersection of  $I$  regarded as left ideal in the rational Weyl algebra  $K(x(1..n)) \langle D(1..n) \rangle$  with the polynomial Weyl algebra  $W$ .  
Reference: (Tsa), Algorithm 2.2.4

**Note:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodloc.lib";
// (OTW), Example 8
ring r = 0, (x,y,z,Dx,Dy,Dz), dp;
def D3 = Weyl();
setring D3;
poly f = x^3-y^2*z^2;
ideal I = f^2*Dx + 3*x^2, f^2*Dy-2*y*z^2, f^2*Dz-2*y^2*z;
// I annihilates exp(1/f)
WeylClosure(I);
↳ _[1]=y*Dy-z*Dz
↳ _[2]=2*y*z^2*Dx+3*x^2*Dy
↳ _[3]=2*y^2*z*Dx+3*x^2*Dz
↳ _[4]=2*z^3*Dx*Dz+3*x^2*Dy^2+2*z^2*Dx
↳ _[5]=4*x^4*Dx^2+12*x^3*z*Dx*Dz+9*x^2*z^2*Dz^2+16*x^3*Dx+27*x^2*z*Dz+12*Dx
↳ _[6]=3*y*z^4*Dz^2-2*x^4*Dx*Dy-6*x^3*z*Dy*Dz+9*y*z^3*Dz-6*Dy
↳ _[7]=3*y^2*z^3*Dz-2*x^4*Dx-6*x^3*z*Dz-6
↳ _[8]=3*z^5*Dz^3-2*x^4*Dx*Dy^2-6*x^3*z*Dy^2*Dz+18*z^4*Dz^2+18*z^3*Dz-6*Dy\
2

```

See also: Section 7.5.14.3 [WeylClosure1], page 522.

### 7.5.14.3 WeylClosure1

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `WeylClosure1(L)`;  $L$  a poly

**Assume:** The basering is the first Weyl algebra  $D=K\langle x,d \mid dx=xd+1 \rangle$  over a field  $K$  of characteristic 0.

**Return:** ideal, the Weyl closure of the principal left ideal generated by  $L$

**Remarks:** The Weyl closure of a left ideal  $I$  in the Weyl algebra  $D$  is defined to be the intersection of  $I$  regarded as left ideal in the rational Weyl algebra  $K(x)\langle d \rangle$  with the polynomial Weyl algebra  $D$ .

Reference: (Tsa), Algorithm 1.2.4

**Note:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0, (x, Dx), dp;
def W = Weyl();
setring W;
poly L = (x^3+2)*Dx-3*x^2;
WeylClosure1(L);
⇨ _[1]=x^3*Dx-3*x^2+2*Dx
⇨ _[2]=x^2*Dx^2-2*x*Dx
⇨ _[3]=x^2*Dx+Dx^2-3*x
⇨ _[4]=x*Dx^2-2*Dx
L = (x^4-4*x^3+3*x^2)*Dx^2+(-6*x^3+20*x^2-12*x)*Dx+(12*x^2-32*x+12);
WeylClosure1(L);
⇨ _[1]=x^4*Dx^2-4*x^3*Dx^2-6*x^3*Dx+3*x^2*Dx^2+20*x^2*Dx+12*x^2-12*x*Dx-32*x+12
⇨ _[2]=x^2*Dx^3-21/10*x^2*Dx^2+3/10*x*Dx^3-6/5*x*Dx^2+63/5*x*Dx-3/5*Dx^2-12/5*Dx-126/5
⇨ _[3]=x^3*Dx^2-43/10*x^2*Dx^2+9/10*x*Dx^3-6*x^2*Dx+12/5*x*Dx^2+109/5*x*Dx-9/5*Dx^2+12*x-36/5*Dx-178/5
⇨ _[4]=x^3*Dx^3-48/5*x^2*Dx^2+9/5*x*Dx^3+24/5*x*Dx^2+198/5*x*Dx-18/5*Dx^2-7/2/5*Dx-336/5
⇨ _[5]=x^3*Dx^4-4*x^2*Dx^4+2*x^2*Dx^3+3*x*Dx^4-69/10*x^2*Dx^2+67/10*x*Dx^3-24/5*x*Dx^2-3*Dx^3+207/5*x*Dx-27/5*Dx^2-18/5*Dx-414/5
⇨ _[6]=x^3*Dx^6+8/3*x^3*Dx^5-4*x^2*Dx^6-2/3*x^2*Dx^5+3*x*Dx^6+16*x^2*Dx^4-2/0*x*Dx^5-92/3*x*Dx^4+12*Dx^5+126/5*x^2*Dx^2-258/5*x*Dx^3-168/5*x*Dx^2+92/3*Dx^3-756/5*x*Dx+356/5*Dx^2+504/5*Dx+1512/5
```

See also: Section 7.5.14.2 [`WeylClosure`], page 521.

### 7.5.14.4 holonomicRank

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `holonomicRank(I[,e])`;  $I$  ideal,  $e$  optional int

**Assume:** The basering is the  $n$ -th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity

$\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Return:** int, the holonomic rank of I

**Remarks:** The holonomic rank of I is defined to be the  $K(x(1..n))$ -dimension of the module  $W/WI$ , where  $W$  is the rational Weyl algebra  $K(x(1..n))\langle D(1..n) \rangle$ . If this dimension is infinite, -1 is returned.

**Note:** If  $e < 0$ , `std` is used for Groebner basis computations, otherwise (and by default) `slimgb` is used. If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodloc.lib";
// (OTW), Example 8
ring r3 = 0, (x,y,z,Dx,Dy,Dz), dp;
def D3 = Weyl();
setring D3;
poly f = x^3-y^2*z^2;
ideal I = f^2*Dx+3*x^2, f^2*Dy-2*y*z^2, f^2*Dz-2*y^2*z;
// I annihilates exp(1/f)
holonomicRank(I);
↪ 1
```

### 7.5.14.5 DsingularLocus

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `DsingularLocus(I)`; I ideal

**Assume:** The basering is the  $n$ -th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Return:** ideal, describing the singular locus of the  $D$ -module  $D/I$

**Note:** If `printlevel>=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed

**Example:**

```
LIB "dmodloc.lib";
// (OTW), Example 8
ring @D3 = 0, (x,y,z,Dx,Dy,Dz), dp;
def D3 = Weyl();
setring D3;
poly f = x^3-y^2*z^2;
ideal I = f^2*Dx + 3*x^2, f^2*Dy-2*y*z^2, f^2*Dz-2*y^2*z;
// I annihilates exp(1/f)
DsingularLocus(I);
↪ _[1]=y^2*z^2-x^3
```

### 7.5.14.6 polSol

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `polSol(I[w,m])`; I ideal, w optional intvec, m optional int

- Assume:** The basering is the  $n$ -th Weyl algebra  $W$  over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ . Moreover, assume that  $I$  is holonomic.
- Return:** ideal, a basis of the polynomial solutions to the given system of linear PDEs with polynomial coefficients, encoded via  $I$
- Remarks:** If  $w$  is given,  $w$  should consist of  $n$  strictly negative entries. Otherwise and by default,  $w$  is set to  $-1:n$ .  
In this case,  $w$  is used as weight vector for the computation of a b-function.  
If  $m$  is given,  $m$  is assumed to be the minimal integer root of the b-function of  $I$  w.r.t.  $w$ . Note that this assumption is not checked.  
Reference: (OTT), Algorithm 2.4
- Note:** If  $\text{printlevel}=1$ , progress debug messages will be printed, if  $\text{printlevel} \geq 2$ , all the debug messages will be printed.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def W = Weyl();
setring W;
poly tx,ty = x*Dx, y*Dy;
ideal I = // Appel F1 with parameters (2,-3,-2,5)
tx*(tx+ty+4)-x*(tx+ty+2)*(tx-3),
ty*(tx+ty+4)-y*(tx+ty+2)*(ty-2),
(x-y)*Dx*Dy+2*Dx-3*Dy;
intvec w = -1,-1;
polSol(I,w);
↪ _[1]=10*x^3*y^2-30*x^3*y-45*x^2*y^2+24*x^3+144*x^2*y+72*x*y^2-126*x^2-252\
*x*y-42*y^2+252*x+168*y-210
```

See also: Section 7.5.14.7 [polSolFiniteRank], page 524; Section 7.5.14.8 [ratSol], page 525.

**7.5.14.7 polSolFiniteRank**

Procedure from library `dmodloc.lib` (see Section 7.5.14 [dmodloc.lib], page 519).

- Usage:** `polSolFiniteRank(I,[w]);`  $I$  ideal,  $w$  optional intvec
- Assume:** The basering is the  $n$ -th Weyl algebra  $W$  over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ . Moreover, assume that  $I$  is of finite holonomic rank.
- Return:** ideal, a basis of the polynomial solutions to the given system of linear PDEs with polynomial coefficients, encoded via  $I$
- Remarks:** If  $w$  is given,  $w$  should consist of  $n$  strictly negative entries. Otherwise and by default,  $w$  is set to  $-1:n$ .  
In this case,  $w$  is used as weight vector for the computation of a b-function.  
Reference: (OTT), Algorithm 2.6
- Note:** If  $\text{printlevel}=1$ , progress debug messages will be printed, if  $\text{printlevel} \geq 2$ , all the debug messages will be printed.

**Example:**

```

LIB "dmodloc.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def W = Weyl();
setring W;
poly tx,ty = x*Dx, y*Dy;
ideal I = // Appel F1 with parameters (2,-3,-2,5)
tx*(tx+ty+4)-x*(tx+ty+2)*(tx-3),
ty*(tx+ty+4)-y*(tx+ty+2)*(ty-2),
(x-y)*Dx*Dy+2*Dx-3*Dy;
intvec w = -1,-1;
polSolFiniteRank(I,w);
↪ _[1]=10*x^3*y^2-30*x^3*y-45*x^2*y^2+24*x^3+144*x^2*y+72*x*y^2-126*x^2-252\
*x*y-42*y^2+252*x+168*y-210

```

See also: Section 7.5.14.6 [polSol], page 523; Section 7.5.14.8 [ratSol], page 525.

**7.5.14.8 ratSol**

Procedure from library `dmodloc.lib` (see Section 7.5.14 [dmodloc.lib], page 519).

**Usage:** `ratSol(I);` I ideal

**Assume:** The basering is the  $n$ -th Weyl algebra  $W$  over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ . Moreover, assume that  $I$  is holonomic.

**Return:** module, a basis of the rational solutions to the given system of linear PDEs with polynomial coefficients, encoded via  $I$ . Note that each entry has two components, the first one standing for the numerator, the second one for the denominator.

**Remarks:** Reference: (OTT), Algorithm 3.10

**Note:** If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```

LIB "dmodloc.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def W = Weyl();
setring W;
poly tx,ty = x*Dx, y*Dy;
ideal I = // Appel F1 with parameters (3,-1,1,1) is a solution
tx*(tx+ty)-x*(tx+ty+3)*(tx-1),
ty*(tx+ty)-y*(tx+ty+3)*(ty+1);
module M = ratSol(I);
// We obtain a basis of the rational solutions to I represented by a
// module / matrix with two rows.
// Each column of the matrix represents a rational function, where
// the first row correspond to the numerator and the second row to
// the denominator.
print(M);
↪ x-y, x,
↪ y^4-3*y^3+3*y^2-y,y

```

See also: Section 7.5.14.6 [polSol], page 523; Section 7.5.14.7 [polSolFiniteRank], page 524.

### 7.5.14.9 bfctBound

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `bfctBound (I,f[,primdec]);` I ideal, f poly, primdec optional string

**Assume:** The basering is the n-th Weyl algebra W over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ . Moreover, assume that I is holonomic.

**Return:** list of roots (of type ideal) and multiplicities (of type intvec) of a multiple of the b-function for  $f^s \cdot u$  at a generic root of f. Here, u stands for [1] in  $D/I$ .

**Remarks:** Reference: (OTT), Algorithm 3.4

**Note:** This procedure requires to compute a primary decomposition in a commutative ring. The optional string primdec can be used to specify the algorithm to do so. It may either be 'GTZ' (Gianni, Trager, Zacharias) or 'SY' (Shimoyama, Yokoyama). By default, 'GTZ' is used.

If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def W = Weyl();
setring W;
poly tx,ty = x*Dx, y*Dy;
ideal I = // Appel F1 with parameters (2,-3,-2,5)
tx*(tx+ty+4)-x*(tx+ty+2)*(tx-3),
ty*(tx+ty+4)-y*(tx+ty+2)*(ty-2),
(x-y)*Dx*Dy+2*Dx-3*Dy;
kill tx,ty;
poly f = x-1;
bfctBound(I,f);
↳ [1]:
↳ _[1]=-1
↳ _[2]=-7
↳ [2]:
↳ 1,1
```

See also: [`bernstein`], page 863; Section 7.5.2.1 [`bfct`], page 371; Section 7.5.2.3 [`bfctAnn`], page 373.

### 7.5.14.10 annRatSyz

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `annRatSyz(f,g[,db,eng]);` f, g polynomials, db,eng optional integers

**Assume:** The basering is commutative and over a field of characteristic 0.

**Return:** ring (a Weyl algebra) containing an ideal 'LD', which is (part of) the annihilator of the rational function  $g/f$  in the corresponding Weyl algebra

**Remarks:** This procedure uses the computation of certain syzygies. One can obtain the full annihilator by computing the Weyl closure of the ideal LD.



**Note:** Activate the output ring with the `setring` command. In the output ring, the ideal 'LD' (in Groebner basis) is (part of) the annihilator of  $g/f$ .  
 If  $db > 0$  is given, operators of order up to  $db$  are considered, otherwise, and by default, a minimal holonomic solution is computed.  
 If  $eng < 0$ , `std` is used for Groebner basis computations, otherwise, and by default, `slimgb` is used.  
 If  $printlevel = 1$ , progress debug messages will be printed, if  $printlevel \geq 2$ , all the debug messages will be printed.

**Example:**

```
LIB "dmodloc.lib";
// printlevel = 3;
ring r = 0,(x,y),dp;
poly f = 2*x*y; poly g = x^2 - y^3;
def A = annRatSyz(f,g); // compute a holonomic solution
setring A; A;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
LD;
↳ LD[1]=3*x*Dx+2*y*Dy+1
↳ LD[2]=y^4*Dy-x^2*y*Dy+2*y^3+x^2
setring r;
def B = annRatSyz(f,g,5); // compute a solution up to degree 5
setring B;
LD; // this is the full annihilator as we will check below
↳ LD[1]=15*y^2*Dx^2*Dy-2*x*Dx*Dy^2-8*y*Dy^3+45*y*Dx^2
↳ LD[2]=3*y^3*Dx^2+15*x^2*Dx^2-8*y^2*Dy^2+30*x*Dx
↳ LD[3]=3*x*Dx+2*y*Dy+1
↳ LD[4]=y^3*Dy^2-x^2*Dy^2+6*y^2*Dy+6*y
↳ LD[5]=y^4*Dy-x^2*y*Dy+2*y^3+x^2
setring r;
def C = annRat(f,g); setring C;
LD; // the full annihilator
↳ LD[1]=3*y^2*Dx^2*Dy+2*x*Dx*Dy^2+9*y*Dx^2+4*Dy^2
↳ LD[2]=3*y^3*Dx^2-10*x*y*Dx*Dy-8*y^2*Dy^2+10*x*Dx
↳ LD[3]=y^3*Dy^2-x^2*Dy^2-6*x*y*Dx+2*y^2*Dy+4*y
↳ LD[4]=3*x*Dx+2*y*Dy+1
↳ LD[5]=y^4*Dy-x^2*y*Dy+2*y^3+x^2
ideal BLD = imap(B,LD);
NF(LD,std(BLD));
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0
↳ _[4]=0
↳ _[5]=0
```

See also: Section 7.5.5.1 [annPoly], page 415; Section 7.5.5.2 [annRat], page 416.

### 7.5.14.11 dmodGeneralAssumptionCheck

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `dmodGeneralAssumptionCheck();`

**Return:** nothing, but checks general assumptions on the basering

**Note:** This procedure checks the following conditions on the basering  $R$  and prints an error message if any of them is violated:

- $R$  is the  $n$ -th Weyl algebra over a field of characteristic 0,
- $R$  is not a qring,
- for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) * \text{var}(i) = \text{var}(i) * \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Example:**

```
LIB "dmodloc.lib";
ring r = 0, (x,D), dp;
dmodGeneralAssumptionCheck(); // prints error message
↳ ? Basing is not a Weyl algebra
↳ ? leaving dmodloc.lib::dmodGeneralAssumptionCheck (0)
def W = Weyl();
setring W;
dmodGeneralAssumptionCheck(); // returns nothing
```

### 7.5.14.12 extendWeyl

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `extendWeyl(S);`  $S$  string or list of strings

**Assume:** The basering is the  $n$ -th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity

$\text{var}(i+n) * \text{var}(i) = \text{var}(i) * \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Return:** ring, Weyl algebra extended by vars given by  $S$

**Example:**

```
LIB "dmodloc.lib";
ring @D2 = 0, (x,y,Dx,Dy), dp;
def D2 = Weyl();
setring D2;
def D3 = extendWeyl("t");
setring D3; D3;
↳ // coefficients: QQ
↳ // number of vars : 6
↳ // block 1 : ordering dp
↳ // : names t x y Dt Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dtt=t*Dt+1
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
list L = "u","v";
```

```

def D5 = extendWeyl(L);
setring D5;
D5;
↳ // coefficients: QQ
↳ // number of vars : 10
↳ // block 1 : ordering dp
↳ // : names u v t x y Du Dv Dt Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Duu=u*Du+1
↳ // Dvv=v*Dv+1
↳ // Dtt=t*Dt+1
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1

```

### 7.5.14.13 polyVars

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `polyVars(f,v)`;  $f$  poly,  $v$  intvec

**Return:** int, 1 if  $f$  contains only variables indexed by  $v$ , 0 otherwise

**Example:**

```

LIB "dmodloc.lib";
ring r = 0,(x,y,z),dp;
poly f = y^2+zy;
intvec v = 1,2;
polyVars(f,v); // does f depend only on x,y?
↳ 0
v = 2,3;
polyVars(f,v); // does f depend only on y,z?
↳ 1

```

### 7.5.14.14 monomialInIdeal

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `monomialInIdeal(I)`;  $I$  ideal

**Return:** ideal consisting of all monomials appearing in generators of ideal

**Example:** `example monomialInIdeal`; shows examples

**Example:**

```

LIB "dmodloc.lib";
ring r = 0,(x,y),dp;
ideal I = x^2+5x^3y^7, x-x^2-6xy;
monomialInIdeal(I);
↳ _[1]=x^3y^7
↳ _[2]=x^2
↳ _[3]=xy
↳ _[4]=x

```

### 7.5.14.15 vars2pars

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `vars2pars(v); v intvec`

**Assume:** The basering is commutative.

**Return:** ring with variables specified by `v` converted into parameters

**Example:**

```

LIB "dmodloc.lib";
ring r = 0,(x,y,z,a,b,c),dp;
intvec v = 4,5,6;
def R = vars2pars(v);
setring R;
R;
↳ // coefficients: QQ(a, b, c)
↳ // number of vars : 3
↳ // block 1 : ordering dp
↳ // : names x y z
↳ // block 2 : ordering C
v = 1,2;
def RR = vars2pars(v);
setring RR;
RR;
↳ // coefficients: QQ(a, b, c, x, y)
↳ // number of vars : 1
↳ // block 1 : ordering dp
↳ // : names z
↳ // block 2 : ordering C

```

### 7.5.14.16 minIntRoot2

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `minIntRoot2(L); L list`

**Assume:** `L` is the output of `bFactor`.

**Return:** int, the minimal integer root in a list of roots

**Example:**

```

LIB "dmodloc.lib";
ring r = 0,x,dp;
poly f = x*(x+1)*(x-2)*(x-5/2)*(x+5/2);
list L = bFactor(f);
minIntRoot2(L);
↳ -1

```

See also: Section 7.5.5.24 [`bFactor`], page 435; Section 7.5.14.17 [`maxIntRoot`], page 530; Section 7.5.4.21 [`minIntRoot`], page 412.

### 7.5.14.17 maxIntRoot

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `maxIntRoot(L); L list`

**Assume:** L is the output of bFactor.

**Return:** int, the maximal integer root in a list of roots

**Example:**

```
LIB "dmodloc.lib";
ring r = 0,x,dp;
poly f = x*(x+1)*(x-2)*(x-5/2)*(x+5/2);
list L = bFactor(f);
maxIntRoot(L);
↪ 2
```

See also: Section 7.5.5.24 [bFactor], page 435; Section 7.5.14.16 [minIntRoot2], page 530.

### 7.5.14.18 dmodAction

Procedure from library `dmodloc.lib` (see Section 7.5.14 [dmodloc.lib], page 519).

**Usage:** `dmodAction(id,f,v)`; id ideal or poly, f poly, v optional intvec

**Assume:** If v is not given, the basering is the n-th Weyl algebra W over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

Otherwise, v is assumed to specify positions of variables, which form a Weyl algebra as a subalgebra of the basering:

If  $\text{size}(v)$  equals  $2 \cdot n$ , then  $\text{bracket}(\text{var}(v[i]), \text{var}(v[j]))$  must equal 1 if and only if j equals  $i+n$ , and 0 otherwise, for all  $1 \leq i, j \leq n$ .

Further, assume that f does not contain any  $D(i)$ .

**Return:** same type as id, the result of the natural D-module action of id on f

**Note:** The assumptions made are not checked.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0,(x,y,z),dp;
poly f = x^2*z - y^3;
def A = annPoly(f);
setring A;
poly f = imap(r,f);
dmodAction(LD,f);
↪ _[1]=0
↪ _[2]=0
↪ _[3]=0
↪ _[4]=0
↪ _[5]=0
↪ _[6]=0
↪ _[7]=0
↪ _[8]=0
↪ _[9]=0
↪ _[10]=0
↪ _[11]=0
↪ _[12]=0
↪ _[13]=0
poly P = y*Dy+3*z*Dz-3;
```

```

dmodAction(P,f);
↳ 0
dmodAction(P[1],f);
↳ -3*y^3

```

### 7.5.14.19 dmodActionRat

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `dmodActionRat(id,w)`; `id` ideal or poly, `f` vector

**Assume:** The basering is the  $n$ -th Weyl algebra  $W$  over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ . Further, assume that  $w$  has exactly two components, second one not 0, and that  $w$  does not contain any  $D(i)$ .

**Return:** same type as `id`, the result of the natural  $D$ -module action of `id` on the rational function  $w[1]/w[2]$

**Example:**

```

LIB "dmodloc.lib";
ring r = 0, (x,y), dp;
poly f = 2*x; poly g = y;
def A = annRat(f,g); setring A;
poly f = imap(r,f); poly g = imap(r,g);
vector v = [f,g]; // represents f/g
// x and y act by multiplication
dmodActionRat(x,v);
↳ _[1]=2*x^2*gen(1)+y*gen(2)
dmodActionRat(y,v);
↳ _[1]=2*x*gen(1)+gen(2)
// Dx and Dy act by partial derivation
dmodActionRat(Dx,v);
↳ _[1]=y*gen(2)+2*gen(1)
dmodActionRat(Dy,v);
↳ _[1]=y^2*gen(2)-2*x*gen(1)
dmodActionRat(x*Dx+y*Dy,v);
↳ _[1]=gen(2)
setring r;
f = 2*x*y; g = x^2 - y^3;
def B = annRat(f,g); setring B;
poly f = imap(r,f); poly g = imap(r,g);
vector v = [f,g];
dmodActionRat(LD,v); // hence LD is indeed the annihilator of f/g
↳ _[1]=gen(2)
↳ _[2]=gen(2)
↳ _[3]=gen(2)
↳ _[4]=gen(2)
↳ _[5]=gen(2)

```

### 7.5.14.20 simplifyRat

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `simplifyRat(v)`;  $v$  vector

**Assume:** Assume that  $v$  has exactly two components, second one not 0.

**Return:** vector, representing simplified rational function  $v[1]/v[2]$

**Note:** Possibly present non-commutative relations of the basering are ignored.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0, (x,y), dp;
vector v = [x2-1,x+1];
simplifyRat(v);
↪ x*gen(1)+gen(2)-gen(1)
simplifyRat(v) - [x-1,1];
↪ 0
```

### 7.5.14.21 addRat

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `addRat(v,w)`;  $v,w$  vectors

**Assume:** Assume that  $v,w$  have exactly two components, second ones not 0.

**Return:** vector, representing rational function  $(v[1]/v[2])+(w[1]/w[2])$

**Note:** Possibly present non-commutative relations of the basering are ignored.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0, (x,y), dp;
vector v = [x,y];
vector w = [y,x];
addRat(v,w);
↪ x2*gen(1)+xy*gen(2)+y2*gen(1)
addRat(v,w) - [x2+y2,xy];
↪ 0
```

### 7.5.14.22 multRat

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `multRat(v,w)`;  $v,w$  vectors

**Assume:** Assume that  $v,w$  have exactly two components, second ones not 0.

**Return:** vector, representing rational function  $(v[1]/v[2])*(w[1]/w[2])$

**Note:** Possibly present non-commutative relations of the basering are ignored.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0, (x,y), dp;
vector v = [x,y];
vector w = [y,x];
multRat(v,w);
↪ gen(2)+gen(1)
multRat(v,w) - [1,1];
↪ 0
```

### 7.5.14.23 diffRat

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `diffRat(v,j)`;  $v$  vector,  $j$  int

**Assume:** Assume that  $v$  has exactly two components, second one not 0.

**Return:** vector, representing rational function derivative of rational function  $(v[1]/v[2])$  w.r.t.  $\text{var}(j)$

**Note:** Possibly present non-commutative relations of the basering are ignored.

**Example:**

```
LIB "dmodloc.lib";
ring r = 0,(x,y),dp;
vector v = [x,y];
diffRat(v,1);
 \mapsto y*gen(2)+gen(1)
diffRat(v,1) - [1,y];
 \mapsto 0
diffRat(v,2);
 \mapsto y2*gen(2)-x*gen(1)
diffRat(v,2) - [-x,y2];
 \mapsto 0
```

### 7.5.14.24 commRing

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `commRing()`;

**Return:** ring, basering without non-commutative relations

**Example:**

```
LIB "dmodloc.lib";
def W = makeWeyl(3);
setring W; W;
 \mapsto // coefficients: QQ
 \mapsto // number of vars : 6
 \mapsto // block 1: ordering dp
 \mapsto // : names x(1) x(2) x(3) D(1) D(2) D(3)
 \mapsto // block 2: ordering C
 \mapsto // noncommutative relations:
 \mapsto // D(1)x(1)=x(1)*D(1)+1
 \mapsto // D(2)x(2)=x(2)*D(2)+1
 \mapsto // D(3)x(3)=x(3)*D(3)+1
def W2 = commRing();
setring W2; W2;
 \mapsto // coefficients: QQ
 \mapsto // number of vars : 6
 \mapsto // block 1: ordering dp
 \mapsto // : names x(1) x(2) x(3) D(1) D(2) D(3)
 \mapsto // block 2: ordering C
ring r = 0,(x,y),dp;
def r2 = commRing(); // same as r
setring r2; r2;
```



```

↳ // coefficients: QQ
↳ // number of vars : 2
↳ // block 1 : ordering dp
↳ // : names x y
↳ // block 2 : ordering C

```

### 7.5.14.25 rightNFWeyl

Procedure from library `dmodloc.lib` (see Section 7.5.14 [`dmodloc.lib`], page 519).

**Usage:** `rightNFWeyl(id,k)`; `id` ideal or poly, `k` int

**Assume:** The basering is the  $n$ -th Weyl algebra over a field of characteristic 0 and for all  $1 \leq i \leq n$  the identity  $\text{var}(i+n) \cdot \text{var}(i) = \text{var}(i) \cdot \text{var}(i+1) + 1$  holds, i.e. the sequence of variables is given by  $x(1), \dots, x(n), D(1), \dots, D(n)$ , where  $D(i)$  is the differential operator belonging to  $x(i)$ .

**Return:** same type as `id`, the right normal form of `id` with respect to the principal right ideal generated by the  $k$ -th variable

**Note:** No Groebner basis computation is used.

**Example:**

```

LIB "dmodloc.lib";
ring r = 0, (x,y,Dx,Dy), dp;
def W = Weyl();
setring W;
ideal I = x^3*Dx^3, y^2*Dy^2, x*Dy, y*Dx;
rightNFWeyl(I,1); // right NF wrt principal right ideal x*W
↳ _[1]=0
↳ _[2]=y^2*Dy^2
↳ _[3]=0
↳ _[4]=y*Dx
rightNFWeyl(I,3); // right NF wrt principal right ideal Dx*W
↳ _[1]=-6
↳ _[2]=y^2*Dy^2
↳ _[3]=x*Dy
↳ _[4]=0
rightNFWeyl(I,2); // right NF wrt principal right ideal y*W
↳ _[1]=x^3*Dx^3
↳ _[2]=0
↳ _[3]=x*Dy
↳ _[4]=0
rightNFWeyl(I,4); // right NF wrt principal right ideal Dy*W
↳ _[1]=x^3*Dx^3
↳ _[2]=2
↳ _[3]=0
↳ _[4]=y*Dx
poly p = x*Dx+1;
rightNFWeyl(p,1); // right NF wrt principal right ideal x*W
↳ 1

```

### 7.5.15 ncfrac.lib

**Library:** `ncfrac.lib`

**Purpose:** object-oriented interface for `olga.lib`

**Author:** Johannes Hoffmann, email: johannes.hoffmann at math.rwth-aachen.de

**Overview:** This library introduces a new type: `ncfrac`.  
 This type wraps the data defining a (non-commutative) fraction in an Ore localization of a G-algebra as in `olga.lib`.  
 An element of type `ncfrac` has five members:  
 - `polys lnum, lden, rnum, rden`  
 - `ncloc loc`

**Operations:**

`string(ncfrac);`  
 give a string representation of the data describing the fraction  
`print(ncfrac);`  
 prints the string representation of the fraction  
`status(ncfrac);`  
 report on the status/validity of the fraction  
`test(ncfrac);`  
 check if the fraction is valid

**Infix operations:**

`ncfrac == ncfrac;`  
 compare two fractions  
`ncfrac != ncfrac;`  
 compare two fractions  
`ncfrac + ncfrac;`  
 add two fractions  
`ncfrac - ncfrac`  
 subtract two fractions  
`ncfrac * ncfrac`  
 multiply two fractions  
`ncfrac / ncfrac`  
 divide two fractions  
`ncfrac = int/number/poly`  
 create a fraction with:  
 - left and right denominator equal to 1  
 - left and right numerator determined by the input - localization data describing the trivial monoidal localization at 1  
`ncfrac = vector`  
 create a fraction from a vector `v` with unspecified localization such that  
`lden,lnum,rnum,rden = v[1],v[2],v[3],v[4]`  
 (note: without specifying a localization afterwards this results is an invalid fraction)  
`ncfrac = list`  
 create a fraction from a list `L` as follows: - try to create a fraction from `L[1]` as above  
 - if `L[2]` is of type `ncloc` set the localization of the fraction to `L[2]`

**Procedures:**

### 7.5.15.1 `hasLeftDenom`

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `hasLeftDenom(frac), ncfrac frac`

**Purpose:** checks if `frac` has a left representation

**Return:** int, 1 if frac has a left representation, 0 otherwise

**Example:**

```
LIB "ncfrac.lib";
⇨ // ** redefining testNcfrac (LIB "ncfrac.lib"); ./examples/hasLeftDenom.s\
 ing:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib"); ncfrac.lib::mod_init:11\
 3
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S;
ncloc loc = ideal(x-3,y+7);
ncfrac noLeft = list([0,0,3*y*Dx,x+2], loc);
hasLeftDenom(noLeft);
⇨ 0
ncfrac left = list([1,Dx,Dx,1], loc);
hasLeftDenom(left);
⇨ 1
```

### 7.5.15.2 hasRightDenom

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `hasRightDenom(frac)`, `ncfrac frac`

**Purpose:** checks if frac has a right representation

**Return:** int, 1 if frac has a right representation, 0 otherwise

**Example:**

```
LIB "ncfrac.lib";
⇨ // ** redefining testNcfrac (LIB "ncfrac.lib"); ./examples/hasRightDenom.\
 sing:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib"); ncfrac.lib::mod_init:11\
 3
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S;
ncloc loc = ideal(x-3,y+7);
ncfrac noRight = list([x+2,3*y*Dx,0,0], loc);
hasRightDenom(noRight);
⇨ 0
ncfrac right = list([1,Dx,Dx,1], loc);
hasRightDenom(right);
⇨ 1
```

### 7.5.15.3 isZeroNcfrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `isZeroNcfrac(frac)`, `ncfrac frac`

**Purpose:** checks if frac is zero

**Return:** int, 1 if frac is zero, 0 otherwise

**Example:**

```

LIB "ncfrac.lib";
⇨ // ** redefining testNcfrac (LIB "ncfrac.lib");) ./examples/isZeroNcfrac.s\
 ing:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib");) ncfrac.lib::mod_init:11\
 3
ring Q = (0,q),(x,y,Qx,Qy),dp;
matrix C[4][4] = UpOneMatrix(4);
C[1,3] = q;
C[2,4] = q;
def ncQ = nc_algebra(C,0);
setring ncQ;
ncloc loc = intvec(2);
ncfrac frac = list([y^2+7*y+1,0,0,0], loc);
isZeroNcfrac(frac);
⇨ 1
frac.lnum = 42*y*Qy+7*Qx+3*x+7;
isZeroNcfrac(frac);
⇨ 0

```

#### 7.5.15.4 isOneNcfrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `isOneNcfrac(frac), ncfrac frac`

**Purpose:** checks if `frac` is one

**Return:** `int`, 1 if `frac` is one, 0 otherwise

**Example:**

```

LIB "ncfrac.lib";
⇨ // ** redefining testNcfrac (LIB "ncfrac.lib");) ./examples/isOneNcfrac.si\
 ng:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib");) ncfrac.lib::mod_init:11\
 3
ring Q = (0,q),(x,y,Qx,Qy),dp;
matrix C[4][4] = UpOneMatrix(4);
C[1,3] = q;
C[2,4] = q;
def ncQ = nc_algebra(C,0);
setring ncQ;
ncloc loc = intvec(2);
ncfrac frac = list([y^2+7*y+1,y^2+7*y+1,0,0], loc);
isOneNcfrac(frac);
⇨ 1
frac.lnum = 42*y*Qy+7*Qx+3*x+7;
isOneNcfrac(frac);
⇨ 0

```

#### 7.5.15.5 zeroNcfrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `zeroNcfrac(loc), ncloc loc`

**Purpose:** returns the zero fraction in the localization `loc`

**Return:** ncffrac

**Example:**

```
LIB "ncfrac.lib";
⇨ // ** redefining testNcffrac (LIB "ncfrac.lib");) ./examples/zeroNcffrac.sin\
 g:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib");) ncffrac.lib::mod_init:11\
 3
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S;
ncloc loc = ideal(x-53,y-7);
zeroNcffrac(loc);
⇨ left repr.: (1,0)
⇨ right repr.: (0,1)
⇨
```

### 7.5.15.6 oneNcffrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** oneNcffrac(`loc`), `ncloc loc`

**Purpose:** returns the one fraction in the localization `loc`

**Return:** ncffrac

**Example:**

```
LIB "ncfrac.lib";
⇨ // ** redefining testNcffrac (LIB "ncfrac.lib");) ./examples/oneNcffrac.sing\
 :1
⇨ // ** redefining testNcloc (LIB "ncloc.lib");) ncffrac.lib::mod_init:11\
 3
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S;
ncloc loc = ideal(x-42,y-17);
oneNcffrac(loc);
⇨ left repr.: (1,1)
⇨ right repr.: (1,1)
⇨
```

### 7.5.15.7 ensureLeftNcffrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** ensureLeftNcffrac(`frac`), `ncfrac frac`

**Purpose:** ensures that `frac` has a left representation (by computing it if not already known)

**Return:** `ncfrac`, a representation of `frac` which has a left representation

**Example:**

```
LIB "ncfrac.lib";
⇨ // ** redefining testNcffrac (LIB "ncfrac.lib");) ./examples/ensureLeftNcfr\
 ac.sing:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib");) ncffrac.lib::mod_init:11\
```

```

3
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x y Dx Dy
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // Dxx=x*Dx+1
⇨ // Dyy=y*Dy+1
// monoidal localization
poly g1 = x+3;
poly g2 = x*y;
list L = g1,g2;
ncloc loc0 = L;
poly g = g1^2*g2;
poly f = Dx;
ncfrac frac0 = [0,0,f,g];
frac0.loc = loc0;
ncfrac rm = ensureLeftNcfrac(frac0);
print(rm);
⇨ left repr.: (x^8*y^4+12*x^7*y^4+54*x^6*y^4+108*x^5*y^4+81*x^4*y^4,x^5*y^3\
 *Dx+6*x^4*y^3*Dx-3*x^4*y^3+9*x^3*y^3*Dx-12*x^3*y^3-9*x^2*y^3)
⇨ right repr.: (Dx,x^3*y+6*x^2*y+9*x*y)
rm.lnum*g-rm.lden*f;
⇨ 0
// geometric localization
ncloc loc1 = ideal(x-1,y-3);
f = Dx;
g = x^2+y;
ncfrac frac1 = [0,0,f,g];
frac1.loc = loc1;
ncfrac rg = ensureLeftNcfrac(frac1);
print(rg);
⇨ left repr.: (x^4+2*x^2*y+y^2,x^2*Dx+y*Dx-2*x)
⇨ right repr.: (Dx,x^2+y)
rg.lnum*g-rg.lden*f;
⇨ 0
// rational localization
intvec rat = 1;
ncloc loc2 = rat;
f = Dx+Dy;
g = x;
ncfrac frac2 = [0,0,f,g];
frac2.loc = loc2;
ncfrac rr = ensureLeftNcfrac(frac2);
print(rr);
⇨ left repr.: (x^2,x*Dx+x*Dy-1)
⇨ right repr.: (Dx+Dy,x)
rr.lnum*g-rr.lden*f;
⇨ 0

```

### 7.5.15.8 ensureRightNcfrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `ensureLeftNcfrac(frac), ncfrac frac`

**Purpose:** ensures that `frac` has a right representation (by computing it if not already known)

**Return:** `ncfrac`, a representation of `frac` which has a right representation

**Example:**

```
LIB "ncfrac.lib";
⇨ // ** redefining testNcfrac (LIB "ncfrac.lib"); ./examples/ensureRightNcf\
 rac.sing:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib"); ncfrac.lib::mod_init:11\
 3
ring R = 0,(x,y,Dx,Dy),dp;
def S = Weyl();
setring S; S;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x y Dx Dy
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // Dxx=x*Dx+1
⇨ // Dyy=y*Dy+1
// monoidal localization
poly g = x;
poly f = Dx;
ncloc loc0 = g;
ncfrac frac0 = [g,f,0,0];
frac0.loc = loc0;
ncfrac rm = ensureRightNcfrac(frac0);
print(rm);
⇨ left repr.: (x,Dx)
⇨ right repr.: (x*Dx+2,x^2)
f*rm.rden-g*rm.rnum;
⇨ 0
// geometric localization
g = x+y;
f = Dx+Dy;
ncloc loc1 = ideal(x-1,y-3);
ncfrac frac1 = [g,f,0,0];
frac1.loc = loc1;
ncfrac rg = ensureRightNcfrac(frac1);
print(rg);
⇨ left repr.: (x+y,Dx+Dy)
⇨ right repr.: (x*Dx+y*Dx+x*Dy+y*Dy+4,x^2+2*x*y+y^2)
f*rg.rden-g*rg.rnum;
⇨ 0
// rational localization
intvec rat = 1;
f = Dx+Dy;
g = x;
```

```

ncloc loc2 = rat;
ncfrac frac2 = [g,f,0,0];
frac2.loc = loc2;
ncfrac rr = ensureRightNcfrac(frac2);
print(rr);
↳ left repr.: (x,Dx+Dy)
↳ right repr.: (x*Dx+x*Dy+2,x^2)
f*rr.rden-g*rr.rnum;
↳ 0

```

### 7.5.15.9 negateNcfrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `negateNcfrac(frac), ncfrac frac`

**Purpose:** compute the negative (i.e. additive inverse) of `frac`

**Return:** `ncfrac`

**Note:** returns  $(-1)*frac$

**Example:**

```

LIB "ncfrac.lib";
↳ // ** redefining testNcfrac (LIB "ncfrac.lib"); ./examples/negateNcfrac.s\
 ing:1
↳ // ** redefining testNcloc (LIB "ncloc.lib"); ncfrac.lib::mod_init:11\
 3
ring R = 0,(x,y,Dx,Dy),dp;
def S = Weyl();
setring S;
poly g = x*y^2+4*x+7*y-98;
ncloc loc = g;
ncfrac frac = list([g, 13*x^2], loc);
frac;
↳ left repr.: (x*y^2+4*x+7*y-98,13*x^2)
↳ right repr.: (0,0)
↳
ncfrac negFrac = negateNcfrac(frac);
negFrac;
↳ left repr.: (x*y^2+4*x+7*y-98,-13*x^2)
↳ right repr.: (0,0)
↳
frac + negFrac;
↳ left repr.: (x*y^2+4*x+7*y-98,0)
↳ right repr.: (0,0)
↳

```

### 7.5.15.10 isInvertibleNcfrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `isInvertibleNcfrac(frac), ncfrac frac`

**Purpose:** checks if `frac` is invertible

**Return:** `int`, 1 if `frac` is invertible, 0 otherwise



**Example:**

```

LIB "ncfrac.lib";
⇨ // ** redefining testNcfrac (LIB "ncfrac.lib";) ./examples/isInvertibleNc\
 frac.sing:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib";) ncfrac.lib::mod_init:11\
 3
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S;
ncloc loc = intvec(2);
ncfrac frac = list([y,y+1,0,0], loc);
isInvertibleNcfrac(frac);
⇨ 1
frac = list([y,x+1,0,0], loc);
isInvertibleNcfrac(frac);
⇨ 0

```

**7.5.15.11 invertNcfrac**

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `invertNcfrac(frac), ncfrac frac`

**Purpose:** compute the inverse of `frac`

**Return:** `ncfrac`

**Note:** returns the zero fraction if `frac` is not invertible

**Example:**

```

LIB "ncfrac.lib";
⇨ // ** redefining testNcfrac (LIB "ncfrac.lib";) ./examples/invertNcfrac.s\
 ing:1
⇨ // ** redefining testNcloc (LIB "ncloc.lib";) ncfrac.lib::mod_init:11\
 3
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S;
ncloc loc = intvec(2);
ncfrac frac1 = list([y,y+1,0,0], loc);
// frac1 is invertible
ncfrac inv = invertNcfrac(frac1);
inv;
⇨ left repr.: (y+1,y)
⇨ right repr.: (0,0)
⇨
ncfrac frac2 = list([y,x+1,0,0], loc);
// frac2 is not invertible
inv = invertNcfrac(frac2);
inv;
⇨ left repr.: (1,0)
⇨ right repr.: (0,1)
⇨

```

### 7.5.15.12 testNcfrac

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `testNcfrac()`

**Purpose:** execute a series of internal testing procedures

**Return:** nothing

**Note:**

### 7.5.15.13 testNcfracExamples

Procedure from library `ncfrac.lib` (see Section 7.5.15 [`ncfrac.lib`], page 535).

**Usage:** `testNcfracExamples()`

**Purpose:** execute the examples of all procedures in this library

**Return:** nothing

**Note:**

## 7.5.16 nchomolog\_lib

Status: experimental

**Library:** `nchomolog.lib`

**Purpose:** Procedures for Noncommutative Homological Algebra

**Authors:** Viktor Levandovskyy [levandov@math.rwth-aachen.de](mailto:levandov@math.rwth-aachen.de),  
Christian Schilli, [christian.schilli@rwth-aachen.de](mailto:christian.schilli@rwth-aachen.de),  
Gerhard Pfister, [pfister@mathematik.uni-kl.de](mailto:pfister@mathematik.uni-kl.de)

**Overview:** In this library we present tools of homological algebra for finitely presented modules over GR-algebras.

**Procedures:**

### 7.5.16.1 ncExt\_R

Procedure from library `nchomolog.lib` (see Section 7.5.16 [`nchomolog.lib`], page 544).

**Usage:** `ncExt_R(i, M)`;  $i$  int,  $M$  module

**Compute:** a presentation of  $\text{Ext}^i(M', R)$ ; for  $M' = \text{coker}(M)$ .

**Return:** right module `Ext`, a presentation of  $\text{Ext}^i(M', R)$

**Example:**

```
LIB "nchomolog.lib";
ring R = 0, (x, y), dp;
poly F = x^2 - y^2;
def A = annfs(F); setring A; // A is the 2nd Weyl algebra
matrix M[1][size(LD)] = LD; // ideal
print(M);
↳ y*Dx + x*Dy, x*Dx + y*Dy + 2, x^2*Dy - y^2*Dy - 2*y
print(ncExt_R(1, M)); // hence the Ext^1 is zero
↳ 1, 0,
```

```

↳ 0,1
module E = ncExt_R(2,M); // define the right module E
print(E); // E is in the opposite algebra
↳ 1, -x, -y,
↳ Dx,y*Dy+1,x*Dy
def Aop = opposite(A); setring Aop;
module Eop = oppose(A,E);
module T1 = ncExt_R(2,Eop);
setring A;
module T1 = oppose(Aop,T1);
print(T1); // this is a left module Ext^2(Ext^2(M,A),A)
↳ y*Dx+x*Dy,x*Dx+y*Dy+2,x^2*Dy-y^2*Dy-2*y
print(M); // it is known that M holonomic implies Ext^2(Ext^2(M,A),A) iso to M
↳ y*Dx+x*Dy,x*Dx+y*Dy+2,x^2*Dy-y^2*Dy-2*y

```

### 7.5.16.2 ncHom

Procedure from library `nchomolog.lib` (see Section 7.5.16 [`nchomolog.lib`], page 544).

**Usage:** `ncHom(M,N)`; M,N modules

**Compute:** A presentation of  $\text{Hom}(M',N')$ ,  $M'=\text{coker}(M)$ ,  $N'=\text{coker}(N)$

**Assume:**  $M'$  is a left module,  $N'$  is a centralizing bimodule

**Note:** `ncHom(M,N)` is a right module, hence a right presentation matrix is returned

**Example:**

```

LIB "nchomolog.lib";
ring A=0,(x,y,z),dp;
matrix M[3][3]=1,2,3,
4,5,6,
7,8,9;
matrix N[2][2]=x,y,
z,0;
module H = ncHom(M,N);
print(H);
↳ 0,0,0,0,y,x,
↳ 0,0,0,z,0,z,
↳ 1,0,0,0,0,0,
↳ 0,1,0,0,0,0,
↳ 0,0,2,1,0,0,
↳ 0,0,1,0,0,0

```

### 7.5.16.3 coHom

Procedure from library `nchomolog.lib` (see Section 7.5.16 [`nchomolog.lib`], page 544).

**Usage:** `coHom(A,k)`; A matrix, k int

**Purpose:** compute the matrix of a homomorphism  $\text{Hom}(R^k,A)$ , where R is the basering. Let A be a matrix defining a map  $F1 \rightarrow F2$  of free R-modules, then the matrix of  $\text{Hom}(R^k,F1) \rightarrow \text{Hom}(R^k,F2)$  is computed.

**Note:** Both A and  $\text{Hom}(A,R^k)$  are matrices for either left or right R-module homomorphisms

**Example:**

```

LIB "nchomolog.lib";
ring A=0,(x,y,z),dp;
matrix M[3][3]=1,2,3,
4,5,6,
7,8,9;
module cM = coHom(M,2);
print(cM);
 ↪ 1,0,2,0,3,0,
 ↪ 0,1,0,2,0,3,
 ↪ 4,0,5,0,6,0,
 ↪ 0,4,0,5,0,6,
 ↪ 7,0,8,0,9,0,
 ↪ 0,7,0,8,0,9

```

#### 7.5.16.4 contraHom

Procedure from library `nchomolog.lib` (see Section 7.5.16 [`nchomolog.lib`], page 544).

**Usage:** `contraHom(A,k)`; A matrix, k int

**Return:** matrix

**Purpose:** compute the matrix of a homomorphism  $\text{Hom}(A, R^k)$ , where R is the basering. Let A be a matrix defining a map  $F_1 \rightarrow F_2$  of free R-modules, then the matrix of  $\text{Hom}(F_2, R^k) \rightarrow \text{Hom}(F_1, R^k)$  is computed.

**Note:** if A is matrix of a left (resp. right) R-module homomorphism, then  $\text{Hom}(A, R^k)$  is a right (resp. left) R-module homomorphism

**Example:**

```

LIB "nchomolog.lib";
ring A=0,(x,y,z),dp;
matrix M[3][3]=1,2,3,
4,5,6,
7,8,9;
module cM = contraHom(M,2);
print(cM);
 ↪ 1,4,7,0,0,0,
 ↪ 2,5,8,0,0,0,
 ↪ 3,6,9,0,0,0,
 ↪ 0,0,0,1,4,7,
 ↪ 0,0,0,2,5,8,
 ↪ 0,0,0,3,6,9

```

#### 7.5.16.5 dmodoublext

Procedure from library `nchomolog.lib` (see Section 7.5.16 [`nchomolog.lib`], page 544).

**Usage:** `dmodoublext(M [,i])`; M module, i optional int

**Compute:** a presentation of  $\text{Ext}^i(\text{Ext}^i(M,D),D)$  for basering D

**Return:** left module

**Note:** by default, i is set to the integer part of the half of number of variables of D for holonomic modules over Weyl algebra, the double ext is known to be holonomic left module

**Example:**

```

LIB "nchomolog.lib";
ring R = 0, (x,y), dp;
poly F = x^3-y^2;
def A = annfs(F);
setring A;
dmodoublext(LD);
↳ _[1]=2*x*Dx+3*y*Dy+6
↳ _[2]=3*x^2*Dy+2*y*Dx
↳ _[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
↳ _[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
LD;
↳ LD[1]=2*x*Dx+3*y*Dy+6
↳ LD[2]=3*x^2*Dy+2*y*Dx
↳ LD[3]=9*x*y*Dy^2-4*y*Dx^2+15*x*Dy
↳ LD[4]=27*y^2*Dy^3+8*y*Dx^3+135*y*Dy^2+105*Dy
// fancier example:
setring A;
ideal I = Dx*(x^2-y^3), Dy*(x^2-y^3);
I = groebner(I);
print(dmodoublext(I,1));
↳ y^3-x^2
print(dmodoublext(I,2));
↳ Dy,
↳ Dx

```

**7.5.16.6 is\_cenBimodule**

Procedure from library `nchomolog.lib` (see Section 7.5.16 [`nchomolog.lib`], page 544).

**Usage:** `is_cenBimodule(M)`;  $M$  module

**Compute:** 1, if a module, presented by  $M$  can be centralizing in the sense of Artin and 0 otherwise

**Note:** only one condition for centralizing factor module can be checked algorithmically

**Example:**

```

LIB "nchomolog.lib";
def A = makeUs12(); setring A;
poly p = 4*e*f + h^2-2*h; // generator of the center
matrix M[2][2] = p, p^2-7, 0, p*(p+1);
is_cenBimodule(M); // M is centralizing
↳ 1
matrix N[2][2] = p, e*f, h, p*(p+1);
is_cenBimodule(N); // N is not centralizing
↳ 0

```

**7.5.16.7 is\_cenSubbimodule**

Procedure from library `nchomolog.lib` (see Section 7.5.16 [`nchomolog.lib`], page 544).

**Usage:** `is_cenSubbimodule(M)`;  $M$  module

**Compute:** 1, if a subbimodule, generated by the columns of  $M$  is centralizing in the sense of Artin and 0 otherwise

**Example:**

```

LIB "nchomolog.lib";
def A = makeUs12(); setring A;
poly p = 4*e*f + h^2-2*h; // generator of the center
matrix M[2][2] = p, p^2-7,0,p*(p+1);
is_cenSubbimodule(M); // M is centralizing subbimodule
 ↪ 1
matrix N[2][2] = p, e*f,h,p*(p+1);
is_cenSubbimodule(N); // N is not centralizing subbimodule
 ↪ 0

```

**7.5.17 ncloc\_lib****Library:** ncloc.lib**Purpose:** Ore-localization in G-Algebras**Author:** Johannes Hoffmann, email: johannes.hoffmann at math.rwth-aachen.de

**Overview:** This library introduces a new type: ncloc.  
 This type wraps the localization data defined as in olga.lib. An element of type ncloc has two members:

- int locType
- def locData

**Operations:**

```

string(ncloc);
give a string representation of the data describing the localization print(ncloc);
prints the string representation of the localization status(ncloc);
report on the status/validity of the localization test(ncloc);
check if the localization is valid

```

**Infix operations:**

```

ncloc == ncloc;
compare two localizations
ncloc != ncloc;
compare two localizations
ncloc = list/poly
create a monoidal localization from the given data ncloc = ideal
create a geometric localization from the given data ncloc = intvec
create a rational localization from the given data

```

**Procedures:****7.5.17.1 isDenom**

Procedure from library `ncloc.lib` (see Section 7.5.17 [`ncloc_lib`], page 548).

**Usage:** isDenom(p, loc), poly a, ncloc loc**Purpose:** check if p is a valid denominator in the localization loc**Return:** int**Note:** returns 1 or 0, depending whether p is a valid denominator**Example:**

```

LIB "ncloc.lib";
⇨ // ** redefining testNcloc (LIB "ncloc.lib"); ./examples/isDenom.sing:1
ring R = 0,(x,y,Dx,Dy),dp;
def S = Weyl();
setring S; S;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x y Dx Dy
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // Dxx=x*Dx+1
⇨ // Dyy=y*Dy+1
// monoidal localization
ncloc loc;
poly g1 = x^2*y+x+2;
poly g2 = y^3+x*y;
list L = g1,g2;
loc = L;
poly g = g1^2*g2;
poly f = g - 1;
isDenom(g, loc);
⇨ 1
isDenom(f, loc);
⇨ 0
// geometrical localization
loc = ideal(x-1,y-3);
g = x^2+y-3;
f = (x-1)*g;
isDenom(g, loc);
⇨ 1
isDenom(f, loc);
⇨ 0
// rational localization
intvec v = 2;
loc = v;
g = y^5+17*y^2-4;
f = x*y;
isDenom(g, loc);
⇨ 1
isDenom(f, loc);
⇨ 0

```

### 7.5.17.2 testNcloc

Procedure from library `ncloc.lib` (see Section 7.5.17 [`ncloc.lib`], page 548).

**Usage:**     `testNcloc()`

**Purpose:**    execute a series of internal testing procedures

**Return:**   nothing

**Note:**

### 7.5.17.3 testNclocExamples

Procedure from library `ncloc.lib` (see Section 7.5.17 [`ncloc.lib`], page 548).

**Usage:** `testNclocExamples()`

**Purpose:** execute the examples of all procedures in this library

**Return:** nothing

**Note:**

### 7.5.18 ncModslimgb\_lib

**Library:** `ncModslimgb.lib`

**Purpose:** A library for computing Groebner bases over G-algebras defined over the rationals using modular techniques.

**Authors:** Wolfram Decker, Christian Eder, Viktor Levandovskyy, and Sharwan K. Tiwari [shrawant@gmail.com](mailto:shrawant@gmail.com)

**References:**

Wolfram Decker, Christian Eder, Viktor Levandovskyy, and Sharwan K. Tiwari, Modular Techniques For Noncommutative Groebner Bases, <https://link.springer.com/article/10.1007/s11786-019-00412-9> and <https://arxiv.org/abs/1704.02852>.

E. A. Arnold, Modular algorithms for computing Groebner bases. *Journal of Symbolic Computation* 35, 403-419 (2003).

N. Idrees, G. Pfister, S. Steidel, Parallelization of Modular Algorithms, *Journal of Symbolic Computation* 46, 672-684 (2011).

**Procedures:**

#### 7.5.18.1 ncmoDSLIMGB

Procedure from library `ncModslimgb.lib` (see Section 7.5.18 [`ncModslimgb.lib`], page 550).

**Usage:** `ncmoDSLIMGB(I[, exactness, ncores]);` I ideal, optional integers exactness and n(umber of )cores

**Return:** ideal

**Purpose:** compute a left Groebner basis of I by modular approach

**Assume:** basering is a G-algebra; base field is prime field Q of rationals.

**Note:** - If the given algebra and ideal are graded (it is not checked by this command), then the computed Groebner basis will be exact. Otherwise, the result will be correct with a very high probability. - The optional parameter 'exactness' justifies, whether the final (expensive) verification step will be performed or not (exactness=0, default value is 1). - The optional parameter 'ncores' (default value is 1) provides an integer to use the number of cores (this must not exceed the number of available cores in the computing machine).

**Example:**





Two, even if the subset of variables in question generates an admissible subalgebra, there might be no admissible elimination ordering, i.e. an elimination ordering which also satisfies the ordering condition for G-algebras.

The difference between the procedure `eliminateNC` provided in this library and the procedure `eliminate (plural)` from the kernel is that `eliminateNC` will always find an admissible elimination if such one exists. Moreover, the use of `slimgb` for performing Groebner basis computations is possible.

As an application of the theory of elimination, the procedure `preimageNC` is provided, which computes the preimage of an ideal under a homomorphism  $f: A \rightarrow B$  between G-algebras A and B. In contrast to the kernel procedure `preimage (plural)`, the assumption that A is commutative is not required.

#### References:

- (BGL) J.L. Bueso, J. Gomez-Torrecillas, F.J. Lobillo: ‘Re-filtering and exactness of the Gelfand-Kirillov dimension’, Bull. Sci. math. 125, 8, 689-715, 2001.  
 (GML) J.I. Garcia Garcia, J. Garcia Miranda, F.J. Lobillo: ‘Elimination orderings and localization in PBW algebras’, Linear Algebra and its Applications 430(8-9), 2133-2148, 2009.  
 (Lev) V. Levandovskyy: ‘Intersection of ideals with non-commutative subalgebras’, ISSAC’06, 212-219, ACM, 2006.

**Procedures:** See also: Section D.4.6 [`elim_lib`], page 812; Section 7.3.21 [`preimage (plural)`], page 349.

#### 7.5.19.1 `eliminateNC`

Procedure from library `ncpreim.lib` (see Section 7.5.19 [`ncpreim.lib`], page 551).

**Usage:** `eliminateNC(I,v,eng)`; I ideal, v intvec, eng optional int

**Return:** ideal, I intersected with the subring defined by the variables not index by the entries of v

**Assume:** The entries of v are in the range  $1..nvars(basering)$  and the corresponding variables generate an admissible subalgebra.

**Remarks:** In order to determine the required elimination ordering, a linear programming problem is solved with the simplex algorithm.

Reference: (GML)

Unlike `eliminate`, this procedure will always find an elimination ordering, if such exists.

**Note:** If `eng<>0`, `std` is used for Groebner basis computations, otherwise (and by default) `slimgb` is used.

If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

#### Example:

```
LIB "ncpreim.lib";
// (Lev): Example 2
ring r = 0, (a,b,x,d), Dp;
matrix D[4][4];
D[1,2] = 3*a; D[1,4] = 3*x^2;
D[2,3] = -x; D[2,4] = d; D[3,4] = 1;
def A = nc_algebra(1,D);
setring A; A;
```

```

⇒ // coefficients: QQ
⇒ // number of vars : 4
⇒ // block 1 : ordering Dp
⇒ // : names a b x d
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // ba=ab+3a
⇒ // da=ad+3x2
⇒ // xb=bx-x
⇒ // db=bd+d
⇒ // dx=xd+1
ideal I = a,x;
// Since d*a-a*d = 3*x^2, any admissible ordering has to satisfy
// x^2 < a*d, while any elimination ordering for {x,d} additionally
// has to fulfil a << x and a << d.
// Hence, the weight (0,0,1,1) is not an elimination weight for
// (x,d) and the call eliminate(I,x*d); will produce an error.
eliminateNC(I,3..4);
⇒ _[1]=a
// This call uses the elimination weight (0,0,1,2), which works.

```

See also: Section 7.3.5 [eliminate (plural)], page 333.

### 7.5.19.2 preimageNC

Procedure from library `ncpreim.lib` (see Section 7.5.19 [ncpreim.lib], page 551).

**Usage:** `preimageNC(A,f,J[,P,eng]);` A ring, f map or ideal, J ideal, P optional string, eng optional int

**Assume:** f defines a map from A to the basering.

**Return:** nothing, instead exports an object 'preim' of type ideal to ring A, being the preimage of J under f.

**Note:** If P is given and not equal to the empty string, the preimage is exported to A under the name specified by P.

Otherwise (and by default), P is set to 'preim'.

If `eng<0`, `std` is used for Groebner basis computations, otherwise (and by default) `slingb` is used.

If `printlevel=1`, progress debug messages will be printed, if `printlevel>=2`, all the debug messages will be printed.

**Remark:** Reference: (Lev)

**Example:**

```

LIB "ncpreim.lib";
def A = makeUgl(3); setring A; A; // universal enveloping algebra of gl_3
⇒ // coefficients: QQ
⇒ // number of vars : 9
⇒ // block 1 : ordering dp
⇒ // : names e_1_1 e_1_2 e_1_3 e_2_1 e_2_2 e_2_3 e_3_1 \
e_3_2 e_3_3
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // e_1_2e_1_1=e_1_1*e_1_2-e_1_2

```

```

⇒ // e_1_3e_1_1=e_1_1*e_1_3-e_1_3
⇒ // e_2_1e_1_1=e_1_1*e_2_1+e_2_1
⇒ // e_3_1e_1_1=e_1_1*e_3_1+e_3_1
⇒ // e_2_1e_1_2=e_1_2*e_2_1-e_1_1+e_2_2
⇒ // e_2_2e_1_2=e_1_2*e_2_2-e_1_2
⇒ // e_2_3e_1_2=e_1_2*e_2_3-e_1_3
⇒ // e_3_1e_1_2=e_1_2*e_3_1+e_3_2
⇒ // e_2_1e_1_3=e_1_3*e_2_1+e_2_3
⇒ // e_3_1e_1_3=e_1_3*e_3_1-e_1_1+e_3_3
⇒ // e_3_2e_1_3=e_1_3*e_3_2-e_1_2
⇒ // e_3_3e_1_3=e_1_3*e_3_3-e_1_3
⇒ // e_2_2e_2_1=e_2_1*e_2_2+e_2_1
⇒ // e_3_2e_2_1=e_2_1*e_3_2+e_3_1
⇒ // e_2_3e_2_2=e_2_2*e_2_3-e_2_3
⇒ // e_3_2e_2_2=e_2_2*e_3_2+e_3_2
⇒ // e_3_1e_2_3=e_2_3*e_3_1-e_2_1
⇒ // e_3_2e_2_3=e_2_3*e_3_2-e_2_2+e_3_3
⇒ // e_3_3e_2_3=e_2_3*e_3_3-e_2_3
⇒ // e_3_3e_3_1=e_3_1*e_3_3+e_3_1
⇒ // e_3_3e_3_2=e_3_2*e_3_3+e_3_2
ring r3 = 0, (x,y,z,Dx,Dy,Dz), dp;
def B = Weyl(); setring B; B; // third Weyl algebra
⇒ // coefficients: QQ
⇒ // number of vars : 6
⇒ // block 1 : ordering dp
⇒ // : names x y z Dx Dy Dz
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Dxx=x*Dx+1
⇒ // Dyy=y*Dy+1
⇒ // Dzz=z*Dz+1
ideal ff = x*Dx,x*Dy,x*Dz,y*Dx,y*Dy,y*Dz,z*Dx,z*Dy,z*Dz;
map f = A,ff; // f: A -> B, e(i,j) |-> x(i)D(j)
ideal J = 0;
preimageNC(A,f,J,"K"); // compute K := ker(f)
setring A;
K;
⇒ K[1]=e_2_3*e_3_2-e_2_2*e_3_3-e_2_2
⇒ K[2]=e_1_3*e_3_2-e_1_2*e_3_3-e_1_2
⇒ K[3]=e_2_3*e_3_1-e_2_1*e_3_3-e_2_1
⇒ K[4]=e_2_2*e_3_1-e_2_1*e_3_2
⇒ K[5]=e_1_3*e_3_1-e_1_1*e_3_3-e_1_1
⇒ K[6]=e_1_2*e_3_1-e_1_1*e_3_2
⇒ K[7]=e_1_3*e_2_2-e_1_2*e_2_3+e_1_3
⇒ K[8]=e_1_3*e_2_1-e_1_1*e_2_3
⇒ K[9]=e_1_2*e_2_1-e_1_1*e_2_2-e_1_1

```

See also: Section 7.3.21 [preimage (plural)], page 349.

### 7.5.19.3 admissibleSub

Procedure from library `ncpreim.lib` (see Section 7.5.19 [ncpreim.lib], page 551).

**Usage:** `admissibleSub(v); v intvec`

**Assume:** The entries of  $v$  are in the range  $1..nvars(\text{basering})$ .

**Return:** int, 1 if the variables indexed by the entries of  $v$  form an admissible subalgebra, 0 otherwise

**Example:**

```
LIB "ncpreim.lib";
ring r = 0,(e,f,h),dp;
matrix d[3][3];
d[1,2] = -h; d[1,3] = 2*e; d[2,3] = -2*f;
def A = nc_algebra(1,d);
setring A; A; // A is U(sl_2)
↪ // coefficients: QQ
↪ // number of vars : 3
↪ // block 1 : ordering dp
↪ // : names e f h
↪ // block 2 : ordering C
↪ // noncommutative relations:
↪ // fe=ef-h
↪ // he=eh+2e
↪ // hf=fh-2f
// the subalgebra generated by e,f is not admissible since [e,f]=h
admissibleSub(1..2);
↪ 0
// but the subalgebra generated by f,h is admissible since [f,h]=2f
admissibleSub(2..3);
↪ 1
```

#### 7.5.19.4 isUpperTriangular

Procedure from library `ncpreim.lib` (see Section 7.5.19 [`ncpreim.lib`], page 551).

**Usage:** `isUpperTriangular(M[,k])`;  $M$  a matrix,  $k$  an optional int

**Return:** int, 1 if the given matrix is upper triangular,  
0 otherwise.

**Note:** If  $k > 0$  is given, it is checked whether  $M$  is strictly upper triangular.

**Example:**

```
LIB "ncpreim.lib";
ring r = 0,x,dp;
matrix M[2][3] =
0,1,2,
0,0,3;
isUpperTriangular(M);
↪ 1
isUpperTriangular(M,1);
↪ 1
M[2,2] = 4;
isUpperTriangular(M);
↪ 1
isUpperTriangular(M,1);
↪ 0
```

### 7.5.19.5 appendWeight2Ord

Procedure from library `ncpreim.lib` (see Section 7.5.19 [`ncpreim.lib`], page 551).

**Usage:** `appendWeight2Ord(w)`;  $w$  an intvec

**Return:** ring, the basering equipped with the ordering  $(a(w), <)$ , where  $<$  is the ordering of the basering.

**Example:**

```
LIB "ncpreim.lib";
ring r = 0, (a,b,x,d), Dp;
intvec w = 1,2,3,4;
def r2 = appendWeight2Ord(w); // for a commutative ring
r2;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering a
↳ // : names a b x d
↳ // : weights 1 2 3 4
↳ // block 2 : ordering Dp
↳ // : names a b x d
↳ // block 3 : ordering C
matrix D[4][4];
D[1,2] = 3*a; D[1,4] = 3*x^2; D[2,3] = -x;
D[2,4] = d; D[3,4] = 1;
def A = nc_algebra(1,D);
setring A; A;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering Dp
↳ // : names a b x d
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // ba=ab+3a
↳ // da=ad+3x2
↳ // xb=bx-x
↳ // db=bd+d
↳ // dx=xd+1
w = 2,1,1,1;
def B = appendWeight2Ord(w); // for a non-commutative ring
setring B; B;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering a
↳ // : names a b x d
↳ // : weights 2 1 1 1
↳ // block 2 : ordering Dp
↳ // : names a b x d
↳ // block 3 : ordering C
↳ // noncommutative relations:
↳ // ba=ab+3a
↳ // da=ad+3x2
↳ // xb=bx-x
```

```

↳ // db=bd+d
↳ // dx=xd+1

```

### 7.5.19.6 elimWeight

Procedure from library `ncpreim.lib` (see Section 7.5.19 [`ncpreim.lib`], page 551).

**Usage:** `elimWeight(v)`;  $v$  an intvec

**Assume:** The basering is a  $G$ -algebra.  
The entries of  $v$  are in the range  $1..nvars(\text{basing})$  and the corresponding variables generate an admissible subalgebra.

**Return:** intvec, say  $w$ , such that the ordering  $(a(w), <)$ , where  $<$  is any admissible global ordering, is an elimination ordering for the subalgebra generated by the variables indexed by the entries of the given intvec.

**Note:** If no such ordering exists, the zero intvec is returned.

**Remark:** Reference: (BGL), (GML)

**Example:**

```

LIB "ncpreim.lib";
// (Lev): Example 2
ring r = 0, (a,b,x,d), Dp;
matrix D[4][4];
D[1,2] = 3*a; D[1,4] = 3*x^2; D[2,3] = -x;
D[2,4] = d; D[3,4] = 1;
def A = nc_algebra(1,D);
setring A; A;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering Dp
↳ // : names a b x d
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // ba=ab+3a
↳ // da=ad+3x2
↳ // xb=bx-x
↳ // db=bd+d
↳ // dx=xd+1
// Since d*a-a*d = 3*x^2, any admissible ordering has to satisfy
// x^2 < a*d, while any elimination ordering for {x,d} additionally
// has to fulfil a << x and a << d.
// Hence neither a block ordering with weights
// (1,1,1,1) nor a weighted ordering with weight (0,0,1,1) will do.
intvec v = 3,4;
elimWeight(v);
↳ 0,0,1,2

```

### 7.5.19.7 extendedTensor

Procedure from library `ncpreim.lib` (see Section 7.5.19 [`ncpreim.lib`], page 551).

**Usage:** `extendedTensor(A,I)`;  $A$  ring,  $I$  ideal

**Return:** ring,  $A+B$  (where  $B$  denotes the basering) extended with non-commutative relations between the vars of  $A$  and  $B$ , which arise from the homomorphism  $A \rightarrow B$  induced by  $I$  in the usual sense, i.e. if the vars of  $A$  are named  $x(i)$  and the vars of  $B$   $y(j)$ , then putting  $q(i)(j) = \text{leadcoef}(y(j)*I[i])/\text{leadcoef}(I[i]*y(j))$  and  $r(i)(j) = y(j)*I[i] - q(i)(j)*I[i]*y(j)$  yields the relation  $y(j)*x(i) = q(i)(j)*x(i)*y(j)+r(i)(j)$ .

**Remark:** Reference: (Lev)

**Example:**

```
LIB "ncpreim.lib";
def A = makeWeyl(2);
setring A; A;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x(1) x(2) D(1) D(2)
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // D(1)x(1)=x(1)*D(1)+1
⇨ // D(2)x(2)=x(2)*D(2)+1
def B = makeUgl(2);
setring B; B;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names e_1_1 e_1_2 e_2_1 e_2_2
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // e_1_2e_1_1=e_1_1*e_1_2-e_1_2
⇨ // e_2_1e_1_1=e_1_1*e_2_1+e_2_1
⇨ // e_2_1e_1_2=e_1_2*e_2_1-e_1_1+e_2_2
⇨ // e_2_2e_1_2=e_1_2*e_2_2-e_1_2
⇨ // e_2_2e_2_1=e_2_1*e_2_2+e_2_1
ideal I = var(1)*var(3), var(1)*var(4), var(2)*var(3), var(2)*var(4);
I;
⇨ I[1]=e_1_1*e_2_1
⇨ I[2]=e_1_1*e_2_2
⇨ I[3]=e_1_2*e_2_1
⇨ I[4]=e_1_2*e_2_2
def C = extendedTensor(A,I);
setring C; C;
⇨ // coefficients: QQ
⇨ // number of vars : 8
⇨ // block 1 : ordering dp
⇨ // : names x(1) x(2) D(1) D(2)
⇨ // block 2 : ordering dp
⇨ // : names e_1_1 e_1_2 e_2_1 e_2_2
⇨ // block 3 : ordering C
⇨ // noncommutative relations:
⇨ // D(1)x(1)=x(1)*D(1)+1
⇨ // e_1_1x(1)=x(1)*e_1_1-e_1_1*e_2_1
⇨ // e_1_2x(1)=x(1)*e_1_2+e_1_1^2-e_1_2*e_2_1-e_1_1*e_2_2
⇨ // e_2_1x(1)=x(1)*e_2_1+e_2_1^2
⇨ // e_2_2x(1)=x(1)*e_2_2+e_1_1*e_2_1
```



```

⇒ // D(2)x(2)=x(2)*D(2)+1
⇒ // e_1_2x(2)=x(2)*e_1_2+e_1_1*e_1_2-e_1_2*e_2_2
⇒ // e_2_1x(2)=x(2)*e_2_1-e_1_1*e_2_1+e_2_1*e_2_2
⇒ // e_1_2D(1)=D(1)*e_1_2+e_1_1*e_1_2-e_1_2*e_2_2-e_1_2
⇒ // e_2_1D(1)=D(1)*e_2_1-e_1_1*e_2_1+e_2_1*e_2_2+e_2_1
⇒ // e_1_1D(2)=D(2)*e_1_1+e_1_2*e_2_2
⇒ // e_1_2D(2)=D(2)*e_1_2+e_1_2^2
⇒ // e_2_1D(2)=D(2)*e_2_1-e_1_2*e_2_1-e_1_1*e_2_2+e_2_2^2
⇒ // e_2_2D(2)=D(2)*e_2_2-e_1_2*e_2_2
⇒ // e_1_2e_1_1=e_1_1*e_1_2-e_1_2
⇒ // e_2_1e_1_1=e_1_1*e_2_1+e_2_1
⇒ // e_2_1e_1_2=e_1_2*e_2_1-e_1_1+e_2_2
⇒ // e_2_2e_1_2=e_1_2*e_2_2-e_1_2
⇒ // e_2_2e_2_1=e_2_1*e_2_2+e_2_1

```

## 7.5.20 nctools\_lib

**Library:** nctools.lib

**Purpose:** General tools for noncommutative algebras

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**Overview:** Support: DFG (Deutsche Forschungsgesellschaft) and Metodos algebraicos y efectivos en grupos cuanticos, BFM2001-3141, MCYT, Jose Gomez-Torrecillas (Main researcher).

**Procedures:**

### 7.5.20.1 Gweights

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `Gweights(r)`; `r` a ring or a square matrix

**Return:** `intvec`

**Purpose:** compute an appropriate weight `int` vector for a `G`-algebra, i.e., such that  $\forall i < j, \text{lm}_w(d_{ij}) <_w x_i x_j$ .  
 the polynomials `d_{ij}` are taken from `r` itself, if it is of the type `ring` or defined by the given square polynomial matrix

**Theory:** `Gweights` returns an integer vector, whose weighting should be used to redefine the `G`-algebra in order to get the same non-commutative structure w.r.t. a weighted ordering. If the input is a matrix and the output is the zero vector then there is not a `G`-algebra structure associated to these relations with respect to the given variables. Another possibility is to use `weightedRing` to obtain directly a `G`-algebra with the new appropriate (weighted) ordering.

**Example:**

```

LIB "nctools.lib";
ring r = (0,q),(a,b,c,d),lp;
matrix C[4][4];

```

```

C[1,2]=q; C[1,3]=q; C[1,4]=1; C[2,3]=1; C[2,4]=q; C[3,4]=q;
matrix D[4][4];
D[1,4]=(q-1/q)*b*c;
def S = nc_algebra(C,D); setring S; S;
↳ // coefficients: QQ(q)
↳ // number of vars : 4
↳ // block 1 : ordering lp
↳ // : names a b c d
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // ba=(q)*ab
↳ // ca=(q)*ac
↳ // da=ad+(q2-1)/(q)*bc
↳ // db=(q)*bd
↳ // dc=(q)*cd
Gweights(S);
↳ 2,1,1,1
def D=fetch(r,D);
Gweights(D);
↳ 2,1,1,1

```

See also: Section 7.5.20.2 [weightedRing], page 560.

### 7.5.20.2 weightedRing

Procedure from library `nctools.lib` (see Section 7.5.20 [nctools.lib], page 559).

**Usage:** `weightedRing(r)`; `r` a ring

**Return:** ring

**Purpose:** equip the variables of the given ring with weights such that the relations of new ring (with weighted variables) satisfies the ordering condition for G-algebras: e.g.  $\forall \text{for all } i < j; \text{lm}_w(d_{ij}) <_w x_i x_j$ .

**Note:** activate this ring with the "setring" command

**Example:**

```

LIB "nctools.lib";
ring r = (0,q),(a,b,c,d),lp;
matrix C[4][4];
C[1,2]=q; C[1,3]=q; C[1,4]=1; C[2,3]=1; C[2,4]=q; C[3,4]=q;
matrix D[4][4];
D[1,4]=(q-1/q)*b*c;
def S = nc_algebra(C,D); setring S; S;
↳ // coefficients: QQ(q)
↳ // number of vars : 4
↳ // block 1 : ordering lp
↳ // : names a b c d
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // ba=(q)*ab
↳ // ca=(q)*ac
↳ // da=ad+(q2-1)/(q)*bc
↳ // db=(q)*bd
↳ // dc=(q)*cd

```

```

def t=weightedRing(S);
setring t; t;
↳ // coefficients: QQ(q)
↳ // number of vars : 4
↳ // block 1 : ordering M
↳ // : names a b c d
↳ // : weights 2 1 1 1
↳ // : weights 0 0 0 1
↳ // : weights 0 0 1 0
↳ // : weights 0 1 0 0
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // ba=(q)*ab
↳ // ca=(q)*ac
↳ // da=ad+(q2-1)/(q)*bc
↳ // db=(q)*bd
↳ // dc=(q)*cd

```

See also: Section 7.5.20.1 [Gweights], page 559.

### 7.5.20.3 ndcond

Procedure from library `nctools.lib` (see Section 7.5.20 [nctools.lib], page 559).

**Usage:** `ndcond()`;

**Return:** ideal

**Purpose:** compute the non-degeneracy conditions of the basering

**Note:** if `printlevel > 0`, the procedure displays intermediate information (by default, `printlevel=0`)

**Example:**

```

LIB "nctools.lib";
ring r = (0,q1,q2),(x,y,z),dp;
matrix C[3][3];
C[1,2]=q2; C[1,3]=q1; C[2,3]=1;
matrix D[3][3];
D[1,2]=x; D[1,3]=z;
def S = nc_algebra(C,D); setring S;
S;
↳ // coefficients: QQ(q1, q2)
↳ // number of vars : 3
↳ // block 1 : ordering dp
↳ // : names x y z
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // yx=(q2)*x*y+x
↳ // zx=(q1)*x*z+z
ideal j=ndcond(); // the silent version
j;
↳ j[1]=(-q2+1)*y*z-z
printlevel=1;
ideal i=ndcond(); // the verbose version
↳ Processing degree : 1

```

```

↳ 1 . 2 . 3 .
↳ failed: (-q2+1)*y*z-z
↳ done
i;
↳ i[1]=(-q2+1)*y*z-z

```

### 7.5.20.4 Weyl

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `Weyl()`

**Return:** ring

**Purpose:** create a Weyl algebra structure on the basering

**Note:** Activate this ring using the command `setring`.  
 Assume the number of variables of a basering is  $2k$ . (if the number of variables is odd, an error message will be returned)  
 by default, the procedure treats first  $k$  variables as coordinates  $x_i$  and the last  $k$  as differentials  $d_i$   
 if a non-zero optional argument is given, the procedure treats  $2k$  variables of a basering as  $k$  pairs  $(x_i, d_i)$ , i.e. variables with odd numbers are treated as coordinates and with even numbers as differentials

**Example:**

```

LIB "nctools.lib";
ring A1=0,(x(1..2),d(1..2)),dp;
def S=Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x(1) x(2) d(1) d(2)
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // d(1)x(1)=x(1)*d(1)+1
↳ // d(2)x(2)=x(2)*d(2)+1
kill A1,S;
ring B1=0,(x1,d1,x2,d2),dp;
def S=Weyl(1);
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x1 d1 x2 d2
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // d1x1=x1*d1+1
↳ // d2x2=x2*d2+1

```

See also: Section 7.5.20.5 [`makeWeyl`], page 562.

### 7.5.20.5 makeWeyl

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `makeWeyl(n,[p]);` n an integer,  $n > 0$ ; p an optional integer (field characteristic)

**Return:** ring

**Purpose:** create the n-th Weyl algebra over the rationals  $\mathbb{Q}$  or  $\mathbb{F}_p$

**Note:** activate this ring with the "setring" command.

The presentation of an n-th Weyl algebra is classical:  $D(i)x(i)=x(i)D(i)+1$ ,

where  $x(i)$  correspond to coordinates and  $D(i)$  to partial differentiations,  $i=1,\dots,n$ .

If p is not prime, the next larger prime number will be used.

**Example:**

```
LIB "nctools.lib";
def a = makeWeyl(3);
setring a;
a;
⇒ // coefficients: QQ
⇒ // number of vars : 6
⇒ // block 1 : ordering dp
⇒ // : names x(1) x(2) x(3) D(1) D(2) D(3)
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // D(1)x(1)=x(1)*D(1)+1
⇒ // D(2)x(2)=x(2)*D(2)+1
⇒ // D(3)x(3)=x(3)*D(3)+1
```

See also: Section 7.5.20.4 [Weyl], page 562.

### 7.5.20.6 makeHeisenberg

Procedure from library `nctools.lib` (see Section 7.5.20 [nctools.lib], page 559).

**Usage:** `makeHeisenberg(n, [p,d]);` int n (setting  $2n+1$  variables), optional int p (field characteristic), optional int d (power of h in the commutator)

**Return:** ring

**Purpose:** create the n-th Heisenberg algebra in the variables  $x(1),y(1),\dots,x(n),y(n),h$  over the rationals  $\mathbb{Q}$  or  $\mathbb{F}_p$  with the relations  $\forall i \in \{1,2,\dots,n\}; y(j)x(i) = x(i)y(j)+h^d$ .

**Note:** activate this ring with the `setring` command

If p is not prime, the next larger prime number will be used.

**Example:**

```
LIB "nctools.lib";
def a = makeHeisenberg(2);
setring a; a;
⇒ // coefficients: QQ
⇒ // number of vars : 5
⇒ // block 1 : ordering lp
⇒ // : names x(1) x(2) y(1) y(2) h
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // y(1)x(1)=x(1)*y(1)+h
⇒ // y(2)x(2)=x(2)*y(2)+h
def H3 = makeHeisenberg(3, 7, 2);
```

```

setring H3; H3;
⇨ // coefficients: ZZ/7
⇨ // number of vars : 7
⇨ // block 1 : ordering lp
⇨ // : names x(1) x(2) x(3) y(1) y(2) y(3) h
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // y(1)x(1)=x(1)*y(1)+h^2
⇨ // y(2)x(2)=x(2)*y(2)+h^2
⇨ // y(3)x(3)=x(3)*y(3)+h^2

```

See also: Section 7.5.20.5 [makeWeyl], page 562.

### 7.5.20.7 Exterior

Procedure from library `nctools.lib` (see Section 7.5.20 [nctools.lib], page 559).

**Usage:** Exterior();

**Return:** qring

**Purpose:** create the exterior algebra of a basering

**Note:** activate this qring with the "setring" command

**Theory:** given a basering, this procedure introduces the anticommutative relations  $x(j)x(i)=-x(i)x(j)$  for all  $j>i$ , moreover, creates a factor algebra modulo the two-sided ideal, generated by  $x(i)^2$  for all  $i$

**Example:**

```

LIB "nctools.lib";
ring R = 0,(x(1..3)),dp;
def ER = Exterior();
setring ER;
ER;
⇨ // coefficients: QQ
⇨ // number of vars : 3
⇨ // block 1 : ordering dp
⇨ // : names x(1) x(2) x(3)
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // x(2)x(1)=-x(1)*x(2)
⇨ // x(3)x(1)=-x(1)*x(3)
⇨ // x(3)x(2)=-x(2)*x(3)
⇨ // quotient ring from ideal
⇨ _[1]=x(3)^2
⇨ _[2]=x(2)^2
⇨ _[3]=x(1)^2

```

### 7.5.20.8 findimAlgebra

Procedure from library `nctools.lib` (see Section 7.5.20 [nctools.lib], page 559).

**Usage:** findimAlgebra(M,[r]); M a matrix, r an optional ring

**Return:** ring

**Purpose:** define a finite dimensional algebra structure on a ring

**Note:** the matrix  $M$  is used to define the relations  $x(i)*x(j) = M[i,j]$  in the basering (by default) or in the optional ring  $r$ .

The procedure equips the ring with the noncommutative structure.

The procedure exports the ideal (not a two-sided Groebner basis!), called `fdQuot`, for further qring definition.

**Theory:** finite dimensional algebra can be represented as a factor algebra of a  $G$ -algebra modulo certain two-sided ideal. The relations of a f.d. algebra are thus naturally divided into two groups: firstly, the relations on the variables of the ring, making it into  $G$ -algebra and the rest of them, which constitute the ideal which will be factored out.

**Example:**

```
LIB "nctools.lib";
ring r=(0,a,b),(x(1..3)),dp;
matrix S[3][3];
S[2,3]=a*x(1); S[3,2]=-b*x(1);
def A=findimAlgebra(S); setring A;
fdQuot = twostd(fdQuot);
qring Qr = fdQuot;
Qr;
↪ // coefficients: QQ(a, b)
↪ // number of vars : 3
↪ // block 1 : ordering dp
↪ // : names x(1) x(2) x(3)
↪ // block 2 : ordering C
↪ // noncommutative relations:
↪ // x(3)x(2)=(-b)/(a)*x(2)*x(3)
↪ // quotient ring from ideal
↪ _[1]=x(3)^2
↪ _[2]=x(2)*x(3)+(-a)*x(1)
↪ _[3]=x(1)*x(3)
↪ _[4]=x(2)^2
↪ _[5]=x(1)*x(2)
↪ _[6]=x(1)^2
```

### 7.5.20.9 superCommutative

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `superCommutative([b,[e, [Q]]]);`

**Return:** qring

**Purpose:** create a super-commutative algebra (as a GR-algebra) over a basering,

**Note:** activate this qring with the "setring" command.

**Note:** if  $b=e$  then the resulting ring is commutative.  
By default,  $b=1$ ,  $e=nvars(basering)$ ,  $Q=0$ .

**Theory:** given a basering, this procedure introduces the anti-commutative relations  $\text{var}(j)\text{var}(i)=-\text{var}(i)\text{var}(j)$  for all  $e \geq j > i \geq b$  and creates the quotient of the anti-commutative algebra modulo the two-sided ideal, generated by  $x(b)^2, \dots, x(e)^2 + Q$

**Display:** If `printlevel > 1`, warning debug messages will be printed

**Example:**

```
LIB "nctools.lib";
ring R = 0,(x(1..4)),dp; // global!
def ER = superCommutative(); // the same as Exterior (b = 1, e = N)
setring ER; ER;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x(1) x(2) x(3) x(4)
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // x(2)x(1)=-x(1)*x(2)
↳ // x(3)x(1)=-x(1)*x(3)
↳ // x(4)x(1)=-x(1)*x(4)
↳ // x(3)x(2)=-x(2)*x(3)
↳ // x(4)x(2)=-x(2)*x(4)
↳ // x(4)x(3)=-x(3)*x(4)
↳ // quotient ring from ideal
↳ _[1]=x(4)^2
↳ _[2]=x(3)^2
↳ _[3]=x(2)^2
↳ _[4]=x(1)^2
"Alternating variables: [" , AltVarStart(), "," , AltVarEnd(), "]" .";
↳ Alternating variables: [1 , 4].
kill R; kill ER;
ring R = 0,(x(1..4)),(lp(1), dp(3)); // global!
def ER = superCommutative(2); // b = 2, e = N
setring ER; ER;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering lp
↳ // : names x(1)
↳ // block 2 : ordering dp
↳ // : names x(2) x(3) x(4)
↳ // block 3 : ordering C
↳ // noncommutative relations:
↳ // x(3)x(2)=-x(2)*x(3)
↳ // x(4)x(2)=-x(2)*x(4)
↳ // x(4)x(3)=-x(3)*x(4)
↳ // quotient ring from ideal
↳ _[1]=x(4)^2
↳ _[2]=x(3)^2
↳ _[3]=x(2)^2
"Alternating variables: [" , AltVarStart(), "," , AltVarEnd(), "]" .";
↳ Alternating variables: [2 , 4].
kill R; kill ER;
ring R = 0,(x, y, z),(ds(1), dp(2)); // mixed!
def ER = superCommutative(2,3); // b = 2, e = 3
setring ER; ER;
↳ // coefficients: QQ
↳ // number of vars : 3
```



```

⇒ // block 1 : ordering ds
⇒ // : names x
⇒ // block 2 : ordering dp
⇒ // : names y z
⇒ // block 3 : ordering C
⇒ // noncommutative relations:
⇒ // zy=-yz
⇒ // quotient ring from ideal
⇒ _[1]=y2
⇒ _[2]=z2
"Alternating variables: [", AltVarStart(), ",", AltVarEnd(), "].";
⇒ Alternating variables: [2 , 3].
x + 1 + z + y; // ordering on variables: y > z > 1 > x
⇒ y+z+1+x
std(x - x*x*x);
⇒ _[1]=x
std(ideal(x - x*x*x, x*x*z + y, z + y*x*x));
⇒ _[1]=y+x2z
⇒ _[2]=z+x2y
⇒ _[3]=x
kill R; kill ER;
ring R = 0,(x, y, z),(ds(1), dp(2)); // mixed!
def ER = superCommutative(2, 3, ideal(x - x*x, x*x*z + y, z + y*x*x)); // b = 2, e =
setring ER; ER;
⇒ // coefficients: QQ
⇒ // number of vars : 3
⇒ // block 1 : ordering ds
⇒ // : names x
⇒ // block 2 : ordering dp
⇒ // : names y z
⇒ // block 3 : ordering C
⇒ // noncommutative relations:
⇒ // zy=-yz
⇒ // quotient ring from ideal
⇒ _[1]=y+x2z
⇒ _[2]=z+x2y
⇒ _[3]=x
⇒ _[4]=y2
⇒ _[5]=z2
"Alternating variables: [", AltVarStart(), ",", AltVarEnd(), "].";
⇒ Alternating variables: [2 , 3].

```

### 7.5.20.10 rightStd

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Purpose:** compute a right Groebner basis of I

**Return:** the same type as input

**Example:**

```

LIB "nctools.lib";
LIB "ncalg.lib";
def A = makeUs1(2);

```

```

setring A;
ideal I = e2,f;
option(redSB);
option(redTail);
ideal LI = std(I);
LI;
↳ LI[1]=f
↳ LI[2]=h2+h
↳ LI[3]=eh+e
↳ LI[4]=e2
ideal RI = rightStd(I);
RI;
↳ RI[1]=f
↳ RI[2]=h2-h
↳ RI[3]=eh+e
↳ RI[4]=e2

```

### 7.5.20.11 rightNF

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `rightNF(I)`;  $v$  a poly/vector,  $M$  an ideal/module

**Purpose:** compute a right normal form of  $v$  w.r.t.  $M$

**Return:** poly/vector (as of the 1st argument)

**Example:**

```

LIB "nctools.lib";
LIB "ncalg.lib";
ring r = 0,(x,d),dp;
def S = nc_algebra(1,1); setring S; // Weyl algebra
ideal I = x; I = std(I);
poly p = x*d+1;
NF(p,I); // left normal form
↳ 0
rightNF(p,I); // right normal form
↳ 1

```

### 7.5.20.12 rightModulo

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `rightModulo(M,N)`;  $M,N$  are ideals/modules

**Purpose:** compute a right representation of the module  $(M+N)/N$

**Return:** module

**Assume:**  $M,N$  are presentation matrices for right modules

**Example:**

```

LIB "nctools.lib";
LIB "ncalg.lib";
def A = makeUs1(2);
setring A;
option(redSB);

```

```

option(redTail);
ideal I = e2,f2,h2-1;
I = twostd(I);
print(matrix(I));
↳ h2-1,fh-f,eh+e,f2,2ef-h-1,e2
ideal E = std(e);
ideal TL = e,h-1; // the result of left modulo
TL;
↳ TL[1]=e
↳ TL[2]=h-1
ideal T = rightModulo(E,I);
T = rightStd(T+I);
T = rightStd(rightNF(T,I)); // make the output canonic
T;
↳ T[1]=h+1
↳ T[2]=e

```

### 7.5.20.13 moduloSlim

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `moduloSlim(A,B)`; A,B module/matrix/ideal

**Return:** module

**Purpose:** compute modulo with `slimgb` as engine

**Example:**

```

LIB "nctools.lib";
LIB "ncalg.lib";
ring r; // first classical example for modulo
ideal h1=x,y,z; ideal h2=x;
module m=moduloSlim(h1,h2);
print(m);
↳ 1,0,0, 0,
↳ 0,0,z, x,
↳ 0,x,-y,0
// now, a noncommutative example
def A = makeUs12(); setring A; // this algebra is U(sl_2)
ideal H2 = e2,f2,h2-1; H2 = twostd(H2);
print(matrix(H2)); // print H2 in a compact form
↳ h2-1,fh-f,eh+e,f2,2ef-h-1,e2
ideal H1 = std(e);
ideal T = moduloSlim(H1,H2);
T = std(NF(std(H2+T),H2));
T;
↳ T[1]=h-1
↳ T[2]=e
// now, a matrix example:
ring r2 = 0,(x,d), (dp);
def R = nc_algebra(1,1); setring R;
matrix M[2][2] = d, 0, 0, d*(x*d);
matrix P[2][1] = (8x+7)*d+9x, (x2+1)*d + 5*x;
module X = moduloSlim(P,M);
print(X);

```

$\mapsto 5x^2d^2-2xd^3-5xd-6d^2+5, xd^5+5xd^3+5d^4+5d^2$

### 7.5.20.14 ncRelations

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `ncRelations(r)`; `r` a ring

**Return:** list `L` with two elements, both elements are of type matrix:  
`L[1]` = matrix of coefficients `C`,  
`L[2]` = matrix of polynomials `D`

**Purpose:** recover the noncommutative relations via matrices `C` and `D` from a noncommutative ring

**Example:**

```
LIB "nctools.lib";
ring r = 0, (x,y,z), dp;
matrix C[3][3]=0,1,2,0,0,-1,0,0,0;
print(C);
 $\mapsto 0,1,2,$
 $\mapsto 0,0,-1,$
 $\mapsto 0,0,0$
matrix D[3][3]=0,1,2y,0,0,-2x+y+1;
print(D);
 $\mapsto 0,1,2y,$
 $\mapsto 0,0,-2x+y+1,$
 $\mapsto 0,0,0$
def S=nc_algebra(C,D);setring S; S;
 \mapsto // coefficients: QQ
 \mapsto // number of vars : 3
 \mapsto // block 1 : ordering dp
 \mapsto // : names x y z
 \mapsto // block 2 : ordering C
 \mapsto // noncommutative relations:
 \mapsto // yx=xy+1
 \mapsto // zx=2xz+2y
 \mapsto // zy=-yz-2x+y+1
def l=ncRelations(S);
print (l[1]);
 $\mapsto 0,1,2,$
 $\mapsto 0,0,-1,$
 $\mapsto 0,0,0$
print (l[2]);
 $\mapsto 0,1,2y,$
 $\mapsto 0,0,-2x+y+1,$
 $\mapsto 0,0,0$
```

See also: Section 7.4.1 [G-algebras], page 360; Section 5.1.135 [ringlist], page 250.

### 7.5.20.15 isCentral

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `isCentral(p)`; `p` poly

**Return:** int, 1 if `p` commutes with all variables and 0 otherwise

**Purpose:** check whether  $p$  is central in a basering (that is, commutes with every generator of the ring)

**Note:** if `printlevel > 0`, the procedure displays intermediate information (by default, `printlevel=0`)

**Example:**

```
LIB "nctools.lib";
ring r=0,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z;
D[1,3]=2*x;
D[2,3]=-2*y;
def S = nc_algebra(1,D); setring S;
S; // this is U(sl_2)
↳ // coefficients: QQ
↳ // number of vars : 3
↳ // block 1 : ordering dp
↳ // : names x y z
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // yx=xy-z
↳ // zx=xz+2x
↳ // zy=yz-2y
poly c = 4*x*y+z^2-2*z;
printlevel = 0;
isCentral(c);
↳ 1
poly h = x*c;
printlevel = 1;
isCentral(h);
↳ Non-central at: y
↳ Non-central at: z
↳ 0
```

### 7.5.20.16 isNC

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `isNC()`;

**Purpose:** check whether a basering is commutative or not

**Return:** `int`, 1 if basering is noncommutative and 0 otherwise

**Example:**

```
LIB "nctools.lib";
def a = makeWeyl(2);
setring a;
isNC();
↳ 1
kill a;
ring r = 17,(x(1..7)),dp;
isNC();
↳ 0
kill r;
```

### 7.5.20.17 isCommutative

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `isCommutative()`;

**Return:** `int`, 1 if basering is commutative, or 0 otherwise

**Purpose:** check whether basering is commutative

**Example:**

```
LIB "nctools.lib";
ring r = 0,(x,y),dp;
isCommutative();
↪ 1
def D = Weyl(); setring D;
isCommutative();
↪ 0
setring r;
def R = nc_algebra(1,0); setring R;
isCommutative();
↪ 1
```

### 7.5.20.18 isWeyl

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `isWeyl()`;

**Return:** `int`, 1 if basering is a Weyl algebra, or 0 otherwise

**Purpose:** check whether basering is a Weyl algebra

**Example:**

```
LIB "nctools.lib";
ring r = 0,(a,b,c,d),dp;
isWeyl();
↪ 0
def D = Weyl(1); setring D; //make from r a Weyl algebra
b*a;
↪ ab+1
isWeyl();
↪ 1
ring t = 0,(Dx,x,y,Dy),dp;
matrix M[4][4]; M[1,2]=-1; M[3,4]=1;
def T = nc_algebra(1,M); setring T;
isWeyl();
↪ 1
```

### 7.5.20.19 UpOneMatrix

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `UpOneMatrix(n)`; `n` an integer

**Return:** `intmat`

**Purpose:** compute an `n x n` matrix with 1's in the whole upper triangle

**Note:** helpful for setting noncommutative algebras with complicated coefficient matrices

**Example:**

```
LIB "nctools.lib";
ring r = (0,q),(x,y,z),dp;
matrix C = UpOneMatrix(3);
C[1,3] = q;
print(C);
↳ 0,1,(q),
↳ 0,0,1,
↳ 0,0,0
def S = nc_algebra(C,0); setring S;
S;
↳ // coefficients: QQ(q)
↳ // number of vars : 3
↳ // block 1 : ordering dp
↳ // : names x y z
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // zx=(q)*xz
```

### 7.5.20.20 AltVarStart

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `AltVarStart()`;

**Return:** `int`

**Purpose:** returns the number of the first alternating variable of basering

**Note:** basering should be a super-commutative algebra constructed by the procedure `superCommutative`, emits an error otherwise

**Example:**

```
LIB "nctools.lib";
ring R = 0,(x(1..4)),dp; // global!
def ER = superCommutative(2); // (b = 2, e = N)
setring ER; ER;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x(1) x(2) x(3) x(4)
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // x(3)x(2)=-x(2)*x(3)
↳ // x(4)x(2)=-x(2)*x(4)
↳ // x(4)x(3)=-x(3)*x(4)
↳ // quotient ring from ideal
↳ _[1]=x(4)^2
↳ _[2]=x(3)^2
↳ _[3]=x(2)^2
"Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "].";
↳ Alternating variables: [2 , 4].
setring R;
"Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "].";
```

```

↳ ? SCA rings are factors by (at least) squares!
↳ ? leaving nctools.lib::AltVarStart (1133)
kill R, ER;
//
ring R = 2,(x(1..4)),dp; // the same in char. = 2!
def ER = superCommutative(2); // (b = 2, e = N)
setring ER; ER;
↳ // coefficients: ZZ/2
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x(1) x(2) x(3) x(4)
↳ // block 2 : ordering C
↳ // quotient ring from ideal
↳ _[1]=x(4)^2
↳ _[2]=x(3)^2
↳ _[3]=x(2)^2
"Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "]" .";
↳ Alternating variables: [4 , 4].
setring R;
"Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "]" .";
↳ ? SCA rings are factors by (at least) squares!
↳ ? leaving nctools.lib::AltVarStart (1133)

```

### 7.5.20.21 AltVarEnd

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `AltVarStart()`;

**Return:** `int`

**Purpose:** returns the number of the last alternating variable of basering

**Note:** basering should be a super-commutative algebra constructed by the procedure `superCommutative`, emits an error otherwise

**Example:**

```

LIB "nctools.lib";
ring R = 0,(x(1..4)),dp; // global!
def ER = superCommutative(2); // (b = 2, e = N)
setring ER; ER;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x(1) x(2) x(3) x(4)
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // x(3)x(2)=-x(2)*x(3)
↳ // x(4)x(2)=-x(2)*x(4)
↳ // x(4)x(3)=-x(3)*x(4)
↳ // quotient ring from ideal
↳ _[1]=x(4)^2
↳ _[2]=x(3)^2
↳ _[3]=x(2)^2
"Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "]" .";

```



```

⇒ Alternating variables: [2 , 4].
setring R;
"Alternating variables: [" , AltVarStart(), "," , AltVarEnd(), "]" .";
⇒ ? SCA rings are factors by (at least) squares!
⇒ ? leaving nctools.lib::AltVarStart (1133)
kill R, ER;
////////////////////////////////////
ring R = 2,(x(1..4)),dp; // the same in char. = 2!
def ER = superCommutative(2); // (b = 2, e = N)
setring ER; ER;
⇒ // coefficients: ZZ/2
⇒ // number of vars : 4
⇒ // block 1 : ordering dp
⇒ // : names x(1) x(2) x(3) x(4)
⇒ // block 2 : ordering C
⇒ // quotient ring from ideal
⇒ _[1]=x(4)^2
⇒ _[2]=x(3)^2
⇒ _[3]=x(2)^2
"Alternating variables: [" , AltVarStart(), "," , AltVarEnd(), "]" .";
⇒ Alternating variables: [4 , 4].
setring R;
"Alternating variables: [" , AltVarStart(), "," , AltVarEnd(), "]" .";
⇒ ? SCA rings are factors by (at least) squares!
⇒ ? leaving nctools.lib::AltVarStart (1133)

```

### 7.5.20.22 IsSCA

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `IsSCA()`;

**Return:** `int`

**Purpose:** returns 1 if basering is a super-commutative algebra and 0 otherwise

**Example:**

```

LIB "nctools.lib";
////////////////////////////////////
ring R = 0,(x(1..4)),dp; // commutative
if(IsSCA())
{ "Alternating variables: [" , AltVarStart(), "," , AltVarEnd(), "]" ."; }
else
{ "Not a super-commutative algebra!!!"; }
⇒ Not a super-commutative algebra!!!
kill R;
////////////////////////////////////
ring R = 0,(x(1..4)),dp;
def S = nc_algebra(1, 0); setring S; S; // still commutative!
⇒ // coefficients: QQ
⇒ // number of vars : 4
⇒ // block 1 : ordering dp
⇒ // : names x(1) x(2) x(3) x(4)
⇒ // block 2 : ordering C
⇒ // noncommutative relations:

```

```

if(IsSCA())
{ "Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "]."; }
else
{ "Not a super-commutative algebra!!!"; }
↳ Not a super-commutative algebra!!!
kill R, S;
//
ring R = 0,(x(1..4)),dp;
list CurrRing = ringlist(R);
def ER = ring(CurrRing);
setring ER; // R;
matrix E = UpOneMatrix(nvars(R));
int i, j; int b = 2; int e = 3;
for (i = b; i < e; i++)
{
for (j = i+1; j <= e; j++)
{
E[i, j] = -1;
}
}
def S = nc_algebra(E,0); setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x(1) x(2) x(3) x(4)
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // x(3)x(2)=-x(2)*x(3)
if(IsSCA())
{ "Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "]."; }
else
{ "Not a super-commutative algebra!!!"; }
↳ Not a super-commutative algebra!!!
kill R, ER, S;
//
ring R = 0,(x(1..4)),dp;
def ER = superCommutative(2); // (b = 2, e = N)
setring ER; ER;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x(1) x(2) x(3) x(4)
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // x(3)x(2)=-x(2)*x(3)
↳ // x(4)x(2)=-x(2)*x(4)
↳ // x(4)x(3)=-x(3)*x(4)
↳ // quotient ring from ideal
↳ _[1]=x(4)^2
↳ _[2]=x(3)^2
↳ _[3]=x(2)^2
if(IsSCA())
{ "This is a SCA! Alternating variables: [" , AltVarStart(), ",", AltVarEnd(), "]."; }

```

```

 ↪ This is a SCA! Alternating variables: [2 , 4].
 else
 { "Not a super-commutative algebra!!!"; }
 kill R, ER;

```

### 7.5.20.23 makeModElimRing

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `makeModElimRing(L)`; L a list

**Return:** ring

**Purpose:** create a copy of a given ring equipped with the elimination ordering for module components  $(c, <)$

**Note:** usually the list argument contains a ring to work with

**Example:**

```

LIB "nctools.lib";
ring r1 = 0, (x,y,z), (C,Dp);
def r2 = makeModElimRing(r1); setring r2; r2; kill r2;
↪ // coefficients: QQ
↪ // number of vars : 3
↪ // block 1 : ordering c
↪ // block 2 : ordering Dp
↪ // : names x y z
ring r3 = 0, (z,t), (wp(2,3),c);
def r2 = makeModElimRing(r3); setring r2; r2; kill r2;
↪ // coefficients: QQ
↪ // number of vars : 2
↪ // block 1 : ordering c
↪ // block 2 : ordering wp
↪ // : names z t
↪ // : weights 2 3
ring r4 = 0, (z,t,u,w), (a(1,2),C,wp(2,3,4,5));
def r2 = makeModElimRing(r4); setring r2; r2;
↪ // coefficients: QQ
↪ // number of vars : 4
↪ // block 1 : ordering c
↪ // block 2 : ordering a
↪ // : names z t
↪ // : weights 1 2
↪ // block 3 : ordering wp
↪ // : names z t u w
↪ // : weights 2 3 4 5

```

### 7.5.20.24 embedMat

Procedure from library `nctools.lib` (see Section 7.5.20 [`nctools.lib`], page 559).

**Usage:** `embedMat(A,m,n)`; A,B matrix/module

**Return:** matrix

**Purpose:** embed A in the left upper corner of  $m \times n$  matrix

**Example:**

```

LIB "nctools.lib";
ring r = 0, (a,b,c,d), dp;
matrix M[2][3]; M[1,1]=a; M[1,2]=b;M[2,2]=d;M[1,3]=c;
print(M);
↳ a,b,c,
↳ 0,d,0
print(embedMat(M,3,4));
↳ a,b,c,0,
↳ 0,d,0,0,
↳ 0,0,0,0
matrix N = M; N[2,2]=0;
print(embedMat(N,3,4));
↳ a,b,c,0,
↳ 0,0,0,0,
↳ 0,0,0,0

```

**7.5.21 olga\_lib****Library:** olga.lib**Purpose:** Ore-localization in G-Algebras**Author:** Johannes Hoffmann, email: johannes.hoffmann at math.rwth-aachen.de**Overview:** Let A be a G-algebra.

Current localization types:

Type 0: monoidal

- represented by a list of polys  $g_1, \dots, g_k$  that have to be contained in a commutative polynomial subring of A generated by a subset of the variables of A

Type 1: geometric

- only for algebras with an even number of variables where the first half induces a commutative polynomial subring B of A

- represented by an ideal p, which has to be a prime ideal in B

- represented by an intvec  $v = [i_1, \dots, i_k]$  in the range  $1..nvars(\text{basering})$

Localization data is an int specifying the type and a def with the corresponding information.

A fraction is represented as a vector with four entries:  $[s,r,p,t]$  Here,  $s^{-1}r$  is the left fraction representation,  $pt^{-1}$  is the right one. If s or t is zero, it means that the corresponding representation is not set. If both are zero, the fraction is not valid.

A detailed description along with further examples can be found in our paper: Johannes Hoffmann, Viktor Levandovskyy:

Constructive Arithmetics in Ore Localizations of Domains  
<https://arxiv.org/abs/1712.01773>

**Procedures:****7.5.21.1 locStatus**

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `locStatus(locType, locData), int locType, list/vector/intvec locData`**Purpose:** determine the status of a set of localization data

**Assume:****Return:** list**Note:** - the first entry is 0 or 1, depending whether the input represents a valid localization  
- the second entry is a string with a status/error message**Example:**

```

LIB "olga.lib";
locStatus(42, list(1));
↳ [1]:
↳ 0
↳ [2]:
↳ invalid localization: type is 42, valid types are:
↳ 0 for a monoidal localization
↳ 1 for a geometric localization
↳ 2 for a rational localization
def undef;
locStatus(0, undef);
↳ [1]:
↳ 0
↳ [2]:
↳ uninitialized or invalid localization: locData has to be defined
string s;
locStatus(0, s);
↳ [1]:
↳ 0
↳ [2]:
↳ for a monoidal localization, locData has to be of type list, but is of\
type string
list L;
locStatus(0, L);
↳ [1]:
↳ 0
↳ [2]:
↳ for a monoidal localization, locData has to be a non-empty list
L = s;
print(L);
↳ [1]:
↳
↳
locStatus(0, L);
↳ [1]:
↳ 0
↳ [2]:
↳ for a monoidal localization, locData has to be a list of polys, ints o\
r numbers, but entry 1 is , which is of type string
ring w = 0, (x,Dx,y,Dy),dp;
def W = Weyl(1);
setring W;
W;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x Dx y Dy

```

```

⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Dxx=x*Dx+1
⇒ // Dyy=y*Dy+1
locStatus(0, list(x, Dx));
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ for a monoidal localization, the variables occurring in the polys in l\
 ocData have to induce a commutative polynomial subring of basering
ring R;
setring R;
R;
⇒ // coefficients: ZZ/32003
⇒ // number of vars : 3
⇒ // block 1 : ordering dp
⇒ // : names x y z
⇒ // block 2 : ordering C
locStatus(1, s);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ for a geometric localization, basering has to have an even number of v\
 ariables
setring W;
locStatus(1, s);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ for a geometric localization, the first half of the variables of baser\
 ing has to induce a commutative polynomial subring of basering
ring t = 0, (x,y,Dx,Dy), dp;
def T = Weyl();
setring T;
T;
⇒ // coefficients: QQ
⇒ // number of vars : 4
⇒ // block 1: ordering dp
⇒ // : names x y Dx Dy
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Dxx=x*Dx+1
⇒ // Dyy=y*Dy+1
locStatus(1, s);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ for a geometric localization, locData has to be of type ideal, but is \
 of type string
locStatus(1, ideal(Dx));
⇒ [1]:
⇒ 0
⇒ [2]:

```

```

⇒ for a geometric localization, locData has to be an ideal generated by \
 polynomials containing only variables from the first half of the variable\
 s
locStatus(2, s);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ for a rational localization, locData has to be of type intvec, but is \
 of type string
intvec v;
locStatus(2, v);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ for a rational localization, locData has to be a non-zero intvec
locStatus(2, intvec(1,2));
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ valid localization

```

### 7.5.21.2 testLocData

Procedure from library `olga.lib` (see Section 7.5.21 [olga.lib], page 578).

**Usage:** `testLocData(locType, locData), int locType,`  
`list/vector/intvec locData`

**Purpose:** test if the given data specifies a denominator set wrt. the checks from `locStatus`

**Assume:**

**Return:** nothing

**Note:** throws error if checks were not successful

**Example:**

```

LIB "olga.lib";
ring R; setring R;
testLocData(0, list(1)); // correct localization, no error
testLocData(42, list(1)); // incorrect localization, results in error
⇒ ? invalid localization: type is 42, valid types are:
⇒ 0 for a monoidal localization
⇒ 1 for a geometric localization
⇒ 2 for a rational localization
⇒ ? leaving olga.lib::testLocData (0)

```

### 7.5.21.3 isInS

Procedure from library `olga.lib` (see Section 7.5.21 [olga.lib], page 578).

**Usage:** `isInS(p, locType, locData(, override)), poly p, int locType, list/vector/intvec locData(,`  
`int override)`

**Purpose:** determine if a polynomial is in a denominator set

**Assume:**

**Return:** int

**Note:** - returns 0 or 1, depending whether  $p$  is in the denominator set specified by `locType` and `locData`  
 - if `override` is set, will not normalize `locData` (use with care)

**Example:**

```
LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x y Dx Dy
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // Dxx=x*Dx+1
⇨ // Dyy=y*Dy+1
// monoidal localization
poly g1 = x^2*y+x+2;
poly g2 = y^3+x*y;
list L = g1,g2;
poly g = g1^2*g2;
poly f = g-1;
isInS(g, 0, L); // g is in the denominator set
⇨ 1
isInS(f, 0, L); // f is NOT in the denominator set
⇨ 0
// geometric localization
ideal p = x-1, y-3;
g = x^2+y-3;
f = (x-1)*g;
isInS(g, 1, p); // g is in the denominator set
⇨ 1
isInS(f, 1, p); // f is NOT in the denominator set
⇨ 0
// rational localization
intvec v = 2;
g = y^5+17*y^2-4;
f = x*y;
isInS(g, 2, v); // g is in the denominator set
⇨ 1
isInS(f, 2, v); // f is NOT in the denominator set
⇨ 0
```

#### 7.5.21.4 fracStatus

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `fracStatus(frac, locType, locData)`, vector `frac`, int `locType`, list/intvec/vector `locData`

**Purpose:** determine if the given vector is a representation of a fraction in the specified localization

**Assume:**



**Return:** list

**Note:** - the first entry is 0 or 1, depending whether the input is valid - the second entry is a string with a status message

**Example:**

```
LIB "olga.lib";
ring r = QQ[x,y,Dx,Dy];
def R = Weyl();
setring R;
fracStatus([1,0,0,0], 42, list(1));
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ invalid localization in fraction: gen(1)
⇒ invalid localization: type is 42, valid types are:
⇒ 0 for a monoidal localization
⇒ 1 for a geometric localization
⇒ 2 for a rational localization
list L = x;
fracStatus([0,7,x,0], 0, L);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ vector is not a valid fraction: no denominator specified in x*gen(3)+7\
*gen(2)
fracStatus([Dx,Dy,0,0], 0, L);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ the left denominator Dx of fraction Dx*gen(1)+Dy*gen(2) is not in the \
denominator set of type 0 given by x
fracStatus([0,0,Dx,Dy], 0, L);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ the right denominator Dy of fraction Dx*gen(3)+Dy*gen(4) is not in the\
denominator set of type 0 given by x
fracStatus([x,Dx,Dy,x], 0, L);
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ left and right representation are not equal in:x*gen(4)+x*gen(1)+Dx*ge\
n(2)+Dy*gen(3)
fracStatus([x,Dx,x*Dx+2,x^2], 0, L);
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ valid fraction
```

### 7.5.21.5 testFraction

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** testFraction(frac, locType, locData), vector frac, int locType, list/intvec/vector locData

**Purpose:** test if the given vector is a representation of a fraction in the specified localization wrt. the checks from fracStatus

**Assume:**

**Return:** nothing

**Note:** throws error if checks were not successful

**Example:**

```
LIB "olga.lib";
ring r = QQ[x,y,Dx,Dy];
def R = Weyl();
setring R;
list L = x;
vector frac = [x,Dx,x*Dx+2,x^2];
testFraction(frac, 0, L); // correct localization, no error
frac = [x,Dx,x*Dx,x^2];
testFraction(frac, 0, L); // incorrect localization, results in error
↳ ? left and right representation are not equal in:x^2*gen(4)+x*Dx*gen(3\
) +x*gen(1)+Dx*gen(2)
↳ ? leaving olga.lib::testFraction (0)
```

### 7.5.21.6 leftOre

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** leftOre(s, r, locType, locData), poly s, r, int locType, list/vector/intvec locData

**Purpose:** compute left Ore data for a given tuple (s,r)

**Assume:** s is in the denominator set determined via locType and locData

**Return:** list

**Note:** - the first entry of the list is a vector [ts,tr] such that  $ts*r=tr*s$  - the second entry of the list is a description of all choices for ts

**Example:**

```
LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
// left Ore
// monoidal localization
poly g1 = x+3;
poly g2 = x*y;
```

```

list L = g1,g2;
poly g = g1^2*g2;
poly f = Dx;
list rm = leftOre(g, f, 0, L);
print(rm[1]);
↳ [x^8*y^4+12*x^7*y^4+54*x^6*y^4+108*x^5*y^4+81*x^4*y^4,x^5*y^3*Dx+6*x^4*y^3*
 3*Dx-3*x^4*y^3+9*x^3*y^3*Dx-12*x^3*y^3-9*x^2*y^3]
rm[2];
↳ [_[1]=x^8*y^4+12*x^7*y^4+54*x^6*y^4+108*x^5*y^4+81*x^4*y^4
rm[1][2]*g-rm[1][1]*f;
↳ 0
// geometric localization
ideal p = x-1, y-3;
f = Dx;
g = x^2+y;
list rg = leftOre(g, f, 1, p);
print(rg[1]);
↳ [x^4+2*x^2*y+y^2,x^2*Dx+y*Dx-2*x]
rg[2];
↳ [_[1]=x^4+2*x^2*y+y^2
rg[1][2]*g-rg[1][1]*f;
↳ 0
// rational localization
intvec rat = 1;
f = Dx+Dy;
g = x;
list rr = leftOre(g, f, 2, rat);
print(rr[1]);
↳ [x^2,x*Dx+x*Dy-1]
rr[2];
↳ [_[1]=x^2
rr[1][2]*g-rr[1][1]*f;
↳ 0

```

### 7.5.21.7 rightOre

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `rightOre(s, r, locType, locData)`, poly  $s, r$ , int `locType`, list/vector/intvec `locData`

**Purpose:** compute right Ore data for a given tuple  $(s,r)$

**Assume:**  $s$  is in the denominator set determined via `locType` and `locData`

**Return:** list

**Note:** - the first entry of the list is a vector  $[ts, tr]$  such that  $r*ts=s*tr$  - the second entry of the list is a description of all choices for  $ts$

**Example:**

```

LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4

```

```

⇒ // block 1 : ordering dp
⇒ // : names x y Dx Dy
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Dxx=x*Dx+1
⇒ // Dyy=y*Dy+1
// monoidal localization
poly g1 = x+3;
poly g2 = x*y;
list L = g1,g2;
poly g = g1^2*g2;
poly f = Dx;
list rm = rightOre(g, f, 0, L);
print(rm[1]);
⇒ [x^8*y^4+12*x^7*y^4+54*x^6*y^4+108*x^5*y^4+81*x^4*y^4,x^5*y^3*Dx+6*x^4*y^3*\
 3*Dx+8*x^4*y^3+9*x^3*y^3*Dx+36*x^3*y^3+36*x^2*y^3]
rm[2];
⇒ _[1]=x^8*y^4+12*x^7*y^4+54*x^6*y^4+108*x^5*y^4+81*x^4*y^4
g*rm[1][2]-f*rm[1][1];
⇒ 0
// geometric localization
ideal p = x-1, y-3;
f = Dx;
g = x^2+y;
list rg = rightOre(g, f, 1, p);
print(rg[1]);
⇒ [x^4+2*x^2*y+y^2,x^2*Dx+y*Dx+4*x]
rg[2];
⇒ _[1]=x^4+2*x^2*y+y^2
g*rg[1][2]-f*rg[1][1];
⇒ 0
// rational localization
intvec rat = 1;
f = Dx+Dy;
g = x;
list rr = rightOre(g, f, 2, rat);
print(rr[1]);
⇒ [x^2,x*Dx+x*Dy+2]
rr[2];
⇒ _[1]=x^2
g*rr[1][2]-f*rr[1][1];
⇒ 0

```

### 7.5.21.8 convertRightToLeftFraction

Procedure from library `olga.lib` (see Section 7.5.21 [olga.lib], page 578).

**Usage:** `convertRightToLeftFraction(frac, locType, locData),`  
vector `frac`, int `locType`, list/vector/intvec `locData`

**Purpose:** determine a left fraction representation of a given fraction

**Assume:**

**Return:** vector

**Note:** - the returned vector contains a repr. of frac as a left fraction - if the left representation of frac is already specified, frac will be returned.

**Example:**

```
LIB "olga.lib";
ring R = 0,(x,y,Dx,Dy),dp;
def S = Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
// monoidal localization
poly g1 = x+3;
poly g2 = x*y;
list L = g1,g2;
poly g = g1^2*g2;
poly f = Dx;
vector fracm = [0,0,f,g];
vector rm = convertRightToLeftFraction(fracm, 0, L);
print(rm);
↳ [x^8*y^4+12*x^7*y^4+54*x^6*y^4+108*x^5*y^4+81*x^4*y^4,x^5*y^3*Dx+6*x^4*y^3*
 3*Dx-3*x^4*y^3+9*x^3*y^3*Dx-12*x^3*y^3-9*x^2*y^3,Dx,x^3*y+6*x^2*y+9*x*y]
rm[2]*g-rm[1]*f;
↳ 0
// geometric localization
ideal p = x-1, y-3;
f = Dx;
g = x^2+y;
vector fracg = [0,0,f,g];
vector rg = convertRightToLeftFraction(fracg, 1, p);
print(rg);
↳ [x^4+2*x^2*y+y^2,x^2*Dx+y*Dx-2*x,Dx,x^2+y]
rg[2]*g-rg[1]*f;
↳ 0
// rational localization
intvec rat = 1;
f = Dx+Dy;
g = x;
vector fracr = [0,0,f,g];
vector rr = convertRightToLeftFraction(fracr, 2, rat);
print(rr);
↳ [x^2,x*Dx+x*Dy-1,Dx+Dy,x]
rr[2]*g-rr[1]*f;
↳ 0
```

### 7.5.21.9 convertLeftToRightFraction

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `convertLeftToRightFraction(frac, locType, locData), vector frac, int locType, list/vector/intvec locData`

**Purpose:** determine a right fraction representation of a given fraction

**Assume:**

**Return:** vector

**Note:** - the returned vector contains a repr. of frac as a right fraction, - if the right representation of frac is already specified, frac will be returned.

**Example:**

```
LIB "olga.lib";
ring R = 0,(x,y,Dx,Dy),dp;
def S = Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
// monoidal localization
poly g = x;
poly f = Dx;
vector fracm = [g,f,0,0];
list L = g;
vector rm = convertLeftToRightFraction(fracm, 0, L);
print(rm);
↳ [x,Dx,x*Dx+2,x^2]
f*rm[4]-g*rm[3];
↳ 0
// geometric localization
g = x+y;
f = Dx+Dy;
vector fracg = [g,f,0,0];
ideal p = x-1, y-3;
vector rg = convertLeftToRightFraction(fracg, 1, p);
print(rg);
↳ [x+y,Dx+Dy,x*Dx+y*Dx+x*Dy+y*Dy+4,x^2+2*x*y+y^2]
f*rg[4]-g*rg[3];
↳ 0
// rational localization
intvec rat = 1;
f = Dx+Dy;
g = x;
vector fracr = [g,f,0,0];
vector rr = convertLeftToRightFraction(fracr, 2, rat);
print(rr);
↳ [x,Dx+Dy,x*Dx+x*Dy+2,x^2]
f*rr[4]-g*rr[3];
↳ 0
```

### 7.5.21.10 addLeftFractions

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `addLeftFractions(a, b, locType, locData(, override)),`  
`vector a, b, int locType, list/vector/intvec locData(, int override)`

**Purpose:** add two left fractions in the specified localization

**Assume:**

**Return:** vector

**Note:** the returned vector is the sum of `a` and `b` as fractions in the localization specified by `locType` and `locData`.

**Example:**

```
LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x y Dx Dy
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // Dxx=x*Dx+1
⇨ // Dyy=y*Dy+1
// monoidal localization
poly g1 = x+3;
poly g2 = x*y+y;
list L = g1,g2;
poly s1 = g1;
poly s2 = g2;
poly r1 = Dx;
poly r2 = Dy;
vector frac1 = [s1,r1,0,0];
vector frac2 = [s2,r2,0,0];
vector rm = addLeftFractions(frac1, frac2, 0, L);
print(rm);
⇨ [x^2*y+4*x*y+3*y,x*y*Dx+y*Dx+x*Dy+3*Dy]
// geometric localization
ideal p = x-1, y-3;
vector rg = addLeftFractions(frac1, frac2, 1, p);
print(rg);
⇨ [x^2*y+4*x*y+3*y,x*y*Dx+y*Dx+x*Dy+3*Dy]
// rational localization
intvec v = 2;
s1 = y^2+y+1;
s2 = y-2;
r1 = Dx;
r2 = Dy;
frac1 = [s1,r1,0,0];
frac2 = [s2,r2,0,0];
vector rr = addLeftFractions(frac1, frac2, 2, v);
```

```
print(rr);
↳ [y^3-y^2-y-2,y^2*Dy+y*Dx+y*Dy-2*Dx+Dy]
```

### 7.5.21.11 multiplyLeftFractions

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `multiplyLeftFractions(a, b, locType, locData(, override))`, vector `a`, `b`, int `locType`, list/vector/intvec `locData`, int `override`

**Purpose:** multiply two left fractions in the specified localization

**Assume:**

**Return:** vector

**Note:** the returned vector is the product of `a` and `b` as fractions in the localization specified by `locType` and `locData`.

**Example:**

```
LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
// monoidal localization
poly g1 = x+3;
poly g2 = x*y+y;
list L = g1,g2;
poly s1 = g1;
poly s2 = g2;
poly r1 = Dx;
poly r2 = Dy;
vector frac1 = [s1,r1,0,0];
vector frac2 = [s2,r2,0,0];
vector rm = multiplyLeftFractions(frac1, frac2, 0, L);
print(rm);
↳ [x^3*y^2+5*x^2*y^2+7*x*y^2+3*y^2,x*y*Dx*Dy+y*Dx*Dy-y*Dy]
// geometric localization
ideal p = x-1, y-3;
vector rg = multiplyLeftFractions(frac1, frac2, 1, p);
print(rg);
↳ [x^3*y+5*x^2*y+7*x*y+3*y,x*Dx*Dy+Dx*Dy-Dy]
// rational localization
intvec v = 2;
s1 = y^2+y+1;
s2 = y-2;
r1 = Dx;
r2 = Dy;
```



```

frac1 = [s1,r1,0,0];
frac2 = [s2,r2,0,0];
vector rr1 = multiplyLeftFractions(frac1, frac2, 2, v);
print(rr1);
↳ [y^3-y^2-y-2,Dx*Dy]
vector rr2 = multiplyLeftFractions(frac2, frac1, 2, v);
print(rr2);
↳ [y^5-y^3-4*y^2-3*y-2,y^2*Dx*Dy+y*Dx*Dy-2*y*Dx+Dx*Dy-Dx]
areEqualLeftFractions(rr1, rr2, 2, v);
↳ 0

```

### 7.5.21.12 areEqualLeftFractions

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `areEqualLeftFractions(a, b, locType, locData), vector a, b, int locType, list/vector/intvec locData`

**Purpose:** check if two given fractions are equal

**Assume:**

**Return:** `int`

**Note:** returns 1 or 0, depending whether  $a=b$  as fractions in the localization specified by `locType` and `locData`

**Example:**

```

LIB "olga.lib";
ring R = 0,(x,y,Dx,Dy),dp;
def S = Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
// monoidal
poly g1 = x*y+3;
poly g2 = y^3;
list L = g1,g2;
poly s1 = g1;
poly s2 = s1*g2;
poly s3 = s2;
poly r1 = Dx;
poly r2 = g2*r1;
poly r3 = s1*r1+3;
vector fracm1 = [s1,r1,0,0];
vector fracm2 = [s2,r2,0,0];
vector fracm3 = [s3,r3,0,0];
areEqualLeftFractions(fracm1, fracm2, 0, L);
↳ 1
areEqualLeftFractions(fracm1, fracm3, 0, L);

```

```

↳ 0
areEqualLeftFractions(fracm2, fracm3, 0, L);
↳ 0

```

### 7.5.21.13 isInvertibleLeftFraction

Procedure from library `olga.lib` (see Section 7.5.21 [olga.lib], page 578).

**Usage:** `isInvertibleLeftFraction(frac, locType, locData), vector frac, int locType, list/vector/intvec locData`

**Purpose:** check if a fraction is invertible in the specified localization

**Assume:**

**Return:** `int`

**Note:** - returns 1, if the numerator of `frac` is in the denominator set, - returns 0, otherwise (NOTE: this does NOT mean that the fraction is not invertible, it just means it could not be determined by the method above).

**Example:**

```

LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
↳ // coefficients: QQ
↳ // number of vars : 4
↳ // block 1 : ordering dp
↳ // : names x y Dx Dy
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // Dxx=x*Dx+1
↳ // Dyy=y*Dy+1
poly g1 = x+3;
poly g2 = x*y;
list L = g1,g2;
vector frac = [g1*g2, 17, 0, 0];
isInvertibleLeftFraction(frac, 0, L);
↳ 1
ideal p = x-1, y;
frac = [g1, x, 0, 0];
isInvertibleLeftFraction(frac, 1, p);
↳ 1
intvec rat = 1,2;
frac = [g1*g2, Dx, 0, 0];
isInvertibleLeftFraction(frac, 2, rat);
↳ 0

```

### 7.5.21.14 invertLeftFraction

Procedure from library `olga.lib` (see Section 7.5.21 [olga.lib], page 578).

**Usage:** `invertLeftFraction(frac, locType, locData), vector frac, int locType, list/vector/intvec locData`

**Purpose:** invert a fraction in the specified localization

**Assume:** frac is invertible in the loc. specified by locType and locData

**Return:** vector

**Note:** - returns the multiplicative inverse of frac in the localization specified by locType and locData,  
- throws error if frac is not invertible (NOTE: this does NOT mean that the fraction is not invertible, it just means it could not be determined by the method listed above).

**Example:**

```
LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
⇨ // coefficients: QQ
⇨ // number of vars : 4
⇨ // block 1 : ordering dp
⇨ // : names x y Dx Dy
⇨ // block 2 : ordering C
⇨ // noncommutative relations:
⇨ // Dxx=x*Dx+1
⇨ // Dyy=y*Dy+1
poly g1 = x+3;
poly g2 = x*y;
list L = g1,g2;
vector frac = [g1*g2, 17, 0, 0];
print(invertLeftFraction(frac, 0, L));
⇨ [17,x^2*y+3*x*y]
ideal p = x-1, y;
frac = [g1, x, 0, 0];
print(invertLeftFraction(frac, 1, p));
⇨ [x,x+3]
intvec rat = 1,2;
frac = [g1*g2, y, 0, 0];
print(invertLeftFraction(frac, 2, rat));
⇨ [y,x^2*y+3*x*y]
```

### 7.5.21.15 isZeroFraction

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `isZeroFraction(frac)`, vector frac

**Purpose:** determine if the vector frac represents zero

**Assume:** frac is a valid fraction

**Return:** int

**Note:** returns 1, if `frac == 0`; 0 otherwise

**Example:**

```
LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
⇨ // coefficients: QQ
```

```

⇒ // number of vars : 4
⇒ // block 1 : ordering dp
⇒ // : names x y Dx Dy
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Dxx=x*Dx+1
⇒ // Dyy=y*Dy+1
isZeroFraction([42,0,0,0]);
⇒ 1
isZeroFraction([0,0,Dx,3]);
⇒ 0
isZeroFraction([1,1,1,1]);
⇒ 0

```

### 7.5.21.16 isOneFraction

Procedure from library `olga.lib` (see Section 7.5.21 [olga.lib], page 578).

**Usage:** `isOneFraction(frac)`, vector `frac`

**Purpose:** determine if the vector `frac` represents one

**Assume:** `frac` is a valid fraction

**Return:** `int`

**Note:** 1, if `frac == 1`; 0 otherwise

**Example:**

```

LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl();
setring S; S;
⇒ // coefficients: QQ
⇒ // number of vars : 4
⇒ // block 1 : ordering dp
⇒ // : names x y Dx Dy
⇒ // block 2 : ordering C
⇒ // noncommutative relations:
⇒ // Dxx=x*Dx+1
⇒ // Dyy=y*Dy+1
isOneFraction([42,42,0,0]);
⇒ 1
isOneFraction([0,0,Dx,3]);
⇒ 0
isOneFraction([1,0,0,1]);
⇒ 0

```

### 7.5.21.17 normalizeMonoidal

Procedure from library `olga.lib` (see Section 7.5.21 [olga.lib], page 578).

**Usage:** `normalizeMonoidal(L)`, list `L`

**Purpose:** compute a normal form of monoidal localization data

**Return:** `list`

**Note:** given a list of polys, returns a list of all unique factors appearing in the given polys

**Example:**

```
LIB "olga.lib";
ring R = 0, (x,y,Dx,Dy), dp;
def S = Weyl(); setring S;
list L = x^2*y^3, (x+1)*(x*y-3*y^2+1);
L = normalizeMonoidal(L);
print(L);
↳ [1]:
↳ x*y-3*y^2+1
↳ [2]:
↳ x+1
↳ [3]:
↳ x
↳ [4]:
↳ y
```

### 7.5.21.18 normalizeRational

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `normalizeRational(v)`, intvec  $v$

**Purpose:** compute a normal form of rational localization data

**Return:** intvec

**Note:** purges double entries and sorts ascendingly

**Example:**

```
LIB "olga.lib";
ring R; setring R;
intvec v = 9,5,9,3,1,5;
v = normalizeRational(v);
v;
↳ 1,3,5,9
```

### 7.5.21.19 testOlga

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `testOlga()`

**Purpose:** execute a series of internal testing procedures

**Return:** nothing

**Note:**

### 7.5.21.20 testOlgaExamples

Procedure from library `olga.lib` (see Section 7.5.21 [`olga.lib`], page 578).

**Usage:** `testOlgaExamples()`

**Purpose:** execute the examples of all procedures in this library

**Return:** nothing

**Note:**

## 7.5.22 perron\_lib

**Library:** perron.lib

**Purpose:** computation of algebraic dependences

**Author:** Oleksandr Motsak U@D, where U={motsak}, D={mathematik.uni-kl.de}

**Procedures:**

### 7.5.22.1 perron

Procedure from library `perron.lib` (see Section 7.5.22 [perron\_lib], page 596).

**Usage:** `perron( L [, D ] )`

**Return:** commutative ring with ideal 'Relations'

**Purpose:** computes polynomial relations ('Relations') between pairwise commuting polynomials of `L` [, up to a given degree bound `D`]

**Note:** the implementation was partially inspired by the Perron's theorem.

**Example:**

```
LIB "perron.lib";
int p = 3;
ring AA = p,(x,y,z),dp;
matrix D[3][3]=0;
D[1,2]=-z; D[1,3]=2*x; D[2,3]=-2*y;
def A = nc_algebra(1,D); setring A; // this algebra is U(sl_2)
ideal I = x^p, y^p, z^p-z, 4*x*y+z^2-2*z; // the center
def RA = perron(I, p);
setring RA;
RA;
↳ // coefficients: ZZ/3
↳ // number of vars : 4
↳ // block 1: ordering dp
↳ // : names F(1) F(2) F(3) F(4)
↳ // block 2: ordering C
Relations; // it was exported from perron to be in the returned ring.
↳ Relations[1]=F(4)^3-F(1)*F(2)-F(3)^2+F(4)^2
// perron can be also used in a commutative case, for example:
ring B = 0,(x,y,z),dp;
ideal J = xy+z^2, z^2+y^2, x^2y^2-2xy^3+y^4;
def RB = perron(J);
setring RB;
Relations;
↳ Relations[1]=F(1)^2-2*F(1)*F(2)+F(2)^2-F(3)
// one more test:
setring A;
map T=RA,I;
T(Relations); // should be zero
↳ _[1]=0
```

### 7.5.23 purityfiltration.lib

Status: experimental

**Library:** purityfiltration.lib

**Purpose:** Algorithms for computing a purity filtration of a given module

**Authors:** Christian Schilli, christian.schilli@rwth-aachen.de  
Viktor Levandovskyy, levandov@math.rwth-aachen.de

**Overview:** Purity is a notion with several meanings. In our context it is equidimensionality of a module (that is all  $M$  is pure iff any nonzero submodule of  $N$  has the same dimension as  $N$ ).  
Notably, one should define purity with respect to a given dimension function. In the context of this library the corresponding function is the homological grade number  $j_A(M)$  of a module  $M$  over an  $K$ -algebra  $A$ .  $j_A(M)$  is the minimal integer  $k$ , such that  $\text{Ext}^k_A(M, A) \neq 0$ .

**References:**

- [AQ] Alban Quadrat: Grade filtration of linear functional systems, INRIA Report 7769 (2010), to appear in Acta Applicanda Mathematica.
- [B93] Jan-Erik Bjoerk: Analytic D-modules and applications, Kluwer Acad. Publ., 1993.
- [MB10] Mohamed Barakat: Purity Filtration and the Fine Structure of Autonomy. Proc. MTNS, 2010.

**Procedures:**

#### 7.5.23.1 projectiveDimension

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `projectiveDimension(R,i,j)`,  $R$  matrix representing the Modul  $M = \text{coker}(R)$   
`int i`, with  $i=0$  or  $i=1$ ,  $j$  a natural number

**Return:** list  $T$ , a projective resolution of  $M$  and its projective dimension

**Purpose:** if  $i=0$  (and by default),  $T[1]$  gives a shortest left resolution of  $M = D^p / D^q(R^t)$  and  $T[2]$  the left projective dimension of  $M$   
if  $i=1$ ,  $T[1]$  gives a shortest right resolution of  $M = D^p / R D^q$  and  $T[2]$  the right projective dimension of  $M$   
in both cases  $T[1][j]$  is the  $(j-1)$ -th syzygy module of  $M$

**Note:** The algorithm is due to A. Quadrat, D. Robertz, Computation of bases of free modules over the Weyl algebras, J.Symb.Comp. 42, 2007.

**Example:**

```
LIB "purityfiltration.lib";
// commutative example
ring D = 0, (x,y,z), dp;
matrix R[6][4] =
0, -2*x, z-2*y-x, -1,
0, z-2*x, 2*y-3*x, 1,
z, -6*x, -2*y-5*x, -1,
0, y-x, y-x, 0,
```

```

y,-x,-y-x,0,
x,-x,-2*x,0;
// compute a left resolution of $M=D^4/D^6R$
list T=projectiveDimension(transpose(R),0);
// so we have the left projective dimension
T[2];
⇒ 3
//we could also compute a right resolution of $M=D^6/RD^4$
list T1=projectiveDimension(R,1);
// and we have right projective dimension
T1[2];
⇒ 1
// check, that a syzygy matrix of R has left inverse:
print(leftInverse(syz(R)));
⇒ 0,-1,0,0
// so lpd(M) must be 1.
// Non-commutative example
ring D1 = 0,(x1,x2,x3,d1,d2,d3),dp;
def S=Weyl(); setring S;
matrix R[3][3]=
1/2*x2*d1, x2*d2+1, x2*d3+1/2*d1,
-1/2*x2*d2-3/2,0,1/2*d2,
-d1-1/2*x2*d3,-d2,-1/2*d3;
list T=projectiveDimension(R,0);
// left projective dimension of coker(R) is
T[2];
⇒ 1
list T1=projectiveDimension(R,1);
// both modules have the same projective dimension, but different resolutions, because
print(T[1][1]);
⇒ 1/2*x2*d1, -1/2*x2*d2-3/2,-1/2*x2*d3-d1,
⇒ x2*d2+1, 0, -d2,
⇒ x2*d3+1/2*d1,1/2*d2, -1/2*d3
// not the same as
print(transpose(T1[1][1]));
⇒ 1/2*x2*d1, -1/2*x2*d2-3/2,-1/2*x2*d3-d1,-1/2*x2,
⇒ x2*d2+1, 0, -d2, 0,
⇒ x2*d3+1/2*d1,1/2*d2, -1/2*d3, 1/2

```

### 7.5.23.2 purityFiltration

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `purityFiltration(S)`,  $S$  matrix with entries of an Auslander regular ring  $D$

**Return:** a list  $T$  of two lists, purity filtration of the module  $M=D^q/D^p(S^t)$

**Purpose:** the first list  $T[1]$  gives a filtration  $\{M_i\}$  of  $M$ ,  
where the  $i$ -th entry of  $T[1]$  gives the representation matrix of  $M_{(i-1)}$ .  
the second list  $T[2]$  gives representations of the factor Modules,  
i.e.  $T[2][i]$  gives the repr. matrix for  $M_{(i-1)}/M_i$

**Example:**

```

LIB "purityfiltration.lib";
ring D = 0,(x1,x2,d1,d2),dp;

```



```

def S=Weyl();
setring S;
int i;
matrix R[3][3]=0,d2-d1,d2-d1,d2,-d1,-d1-d2,d1,-d1,-2*d1;
print(R);
↳ 0, -d1+d2,-d1+d2,
↳ d2,-d1, -d1-d2,
↳ d1,-d1, -2*d1
list T=purityFiltration(transpose(R));
// the purity filtration of coker(M)
print(T[1][1]);
↳ 0, -d1+d2,-d1+d2,
↳ d2,-d1, -d1-d2,
↳ d1,-d1, -2*d1
print(T[1][2]);
↳ d2, d2,
↳ d1-d2,0,
↳ d2, d1
print(T[1][3]);
↳ 1,0,
↳ 0,d2,
↳ 0,d1
// factor modules of the filtration
print(T[2][1]);
↳ 0, 1,1,
↳ -1,0,1
print(T[2][2]);
↳ 1, 1,
↳ d1-d2,0
print(T[2][3]);
↳ 1,0,
↳ 0,d2,
↳ 0,d1

```

### 7.5.23.3 purityTriang

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `purityTriang(S)`,  $S$  matrix with entries of an Auslander regular ring  $D$

**Return:** a matrix  $T$

**Purpose:** compute a triangular block matrix  $T$ , such that  $M=D^p/D^q(S^t)$  is isomorphic to  $M'=D^p/D^q(T^t)$

**Example:**

```

LIB "purityfiltration.lib";
ring D = 0,(x1,x2,d1,d2),dp;
def S=Weyl();
setring S;
int i;
matrix R[3][3]=0,d2-d1,d2-d1,d2,-d1,-d1-d2,d1,-d1,-2*d1;
print(R);
↳ 0, -d1+d2,-d1+d2,
↳ d2,-d1, -d1-d2,

```

```

↳ d1,-d1, -2*d1
matrix T=purityTriang(transpose(R));
// a triangular blockmatrix representing the module coker(R)
print(T);
↳ 0, 1,1,-1, 0, 0, 0,
↳ -1,0,1,0, -1, 0, 0,
↳ 0, 0,0,-d1,-d2,-1,0,
↳ 0, 0,0,-1, -1, 0, -1,
↳ 0, 0,0,0, 0, 1, -d2,
↳ 0, 0,0,0, 0, 1, 0,
↳ 0, 0,0,0, 0, 0, d1

```

### 7.5.23.4 gradeNumber

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `gradeNumber(R)`, `R` matrix, representing  $M=D^p/D^q(R^t)$  over a ring `D`

**Return:** `int`, grade number of `M`

**Purpose:** computes the grade number of `M`, i.e. the first `i`, with  $\text{ext}^i(M,D) \neq 0$   
returns -1 if  $M=0$

**Example:**

```

LIB "purityfiltration.lib";
// trivial example
ring D=0,(x,y,z),dp;
matrix R[2][1]=1,x;
gradeNumber(R);
↳ 0
// R has left inverse, so M=D/D^2R=0
gradeNumber(transpose(R));
↳ -1
print(ncExt_R(0,R));
↳ 0
// so, ext^0(coker(R),D) != 0
//
// a little bit more complex
matrix R1[3][1]=x,-y,z;
gradeNumber(transpose(R1));
↳ 3
print(ncExt_R(0,transpose(R1)));
↳ 1
print(ncExt_R(1,transpose(R1)));
↳ 1
print(ncExt_R(2,transpose(R1)));
↳ 1,0,0,
↳ 0,1,0,
↳ 0,0,1
// ext^i are zero for i=0,1,2
matrix ext3=ncExt_R(3,transpose(R1));
print(ext3);
↳ z,y,x
// not zero
is_zero(ext3);

```

↪ 0

### 7.5.23.5 showgrades

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `showgrades(T)`, `T` list, which includes representation matrices of modules

**Return:** list, gradenumbers of the entries in `T`

**Purpose:** computes a list `L` with  $L[i]=\text{gradenumber}(M)$ ,  $M=D^p/D^qT[i]$

**Example:**

```
LIB "purityfiltration.lib";
ring D = 0, (x,y,z), dp;
matrix R[6][4]=
0, -2*x, z-2*y-x, -1,
0, z-2*x, 2*y-3*x, 1,
z, -6*x, -2*y-5*x, -1,
0, y-x, y-x, 0,
y, -x, -y-x, 0,
x, -x, -2*x, 0;
list T=purityFiltration(transpose(R))[2];
showgrades(T);
↪ [1]:
↪ 0
↪ [2]:
↪ -1
↪ [3]:
↪ 2
↪ [4]:
↪ 3
// T[i] are i-1 pure (i=1,3,4) or zero (i=2)
```

### 7.5.23.6 allExtOfLeft

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `allExtOfLeft(M)`,

**Return:** list, entries are ext-modules

**Assume:** `M` presents a left module of finite left projective dimension `n`

**Purpose:** For a left module presented by `M` over the basering `D`, compute a list `T`, whose entry `T[i+1]` is a matrix, presenting the right module  $\text{Ext}^i_D(M,D)$  for  $i=0..n$

**Example:**

```
LIB "purityfiltration.lib";
ring D = 0, (x,y,z), dp;
matrix R[6][4]=
0, -2*x, z-2*y-x, -1,
0, z-2*x, 2*y-3*x, 1,
z, -6*x, -2*y-5*x, -1,
0, y-x, y-x, 0,
y, -x, -y-x, 0,
```

```

x,-x,-2*x,0;
// coker(R) consider the left module M=D^6/D^4R
list T=allExtOfLeft(transpose(R));
print(T[1]);
↳ 0
print(T[2]);
↳ 1,0,0,
↳ 0,1,0,
↳ 0,0,1
print(T[3]);
↳ 0,0, z,4y-z,4x-z,
↳ 0,-2,1,0, 0,
↳ 1,0, 0,0, 0,
↳ 0,1, 0,0, 0
print(T[4]);
↳ z,y,x
// right modules coker(T[i].)!!

```

### 7.5.23.7 allExtOfRight

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `allExtOfRight(R)`,  $R$  matrix representing the right Module  $M=D^q/RD^p$  over a ring  $D$   
 $M$  module with finite right projective dimension  $n$

**Return:** list, entries are ext-modules

**Purpose:** computes a list  $T$ , which entries are representations of the left modules  $\text{ext}^i(M,D)$   
 $T[i]$  gives the repr. matrix of  $\text{ext}^{i-1}(M,D)$ ,  $i=1,\dots,n+1$

**Example:**

```

LIB "purityfiltration.lib";
ring D = 0,(x,y,z),dp;
matrix R[6][4]=
0,-2*x,z-2*y-x,-1,
0,z-2*x,2*y-3*x,1,
z,-6*x,-2*y-5*x,-1,
0,y-x,y-x,0,
y,-x,-y-x,0,
x,-x,-2*x,0;
// coker(R) considered as right module
projectiveDimension(R,1)[2];
↳ 1
list T=allExtOfRight(R);
print(T[1]);
↳ 4x,
↳ -4y,
↳ -z,
↳ z
print(T[2]);
↳ 1,0,0, 0,0, 0,
↳ 0,0,4y-z,0,4x-z,0,
↳ 0,z,0, y,0, x
// left modules coker(.T[i])!!

```

### 7.5.23.8 doubleExt

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `doubleExt(R,i)`,  $R$  matrix representing the left Module  $M=D^p/D^q(R^t)$  over a ring  $D$   
`int i`, less or equal the left projective dimension of  $M$

**Return:** matrix  $P$ , representing the double ext module

**Purpose:** computes a matrix  $P$ , which represents the left module  $\text{ext}^i(\text{ext}^i(M,D))$

**Example:**

```
LIB "purityfiltration.lib";
ring D = 0, (x,y,z), dp;
matrix R[7][3]=
0 ,0,1,
1 , -4*x+z, -z,
-1, 8*x-2*z, z,
1 ,0 ,0,
0 ,x-y,0,
0 ,x-y,y,
0 ,0 ,x;
// coker(R) is 2-pure, so all doubleExt are zero
print(doubleExt(transpose(R),0));
↪ 1
print(doubleExt(transpose(R),1));
↪ 1,0,0,
↪ 0,1,0,
↪ 0,0,1
print(doubleExt(transpose(R),3));
↪ 1
// except of the second
print(doubleExt(transpose(R),2));
↪ 4y-z,4x-z
```

### 7.5.23.9 allDoubleExt

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `allDoubleExt(R)`,  $R$  matrix representing the left Module  $M=D^p/D^q(R^t)$  over a ring  $D$

**Return:** list  $T$ , double indexed, which include all double-ext modules

**Purpose:** computes all double ext-modules  
 $T[i][j]$  gives a representation matrix of  $\text{ext}^{(j-1)}(\text{ext}^{(i-1)}(M,D))$

**Example:**

```
LIB "purityfiltration.lib";
ring D = 0, (x1,x2,x3,d1,d2,d3), dp;
def S=Weyl();
setring S;
matrix R[6][4]=
0, -2*d1, d3-2*d2-d1, -1,
0, d3-2*d1, 2*d2-3*d1, 1,
```

```

d3,-6*d1,-2*d2-5*d1,-1,
0,d2-d1,d2-d1,0,
d2,-d1,-d2-d1,0,
d1,-d1,-2*d1,0;
list T=allDoubleExt(transpose(R));
// left projective dimension of M=coker(R) is 3
// ext^i(ext^0(M,D)), i=0,1,2,3
print(T[1][1]);
↳ 0,
↳ d1,
↳ d3,
↳ -d2
print(T[1][2]);
↳ d3,d3,d2,d1
print(T[1][3]);
↳ 1
print(T[1][4]);
↳ 1
// ext^i(ext^1(M,D)), i=0,1,2,3
print(T[2][1]);
↳ 1
print(T[2][2]);
↳ 1,0,0,
↳ 0,1,0,
↳ 0,0,1
print(T[2][3]);
↳ 0,0,0,4*d2-d3,4*d1-d3,
↳ 1,0,0,0, 0,
↳ 0,1,0,0, 0,
↳ 0,0,1,0, 0
print(T[2][4]);
↳ d3,d2,d1
// ext^i(ext^2(M,D)), i=0,1,2,3 (all zero)
print(T[3][1]);
↳ 1
print(T[3][2]);
↳ 1
print(T[3][3]);
↳ 1
print(T[3][4]);
↳ 1
// ext^i(ext^3(M,D)), i=0,1,2,3 (all zero)
print(T[4][1]);
↳ 1
print(T[4][2]);
↳ 1
print(T[4][3]);
↳ 1
print(T[4][4]);
↳ 1

```

### 7.5.23.10 is\_pure

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `is_pure(R)`,  $R$  representing the module  $M=D^p/D^q(R^t)$

**Return:** int, 0 or 1

**Purpose:** checks pureness of  $M$ .  
returns 1, if  $M$  is pure, or 0, if it's not  
remark: if  $M$  is zero, `is_pure` returns 1

**Example:**

```
LIB "purityfiltration.lib";
ring D = 0, (x,y,z), dp;
matrix R[3][2]=y,-z,x,0,0,x;
list T=purityFiltration(transpose(R));
print(transpose(std(transpose(T[2][2]))));
↳ y,-z,
↳ x,0,
↳ 0,x
// so the purity filtration of coker(R) is trivial,
// i.e. coker(R) is already pure
is_pure(transpose(R));
↳ 1
// we can also have non-pure modules:
matrix R2[6][4]=
0,-2*x,z-2*y-x,-1,
0,z-2*x,2*y-3*x,1,
z,-6*x,-2*y-5*x,-1,
0,y-x,y-x,0,
y,-x,-y-x,0,
x,-x,-2*x,0;
is_pure(transpose(R2));
↳ 0
```

### 7.5.23.11 purelist

Procedure from library `purityfiltration.lib` (see Section 7.5.23 [`purityfiltration.lib`], page 597).

**Usage:** `purelist(T)`,  $T$  list, in which the  $i$ -th entry  $R=T[i]$  represents  $M=D^p/D^q(R^t)$

**Return:** list  $M$ , entries of  $M$  are 0 or 1

**Purpose:** if  $T[i]$  is pure,  $M[i]$  is 1, else  $M[i]$  is 0

**Example:**

```
LIB "purityfiltration.lib";
ring D = 0, (x,y,z), dp;
matrix R[6][4]=
0,-2*x,z-2*y-x,-1,
0,z-2*x,2*y-3*x,1,
z,-6*x,-2*y-5*x,-1,
0,y-x,y-x,0,
y,-x,-y-x,0,
x,-x,-2*x,0;
is_pure(transpose(R));
```

```

⇒ 0
// R is not pure, so we do the purity filtration
list T=purityFiltration(transpose(R));
// all Elements of T[2] are either zero or pure
purelist(T[2]);
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ 1
⇒ [3]:
⇒ 1
⇒ [4]:
⇒ 1

```

### 7.5.24 qmatrix\_lib

**Library:** qmatrix.lib

**Purpose:** Quantum matrices, quantum minors and symmetric groups

**Authors:** Lobillo, F.J., jlobillo@ugr.es  
Rabelo, C., crabelo@ugr.es

**Support:** 'Metodos algebraicos y efectivos en grupos cuanticos', BFM2001-3141, MCYT, Jose Gomez-Torrecillas (Main researcher).

**Procedures:**

#### 7.5.24.1 quantMat

Procedure from library `qmatrix.lib` (see Section 7.5.24 [`qmatrix_lib`], page 606).

**Usage:** `quantMat(n [, p]);` `n` integer ( $n > 1$ ), `p` an optional integer

**Return:** ring (of quantum matrices). If `p` is specified, the quantum parameter `q` will be specialized at the `p`-th root of unity

**Purpose:** compute the quantum matrix ring of order `n`

**Note:** activate this ring with the "setring" command.  
The usual representation of the variables in this quantum algebra is not used because double indexes are not allowed in the variables. Instead the variables are listed by reading the rows of the usual matrix representation, that is, there will be  $n \cdot n$  variables (one for each entry an  $n \cdot N$  generic matrix), listed row-wise

**Example:**

```

LIB "qmatrix.lib";
def r = quantMat(2); // generate O_q(M_2) at q generic
setring r; r;
⇒ // coefficients: QQ(q)
⇒ // number of vars : 4
⇒ // block 1 : ordering Dp
⇒ // : names y(1) y(2) y(3) y(4)
⇒ // block 2 : ordering C
⇒ // noncommutative relations:

```



```

↳ // y(2)y(1)=1/(q)*y(1)*y(2)
↳ // y(3)y(1)=1/(q)*y(1)*y(3)
↳ // y(4)y(1)=y(1)*y(4)+(-q^2+1)/(q)*y(2)*y(3)
↳ // y(4)y(2)=1/(q)*y(2)*y(4)
↳ // y(4)y(3)=1/(q)*y(3)*y(4)
kill r;
def r = quantMat(2,5); // generate 0_q(M_2) at q^5=1
setring r; r;
↳ // coefficients: QQ[q]/(q^4+q^3+q^2+q+1)
↳ // number of vars : 4
↳ // block 1 : ordering Dp
↳ // : names y(1) y(2) y(3) y(4)
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // y(2)y(1)=(-q^3-q^2-q-1)*y(1)*y(2)
↳ // y(3)y(1)=(-q^3-q^2-q-1)*y(1)*y(3)
↳ // y(4)y(1)=y(1)*y(4)+(-q^3-q^2-2*q-1)*y(2)*y(3)
↳ // y(4)y(2)=(-q^3-q^2-q-1)*y(2)*y(4)
↳ // y(4)y(3)=(-q^3-q^2-q-1)*y(3)*y(4)

```

See also: Section 7.5.24.2 [qminor], page 607.

### 7.5.24.2 qminor

Procedure from library `qmatrix.lib` (see Section 7.5.24 [qmatrix.lib], page 606).

**Usage:** `qminor(I,J,n)`; I,J intvec, n int

**Return:** poly, the quantum minor of a generic  $n*n$  quantum matrix

**Assume:** I is the ordered list of the rows to consider in the minor,  
 J is the ordered list of the columns to consider in the minor,  
 I and J must have the same number of elements,  
 n is the order of the quantum matrix algebra you are working with (`quantMat(n)`).  
 The base ring should be constructed using `quantMat`.

**Example:**

```

LIB "qmatrix.lib";
def r = quantMat(3); // let r be a quantum matrix of order 3
setring r;
intvec u = 1,2;
intvec v = 2,3;
intvec w = 1,2,3;
qminor(w,w,3);
↳ y(1)*y(5)*y(9)+(-q)*y(1)*y(6)*y(8)+(-q)*y(2)*y(4)*y(9)+(q^2)*y(2)*y(6)*y(\
 7)+(q^2)*y(3)*y(4)*y(8)+(-q^3)*y(3)*y(5)*y(7)
qminor(u,v,3);
↳ y(2)*y(6)+(-q)*y(3)*y(5)
qminor(v,u,3);
↳ y(4)*y(8)+(-q)*y(5)*y(7)
qminor(u,u,3);
↳ y(1)*y(5)+(-q)*y(2)*y(4)

```

See also: Section 7.5.24.1 [quantMat], page 606.

### 7.5.24.3 SymGroup

Procedure from library `qmatrix.lib` (see Section 7.5.24 [`qmatrix.lib`], page 606).

**Usage:** `SymGroup(n)`;  $n$  an integer (positive)

**Return:** `intmat`

**Purpose:** represent the symmetric group  $S(n)$  via integer vectors (permutations)

**Note:** each row of the output integer matrix is an element of  $S(n)$

**Example:**

```
LIB "qmatrix.lib";
// "S(3)={(1,2,3),(1,3,2),(3,1,2),(2,1,3),(2,3,1),(3,2,1)}";
SymGroup(3);
↪ 1,2,3,
↪ 1,3,2,
↪ 3,1,2,
↪ 2,1,3,
↪ 2,3,1,
↪ 3,2,1
```

See also: Section 7.5.24.5 [`LengthSym`], page 608; Section 7.5.24.4 [`LengthSymElement`], page 608.

### 7.5.24.4 LengthSymElement

Procedure from library `qmatrix.lib` (see Section 7.5.24 [`qmatrix.lib`], page 606).

**Usage:** `LengthSymElement(v)`;  $v$  `intvec`

**Return:** `int`

**Purpose:** determine the length of the permutation given by  $v$  in some  $S(n)$

**Assume:**  $v$  represents an element of  $S(n)$ ; otherwise the output may have no sense

**Example:**

```
LIB "qmatrix.lib";
intvec v=1,3,4,2,8,9,6,5,7,10;
LengthSymElement(v);
↪ 9
```

See also: Section 7.5.24.5 [`LengthSym`], page 608; Section 7.5.24.3 [`SymGroup`], page 608.

### 7.5.24.5 LengthSym

Procedure from library `qmatrix.lib` (see Section 7.5.24 [`qmatrix.lib`], page 606).

**Usage:** `LengthSym(M)`;  $M$  an `intmat`

**Return:** `intvec`

**Purpose:** determine a vector, where the  $i$ -th element is the length of the permutation of  $S(n)$  given by the  $i$ -th row of  $M$

**Assume:**  $M$  represents a subset of  $S(n)$  (each row must be an element of  $S(n)$ ); otherwise, the output may have no sense

**Example:**

```

LIB "qmatrix.lib";
def M = SymGroup(3); M;
 ↪ 1,2,3,
 ↪ 1,3,2,
 ↪ 3,1,2,
 ↪ 2,1,3,
 ↪ 2,3,1,
 ↪ 3,2,1
LengthSym(M);
 ↪ 0,1,2,1,2,3

```

See also: Section 7.5.24.4 [LengthSymElement], page 608; Section 7.5.24.3 [SymGroup], page 608.

### 7.5.25 ratgb\_lib

Status: experimental

**Library:** ratgb.lib

**Purpose:** Groebner bases in Ore localizations of noncommutative G-algebras

**Author:** Viktor Levandovskyy, levandov@risc.uni-linz.ac.at

**Overview:** Theory: Let  $A$  be an operator algebra with  $R = K[x_1, \dots, x_N]$  as subring. The operators are usually denoted by  $d_1, \dots, d_M$ .

Assume, that  $A$  is a  $G$ -algebra, then the set  $S=R-0$  is multiplicatively closed Ore set in  $A$ . That is, for any  $s$  in  $S$  and  $a$  in  $A$ , there exist  $t$  in  $S$  and  $b$  in  $A$ , such that  $sa=bt$ . In other words, one can transform any left fraction into a right fraction. The algebra  $A_S$  is called an Ore localization of  $A$  with respect to  $S$ .

This library provides Groebner basis procedure for  $A_S$ , performing polynomial (that is fraction-free) computations only. Note, that there is ongoing development of the subsystem called Singular:Locapal, which will provide yet another approach to Groebner bases over such Ore localizations.

Assumptions: in order to treat such localizations constructively, some care need to be taken. We will assume that the variables  $x_1, \dots, x_N$  from above (which will become invertible in the localization) come as the first block among the variables of the basering. Moreover, the ordering on the basering must be an antiblock ordering, that is its matrix form has the left upper  $N \times N$  block zero. Here is a recipe to create such an ordering easily: use 'a(w)' definitions of the ordering  $N$  times with intvecs  $w_i$  of the following form:  $w_i$  has first  $N$  components zero. The rest entries need to be positive and such, that  $w_1, \dots, w_N$  are linearly independent (see an example below).

Guide: with this library, it is possible

- to compute a Groebner basis of an ideal or a submodule in the 'rational' Ore localization  $D = A_S$
- to compute a dimension of associated graded submodule (called  $D$ -dimension) - to compute a vector space dimension over  $\text{Quot}(R)$  of a submodule of  $D$ -dimension 0 (so called  $D$ -finite submodule)
- to compute a basis over  $\text{Quot}(R)$  of a  $D$ -finite submodule

**Procedures:** See also: Section D.11.3 [jacobson.lib], page 889; Section 7.5.21 [olga.lib], page 578.

#### 7.5.25.1 ratstd

Procedure from library `ratgb.lib` (see Section 7.5.25 [ratgb.lib], page 609).

**Usage:** `ratstd(I, n [,eng]);` I an ideal/module, n an integer, eng an optional integer

**Return:** ring

**Purpose:** compute the Groebner basis of I in the Ore localization of the basering with respect to the subalgebra, generated by first n variables

**Assume:** the variables of basering are organized in two blocks and - the first block of length n contains the elements with respect to which one localizes, - the basering is equipped with anti-block ordering, giving block dominance for the variables in the second block

**Note:** the output ring C is commutative. The ideal `rGBid` in C represents the rational form of the output ideal `pGBid` in the basering. - During the computation, the D-dimension of I, `Ratgb::Ddim` and the corresponding dimension as  $K(x)$ -vector space of I (`Ratgb::KXdim`, if `Ratgb::Ddim=0`) are computed and exported. - Setting optional integer `eng` to 1, `std` is taken as Groebner engine; default is `slimgb`.

**Display:** In order to see the steps of the computation, set `printlevel` to  $\geq 2$

**Example:**

```
LIB "ratgb.lib";
ring r = (0,c),(x,y,Dx,Dy),(a(0,0,1,1),a(0,0,1,0),dp);
// this ordering is an antiblock ordering, as it must be
def S = Weyl(); setring S;
// the ideal I below annihilates parametric Appel F4 function
// where we set parameters to a=-2, b=-1 and d=0
ideal I =
x*Dx*(x*Dx+c-1) - x*(x*Dx+y*Dy-2)*(x*Dx+y*Dy-1),
y*Dy*(y*Dy-1) - y*(x*Dx+y*Dy-2)*(x*Dx+y*Dy-1);
int is = 2; // hence 1st and 2nd variables, that is x and y
// will become invertible in the localization
def A = ratstd(I,2); // main call
pGBid; // polynomial form of the basis in the localized ring
↳ pGBid[1]=2*x*y*Dx*Dy+x*y*Dy^2+y^2*Dy^2-y*Dy^2+(-c-2)*x*Dx-2*y*Dy+2
↳ pGBid[2]=x*Dx^2-y*Dy^2+(c)*Dx
↳ pGBid[3]=2*x^2*y*Dy^3-4*x*y^2*Dy^3+2*y^3*Dy^3-4*x*y*Dy^3-4*y^2*Dy^3+2*y*D\
y^3+(-2*c)*x^2*Dx*Dy+(2*c)*x*Dx*Dy+2*x^2*Dy^2+(c-2)*x*y*Dy^2+(-3*c)*y^2*D\
y^2-4*x*Dy^2+(3*c-2)*y*Dy^2+2*Dy^2+(-c^2+2*c)*x*Dx+(6*c)*y*Dy+(-6*c)
setring A; // A is a commutative ring used for presentation
rGBid; // "rational" or "localized" form of the basis
↳ rGBid[1]=(2*x*y)*Dx*Dy+(x*y+y^2-y)*Dy^2+(-c*x-2*x)*Dx+(-2*y)*Dy+2
↳ rGBid[2]=(x)*Dx^2+(-y)*Dy^2+(c)*Dx
↳ rGBid[3]=(2*x^2*y-4*x*y^2-4*x*y+2*y^3-4*y^2+2*y)*Dy^3+(-2*c*x^2+2*c*x)*Dx\
*Dy+(c*x*y-3*c*y^2+3*c*y+2*x^2-2*x*y-4*x-2*y+2)*Dy^2+(-c^2*x+2*c*x)*Dx+(6\
*c*y)*Dy+(-6*c)
Ratgb::Ddim; // the Krull-like dimension of A/I
↳ 0
Ratgb::KXdim; // the dimension of A/I as a left K(x,y)-vector space
↳ 4
//--- Now, let us compute a K(x,y) basis explicitly
print(matrix(kbase(rGBid)));
↳ // ** rGBid is no standard basis
↳ Dy^2,Dy,Dx,1
```

## 7.6 Graded commutative algebras (SCA)

This section describes basic mathematical notions, definition, and a little bit the implementation of the experimental non-commutative kernel extension SCA of SINGULAR which improves performance of many algorithms in graded commutative algebras.

In order to improve performance of SINGULAR in specific non-commutative algebras one can extend the internal implementation for them in a virtual-method-overloading-like manner. At the moment graded commutative algebras (SCA) and in particular exterior algebras are implemented this way.

Note that graded commutative algebras require no special user actions apart from defining an appropriate non-commutative GR-algebra in SINGULAR. Upon doing that, the super-commutative structure will be automatically detected and special multiplication will be used. Moreover, in most SCA-aware (e.g. `std`) algorithms special internal improvements will be used (otherwise standard generic non-commutative implementations will be used).

All considered algebras are assumed to be associative  $K$ -algebras for some ground field  $K$ .

Definition

**Polynomial graded commutative algebras** are factors of tensor products of commutative algebras with an exterior algebra over a ground field  $K$ .

These algebras can be naturally endowed with a  $\mathbb{Z}/2\mathbb{Z}$ -grading, where anti-commutative algebra generators have degree 1 and commutative algebra generators (and naturally scalars) have degree 0. In this particular case they may be considered as super-commutative algebras.

GR-algebra representation

A graded commutative algebra with  $n$  commutative and  $m$  anti-commutative algebra generators can be represented as factors of the following GR-algebra by some two-sided ideal:

$$K \langle x_1, \dots, x_n; y_1, \dots, y_m \mid y_j * y_i = -y_i y_j, i < j \rangle / \langle y_1^2, \dots, y_m^2 \rangle.$$

Distinctive features

Graded commutative algebras are Noetherian.

Graded commutative algebras have zero divisors if and only if  $m > 0 : y_i * y_i = 0$ .

Unlike other non-commutative algebras one may use any monomial ordering where only the non-commutative variables are required to be global. In particular, commutative variables are allowed to be local. This means that one can work in tensor products of any commutative ring with an exterior algebra.

Example of defining graded commutative algebras in SINGULAR:SCA and computing with them

Given a commutative polynomial ring  $r$ , super-commutative structure on it can be introduced as follows:

```
LIB "nctools.lib";
ring r = 0,(a, b, x,y,z, Q, W),(lp(2), dp(3), Dp(2));
// Let us make variables x = var(3), ..., z = var(5) to be anti-commutative
// and add additionally a quotient ideal:
def S = superCommutative(3, 5, ideal(a*W + b*Q*x + z)); setring S; S;
⇨ // coefficients: QQ
⇨ // number of vars : 7
⇨ // block 1 : ordering lp
⇨ // : names a b
⇨ // block 2 : ordering dp
⇨ // : names x y z
⇨ // block 3 : ordering Dp
```

```

⇒ // : names Q W
⇒ // block 4 : ordering C
⇒ // noncommutative relations:
⇒ // yx=-xy
⇒ // zx=-xz
⇒ // zy=-yz
⇒ // quotient ring from ideal
⇒ _[1]=xz
⇒ _[2]=bxyQ-yz
⇒ _[3]=aW+bxQ+z
⇒ _[4]=z2
⇒ _[5]=y2
⇒ _[6]=x2
ideal I = a*x*y + z*Q + b, y*Q + a; I;
⇒ I[1]=axy+b+zQ
⇒ I[2]=a+yQ
std(I); // Groebner basis is used here since > is global
⇒ _[1]=yQW-z
⇒ _[2]=yz
⇒ _[3]=b+zQ
⇒ _[4]=a+yQ
kill r, S;
// Let's do the same but this time with some local commutative variables:
ring r = 0,(a, b, x,y,z, Q, W),(dp(1), ds(1), lp(3), ds(2));
def S = superCommutative(3, 5, ideal(a*W + b*Q*x + z)); setring S; S;
⇒ // coefficients: QQ
⇒ // number of vars : 7
⇒ // block 1 : ordering dp
⇒ // : names a
⇒ // block 2 : ordering ds
⇒ // : names b
⇒ // block 3 : ordering lp
⇒ // : names x y z
⇒ // block 4 : ordering ds
⇒ // : names Q W
⇒ // block 5 : ordering C
⇒ // noncommutative relations:
⇒ // yx=-xy
⇒ // zx=-xz
⇒ // zy=-yz
⇒ // quotient ring from ideal
⇒ _[1]=xz
⇒ _[2]=yz-bxyQ
⇒ _[3]=aW+z+bxQ
⇒ _[4]=x2
⇒ _[5]=y2
⇒ _[6]=z2
ideal I = a*x*y + z*Q + b, y*Q + a; I;
⇒ I[1]=axy+zQ+b
⇒ I[2]=a+yQ
std(I);
⇒ _[1]=yQW-z-bxQ
⇒ _[2]=zQ+b

```

```

↳ _[3]=bx
↳ _[4]=by
↳ _[5]=bz
↳ _[6]=b2
↳ _[7]=a+yQ

```

See example of Section 7.5.20.9 [superCommutative], page 565 from the library `nctools.lib`.

Reference: Ph.D thesis by Oleksandr Motsak (2010), <https://nbn-resolving.org/urn:nbn:de:hbz:386-kluedo-26479>.

## 7.7 LETTERPLACE

This section describes mathematical notions and definitions used in the LETTERPLACE subsystem of SINGULAR.

All algebras are assumed to be associative  $K$ -algebras for some field  $K$ .

### What is and what does LETTERPLACE?

What is LETTERPLACE? It is a subsystem of SINGULAR, providing the manipulations and computations within free associative algebras over rings  $R$

$\langle x_1, \dots, x_n \rangle$ , where the coefficient domain  $R$  is either a ring  $Z$  or a field, supported by SINGULAR.

LETTERPLACE can perform computations also in the factor-algebras of the above (via data type `qring`) by two-sided ideals.

Free algebras are internally represented in SINGULAR as so-called Letterplace rings.

Each such ring is constructed from a commutative ring  $R[x_1, \dots, x_n]$  and a **degree (length) bound**  $d$ .

This encodes a sub- $K$ -vector space (also called a filtered part) of  $K$

$\langle x_1, \dots, x_n \rangle$ , spanned by all monomials of **length** at most  $d$ . Analogously for free  $R$ -submodules of a free  $R$ -module.

Within such a construction we offer the computations of Groebner (also known as Groebner-Shirshov) bases, normal forms, syzygies and many more. We address both ideals and submodules of the free bimodule of the fixed rank.

A variety of monomial and module orderings is supported, including **elimination** orderings for both variables and bimodule components. A monomial ordering has to be a well-ordering.

LETTERPLACE works with every field, supported by SINGULAR, and with the coefficient ring  $Z$ . Note, that the elements of the coefficient field (or a ring) mutually commute with all variables.

### 7.7.1 Examples of use of LETTERPLACE

First, define a commutative ring  $K[X]$  in SINGULAR, equipped with a monomial well-ordering and call it, say, `r`.

Then, decide what should be the degree (length) bound  $d$ , that is how long may the words (monomials in the free algebra) become and run the procedure `freeAlgebra(r, d)`.

This procedure creates free algebra  $K \langle X \rangle$  with a monomial ordering, corresponding to the one in the original commutative ring  $K[X]$ , see Section 7.9.2 [Monomial orderings on free algebras], page 625.

Polynomial arithmetics in this  $K$ -algebra is the usual one:  $+$ ,  $-$ ,  $*$  while of course,  $x*y$  and  $y*x$  are different monomials while  $x*7=7*x$ .

Let us define an ideal  $I$  as a list of polynomials in the free algebra and run, for example, `twostd` (see Section 7.8.9 [twostd (letterplace)], page 624). The answer is a two-sided Groebner basis  $J$  of the two-sided ideal  $I$

up to the length bound  $d$ .

Then, we want to compute the two-sided normal form of  $xyzy$  with respect to  $J$  using the function `reduce` (see Section 7.8.5 [reduce (letterplace)], page 622).

We illustrate the approach with the following example:

```
LIB "freegb.lib";
ring r = 0,(x,y,z),dp; // the ordering on the free algebra will be degree right lex
ring R = freeAlgebra(r, 4); // 4 the is degree (length) bound;
ideal I = x*y + y*z, x*x + x*y - z; // define a non-graded ideal
ideal J = twostd(I);
J;
↳ J[1]=x*y+y*z
↳ J[2]=x*x-y*z-z
↳ J[3]=y*z*y-y*z*z+z*y
↳ J[4]=y*z*x+y*z*z+z*x-x*z
↳ J[5]=y*z*z*y-y*z*z*z-x*z*y
↳ J[6]=y*z*z*x+y*z*z*z-x*z*x+y*z*z+z*z
poly p = reduce(x*y*z*y,J);
p; // since p!=0, x*y*z*y is not contained in J
↳ -y*z*z*z-x*z*y

// Now, we introduce a factor algebra K<x,y,z>/J} of type qring,
// and demonstrate the functions reduce and rightstd:

qring Q = J; // J is a Groebner basis
poly p = reduce(x*x, twostd(0)); // the canonical representative of x*x in Q
p;
↳ y*z+z
rightstd(ideal(p)); // right Groebner basis of the right ideal, generated by p in Q
↳ _[1]=z*z
↳ _[2]=y*z+z
↳ _[3]=x*z
```

See Section 7.8 [Functions (letterplace)], page 619 for the list of all available kernel functions.

There are various conversion routines in the library `freegb_lib` (see Section 7.10.4 [freegb\_lib], page 652). Many algebras are predefined in the library `fpalgebras_lib` (see Section 7.10.2 [fpalgebras\_lib], page 634). Important ring-theoretic properties can be established with the help of the library `fpaprops_lib` (see Section 7.10.3 [fpaprops\_lib], page 649), while  $K$ -dimension and monomial bases and Hilbert data - with the help of the library `fpadim_lib` (see Section 7.10.1 [fpadim\_lib], page 629). We work further on implementing more algorithms for non-commutative ideals and modules over free associative algebra.

## 7.7.2 Example of use of LETTERPLACE over $Z$

Consider the following paradigmatic example:

```
LIB "freegb.lib";
ring r = integer,(x,y),Dp;
```



```

ring R = freeAlgebra(r,5); // length bound is 5
ideal I = 2*x, 3*y;
I = twostd(I);
print(matrix(I)); // pretty prints the generators
↳ 3*y,2*x,y*x,x*y

```

As we can see, over  $Z \langle x, y \rangle$  the ideal  $\langle 2x, 3y \rangle$  has a finite Groebner basis and indeed

```

Z < x, y > / < 2x, 3y > =
Z < x, y > / < 2x, 3y, yx, xy > =
Z < x, y > / < 2x, 3y, yx - xy, xy > =
Z[x, y] / < 2x, 3y, xy > holds.

```

Now, we analyze the same ideal in the ring with one more variable  $z$  :

```

LIB "freegb.lib";
ring r = integer, (x,y,z), Dp;
ring R = freeAlgebra(r,5,2); // length bound is 5
ideal I = 2*x, 3*y;
I = twostd(I);
print(matrix(I)); // pretty prints the generators
↳ 3*y,2*x,y*x,x*y,y*z*x,x*z*y,y*z*z*x,x*z*z*y,y*z*z*z*x,x*z*z*z*y

```

Now we see, that this Groebner basis is potentially infinite and the following argument delivers a proof. Namely,  $y * z^i * x$  and

$x * z^i * y$  are present in the ideal for all  $i \geq 0$ . How can we do this? We wish to express  $y * z^i * x$  and

$x * z^i * y$  via the original generators:

```

LIB "freegb.lib";
ring r = integer, (x,y,z), Dp;
ring R = freeAlgebra(r,5,2); // length bound is 5, rank of the free bimodule is 2
ideal I = 2*x, 3*y;
matrix T1 = lift(I, ideal(y*z*x,x*z*y));
print(T1);
↳ -y*z*ncgen(1),-ncgen(1)*z*y,
↳ ncgen(2)*z*x, x*z*ncgen(2)
-y*z*I[1] + I[2]*z*x; // gives y*z*x
↳ y*z*x
matrix T2 = lift(I, ideal(y*z^2*x,x*z^2*y));
print(T2);
↳ -y*z*z*ncgen(1),-ncgen(1)*z*z*y,
↳ ncgen(2)*z*z*x, x*z*z*ncgen(2)
-y*z^2*I[1] + I[2]*z^2*x; // gives y*z^2*x
↳ y*z*z*x

```

The columns of matrices, returned by `lift`, encode the presentation of new elements in terms of generators. From this we conjecture, that in particular

$-y * z^i * (2x) + (3y) * z^i * x = y * z^i * x$  holds for all  $i \geq 0$

and indeed, confirm it via a routine computation by hands.

**Comparing computations over  $\mathbb{Q}$  with computations over  $\mathbb{Z}$ .** In the next example, we first compute over  $\mathbb{Q}$  and a bit later compare the result with computations over  $\mathbb{Z}$ .

```

LIB "freegb.lib"; // initialization of free algebras
ring r = 0, (z,y,x), Dp; // degree left lex ord on z>y>x
ring R = freeAlgebra(r,7); // length bound is 7

```

```

ideal I = y*x - 3*x*y - 3*z, z*x - 2*x*z + y, z*y-y*z-x;
option(redSB); option(redTail); // for minimal reduced GB
option(intStrategy); // avoid divisions by coefficients
ideal J = twostd(I); // compute a two-sided GB of I
J; // prints generators of J
↪ J[1]=4*x*y+3*z
↪ J[2]=3*x*z-y
↪ J[3]=4*y*x-3*z
↪ J[4]=2*y*y-3*x*x
↪ J[5]=2*y*z+x
↪ J[6]=3*z*x+y
↪ J[7]=2*z*y-x
↪ J[8]=3*z*z-2*x*x
↪ J[9]=4*x*x*x+x
LIB "fpadim.lib"; // load the library for K-dimensions
lpMonomialBasis(7,0,J); // compute all monomials
↪ _[1]=1
↪ _[2]=z
↪ _[3]=y
↪ _[4]=x
↪ _[5]=x*x
// of length up to 7 in $\mathbb{Q}\langle x,y,z \rangle/J$

```

As we see, we obtain a nice finite Groebner basis  $J$ . Moreover, from the form of its leading monomials, we conjecture that

$\mathbb{Q}\langle x,y,z \rangle/J$  is finite dimensional  $\mathbb{Q}$ -vector space. We check it with `lpMonomialBasis` and obtain an affirmative answer.

Now, for doing similar computations over  $Z$  one needs to change only the initialization of the ring, the rest stays the same

```

LIB "freegb.lib"; // initialization of free algebras
ring r = integer,(z,y,x),Dp; // Z and deg left lex ord on z>y>x
ring R = freeAlgebra(r,7); // length bound is 7
ideal I = y*x - 3*x*y - 3*z, z*x - 2*x*z + y, z*y-y*z-x;
option(redSB); option(redTail); // for minimal reduced GB
option(intStrategy); // avoid divisions by coefficients
ideal J = twostd(I); // compute a two-sided GB of I
J; // prints generators of J
↪ J[1]=12*x*y+9*z
↪ J[2]=9*x*z-3*y
↪ J[3]=y*x-3*x*y-3*z
↪ J[4]=6*y*y-9*x*x
↪ J[5]=6*y*z+3*x
↪ J[6]=z*x-2*x*z+y
↪ J[7]=z*y-y*z-x
↪ J[8]=3*z*z+2*y*y-5*x*x
↪ J[9]=6*x*x*x-3*y*z
↪ J[10]=4*x*x*y+3*x*z
↪ J[11]=3*x*x*z+3*x*y+3*z
↪ J[12]=2*x*y*y+75*x*x*x+39*y*z+39*x
↪ J[13]=3*x*y*z+3*y*y-3*x*x
↪ J[14]=2*y*y*y+x*x*y+3*x*z
↪ J[15]=2*x*x*x*x+y*y-x*x
↪ J[16]=2*x*x*x*y+3*y*y*z+3*x*y+3*z

```

```

↳ J[17]=x*x*y*z+x*y*y-x*x*x
↳ J[18]=x*y*y*z-y*y*y+x*x*y
↳ J[19]=x*x*x*x*x-y*y*y*z-x*y*y+x*x*x
↳ J[20]=x*x*x*x*z+x*x*x*y+2*y*y*z+x*x*z+3*x*y+3*z
↳ J[21]=x*y*y*y*z-y*y*y*y+x*x*x*x-y*y+x*x
↳ J[22]=y*y*y*z*z-x*x*x*x*y
↳ J[23]=x*y*y*y*y*z-y*y*y*y*y+x*x*y*y*y
↳ J[24]=x*y*y*y*y*y*z-y*y*y*y*y*y+x*x*x*x*y*y+y*y*y*y+x*x*x*x+2*y*y-2*x*x

```

The output has plenty of elements in each degree (which is the same as length because of the degree ordering), what hints at potentially infinite Groebner basis. Indeed, one can show that for every  $i \geq 2$  the ideal  $J$  contains an element with the leading monomial  $xy^i z$ .

### 7.7.3 Functionality and release notes of LETTERPLACE

With the present functionality it is possible to compute two-sided Groebner basis of an arbitrary two-sided ideal in a free associative algebra up to a given degree.

The weights of variables are nonnegative and are determined by the current monomial ordering.

Restrictions/conventions of the LETTERPLACE subsystem:

Since free algebra is not Noetherian, one has to work with explicitly fixed degree (length) bound, up to which a partial Groebner basis will be computed. The initialization routine `freeAlgebra` (`letterplace`) constructs the ring with this bound. For increasing the length bound one needs to define another ring and to use `imap` for mapping the objects over.

All the computations happen up to the explicitly fixed length bound.

The options `redSB`, `redTail` are effective for computations involving Groebner bases,

The options `prot`, `mem` are effective for the whole LETTERPLACE subsystem.

For monomial orderings, which are not compatible with the length, the following error message might appear: `degree bound of Letterplace ring is 11, but at least 12 is needed for this multiplication` In such a situation, activating option(`redSB`), option(`redTail`) and increasing the length (degree) bound might help. Though there are situations, where nothing leads to a finite computation.

Operations for polynomials in Letterplace rings are the usual ones: `+` (addition), `-` (subtraction), `*` (multiplication) and `^` (power).

The functions Section 7.3.2 [bracket], page 330, Section 5.1.88 [maxideal], page 216 and Section 5.1.149 [std], page 266 (an alias for Section 7.8.9 [twostd (letterplace)]), page 624) also work within letterplace rings:

```

LIB "freegb.lib";
ring r = 0,(x,y,z),dp; // the ordering will be degree right lex
ring R = freeAlgebra(r, 5); // degree (length) bound is 5
// maxideal in a letterplace ring:
print(matrix(maxideal(2))); // all monomials of length 2
↳ x*x,y*x,z*x,x*y,y*y,z*y,x*z,y*z,z*z
// bracket in a letterplace ring:
bracket(x,y);
↳ -y*x+x*y
poly f = x*x + x*y - z;
bracket(f,x);
↳ x*y*x-x*x*y-z*x+x*z
bracket(f,x,2);
↳ -x*y*x*x*x+x*x*y*x*x+x*x*x*y*x+x*y*x*y*x-x*x*x*x*y-2*x*y*x*x*y+x*x*y*x*y+\

```

```
z*x*x*x-x*z*x*x-z*x*y*x-x*x*z*x-x*y*z*x+2*z*x*x*y-x*z*x*y+x*x*x*z+2*x*y*x\
*z-x*x*y*z+z*z*x-2*z*x*z+x*z*z
```

Further functionality is provided in the libraries for the LETTERPLACE subsystem: see Section 7.10 [LETTERPLACE libraries], page 629 for details.

In the Section 7.10.4 [freegb.lib], page 652 one finds e.g. Letterplace initialization together with legacy, conversion and convenience tools.

The Section 7.10.1 [fpadim.lib], page 629 contains procedures for computations with vector space basis of a factor algebra including finiteness check and dimension computation.

The Section 7.10.3 [fpaprops.lib], page 649 contains procedures for determining important ring-theoretic properties including Gelfand-Kirillov dimension.

The Section 7.10.2 [fpalgebras.lib], page 634 contains procedures for the generation of various algebras, including group algebras of finitely presented groups in the Letterplace ring.

The Section 7.5.12 [ncfactor.lib], page 480 contains the procedure `ncfactor` for factorizing polynomials in the Letterplace ring.

See Section 7.3.2 [bracket], page 330; Section 5.1.88 [maxideal], page 216; Section 7.8.5 [reduce (letterplace)], page 622; Section 7.8.6 [rightstd (letterplace)], page 623; Section 7.8.7 [std (letterplace)], page 623; Section 7.8.9 [twostd (letterplace)], page 624.

#### 7.7.4 References and history of LETTERPLACE

LETTERPLACE has undergone several stages of development.

The first one, the pure Letterplace implementation for homogeneous ideals, was created by V. Levandovskyy and H. Schoenemann in 2007-2009.

Later in 2010-2014, experiments with advanced (among other, with shift-invariant) data structures were performed by V. Levandovskyy, B. Schnitzler and G. Studzinski, and new libraries for  $K$ -dimension,  $K$ -bases, and Ufnarovskij graph were written.

The next stage started in 2017, when K. Abou Zeid joined the team of H. Schoenemann and V. Levandovskyy. Those recent activities led to the change of interface to the one, usual in the free algebra. The Letterplace data structure is still at heart of the implementation, though not explicitly visible by default. It has been generalized to support  $Z$  as coefficient ring (together with T. Metzloff (RWTH Aachen and INRIA Sophia Antipolis)); to support bimodules and compute syzygies and lifts, to name a few. We are grateful to L. Schmitz (RWTH Aachen) for his contributions to the development.

References:

[LL09]: Roberto La Scala and Viktor Levandovskyy, "Letterplace ideals and non-commutative Groebner bases", Journal of Symbolic Computation, Volume 44, Issue 10, October 2009, Pages 1374-1393, see <http://dx.doi.org/10.1016/j.jsc.2009.03.002>.

[LL13]: Roberto La Scala and Viktor Levandovskyy, "Skew polynomial rings, Groebner bases and the letterplace embedding of the free associative algebra", Journal of Symbolic Computation, Volume 48, Issue 1, January 2013, Pages 1374-1393, see <http://dx.doi.org/10.1016/j.jsc.2012.05.003> and also <http://arxiv.org/abs/1009.4152>.

[LSS13]: Viktor Levandovskyy, Grisha Studzinski and Benjamin Schnitzler, "Enhanced Computations of Groebner Bases in Free Algebras as a New Application of the Letterplace Paradigm", Proc. ISSAC 2013, ACM Press, 259-266, see <https://doi.org/10.1145/2465506.2465948>.

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[Mora16]: Teo Mora, "Solving Polynomial Equation Systems IV: Volume 4, Buchberger Theory and Beyond.", Cambridge University Press, 2016.

[LMZ20]: Viktor Levandovskyy, Tobias Metzloff and Karim Abou Zeid, "Computation of free non-commutative Groebner Bases over  $Z$  with SINGULAR:LETTERPLACE", Proc. ISSAC 2020, accepted

[LSZ20]: Viktor Levandovskyy, Hans Schoenemann and Karim Abou Zeid, "LETTERPLACE - a Subsystem of SINGULAR for computations with free algebras via Letterplace Embedding", Proc. ISSAC 2020, accepted

[LMSZ20]: Viktor Levandovskyy, Tobias Metzloff, Hans Schoenemann and Karim Abou Zeid, "Groebner Bases over  $K < X >$  and  $Z < X >$  in theory and practice", Proc. ICMS 2020, accepted

## 7.8 Functions (letterplace)

This chapter gives a complete reference of all functions and commands of the LETTERPLACE kernel, i.e. all built-in commands (for the numerous LETTERPLACE libraries see Section 7.10 [LETTERPLACE libraries], page 629).

The general syntax of a function is

```
[target =] function_name (<arguments>);
```

Note, that both **Control structures** and **System variables** of LETTERPLACE are the same as of SINGULAR (see Section 5.2 [Control structures], page 285, Section 5.3 [System variables], page 297).

### 7.8.1 freeAlgebra (letterplace)

#### Syntax:

```
freeAlgebra(ring_expression r, int_expression d)
```

**Type:** ring

**Purpose:** Creates a free (letterplace) ring with the variables of the ring  $r$  up to the degree (length) bound  $d$ , with the monomial ordering, determined by those on the ring  $r$ .

**Note:** A letterplace ring has an attribute called `isLetterplaceRing`, which is zero for non-letterplace rings and contains the number of variables of the free algebra it encodes, otherwise.

#### Example:

```
LIB "freegb.lib";
ring r = 0,(x,y,z),dp;
ring R = freeAlgebra(r, 7); // this ordering is degree right lex
R;
↳ // coefficients: QQ
↳ // number of vars : 21
↳ // block 1 : ordering dp
↳ // : names x y z x y z x y z x y z x y z x y z x y z
↳ // block 2 : ordering C
↳ // letterplace ring (block size 3, ncgen count 0)
attrib(R,"isLetterplaceRing");
↳ 3
ring r2 = 0,(x,y,z),lp;
ring R2 = freeAlgebra(r2, 5); // note, that this ordering is NOT left or r
R2;
↳ // coefficients: QQ
```

```

↳ // number of vars : 15
↳ // block 1 : ordering a
↳ // : names x y z x y z x y z x y z x y z
↳ // : weights 1 0 0 1 0 0 1 0 0 1 0 0 1 0 0
↳ // block 2 : ordering a
↳ // : names x y z x y z x y z x y z x y z
↳ // : weights 0 1 0 0 1 0 0 1 0 0 1 0 0 1 0
↳ // block 3 : ordering a
↳ // : names x y z x y z x y z x y z x y z
↳ // : weights 0 0 1 0 0 1 0 0 1 0 0 1 0 0 1
↳ // block 4 : ordering lp
↳ // : names x y z x y z x y z x y z x y z
↳ // block 5 : ordering C
↳ // letterplace ring (block size 3, nongen count 0)
attrib(R2,"isLetterplaceRing");
↳ 3

```

See Section 7.9.2 [Monomial orderings on free algebras], page 625.

### 7.8.2 lift (letterplace)

**Syntax:** lift ( ideal\_expression, subideal\_expression )  
lift ( module\_expression, submodule\_expression )

**Type:** matrix

**Purpose:** computes the transformation matrix which expresses the generators of a subbimodule in terms of the generators of a bimodule.

More precisely, if  $m$  is the module (or ideal),  $sm$  the submodule (or ideal), and  $T$  the transformation matrix returned by lift, then the substitution of each  $ncgen(i)$  in  $T$  by the  $m[i]$  delivers a matrix, say  $N$ . The  $i$ -th generator of  $sm$  is equal to the sum of elements in the  $i$ -th column of  $N$ .

**Note:** Gives a warning if  $sm$  is not a submodule.

**Note:** The procedure Section 7.10.4.12 [testLift], page 659 can be used for testing the result.

**Example:**

```

LIB "freegb.lib";
ring r = 0, (x,y), (c,Dp);
ring R = freeAlgebra(r, 7, 2);
ideal I = std(x*y*x + 1);
print(matrix(I));
↳ x*y-y*x,y*x*x+1
ideal SI = x*I[1]*y + y*x*I[2], I[1]*y*x + I[2]*y;
matrix T = lift(I, SI);
print(T);
↳ y*ncgen(1)*x*x+x*ncgen(1)*y,y*x*ncgen(1)+y*ncgen(1)*x+ncgen(1)*y*x,
↳ y*ncgen(2)*x, y*ncgen(2)
print(matrix(SI)); // the original generators
↳ y*x*y*x*x+x*x*y*y-x*y*x*y+y*x,x*y*y*x+y*x*x*y-y*x*y*x+y
print(matrix(testLift(I,T))); // test for the result of lift
↳ y*x*y*x*x+x*x*y*y-x*y*x*y+y*x,x*y*y*x+y*x*x*y-y*x*y*x+y

```

See Section 4.5 [ideal], page 78; Section 7.8.3 [liftstd (letterplace)], page 621; Section 7.8.8 [syz (letterplace)], page 623; Section 7.8.9 [twostd (letterplace)], page 624.

### 7.8.3 liftstd (letterplace)

**Syntax:** liftstd ( ideal\_expression, matrix\_name )  
 liftstd ( module\_expression, matrix\_name )  
 liftstd ( ideal\_expression, matrix\_name, module\_name )  
 liftstd ( module\_expression, matrix\_name, module\_name )

**Type:** ideal or module

**Purpose:** returns a Groebner basis of a two-sided ideal or a bimodule and the transformation matrix from the given ideal, resp. module, to the Groebner basis from the output. That is, if  $m$  is the module (or ideal),  $sm$  the submodule (or ideal), and  $T$  the transformation matrix returned by lift, then the substitution of each  $ncgen(i)$  in  $T$  by the  $m[i]$  delivers a matrix, say  $N$ . The  $i$ -th generator of  $sm$  is equal to the sum of elements in the  $i$ -th column of  $N$ .  
 In an optional third argument the syzygy bimodule will be returned.

**Example:**

```
LIB "freegb.lib";
ring r = 0, (x,y), (c,Dp);
ring R = freeAlgebra(r, 8, 2);
ideal I = x*y*x + 1;
matrix T; module S;
ideal SI = liftstd(I,T,S);
print(matrix(SI));
↳ x*y-y*x,y*x*x+1
print(matrix(testLift(I,T))); // test for the result of lift
↳ x*y-y*x,y*x*x+1
S; // the bisyzygy module of I
↳ S[1]=[x*y*ncgen(1)*x*y+y*x*x*y*ncgen(1)-y*x*ncgen(1)*y*x-ncgen(1)*y*x*x
+y*ncgen(1)-ncgen(1)*y]
↳ S[2]=[x*x*y*ncgen(1)*x-x*ncgen(1)*y*x*x-x*ncgen(1)+ncgen(1)*x]
↳ S[3]=[x*y*ncgen(1)*x*x*y+y*x*x*x*y*ncgen(1)-y*x*x*ncgen(1)*y*x-ncgen(1)
*x*x*x*y+x*y*ncgen(1)+y*x*ncgen(1)-ncgen(1)*x*y-ncgen(1)*y*x]
↳ S[4]=[x*y*x*y*ncgen(1)*x-ncgen(1)*y*x*y*x*x+y*ncgen(1)*x-ncgen(1)*y*x]
testSyz(I,S);
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0
↳ _[4]=0
```

See Section 4.5 [ideal], page 78; Section 7.8.2 [lift (letterplace)], page 620; Section 7.8.8 [syz (letterplace)], page 623; Section 7.8.9 [twostd (letterplace)], page 624.

### 7.8.4 ncgen

**Syntax:** ncgen ( int\_expression )

**Type:** poly

**Purpose:** returns the  $i$ -th free non-commutative generator of a free bimodule.

**Note:** `ncgen` in bimodules is used together with the commutative `gen`, which encodes the component or the position in a vector.

**Example:**

```

LIB"freegb.lib";
ring r= 0,(x,y),dp;
ring R = freeAlgebra(r,4,3); // R supports free bimodule up to rank 3
typeof(ncgen(2));
↳ poly
vector v = [ncgen(1)*x, x*ncgen(2), y*ncgen(3)*x];
v;
↳ y*ncgen(3)*x*gen(3)+ncgen(1)*x*gen(1)+x*ncgen(2)*gen(2)
print(v*x);
↳ [ncgen(1)*x*x,x*ncgen(2)*x,y*ncgen(3)*x*x]
print(x*v);
↳ [x*ncgen(1)*x,x*x*ncgen(2),x*y*ncgen(3)*x]

```

See Section 7.8.1 [freeAlgebra (letterplace)], page 619; Section 4.13 [module], page 110; Section 4.22 [vector], page 131.

### 7.8.5 reduce (letterplace)

**Syntax:** reduce ( poly\_expression, ideal\_expression )  
 reduce ( poly\_expression, ideal\_expression, int\_expression )  
 reduce ( vector\_expression, ideal\_expression )  
 reduce ( vector\_expression, ideal\_expression, int\_expression )  
 reduce ( vector\_expression, module\_expression, int\_expression )  
 reduce ( ideal\_expression, ideal\_expression )  
 reduce ( ideal\_expression, ideal\_expression, int\_expression )

**Type:** the type of the first argument

**Purpose:** reduces a polynomial, vector, or ideal (the first argument) to its **two-sided** normal form with respect to the second argument, meant to be an ideal, represented by its two-sided Groebner basis (otherwise, the result may have no meaning).

returns 0 if and only if the polynomial (resp. vector, ideal) is an element (resp. subideal) of the ideal.

The third (optional) argument of type int modifies the behavior:

0 default

1 consider only the leading term and do no tail reduction.

2 tail reduction: in the local/mixed ordering case: reduce also with bad ecart

4 reduce without division, return possibly a non-zero constant multiple of the remainder

**Note:** The commands `reduce` and `NF` are synonymous.

**Note:** A two-sided Groebner presentation of a polynomial with respect to a two-sided ideal can be computed by the procedure Section 7.10.4.5 [lpDivision], page 654 from Section 7.10.4 [freegb-lib], page 652.

**Example:**

```

LIB "freegb.lib";
ring r = 0,(x,y),dp;
ring R = freeAlgebra(r,5);
ideal I = x*x + y*y - 1; // 2D sphere
ideal J = twostd(I); // computes a two-sided Groebner basis
J; // it is finite and nice
↳ J[1]=x*x+y*y-1

```



```

↳ J[2]=y*y*x-x*y*y
poly g = x*y*y - y*y*x;
reduce(g,J); // 0, hence g belongs to J
↳ 0
poly h = x*y*y*x - y*x*x;
reduce(h,J); // the rest of two-sided division of h by J
↳ -y*y*y*y+y*y*y+y*y-y
qring Q = J; // swith to K<x,y>/J
reduce(x*y*y - y*y*x,twostd(0)); //image of g above
↳ 0
reduce(x*y*y*x - y*x*x,std(0)); //image of h above
↳ -y*y*y*y+y*y*y+y*y-y

```

See also Section 7.8.2 [lift (letterplace)], page 620.

### 7.8.6 rightstd (letterplace)

**Syntax:** rightstd( ideal\_expression); rightstd( module\_expression);

**Type:** ideal or module

**Purpose:** Compute a right Groebner basis of the set of generators of the input ideal/module.

**Note:** It is effective also in factor rings.

**Example:**

```

LIB "freedb.lib";
ring r = 0,(x,z),dp;
ring R = freeAlgebra(r,7);
ideal I = z, x*z, x*x*z;
rightstd(I); // a right GB of I in K<x,z>
↳ _[1]=z
↳ _[2]=x*z
↳ _[3]=x*x*z
qring Q = twostd(x*z); // now we change to the factor algebra modulo x*z
ideal I = imap(R,I);
rightstd(I); // a right GB in a factor algebra
↳ _[1]=z
reduce(I,twostd(0)); // an explanation for the latter
↳ _[1]=z
↳ _[2]=0
↳ _[3]=0

```

### 7.8.7 std (letterplace)

**Syntax:** std( ideal\_expression); std( module\_expression);

**Type:** ideal or module

**Purpose:** Alias to Section 7.8.9 [twostd (letterplace)], page 624.

### 7.8.8 syz (letterplace)

**Syntax:** syz ( ideal\_expression )  
syz ( module\_expression )

**Type:** module

**Purpose:** computes the first syzygy (i.e., the module of relations of the given generators) bimodule of the ideal, resp. module.

If `option(returnSB)` is set, a Groebner basis is returned, otherwise a generating set.

**Example:**

```
LIB "freegb.lib";
ring r = 0,(x,y),(c,Dp);
ring R = freeAlgebra(r, 7, 2);
ideal I = twostd(x*y*x + 1);
I;
↳ I[1]=x*y-y*x
↳ I[2]=y*x*x+1
module S = syz(I);
print(S);
↳ ncgen(1)*x*x,S[1,2],S[1,3],S[1,4],S[1,5],
↳ S[2,1], S[2,2],S[2,3],S[2,4],S[2,5]
testSyz(I,S);
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0
↳ _[4]=0
↳ _[5]=0
```

See Section 4.5 [ideal], page 78; Section 7.8.2 [lift (letterplace)], page 620; Section 7.8.3 [liftstd (letterplace)], page 621; Section 4.13 [module], page 110; Section 7.8.4 [ncgen], page 621; Section 5.1.110 [option], page 230.

### 7.8.9 twostd (letterplace)

**Syntax:** `twostd(ideal_expression); twostd(module_expression);`

**Type:** ideal

**Purpose:** returns a two-sided Groebner basis of the two-sided ideal, generated by the input, which is treated as a set of two-sided generators.

**Example:**

```
LIB "freegb.lib";
ring r = 3,(x,d),dp; // notice: we work over Z/3Z
ring R = freeAlgebra(r,5);
ideal I = x^4, d^3, d*x - x*d - 1;
twostd(I); // a proper ideal, note x^3 as a generator
↳ _[1]=d*x-x*d-1
↳ _[2]=d*d*d
↳ _[3]=x*x*x
ideal J = x^2, d^3, d*x - x*d - 1;
twostd(J); // the whole ring
↳ _[1]=1
ideal T = twostd(ideal(d*x - x*d - 1));
T;
↳ T[1]=d*x-x*d-1
qring Q = T; // thus Q is the Weyl algebra over Z/3z
ideal I = x^4, d^3;
```

```

twostd(I);
↳ _[1]=d*d*d
↳ _[2]=x*x*x
ideal J = x^2, d^3;
twostd(J);
↳ _[1]=1

```

See Section 4.5 [ideal], page 78; Section 7.8.2 [lift (letterplace)], page 620; Section 7.8.3 [liftstd (letterplace)], page 621; Section 4.13 [module], page 110; Section 7.8.4 [ncgen], page 621; Section 5.1.110 [option], page 230; Section 7.8.6 [rightstd (letterplace)], page 623; Section 7.8.8 [syz (letterplace)], page 623.

## 7.9 Mathematical background (letterplace)

### 7.9.1 Free associative algebras

Let  $V$  be a  $K$ -vector space, spanned by the symbols  $x_1, \dots, x_n$ . A free associative algebra in  $x_1, \dots, x_n$  over  $K$ , denoted by  $K\langle x_1, \dots, x_n \rangle$

$\langle x_1, \dots, x_n \rangle$

is also known as the tensor algebra  $T(V)$  of  $V$ ; it is also the monoid  $K$ -algebra of the free monoid  $\langle x_1, \dots, x_n \rangle$ . The elements of this free monoid constitute an infinite  $K$ -basis of  $K\langle x_1, \dots, x_n \rangle$ , where the identity element (the empty word) of the free monoid is identified with the 1 in  $K$ . Yet in other words, the monomials of  $K\langle x_1, \dots, x_n \rangle$  are the words of finite length in the finite alphabet  $\{x_1, \dots, x_n\}$ .

The algebra  $K\langle x_1, \dots, x_n \rangle$  is an integral domain, which is not (left, right, weak or two-sided) Noetherian for  $n > 1$ ; hence, a Groebner basis of a finitely generated ideal might be infinite. Therefore, a general computation takes place **up to an explicit degree (length) bound**, provided by the user. The free associative algebra can be regarded as a graded algebra in a natural way.

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**Definition.** An associative algebra  $A$  is called **finitely presented (f.p.)**, if it is isomorphic to  $K\langle x_1, \dots, x_n \rangle / I$ , where  $I$  is a two-sided ideal.

$A$  is called **standard finitely presented (s.f.p.)**, if there exists a monomial ordering, such that  $I$  is given via its **finite** Groebner basis  $G$ .

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### 7.9.2 Monomial orderings on free algebras

We provide many types of orderings for non-commutative Groebner bases up to a degree (length) bound. In general it is not clear, whether a given generating set has a finite Groebner bases with respect to some ordering.

Let  $X = \{x_1, \dots, x_n\}$  be a set of symbols. A total ordering  $<$  on the free monoid  $\langle X \rangle$  with 1 as the neutral element is called a **monomial ordering** if

- it is a well-ordering, i.e., every non empty subset has a least element with respect to  $<$ , and
- it is compatible with multiplication, that is  $u < v$  implies  $aub < avb$  for all  $u, v, a$  and  $b$  in  $\langle X \rangle$ .

Note that the latter implies  $1 \leq m$  for all  $m$  in  $\langle X \rangle$ .

The **left lexicographical ordering** on  $\langle X \rangle$  with  $x_1 > x_2 > \dots > x_n$  is defined as follows: For arbitrary  $a, b$  in  $\langle X \rangle$  we say that  $a < b$ , if

$\exists u \in \langle X \rangle \setminus \{1\} : au = b$  or

$\exists u, v, w \in \langle X \rangle \exists x_i, x_j \in X : a = ux_iv, b = ux_jw$  and  $i < j$  holds.

**Note:** left lex is **not** a monomial ordering, though it is a natural choice to break ties after, say, comparing elements by the total degree.

In a similar manner one can define the **right lexicographical ordering**.

On the monoid  $(N_0, +)$  define the **weight** homomorphism  $w : \langle X \rangle \rightarrow N_0$ , uniquely determined by  $w(x_i) = w_i$  in  $N_0$  for  $1 \leq i \leq n$ .

As a special case, define the **length**  $\text{len} : \langle X \rangle \rightarrow N_0$  by  $\text{len}(x_i) = 1$  for  $1 \leq i \leq n$ .

For any ordering  $\ll$  on  $\langle X \rangle$  and any weight  $w : \langle X \rangle \rightarrow N_0$  define an ordering  $<$ , called the  **$w$ -weight extension of  $\ll$**  as follows: For arbitrary  $a, b$  in  $\langle X \rangle$  we say that  $a < b$  if

$w(a) < w(b)$  or

$w(a) = w(b)$  and  $a \ll b$  holds.

An ordering  $<$  on  $\langle X \rangle$  **eliminates** a certain subset  $\emptyset \neq Y \subset X$  if for all  $f \in K\langle X \rangle \setminus \{0\}$  one has  $\text{lm}(f) \in K\langle X \setminus Y \rangle \Rightarrow f \in K\langle X \setminus Y \rangle \subseteq K\langle X \rangle$ .

In a ring declaration, LETTERPLACE supports the following monomial orderings.

We illustrate each of the available choices by an example on the free monoid  $\langle x_1, x_2, x_3 \rangle$ , where we order the monomials

$x_1x_1x_1, x_3x_2x_1, x_1x_2x_3, x_3x_3x_3, x_3x_1, x_2x_2, x_1x_3, x_2x_3, x_1, x_2$  and  $x_3$  correspondingly.

‘dp’ The **degree right lexicographical ordering** is the length-weight extension of the right lexicographical ordering.

With respect to the ordering ‘dp’, the test monomials are ordered as follows:

$x_1x_1x_1 > x_3x_2x_1 > x_1x_2x_3 > x_3x_3x_3 > x_3x_1 > x_2x_2 > x_1x_3 > x_2x_3 > x_1 > x_2 > x_3$

‘Dp’ The **degree left lexicographical ordering** is the length-weight extension of the left lexicographical ordering.

With respect to the ordering ‘Dp’, the test monomials are ordered as follows:

$x_1x_1x_1 > x_1x_2x_3 > x_3x_2x_1 > x_3x_3x_3 > x_1x_3 > x_2x_2 > x_2x_3 > x_3x_1 > x_1 > x_2 > x_3$

‘Wp(w) for intvec w’

The **weighted degree left lexicographical ordering** is the  $w$ -weight extension of the left lexicographical ordering with weight  $w : \langle X \rangle \rightarrow N_0$  uniquely determined by strict positive  $w(x_i) = w_i > 0$ .

With respect to the ordering ‘Wp(1, 2, 1)’, the test monomials are ordered as follows:

$x_1x_2x_3 > x_2x_2 > x_3x_2x_1 > x_1x_1x_1 > x_2x_3 > x_3x_3x_3 > x_1x_3 > x_2 > x_3x_1 > x_1 > x_3$

‘lp’ Let  $w^{(i)}$  be weights uniquely determined by  $w^{(i)}(x_j) = \delta_{i,j}$  for  $1 \leq i, j \leq n$  where  $\delta$  denotes the Kronecker delta. Let  $<_n$  be the  $w^{(n)}$ -weight extension of the left lexicographical ordering on  $\langle X \rangle$  and inductively  $<_i$  be the  $w^{(i)}$ -weight extension of  $<_{i+1}$  for all  $1 \leq i < n$ . The monomial ordering lp corresponds to  $<_1$  and eliminates  $x_1, \dots, x_j$  for all  $1 \leq j < n$ .

The monomial ordering ‘lp’ corresponds to  $<_1$  and eliminates  $\{x_1, \dots, x_j\}$  for all  $1 \leq j < n$ .

With respect to the ordering ‘lp’, the test monomials are ordered as follows:

$x_1x_1x_1 > x_1x_2x_3 > x_3x_2x_1 > x_1x_3 > x_3x_1 > x_1 > x_2x_2 > x_2x_3 > x_2 > x_3x_3x_3 > x_3$

‘**rp**’ Let  $w^{(i)}$  be weights uniquely determined by  $w^{(i)}(x_j) = \delta_{i,j}$  for  $1 \leq i, j \leq n$  where  $\delta$  denotes the Kronecker delta. Let  $<_1$  be the  $w^{(1)}$ -weight extension of the left lexicographical ordering on  $\langle X \rangle$  and inductively  $<_i$  be the  $w^{(i)}$ -weight extension of  $<_{i-1}$  for all  $1 < i \leq n$ . The monomial ordering **rp** corresponds to  $<_n$  and eliminates  $\{x_j, \dots, x_n\}$  for all  $1 < j \leq n$ .

The monomial ordering ‘**rp**’ corresponds to  $<_n$  and eliminates  $\{x_j, \dots, x_n\}$  for all  $1 < j \leq n$ .

With respect to the ordering ‘**rp**’, the test monomials are ordered as follows:

$$x_3x_3x_3 > x_1x_2x_3 > x_3x_2x_1 > x_2x_3 > x_1x_3 > x_3x_1 > x_3 > x_2x_2 > x_2 > x_1x_1x_1 > x_1$$

‘**(a(v), ordering)** for intvec v’

For weight  $v : \langle X \rangle \rightarrow N_0$  determined by  $v(x_i) = v_i \in N_0$  with  $1 \leq i \leq n$  and monomial ordering  $\prec$  on  $\langle X \rangle$ , the  $v$ -weight extension of  $\prec$  corresponds to  $(\mathbf{a}(v), \mathbf{o})$ . As a choice for  $\prec$  there are currently two options implemented, which are **dp** and **Dp**. Notice that this ordering eliminates  $\{x_i \in X \mid v(x_i) \neq 0\}$ .

With respect to the ordering ‘ $(\mathbf{a}(1, 0, 0), \mathbf{Dp})$ ’, the test monomials are ordered as follows:

$$x_1x_1x_1 > x_1x_2x_3 > x_3x_2x_1 > x_1x_3 > x_3x_1 > x_1 > x_3x_3x_3 > x_2x_2 > x_2x_3 > x_2 > x_3$$

With ordering ‘ $(\mathbf{a}(1, 1, 0), \mathbf{Dp})$ ’ one obtains:

$$x_1x_1x_1 > x_1x_2x_3 > x_3x_2x_1 > x_2x_2 > x_1x_3 > x_2x_3 > x_3x_1 > x_1 > x_2 > x_3x_3x_3 > x_3$$

The examples are generated by the following code but with customized orderings denoted above.

```
LIB "freegb.lib";
ring r = 0, (x1,x2,x3),Dp; // variate ordering here
ring R = freeAlgebra(r, 4);
poly wr = x1*x1*x1+x3*x3*x3+x1*x2*x3+x3*x2*x1+x2*x2+x2*x3+x1*x3+x3*x1+x1+x2+x3;
wr; // polynomial will be automatically ordered according to the ordering on R
↳ x1*x1*x1+x1*x2*x3+x3*x2*x1+x3*x3*x3+x1*x3+x2*x2+x2*x3+x3*x1+x1+x2+x3
```

### 7.9.3 Groebner bases for two-sided ideals in free associative algebras

We say that a monomial  $v$  divides (two-sided or bilaterally) a monomial  $w$ , if there exist monomials  $p, s \in X$ , such that  $w = p \cdot v \cdot s$ , in other words  $v$  is a subword of  $w$ .

For a subset  $G \subset K\langle x_1, \dots, x_n \rangle =: T$ , define the **leading ideal of  $G$  to be the two-sided ideal**  $LM(G) = {}_T\langle \{lm(g) \mid g \in G \setminus \{0\}\} \rangle_T \subseteq T$ .

**Let  $<$  be a fixed monomial ordering on  $T$ . We say that a subset  $G \subset I$  is a (two-sided) Groebner basis for the ideal  $I$  with respect to  $<$ , if  $LM(G) = LM(I)$ . That is  $\forall f \in I \setminus \{0\}$  there exists  $g \in G$ , such that  $lm(g)$  divides  $lm(f)$ .**

The notion of **Groebner-Shirshov** basis applies to more general algebraic structures, but means the same as Groebner basis for associative algebras.

Suppose, that the weights of the ring variables are strictly positive. We can interpret these weights as defining a non-standard grading on the ring. If the set of input polynomials is weighted homogeneous with respect to the given weights of the ring variables, then computing up to a weighted degree (and thus, also length) bound  $d$

results in the **truncated Groebner basis  $G(d)$** . In other words, by trimming elements of degree exceeding  $d$  from the complete Groebner basis  $G$ , one obtains precisely  $G(d)$ .

In general, given a set  $G(d)$ , which is the result of Groebner basis computation up to weighted degree bound  $d$ , then it is the complete finite Groebner basis, if and only if  $G(2d - 1) = G(d)$  holds.

**Note:** If the set of input polynomials is **not** weighted homogeneous with respect to the weights of the ring variables, and a Groebner is **not** finite, then actually not much can be said precisely on the properties of the given ideal. By increasing the length bound bigger generating sets will be computed, but in contrast to the weighted homogeneous case some polynomials in of small length first enter the basis after computing up to a much higher length bound.

### 7.9.4 Bimodules and syzygies and lifts

Let  $A = K$

$\langle x_1, \dots, x_n \rangle$  be the free algebra. A free bimodule of rank  $r$  over  $A$  is  $Ae_1A \oplus \dots \oplus Ae_rA$ , where  $e_i$  are the generators of the free bimodule.

NOTE: these  $e_i$  are freely non-commutative with respect to elements of  $A$  except constants from the ground field  $K$ .

The free bimodule of rank 1  $AeA$  surjects onto the algebra  $A$  itself. A two-sided ideal of the algebra  $A$  can be converted to a subbimodule of  $AeA$ .

The **syzygy bimodule** or even **module of bisyzygies** of the given finitely generated subbimodule  $N = \langle g_1, \dots, g_m \rangle \subset \bigoplus_{i=1}^r Ae_iA$  is the kernel of the natural homomorphism of  $A$ -bimodules  $\bigoplus_{j=1}^m Ae_jA \rightarrow \bigoplus_{i=1}^r Ae_iA$ ,  $\epsilon_j \mapsto g_j$ , that is  $\sum_{j=1}^m \sum_k \ell_{jk} \epsilon_j r_{jk} \mapsto \sum_{j=1}^m \sum_k \ell_{jk} g_j r_{jk}$ .

The syzygy bimodule is in general not finitely generated. Therefore as a bimodule, both the set of generators of the syzygy bimodule and its Groebner basis are computed up to a specified length bound.

Given a subbimodule  $N$  of a bimodule  $M$ , the **lift(ing)** process returns a matrix, which encodes the expression of generators  $N_1, \dots, N_s$

in terms of generators of  $M_1, \dots, M_m$  like this:  $N_i = \sum_{j=1}^m \sum_k \ell_{jk} M_j r_{jk} = \sum_{j=1}^m T_{ij} M_j$ ,

where  $T_{ij}$  are elements from the enveloping algebra  $R\langle X \rangle \otimes R\langle X \rangle$ , encoded as elements of the free bimodule of rank  $m$ , namely by using the non-commutative generators of the free bimodule which we call **ncgen**.

### 7.9.5 Letterplace correspondence

The name letterplace has been inspired by the work of Rota and, independently, Feynman.

Already Feynman and Rota encoded the monomials (words) of the free algebra  $x_{i_1} x_{i_2} \dots x_{i_m} \in K\langle x_1, \dots, x_n \rangle$  via the double-indexed letterplace (that is encoding the letter (= variable) and its place in the word) monomials  $x(i_1|1)x(i_2|2) \dots x(i_m|m) \in K[X \times N]$ , where  $X = \{x_1, \dots, x_n\}$  and  $N$  is the semigroup of natural numbers, starting with 1 as the first possible place. Note, that the letterplace algebra  $K[X \times N]$  is an infinitely generated commutative polynomial  $K$ -algebra. Since  $K\langle x_1, \dots, x_n \rangle$  is not Noetherian, it is common to perform the computations with its ideals and modules up to a given degree bound.

Subject to the given degree (length) bound  $d$ , the truncated letterplace algebra  $K[X \times (1, \dots, d)]$  is finitely generated commutative polynomial  $K$ -algebra.

In [LL09] a natural shifting on letterplace polynomials was introduced and used. Indeed, there is 1-to-1 correspondence between two-sided ideals of a free algebra and so-called letterplace ideals in the letterplace algebra, see [LL09], [LL13], [LSS13] and [L14] for details. Note, that first this correspondence was established for graded ideals, but holds more generally for arbitrary ideals and subbimodules of a free bimodule of a finite rank. All the computations internally take place in the Letterplace algebra.

A letterplace monomial of length  $m$  is a monomial of a letterplace algebra, such that its  $m$  places are exactly  $1, 2, \dots, m$ . In particular, such monomials are multilinear with respect to places (i.e.

no place, smaller than the length is omitted or filled more than with one letter). A letterplace polynomial is an element of the  $K$ -vector space, spanned by letterplace monomials. A letterplace ideal is generated by letterplace polynomials subject to two kind of operations:

the  $K$ -algebra operations of the letterplace algebra **and simultaneous shifting of places by any natural number  $n$** .

**Note:** Letterplace correspondence naturally extends to the correspondence over

$R <$

$x_1, \dots, x_n$

$>$ , where  $R$  is a commutative unital ring. The case  $R = Z$  is implemented, in addition to  $R$  being a field.

## 7.10 LETTERPLACE libraries

The content of libraries, created for LETTERPLACE is described in the following subsections.

Use the LIB command for loading of single libraries.

See also Section 7.5.12 [ncfactor\_lib], page 480 for the factorization of polynomials in noncommutative algebras.

### 7.10.1 fpadim\_lib

**Library:** fpadim.lib

**Purpose:** Vector space dimension, basis and Hilbert series for finitely presented algebras (Letterplace)

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**Note:** - basering is a Letterplace ring  
- all intvecs correspond to Letterplace monomials  
- if a degree bound  $d$  is specified,  $d \leq \text{attrib}(\text{basing}, \text{uptodeg})$  holds

In the procedures below, 'iv' stands for intvec representation and 'lp' for the letterplace representation of monomials

**Overview:** Given the free associative algebra  $A = K\langle x_1, \dots, x_n \rangle$  and a (finite or truncated) Groebner basis GB, one is interested in the following data:

- the  $K$ -dimension of  $A/\langle \text{GB} \rangle$  (check for finiteness or explicit value) - the Hilbert series of  $A/\langle \text{GB} \rangle$   
- the explicit monomial  $K$ -basis of  $A/\langle \text{GB} \rangle$

In order to determine these, we need

- the Ufnarovskij graph induced by GB  
- the mistletoes of  $A/\langle \text{GB} \rangle$  (which are special monomials in a basis)

The Ufnarovskij graph is used to determine whether  $A/\langle \text{GB} \rangle$  has finite  $K$ -dimension. One has to check if the graph contains cycles. For the whole theory we refer to [Ufn].

Given a reduced set of monomials  $GB$  one can define the basis tree, whose vertex set  $V$  consists of all normal monomials w.r.t.  $GB$ . For every two monomials  $m_1, m_2$  in  $V$  there is a direct edge from  $m_1$  to  $m_2$ , if and only if there exists  $x_k$  in  $\{x_1, \dots, x_n\}$ , such that  $m_1 * x_k = m_2$ . The set  $M = \{m \text{ in } V \mid \text{there is no edge from } m \text{ to another monomial in } V\}$  is called the set of mistletoes. As one can easily see it consists of the endpoints of the graph. Since there is a unique path to every monomial in  $V$ , the whole graph can be described only from the knowledge of the mistletoes. Note that  $V$  corresponds to a basis of  $A/\langle GB \rangle$ , so knowing the mistletoes we know a  $K$ -basis. The name mistletoes was given to those points because of these miraculous value and the algorithm is named sickle, because a sickle is the tool to harvest mistletoes. For more details see [Stu]. This package uses the Letterplace format introduced by [LL09]. The algebra can either be represented as a Letterplace ring or via integer vectors: Every variable will only be represented by its number, so variable one is represented as 1, variable two as 2 and so on. The monomial  $x_1 * x_3 * x_2$  for example will be stored as (1,3,2). Multiplication is concatenation. Note that the approach in this library does not need an algorithm for computing the normal form. Note that fpa is an acronym for Finitely Presented Algebra.

#### References:

[Ufn] V. Ufnarovskij: Combinatorial and asymptotic methods in algebra, 1990. [LL09] R. La Scala, V. Levandovskyy: Letterplace ideals and non-commutative Groebner bases, Journal of Symbolic Computation, 2009. [Stu] G. Studzinski: Dimension computations in non-commutative, associative algebras, Diploma thesis, RWTH Aachen, 2010.

**Procedures:** See also: Section 7.10.3 [fpaprops\_lib], page 649; Section 7.10.4 [freegb\_lib], page 652; Section 7.10.5 [ncHilb\_lib], page 665.

#### 7.10.1.1 lpKDimCheck

Procedure from library `fpadim.lib` (see Section 7.10.1 [fpadim.lib], page 629).

**Usage:** `lpKDimCheck(G);`

**Return:** `int`, 1 if  $K$ -dimension of the factor algebra is infinite, 0 otherwise

**Purpose:** Checking a factor algebra for finiteness of the  $K$ -dimension

**Assume:** - basering is a Letterplace ring.

#### Example:

```
LIB "fpadim.lib";
ring r = 0,(x,y),dp;
def R = freeAlgebra(r, 5); // constructs a Letterplace ring
setring R; // sets basering to Letterplace ring
ideal G = x*x, y*y,x*y*x;
// Groebner basis
ideal I = x*x, y*x*y, x*y*x;
// Groebner basis
lpKDimCheck(G); // invokes procedure, factor algebra is of finite K-dimension
↳ 0
lpKDimCheck(I); // invokes procedure, factor algebra is of infinite Kdimension
↳ 1
```



### 7.10.1.2 lpKDim

Procedure from library `fpadim.lib` (see Section 7.10.1 [`fpadim.lib`], page 629).

**Usage:** `lpKDim(G[,degbound, n]);` `G` an ideal, `degbound`, `n` optional integers

**Return:** `int`, the  $K$ -dimension of the factor algebra

**Purpose:** Compute the  $K$ -dimension of a factor algebra, given via an ideal

**Assume:** - basering is a Letterplace ring  
 - if you specify a different degree bound `degbound`,  
`degbound <= attrib(basering,uptodeg)` holds.

**Note:** - If `degbound` is set, there will be a degree bound added. 0 means no degree bound. Default: `attrib(basering, uptodeg)`.  
 - `n` is the number of variables, which can be set to a different number. Default: `attrib(basering, IV)`.  
 - If the  $K$ -dimension is known to be infinite, a degree bound is needed

**Example:**

```
LIB "fpadim.lib";
ring r = 0,(x,y),dp;
def R = freeAlgebra(r, 5); // constructs a Letterplace ring
setring R; // sets basering to Letterplace ring
ideal G = x*x, y*y,x*y*x;
// ideal G contains a Groebner basis
lpKDim(G); //procedure invoked with ring parameters
↳ 6
// the factor algebra is finite, so the degree bound given by the Letterplace
// ring is not necessary
lpKDim(G,0); // procedure without any degree bound
↳ 6
```

### 7.10.1.3 lpMonomialBasis

Procedure from library `fpadim.lib` (see Section 7.10.1 [`fpadim.lib`], page 629).

**Usage:** `lpMonomialBasis(d, donly, J);` `d`, `donly` integers, `J` an ideal

**Return:** ideal

**Purpose:** computes a list of free monomials in a Letterplace basering `R` of degree at most `d` and not contained in  $\langle LM(J) \rangle$  if `donly <> 0`, only monomials of degree `d` are returned

**Assume:** - basering is a Letterplace ring.  
 - `d <= attrib(basering,uptodeg)` holds.  
 - `J` is a Groebner basis

**Note:** will be replaced with `reduce(maxideal(d), J)`; soon

**Example:**

```
LIB "fpadim.lib";
ring r = 0,(x,y),dp;
def R = freeAlgebra(r, 7); setring R;
ideal J = x*y*x - y*x*y;
option(redSB); option(redTail);
```

```

J = letplaceGBasis(J);
J;
↳ J[1]=x*y*x-y*x*y
↳ J[2]=y*x*y*y*x-x*y*y*x*y
↳ J[3]=y*x*y*y*y*x-x*x*y*y*x*y
↳ J[4]=y*x*y*y*y*y*x-x*x*x*y*y*x*y
//monomials of degree 2 only in K<x,y>:
lpMonomialBasis(2,1,ideal(0));
↳ _[1]=x*x
↳ _[2]=y*x
↳ _[3]=x*y
↳ _[4]=y*y
//monomials of degree <=2 in K<x,y>
lpMonomialBasis(2,0,ideal(0));
↳ _[1]=1
↳ _[2]=x
↳ _[3]=y
↳ _[4]=x*x
↳ _[5]=y*x
↳ _[6]=x*y
↳ _[7]=y*y
//monomials of degree 3 only in K<x,y>/J
lpMonomialBasis(3,1,J);
↳ _[1]=x*x*x
↳ _[2]=y*x*x
↳ _[3]=y*y*x
↳ _[4]=x*x*y
↳ _[5]=y*x*y
↳ _[6]=x*y*y
↳ _[7]=y*y*y
//monomials of degree <=3 in K<x,y>/J
lpMonomialBasis(3,0,J);
↳ _[1]=1
↳ _[2]=x
↳ _[3]=y
↳ _[4]=x*x
↳ _[5]=y*x
↳ _[6]=x*y
↳ _[7]=y*y
↳ _[8]=x*x*x
↳ _[9]=y*x*x
↳ _[10]=y*y*x
↳ _[11]=x*x*y
↳ _[12]=y*x*y
↳ _[13]=x*y*y
↳ _[14]=y*y*y

```

#### 7.10.1.4 lpHilbert

Procedure from library `fpadim.lib` (see Section 7.10.1 [`fpadim.lib`], page 629).

**Usage:** `lpHilbert(G[,degbound,n]);`  $G$  an ideal, `degbound`,  $n$  optional integers

**Return:** `intvec`, containing the coefficients of the Hilbert series

**Purpose:** Compute the truncated Hilbert series of  $K\langle X \rangle / \langle G \rangle$  up to a degree bound

**Assume:** - basering is a Letterplace ring.  
- if you specify a different degree bound degbound, degbound  $\leq$  attrib(basering,uptodeg) holds.

**Theory:** Hilbert series of an algebra  $K\langle X \rangle / \langle G \rangle$  is  $\sum_{i \geq 0} h_i t^i$ , where  $h_i$  is the K-dimension of the space of monomials of degree  $i$ , not contained in  $\langle G \rangle$ . For finitely presented algebras Hilbert series NEED NOT be a rational function, though it happens often. Therefore in general there is no notion of a Hilbert polynomial.

**Note:** - If degbound is set, there will be a degree bound added. 0 means no degree bound. Default: attrib(basering,uptodeg).  
-  $n$  is the number of variables, which can be set to a different number. Default: attrib(basering, IV).  
- In the output intvec  $I$ ,  $I[k]$  is the  $(k-1)$ -th coefficient of the Hilbert series, i.e.  $h_{(k-1)}$  as above.

**Example:**

```
LIB "fpadim.lib";
ring r = 0, (x,y), dp;
def R = freeAlgebra(r, 5); // constructs a Letterplace ring
setring R; // sets basering to Letterplace ring
ideal G = y*y, x*y*x; // G is a Groebner basis
lpHilbert(G); // procedure with default parameters
 ↪ 1,2,3,4,4,4
lpHilbert(G,3,2); // invokes procedure with degree bound 3 and (same) 2 variables
 ↪ 1,2,3,4
```

See also: Section 7.10.5 [ncHilb\_lib], page 665.

### 7.10.1.5 lpSickleDim

Procedure from library `fpadim.lib` (see Section 7.10.1 [fpadim.lib], page 629).

**Usage:** `lpSickleDim(G[,degbound,n]);`  $G$  an ideal, degbound,  $n$  optional integers

**Return:** list

**Purpose:** Compute the K-dimension and the mistletoes of  $K\langle X \rangle / \langle G \rangle$

**Assume:** - basering is a Letterplace ring.  
- if you specify a different degree bound degbound, degbound  $\leq$  attrib(basering,uptodeg) holds.

**Note:** - If  $L$  is the list returned, then  $L[1]$  is an integer, the K-dimension,  $L[2]$  is an ideal, the mistletoes.  
- If degbound is set, there will be a degree bound added. 0 means no degree bound. Default: attrib(basering,uptodeg).  
-  $n$  is the number of variables, which can be set to a different number. Default: attrib(basering, IV).  
- If the K-dimension is known to be infinite, a degree bound is needed

**Example:**

```
LIB "fpadim.lib";
ring r = 0, (x,y), dp;
def R = freeAlgebra(r, 5); // constructs a Letterplace ring
```

```

setring R; // sets basering to Letterplace ring
ideal G = x*x, y*y,x*y*x; // G is a monomial Groebner basis
lpSickleDim(G); // invokes the procedure with ring parameters
↳ [1]:
↳ 6
↳ [2]:
↳ _[1]=x*y
↳ _[2]=y*x*y
// the factor algebra is finite, so the degree bound, given
// by the Letterplace ring is not necessary
lpSickleDim(G,0); // procedure without any degree bound
↳ [1]:
↳ 6
↳ [2]:
↳ _[1]=x*y
↳ _[2]=y*x*y

```

## 7.10.2 fpalgebras.lib

**Library:** fpalgebras.lib

**Purpose:** Definitions of some finitely presented algebras and groups (Letterplace)

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Support: Project II.6 in the transregional collaborative research centre SFB-TRR 195 'Symbolic Tools in Mathematics and their Application' of the German DFG

**Overview:** Generation of various algebras, including group algebras of finitely presented groups in the Letterplace ring. FPA stands for finitely presented algebra.

**Procedures:** See also: Section 7.7 [LETTERPLACE], page 613; Section 7.10.1 [fpadim.lib], page 629; Section 7.10.3 [fpaprops.lib], page 649; Section 7.10.4 [freegb.lib], page 652.

### 7.10.2.1 operatorAlgebra

Procedure from library fpalgebras.lib (see Section 7.10.2 [fpalgebras.lib], page 634).

**Usage:** operatorAlgebra(a,d); a a string, d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - a gives the name of the algebra

- d gives the degreebound for the Letterplace ring

a must be one of the following:

integrodif3

toeplitz

weyl1

usl2

usl2h

shift1inverse

exterior2

quadrowmm

```

shift1
weyllinverse
This is a collection of common algebras.

```

**Example:**

```

LIB "fpalgebras.lib";
def R = operatorAlgebra("integrodifff3",5); setring R;
I; //relations of the algebra
⇨ I[1]=-x*D+D*x-1
⇨ I[2]=II*II-x*II+II*x
⇨ I[3]=D*II-1

```

**7.10.2.2 serreRelations**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `serreRelations(A,z)`; A an intmat, z an int

**Return:** ideal

**Assume:** basering has a letterplace ring structure and  
A is a generalized Cartan matrix with integer entries

**Purpose:** compute the ideal of Serre's relations associated to A

**Example:**

```

LIB "fpalgebras.lib";
intmat A[3][3] =
2, -1, 0,
-1, 2, -3,
0, -1, 2; // G^1_2 Cartan matrix
ring r = 0,(f1,f2,f3),dp;
int uptodeg = 5;
def R = freeAlgebra(r, uptodeg);
setring R;
ideal I = serreRelations(A,1); I = simplify(I,1+2+8);
I;
⇨ I[1]=f2*f2*f1-2*f2*f1*f2+f1*f2*f2
⇨ I[2]=f3*f1-f1*f3
⇨ I[3]=f2*f1*f1-2*f1*f2*f1+f1*f1*f2
⇨ I[4]=f3*f3*f3*f3*f2-4*f3*f3*f3*f2*f3+6*f3*f3*f2*f3*f3-4*f3*f2*f3*f3*f3+f2\
*f3*f3*f3*f3
⇨ I[5]=f3*f2*f2-2*f2*f3*f2+f2*f2*f3

```

**7.10.2.3 fullSerreRelations**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `fullSerreRelations(A,N,C,P,d)`; A an intmat, N,C,P ideals, d an int

**Return:** ring (and ideal)

**Purpose:** compute the inhomogeneous Serre's relations associated to A in given variable names

**Assume:** three ideals in the input are of the same sizes and contain merely variables which are interpreted as follows: N resp. P stand for negative resp. positive roots, C stand for Cartan elements. d is the degree bound for letterplace ring, which will be returned.

The matrix A is a generalized Cartan matrix with integer entries The result is the ideal called 'fsRel' in the returned ring.

**Example:**

```
LIB "fpalgebras.lib";
intmat A[2][2] =
2, -1,
-1, 2; // A_2 = sl_3 Cartan matrix
ring r = 0,(f1,f2,h1,h2,e1,e2),dp;
ideal negroots = f1,f2; ideal cartans = h1,h2; ideal posroots = e1,e2;
int uptodeg = 5;
def RS = fullSerreRelations(A,negroots,cartans,posroots,uptodeg);
setring RS; fsRel;
↪ fsRel[1]=f2*f2*f1-2*f2*f1*f2+f1*f2*f2
↪ fsRel[2]=f2*f1*f1-2*f1*f2*f1+f1*f1*f2
↪ fsRel[3]=e2*e2*e1-2*e2*e1*e2+e1*e2*e2
↪ fsRel[4]=e2*e1*e1-2*e1*e2*e1+e1*e1*e2
↪ fsRel[5]=e1*f2-f2*e1
↪ fsRel[6]=e2*f1-f1*e2
↪ fsRel[7]=e1*f1-f1*e1-h1
↪ fsRel[8]=e2*f2-f2*e2-h2
↪ fsRel[9]=h2*h1-h1*h2
↪ fsRel[10]=e1*h1-h1*e1+2*e1
↪ fsRel[11]=h1*f1-f1*h1+2*f1
↪ fsRel[12]=e2*h1-h1*e2-e2
↪ fsRel[13]=h1*f2-f2*h1-f2
↪ fsRel[14]=e1*h2-h2*e1-e1
↪ fsRel[15]=h2*f1-f1*h2-f1
↪ fsRel[16]=e2*h2-h2*e2+2*e2
↪ fsRel[17]=h2*f2-f2*h2+2*f2
```

**7.10.2.4 ademRelations**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `ademRelations(i,j); i,j int`

**Return:** ring (and exports ideal)

**Purpose:** compute the ideal of Adem relations for  $i < 2j$  in characteristic 0 the ideal is exported under the name `AdemRel` in the output ring

**Example:**

```
LIB "fpalgebras.lib";
def A = ademRelations(2,5);
setring A;
AdemRel;
↪ 6*s(7)*s(0)+s(6)*s(1)
```

**7.10.2.5 baumslagSolitar**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `baumslagSolitar(m,n,d,[IsGroup]); n an integer, m an integer, d an integer, IsGroup an optional integer`

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - in the group case:  
 $A = a^{-1}$ ,  $B = b^{-1}$   
 - negative input is only allowed in the group case!  
 - d gives a degreebound and must be  $> m, n$   
 - varying n and m produces a family of examples

**Example:**

```
LIB "fpalgebras.lib";
def R = baumslagSolitar(2,3,4); setring R;
I;
↪ I[1]=-b*a*a*a+a*a*b
↪ I[2]=a*A-1
↪ I[3]=b*B-1
↪ I[4]=a*A-A*a
↪ I[5]=b*B-B*b
```

### 7.10.2.6 baumslagGroup

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `baumslagGroup(m,n,d)`; m an integer, n an integer, d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - Baumslag group with the following presentation  
 $\langle a, b \mid a^m = b^n = 1 \rangle$   
 -d gives the degreebound for the Letterplace ring  
 - varying n and m produces a family of examples

**Example:**

```
LIB "fpalgebras.lib";
def R = baumslagGroup(2,3,4); setring R;
I;
↪ I[1]=a*a-1
↪ I[2]=b*b*b-1
```

### 7.10.2.7 crystallographicGroupP1

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP1(d)`; d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - p1 group with the following presentation  
 $\langle x, y \mid [x, y] = 1 \rangle$   
 -d gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP1(5); setring R;
I;
↪ I[1]=y*x+x*y+1
```

```

↳ I[2]=X*x+1
↳ I[3]=x*X+1
↳ I[4]=y*Y+1
↳ I[5]=Y*y+1

```

### 7.10.2.8 crystallographicGroupPM

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupPM(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations - pm group with the following presentation  
 $\langle x, y, m \mid [x, y] = m^2 = 1, m^{(-1)} * x * m = x, m^{(-1)} * y * m = y^{(-1)} \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```

LIB "fpalgebras.lib";
def R = crystallographicGroupPM(5); setring R;
I;
↳ I[1]=y*x+x*y+1
↳ I[2]=y*x+x*y+m*m
↳ I[3]=m*m+1
↳ I[4]=m*x*m+x
↳ I[5]=m*y*m+Y
↳ I[6]=x*X+1
↳ I[7]=X*x+1
↳ I[8]=Y*y+1
↳ I[9]=y*Y+1

```

### 7.10.2.9 crystallographicGroupPG

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupPG(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations - pg group with the following presentation  
 $\langle x, y, t \mid [x, y] = 1, t^2 = x, t^{(-1)} * y * t = y^{(-1)} \rangle$   
-  $d$  gives the degreebound for the Letterplace ring

**Example:**

```

LIB "fpalgebras.lib";
def R = crystallographicGroupPG(5); setring R;
I;
↳ I[1]=y*x+x*y+1
↳ I[2]=t*t+x
↳ I[3]=T*y*t+Y
↳ I[4]=X*x+1
↳ I[5]=x*X+1
↳ I[6]=Y*y+1
↳ I[7]=y*Y+1
↳ I[8]=t*T+1
↳ I[9]=T*t+1

```



### 7.10.2.10 crystallographicGroupP2MM

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP2MM(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations -  $p2mm$  group with the following presentation  
 $\langle x, y, p, q \mid [x, y] = [p, q] = p^2 = q^2 = 1, p^{(-1)}*x*p = x, q^{(-1)}*x*q = x^{(-1)}, p^{(-1)}*y*p = y^{(-1)}, q^{(-1)}*y*q = y \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP2MM(5); setring R;
I;
↪ I[1]=y*x+x*y+1
↪ I[2]=q*p+p*q+1
↪ I[3]=p*p+1
↪ I[4]=q*q+1
↪ I[5]=p*y*p+Y
↪ I[6]=p*x*p+x
↪ I[7]=q*y*q+y
↪ I[8]=q*x*q+X
↪ I[9]=X*x+1
↪ I[10]=x*X+1
↪ I[11]=Y*y+1
↪ I[12]=y*Y+1
↪ I[13]=y*x+x*y+p*p
↪ I[14]=y*x+x*y+q*q
↪ I[15]=p*p+q*q
```

### 7.10.2.11 crystallographicGroupP2

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP2(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations -  $p2$  group with the following presentation  
 $\langle x, y, m, t \mid [x, y] = t^2 = 1, m^2 = y, t^{(-1)}*x*t = x, m^{(-1)}*x*m = x^{(-1)}, t^{(-1)}*y*t = y^{(-1)}, t^{(-1)}*m*t = m^{(-1)} \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP2(5); setring R;
I;
↪ I[1]=y*x+x*y+1
↪ I[2]=y*x+x*y+t*t
↪ I[3]=m*m+y
↪ I[4]=t*t+1
```

```

↳ I[5]=t*x*t+x
↳ I[6]=M*x*m+X
↳ I[7]=t*y*t+Y
↳ I[8]=t*m*t+M
↳ I[9]=X*x+1
↳ I[10]=x*X+1
↳ I[11]=Y*y+1
↳ I[12]=y*Y+1
↳ I[13]=m*M+1
↳ I[14]=M*m+1

```

### 7.10.2.12 crystallographicGroupP2GG

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP2GG(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations - p2gg group with the following presentation  
 $\langle x, y, u, v \mid [x, y] = (u*v)^2 = 1, u^2 = x, v^2 = y, v^{(-1)}*x*v = x^{(-1)}, u^{(-1)}*y*u = y^{(-1)} \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```

LIB "fpalgebras.lib";
def R = crystallographicGroupP2GG(5); setring R;
I;
↳ I[1]=y*x+x*y+1
↳ I[2]=u*v*u*v+y*x+x*y
↳ I[3]=u*v*u*v+1
↳ I[4]=u*u+x
↳ I[5]=v*v+y
↳ I[6]=V*x*v+X
↳ I[7]=U*y*u+Y
↳ I[8]=X*x+1
↳ I[9]=x*X+1
↳ I[10]=Y*y+1
↳ I[11]=y*Y+1
↳ I[12]=u*U+1
↳ I[13]=U*u+1
↳ I[14]=v*V+1
↳ I[15]=V*v+1

```

### 7.10.2.13 crystallographicGroupCM

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupCM(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations - cm group with the following presentation  
 $\langle x, y, t \mid [x, y] = t^2 = 1, t^{(-1)}*x*t = x*y, t^{(-1)}*y*t = y^{(-1)} \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```

LIB "fpalgebras.lib";
def R = crystallographicGroupCM(5); setring R;
I;
↪ I[1]=y*x+x*y+1
↪ I[2]=y*x+x*y+t*t
↪ I[3]=t*t+1
↪ I[4]=t*x*t+x*y
↪ I[5]=t*y*t+Y
↪ I[6]=X*x+1
↪ I[7]=x*X+1
↪ I[8]=Y*y+1
↪ I[9]=y*Y+1

```

**7.10.2.14 crystallographicGroupC2MM**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupC2MM(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations -  $c2mm$  group with the following presentation  
 $\langle x, y, m, r \mid [x, y] = m^2 = r^2 = 1, m^{-1}y^*m = y^*(-1), m^{-1}x^*m = x^*y, r^{-1}y^*r = y^*(-1), r^{-1}x^*r = x^*(-1), m^{-1}r^*m = r^*(-1) \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```

LIB "fpalgebras.lib";
def R = crystallographicGroupC2MM(5); setring R;
I;
↪ I[1]=y*x+x*y+1
↪ I[2]=y*x+x*y+m*m
↪ I[3]=y*x+x*y+r*r
↪ I[4]=m*m+1
↪ I[5]=r*r+1
↪ I[6]=m*m+r*r
↪ I[7]=m*y*m+Y
↪ I[8]=m*x*m+x*y
↪ I[9]=r*y*r+Y
↪ I[10]=r*x*r+X
↪ I[11]=m*r*m+r
↪ I[12]=X*x+1
↪ I[13]=x*X+1
↪ I[14]=Y*y+1
↪ I[15]=y*Y+1

```

**7.10.2.15 crystallographicGroupP4**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP4(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - p4 group with the following presentation  
 $\langle x, y, r \mid [x, y] = r^4 = 1, r^{(-1)}*x*r = x^{(-1)}, r^{(-1)}*x*r = y \rangle$  - d gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP4(5); setring R;
I;
⇒ I[1]=y*x+x*y+1
⇒ I[2]=r*r*r*r+y*x+x*y
⇒ I[3]=r*r*r*r+1
⇒ I[4]=r*r*r*x*r+X
⇒ I[5]=r*r*r*x*r+y
⇒ I[6]=X*x+1
⇒ I[7]=x*X+1
⇒ I[8]=Y*y+1
⇒ I[9]=y*Y+1
```

### 7.10.2.16 crystallographicGroupP4MM

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP4MM(d)`; d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - p4mm group with the following presentation  
 $\langle x, y, r, m \mid [x, y] = r^4 = m^2 = 1, r^{(-1)}*y*r = x^{(-1)}, r^{(-1)}*x*r = y, m^{(-1)}*x*m = y, m^{(-1)}*r*m = r^{(-1)} \rangle$  - d gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP4MM(5); setring R;
I;
⇒ I[1]=y*x+x*y+1
⇒ I[2]=r*r*r*r+y*x+x*y
⇒ I[3]=r*r*r*r+1
⇒ I[4]=r*r*r*x*r+X
⇒ I[5]=r*r*r*x*r+y
⇒ I[6]=X*x+1
⇒ I[7]=x*X+1
⇒ I[8]=Y*y+1
⇒ I[9]=y*Y+1
```

### 7.10.2.17 crystallographicGroupP4GM

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP4GM(d)`; d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - p4gm group with the following presentation

$\langle x, y, r, t \mid [x, y] = r^4 = t^2 = 1, r^{(-1)}*y*r = x^{(-1)}, r^{(-1)}*x*r = y, t^{(-1)}*x*t = y, t^{(-1)}*r*t = x^{(-1)}*r^{(-1)} \rangle$  - d gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP4GM(5); setring R;
I;
→ I[1]=y*x+x*y+1
→ I[2]=r*r*r*r+y*x+x*y
→ I[3]=r*r*r*r+1
→ I[4]=y*x+x*y+t*t
→ I[5]=t*t+1
→ I[6]=r*r*r*r+t*t
→ I[7]=r*r*r*y*r+X
→ I[8]=r*r*r*x*r+y
→ I[9]=X*r*r*r+t*r*t
→ I[10]=X*x+1
→ I[11]=x*X+1
→ I[12]=Y*y+1
→ I[13]=y*Y+1
```

**7.10.2.18 crystallographicGroupP3**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP3(d)`; d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - p3 group with the following presentation

$\langle x, y, r \mid [x, y] = r^3 = 1, r^{(-1)}*x*r = x^{(-1)}*y, r^{(-1)}*y*r = x^{(-1)} \rangle$  - d gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP3(5); setring R;
I;
→ I[1]=y*x+x*y+1
→ I[2]=r*r*r+y*x+x*y
→ I[3]=r*r*r+1
→ I[4]=r*r*x*r+X*y
→ I[5]=r*r*y*r+X
→ I[6]=X*x+1
→ I[7]=x*X+1
→ I[8]=Y*y+1
→ I[9]=y*Y+1
```

**7.10.2.19 crystallographicGroupP31M**

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP31M(d)`; d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - p31m group with the following presentation  
 $\langle x, y, r, t \mid [x, y] = r^2 = t^2 = (t^*r)^3 = 1, r^{(-1)}x^*r = x, t^{(-1)}y^*t = y, t^{(-1)}x^*t = x^{(-1)}y, r^{(-1)}y^*r = x^*y^{(-1)} \rangle$  - d gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP31M(6); setring R;
I;
⇨ I[1]=y*x+x*y+1
⇨ I[2]=y*x+x*y+r*r
⇨ I[3]=y*x+x*y+t*t
⇨ I[4]=r*r+1
⇨ I[5]=t*t+1
⇨ I[6]=t*r*t*r*t*r+1
⇨ I[7]=r*r+t*t
⇨ I[8]=t*r*t*r*t*r+y*x+x*y
⇨ I[9]=t*r*t*r*t*r+r*r
⇨ I[10]=t*r*t*r*t*r+t*t
⇨ I[11]=r*x*r+x
⇨ I[12]=t*y*t+y
⇨ I[13]=t*x*t+X*y
⇨ I[14]=r*y*r+x*Y
⇨ I[15]=X*x+1
⇨ I[16]=x*X+1
⇨ I[17]=Y*y+1
⇨ I[18]=y*Y+1
```

### 7.10.2.20 crystallographicGroupP3M1

Procedure from library fpalgebras.lib (see Section 7.10.2 [fpalgebras.lib], page 634).

**Usage:** crystallographicGroupP3M1(d); d an integer

**Return:** ring

**Note:** - the ring contains the ideal I, which contains the required relations - p3m1 group with the following presentation  
 $\langle x, y, r, m \mid [x, y] = r^3 = m^2 = 1, m^{(-1)}r^*m = r^2, r^{(-1)}x^*r = x^{(-1)}y, r^{(-1)}y^*r = x^{(-1)}, m^{(-1)}x^*m = x^{(-1)}, m^{(-1)}y^*m = x^{(-1)}y \rangle$  - d gives the degreebound for the Letterplace ring

**Example:**

```
LIB "fpalgebras.lib";
def R = crystallographicGroupP3M1(5); setring R;
I;
⇨ I[1]=y*x+x*y+1
⇨ I[2]=r*r*r+y*x+x*y
⇨ I[3]=y*x+x*y+m*m
⇨ I[4]=r*r*r+1
⇨ I[5]=m*m+1
⇨ I[6]=r*r*r+m*m
⇨ I[7]=m*r*m+r*r
⇨ I[8]=r*r*x*r+X*y
⇨ I[9]=r*r*y*r+X
```

```

↳ I[10]=m*x*m+X
↳ I[11]=m*y*m+X*y
↳ I[12]=X*x+1
↳ I[13]=x*X+1
↳ I[14]=Y*y+1
↳ I[15]=y*Y+1

```

### 7.10.2.21 crystallographicGroupP6

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP6(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations -  $p_6$  group with the following presentation

$\langle x, y, r \mid [x, y] = r^6 = 1, r^{(-1)}*x*r = y, r^{(-1)}*y*r = x^{(-1)}*y \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```

LIB "fpalgebras.lib";
def R = crystallographicGroupP6(7); setring R;
I;
↳ I[1]=y*x+x*y+1
↳ I[2]=r*r*r*r*r*r+y*x+x*y
↳ I[3]=r*r*r*r*r*r+1
↳ I[4]=r*r*r*r*r*r*x*r+y
↳ I[5]=r*r*r*r*r*r*y*r+X*y
↳ I[6]=X*x+1
↳ I[7]=x*X+1
↳ I[8]=Y*y+1
↳ I[9]=y*Y+1

```

### 7.10.2.22 crystallographicGroupP6MM

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `crystallographicGroupP6MM(d)`;  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations -  $p_6mm$  group with the following presentation

$\langle x, y, r, m \mid [x, y] = r^6 = m^2 = 1, r^{(-1)}*y*r = x^{(-1)}*y, r^{(-1)}*x*r = y, m^{(-1)}*x*m = x^{(-1)}, m^{(-1)}*y*m = x^{(-1)}*y, m^{(-1)}*r*m = r^{(-1)}*y \rangle$  -  $d$  gives the degreebound for the Letterplace ring

**Example:**

```

LIB "fpalgebras.lib";
def R = crystallographicGroupP6MM(7); setring R;
I;
↳ I[1]=y*x+x*y+1
↳ I[2]=r*r*r*r*r*r+y*x+x*y
↳ I[3]=r*r*r*r*r*r+1
↳ I[4]=y*x+x*y+m*m

```

```

↳ I[5]=r*r*r*r*r*r+m*m
↳ I[6]=m*m+1
↳ I[7]=m*x*m+X
↳ I[8]=m*y*m+X*y
↳ I[9]=r*r*r*r*r*x*r+y
↳ I[10]=r*r*r*r*r*y*r+X*y
↳ I[11]=r*r*r*r*r*y+m*r*m
↳ I[12]=X*x+1
↳ I[13]=x*X+1
↳ I[14]=Y*y+1
↳ I[15]=y*Y+1

```

### 7.10.2.23 dyckGroup1

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `dyckGroup1(n,d,P)`;  $n$  an integer,  $d$  an integer,  $P$  an intvec

**Return:** ring

**Note:**

- the ring contains the ideal  $I$ , which contains the required relations - The Dyck group with the following presentation
- $\langle x_1, x_2, \dots, x_n \mid (x_1)^{p_1} = (x_2)^{p_2} = \dots = (x_n)^{p_n} = x_1 * x_2 * \dots * x_n = 1 \rangle$  - negative exponents are allowed
- representation in the form  $x_i^{p_i} - x_{(i+1)}^{p_{(i+1)}}$
- $d$  gives the degreebound for the Letterplace ring
- varying  $n$  and  $P$  produces a family of examples

**Example:**

```

LIB "fpalgebras.lib";
intvec P = 1,2,3;
def R = dyckGroup1(3,5,P); setring R;
I;
↳ I[1]=x(2)*x(2)+x(1)
↳ I[2]=x(3)*x(3)*x(3)+x(2)*x(2)
↳ I[3]=x(1)*x(2)*x(3)+x(3)*x(3)*x(3)
↳ I[4]=x(1)*x(2)*x(3)+1

```

### 7.10.2.24 dyckGroup2

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `dyckGroup2(n,d,P)`;  $n$  an integer,  $d$  an integer,  $P$  an intvec

**Return:** ring

**Note:**

- the ring contains the ideal  $I$ , which contains the required relations - The Dyck group with the following presentation
- $\langle x_1, x_2, \dots, x_n \mid (x_1)^{p_1} = (x_2)^{p_2} = \dots = (x_n)^{p_n} = x_1 * x_2 * \dots * x_n = 1 \rangle$  - negative exponents are allowed
- representation in the form  $x_i^{p_i} - 1$
- $d$  gives the degreebound for the Letterplace ring
- varying  $n$  and  $P$  produces a family of examples

**Example:**



```

LIB "fpalgebras.lib";
intvec P = 1,2,3;
def R = dyckGroup2(3,5,P); setring R;
I;
↪ I[1]=x(1)+1
↪ I[2]=x(2)*x(2)+1
↪ I[3]=x(3)*x(3)*x(3)+1
↪ I[4]=x(1)*x(2)*x(3)+1

```

### 7.10.2.25 dyckGroup3

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `dyckGroup2(n,d,P)`;  $n$  an integer,  $d$  an integer,  $P$  an intvec

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations - The Dyck group with the following presentation  $\langle x_1, x_2, \dots, x_n \mid (x_1)^{p_1} = (x_2)^{p_2} = \dots = (x_n)^{p_n} = x_1 * x_2 * \dots * x_n = 1 \rangle$  - only positive exponents are allowed  
 - no inverse generators needed  
 -  $d$  gives the degreebound for the Letterplace ring  
 - varying  $n$  and  $P$  produces a family of examples

**Example:**

```

LIB "fpalgebras.lib";
intvec P = 1,2,3;
def R = dyckGroup3(3,5,P); setring R;
I;
↪ I[1]=x(1)+1
↪ I[2]=x(2)*x(2)+1
↪ I[3]=x(3)*x(3)*x(3)+1
↪ I[4]=x(1)*x(2)*x(3)+1

```

### 7.10.2.26 fibonacciGroup

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `fibonacciGroup(m,d)`;  $m$  an integer,  $d$  an integer

**Return:** ring

**Note:** - the ring contains the ideal  $I$ , which contains the required relations - The Fibonacci group  $F(2, m)$  with the following presentation  $\langle x_1, x_2, \dots, x_m \mid x_i * x_{(i+1)} = x_{(i+2)} \rangle$   
 -  $d$  gives the degreebound for the Letterplace ring  
 - varying  $m$  produces a family of examples

**Example:**

```

LIB "fpalgebras.lib";
def R = fibonacciGroup(3,5); setring R;
I;
↪ I[1]=x(1)*x(2)+x(3)
↪ I[2]=x(1)*Y(1)+1
↪ I[3]=Y(1)*x(1)+1

```

```

↳ I[4]=x(2)*Y(2)+1
↳ I[5]=Y(2)*x(2)+1
↳ I[6]=x(3)*Y(3)+1
↳ I[7]=Y(3)*x(3)+1

```

### 7.10.2.27 tetrahedronGroup

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `tetrahedronGroup(g,d)`; `g` an integer, `d` an integer

**Return:** ring

**Note:** - the ring contains the ideal `I`, which contains the required relations - `g` gives the number of the example (1 - 5)  
 - `d` gives the degreebound for the Letterplace ring  
 - varying `g` produces a family of examples

The examples are found in “Classification of the finite generalized tetrahedron groups” by Gerhard Rosenberger and Martin Scheer.

The 5 examples originate from Proposition 1.9 and describe finite generalized tetrahedron group in the Tsaranov-case, which are not equivalent to a presentation for an ordinary tetrahedron group.

**Example:**

```

LIB "fpalgebras.lib";
def R = tetrahedronGroup(3,5); setring R;
I;
↳ I[1]=x*x*x+1
↳ I[2]=y*y*y+1
↳ I[3]=z*z*z+1
↳ I[4]=x*y*x*y+1
↳ I[5]=x*z*x*z+1
↳ I[6]=y*z*y*z+1

```

### 7.10.2.28 triangularGroup

Procedure from library `fpalgebras.lib` (see Section 7.10.2 [`fpalgebras.lib`], page 634).

**Usage:** `triangularGroup(g,d)`; `g` an integer, `d` an integer

**Return:** ring

**Note:** - the ring contains the ideal `I`, which contains the required relations - `g` gives the number of the example (1 - 14)  
 - `d` gives the degreebound for the Letterplace ring  
 - varying `g` produces a family of examples

The examples are found in

Classification of the finite generalized tetrahedron groups by Gerhard Rosenberger and Martin Scheer.

The 14 examples are denoted in theorem 2.12

**Example:**

```

LIB "fpalgebras.lib";
def R = triangularGroup(3,10); setring R;
I;

```

```

↳ I[1]=a*a*a+1
↳ I[2]=b*b*b+1
↳ I[3]=a*b*a*b*b*a*b*a*b*b+1

```

### 7.10.3 fpaprops\_lib

**Library:** fpaprops.lib

**Purpose:** Algorithmic ring-theoretic properties of finitely presented algebras (Letterplace)

**Authors:** Karim Abou Zeid, karim.abou.zeid at rwth-aachen.de

Support: Project II.6 in the transregional collaborative research centre SFB-TRR 195 'Symbolic Tools in Mathematics and their Application' of the German DFG

**Overview:** In this library, algorithms for computing various ring-theoretic properties of finitely presented algebras are implemented.

Applicability: Letterplace rings.

**References:**

Huishi Li: Groebner bases in ring theory. World Scientific, 2010.

**Procedures:** See also: Section 7.10.1 [fpadim.lib], page 629; Section 7.10.4 [freegb.lib], page 652.

#### 7.10.3.1 lpNoetherian

Procedure from library fpaprops.lib (see Section 7.10.3 [fpaprops.lib], page 649).

**Usage:** lpNoetherian(G); G an ideal in a Letterplace ring

**Return:** int  
 0 not Noetherian  
 1 left Noetherian  
 2 right Noetherian  
 3 Noetherian  
 4 weak Noetherian

**Purpose:** Check whether the monomial algebra  $A/\langle LM(G) \rangle$  is (left/right) noetherian

**Assume:** - basering is a Letterplace ring  
 - G is a Groebner basis

**Theory:** lpNoetherian works with the monomial algebra  $A/\langle LM(G) \rangle$ . If it gives an affirmative answer for one of the properties, then it holds for both  $A/\langle LM(G) \rangle$  and  $A/\langle G \rangle$ . However, a negative answer applies only to  $A/\langle LM(G) \rangle$  and not necessarily to  $A/\langle G \rangle$ .

**Note:** Weak Noetherian means that two-sided ideals in  $A/\langle LM(G) \rangle$  satisfy the acc (ascending chain condition).

**Example:**

```

LIB "fpaprops.lib";
ring r = 0, (x,y), dp;
def R = freeAlgebra(r, 5);
setring R;
ideal G = x*x, y*x; // K<x,y>/<xx,yx> is right noetherian
lpNoetherian(G);
↳ 2

```

### 7.10.3.2 lpIsSemiPrime

Procedure from library `fpaprops.lib` (see Section 7.10.3 [`fpaprops.lib`], page 649).

**Usage:** `lpIsSemiPrime(G)`;  $G$  an ideal in a Letterplace ring

**Return:** boolean

**Purpose:** Check whether  $A/\langle LM(G) \rangle$  is semi-prime ring, alternatively whether  $\langle LM(G) \rangle$  is a semi-prime ideal in  $A$ .

**Assume:** - basering is a Letterplace ring  
-  $G$  is a Groebner basis

**Theory:** A (two-sided) ideal  $I$  in the ring  $A$  is semi-prime, if for any  $a$  in  $A$  one has  $aAa \subseteq I$  implies  $a \in I$ .

**Note:** `lpIsSemiPrime` works with the monomial algebra  $A/\langle LM(G) \rangle$ . A positive answer holds for both  $A/\langle LM(G) \rangle$  and  $A/\langle G \rangle$ , while a negative answer applies only to  $A/\langle LM(G) \rangle$  and not necessarily to  $A/\langle G \rangle$ .

**Example:**

```
LIB "fpaprops.lib";
ring r = 0, (x1, x2), dp;
def R = freeAlgebra(r, 5);
setring R;
ideal G = x1*x2, x2*x1; // K<x1,x2>/<x1*x2,x2*x1> is semi prime
lpIsSemiPrime(G);
↪ 1
```

### 7.10.3.3 lpIsPrime

Procedure from library `fpaprops.lib` (see Section 7.10.3 [`fpaprops.lib`], page 649).

**Usage:** `lpIsPrime(G)`;  $G$  an ideal in a Letterplace ring

**Return:** boolean

**Purpose:** Check whether  $A/\langle LM(G) \rangle$  is prime ring, alternatively whether  $\langle LM(G) \rangle$  is a prime ideal in  $A$ .

**Assume:** - basering is a Letterplace ring  
-  $G$  is a Groebner basis

**Theory:** A (two-sided) ideal  $I$  in the ring  $A$  is prime, if for any  $a, b$  in  $A$  one has  $aAb \subseteq I$  implies  $a \in I$  or  $b \in I$ .

**Note:** `lpIsPrime` works with the monomial algebra  $A/\langle LM(G) \rangle$ . A positive answer holds for both  $A/\langle LM(G) \rangle$  and  $A/\langle G \rangle$ , while a negative answer applies only to  $A/\langle LM(G) \rangle$  and not necessarily to  $A/\langle G \rangle$ .

**Example:**

```
LIB "fpaprops.lib";
ring r = 0, (x, y), dp;
def R = freeAlgebra(r, 5);
setring R;
ideal G = x*x, y*y; // K<x,y>/<xx,yy> is prime
lpIsPrime(G);
↪ 1
```

### 7.10.3.4 lpGldimBound

Procedure from library `fpaprops.lib` (see Section 7.10.3 [`fpaprops.lib`], page 649).

**Usage:** `lpGldimBound(I)`; I an ideal

**Return:** int, an upper bound for the global dimension, -1 means infinity

**Purpose:** computing an upper bound for the global dimension

**Assume:** - basering is a Letterplace ring, G is a reduced Groebner Basis

**Note:** if  $I = \text{LM}(I)$ , then the global dimension is equal the Gelfand Kirillov dimension if it is finite

Global dimension should be 0 for  $A/G = K$  and 1 for  $A/G = K\langle x_1 \dots x_n \rangle$

**Example:**

```
LIB "fpaprops.lib";
ring r = 0, (x,y), dp;
def R = freeAlgebra(r, 5); // constructs a Letterplace ring
setring R; // sets basering to Letterplace ring
ideal G = x*x, y*y, x*y*x; // it is a monomial Groebner basis
lpGldimBound(G);
⇒ // ** G is no standard basis
⇒ 0
ideal H = y*x - x*y; H = std(H); // H is a Groebner basis
lpGldimBound(H); // gl dim of K[x,y] is 2, as expected
⇒ 2
```

### 7.10.3.5 lpSubstitute

Procedure from library `fpaprops.lib` (see Section 7.10.3 [`fpaprops.lib`], page 649).

**Usage:** `lpSubstitute(f,s1,s2[,G])`; f poly, s1 list (ideal) of variables to replace, s2 list (ideal) of polynomials to replace with, G optional ideal to reduce with.

**Return:** poly, the substituted polynomial

**Assume:** - basering is a Letterplace ring  
 - s1 contains a subset of the set of variables  
 - s2 and s1 are of the same size  
 - G is a Groebner basis,  
 - the current ring has a sufficient degbound (which also can be calculated with `lpCalcSubstDegBound()`)

**Note:** the procedure implements the image of a polynomial f under an endomorphism of a free algebra, defined by s1 and s2: variables, not present in s1, are left unchanged;

variable  $s1[k]$  is mapped to a polynomial  $s2[k]$ .

- An optional ideal G extends the endomorphism as above to the morphism into the factor algebra  $K\langle X \rangle / G$ .

**Example:**

```
LIB "fpaprops.lib";
ring r = 0, (x,y,z), dp;
def R = freeAlgebra(r, 4);
setring R;
```

```

ideal G = x*y; // optional
poly f = 3*x*x+y*x;
ideal s1 = x, y;
ideal s2 = y*z*z, x; // i.e. x --> yzz and y --> x
// the substitution probably needs a higher degbound
int minDegBound = lpCalcSubstDegBound(f,s1,s2);
minDegBound; // thus the bound needs to be increased
↳ 9
setring r; // back to original r
def R1 = freeAlgebra(r, minDegBound);
setring R1;
lpSubstitute(imap(R,f), imap(R,s1), imap(R,s2));
↳ 3*y*z*z*y*z*z+x*y*z*z
// the last parameter is optional; above it was G=<xy>
// the output will be reduced with respect to G
lpSubstitute(imap(R,f), imap(R,s1), imap(R,s2), imap(R,G));
↳ 3*y*z*z*y*z*z

```

### 7.10.3.6 lpCalcSubstDegBound

Procedure from library `fpaprops.lib` (see Section 7.10.3 [`fpaprops.lib`], page 649).

**Usage:** `lpCalcSubstDegBound(I,s1,s2)`; I ideal of polynomials, s1 ideal of variables to replace, s2 ideal of polynomials to replace with

**Return:** int, the min degbound required to perform all of the substitutions

**Assume:** - basering is a Letterplace ring

**Note:** convenience method

**Example:**

```

LIB "fpaprops.lib";
ring r = 0, (x,y,z), dp;
def R = freeAlgebra(r, 4);
setring R;
ideal I = 3*x*x+y*x, x*y*x - z;
ideal s1 = x, y; // z --> z
ideal s2 = y*z*z, x; // i.e. x --> yzz and y --> x
// the substitution probably needs a higher degbound
lpCalcSubstDegBound(I,s1,s2);
↳ 10
lpCalcSubstDegBound(I[1],s1,s2);
↳ 9

```

### 7.10.4 freegb\_lib

**Library:** `freegb.lib`

**Purpose:** Two-sided Groebner bases in free algebras and tools via Letterplace approach

**Authors:** Viktor Levandovskyy, viktor.levandovskyy at math.rwth-aachen.de  
Karim Abou Zeid, karim.abou.zeid at rwth-aachen.de  
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**Overview:** For the theory, see chapter 'Letterplace' in the Singular Manual.

This library provides access to kernel functions and also contains legacy code (partially as static procedures) for compatibility reasons.

Support: Joint projects LE 2697/2-1 and KR 1907/3-1 of the Priority Programme SPP 1489: 'Algorithmische und Experimentelle Methoden in Algebra, Geometrie und Zahlentheorie' of the German DFG and Project II.6 of the transregional collaborative research centre SFB-TRR 195 'Symbolic Tools in Mathematics and their Application' of the German DFG

**Procedures:** See also: Section 7.7 [LETTERPLACE], page 613; Section 7.10.1 [fpadim.lib], page 629; Section 7.10.2 [fpalgebras.lib], page 634; Section 7.10.3 [fpaprops.lib], page 649.

#### 7.10.4.1 isFreeAlgebra

Procedure from library `freegb.lib` (see Section 7.10.4 [freegb.lib], page 652).

**Usage:** `isFreeAlgebra(r)`; `r` a ring

**Return:** boolean

**Purpose:** check whether `R` is a letterplace ring (free algebra)

**Example:**

```
LIB "freegb.lib";
ring r = 0, (x,y,z), dp;
isFreeAlgebra(r);
↳ 0
ring R = freeAlgebra(r, 7);
isFreeAlgebra(R);
↳ 1
```

#### 7.10.4.2 lpDegBound

Procedure from library `freegb.lib` (see Section 7.10.4 [freegb.lib], page 652).

**Usage:** `lpDegBound(R)`; `R` a letterplace ring

**Return:** int

**Purpose:** returns the degree bound of the letterplace ring

**Example:**

```
LIB "freegb.lib";
ring r = 0, (x,y,z), dp;
def R = freeAlgebra(r, 7);
lpDegBound(R);
↳ 7
```

#### 7.10.4.3 lpVarBlockSize

Procedure from library `freegb.lib` (see Section 7.10.4 [freegb.lib], page 652).

**Usage:** `lpVarBlockSize(R)`; `R` a letterplace ring

**Return:** int

**Purpose:** returns the variable block size of the letterplace ring, that is the number of variables of the original ring.

**Example:**

```
LIB "freegb.lib";
ring r = 0, (x,y,z), dp;
ring R = freeAlgebra(r, 7);
lpVarBlockSize(R);
↪ 3
```

**7.10.4.4 lpNcgenCount**

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `lpNcgenCount(R)`;  $R$  a letterplace ring

**Return:** `int`

**Purpose:** returns the number of `ncgen` variables in the letterplace ring.

**Example:**

```
LIB "freegb.lib";
ring r = 0, (x,y,z), dp;
ring R = freeAlgebra(r, 7, 3);
lpNcgenCount(R); // should be 3
↪ 3
```

**7.10.4.5 lpDivision**

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `lpDivision(p,G)`; poly  $p$ , ideal  $G$

**Purpose:** compute a two-sided division with remainder of  $p$  wrt  $G$ ; two-sided noncommutative analogue of the procedure `division`

**Assume:**  $G = \{g_1, \dots, g_N\}$  is a Groebner basis, the original ring of the Letterplace ring has the name `r` and no variable is called `'tag_i'` for  $i$  in  $1 \dots N$

**Return:** `list L`

**Note:**

- `L[1]` is  $NF(p,I)$
- `L[2]` is the list of expressions `[i, l_(ij), r_(ij)]` with  $\sum_{(i,j)} l_{(ij)} g_i r_{(ij)} = p - NF(p,I)$
- procedure `lpGBPres2Poly`, applied to `L`, reconstructs  $p$

**Example:**

```
LIB "freegb.lib";
ring r = 0, (x,y), dp;
ring R = freeAlgebra(r, 4);
ideal I = x*x + y*y - 1; // 2D sphere
ideal J = twostd(I); // compute a two-sided Groebner basis
J; // it is finite and nice
↪ J[1]=x*x+y*y-1
↪ J[2]=y*y*x-x*y*y
poly h = x*x*y-y*x*x+x*y;
list L = lpDivision(h,J); L; // what means that the NF of h wrt J is x*y
↪ [1]:
↪ x*y
↪ [2]:
↪ [1]:
```



```

↳ [1]:
↳ 1
↳ [2]:
↳ 1
↳ [3]:
↳ y
↳ [2]:
↳ [1]:
↳ 1
↳ [2]:
↳ -y
↳ [3]:
↳ 1
h - lpNF(h,J); // and this poly has the following two-sided Groebner presentation:
↳ -y*x*x+x*x*y
-y*J[1] + J[1]*y;
↳ -y*x*x+x*x*y
lpGBPres2Poly(L,J); // reconstructs the above automatically
↳ -y*x*x+x*x*y+x*y

```

#### 7.10.4.6 lpGBPres2Poly

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `lpGBPres2Poly(p,G)`; poly `p`, ideal `G`

**Assume:** `L` is a valid Groebner presentation like the result of `lpDivision`

**Return:** poly

**Note:**  $\text{assembles } p = \sum_{(i,j)} L_{(ij)} g_i r_{(ij)} + \text{NF}(p,I) = \sum_{(i)} L_{[2][i][2]} I_{[L[2][i][1]]} L_{[2][i][3]} + L_{[1]}$

**Example:**

```

LIB "freegb.lib";
ring r = 0,(x,y),dp;
ring R = freeAlgebra(r, 4);
ideal I = x*x + y*y - 1; // 2D sphere
ideal J = twostd(I); // compute a two-sided Groebner basis
J; // it is finite and nice
↳ J[1]=x*x+y*y-1
↳ J[2]=y*y*x-x*y*y
poly h = x*x*y-y*x*x+x*y;
list L = lpDivision(h,J);
L[1]; // what means that the normal form (or the remainder) of h wrt J is x*y
↳ x*y
lpGBPres2Poly(L,J); // we see, that it is equal to h from above
↳ -y*x*x+x*x*y+x*y

```

#### 7.10.4.7 isOrderingShiftInvariant

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `isOrderingShiftInvariant(b)`; `b` an integer interpreted as a boolean

**Return:** int

**Note:** Tests whether the ordering of the current ring is shift invariant, which is the case, when  $LM(p) > LM(p')$  for all  $p$  and  $p'$  where  $p'$  is  $p$  shifted by any number of places.

If withHoles != 0 even Letterplace polynomials with holes (eg.  $x(1)*y(4)$ ) are considered.

**Assume:** - basering is a Letterplace ring.

**Example:**

```
LIB "freegb.lib";
ring r = 0,(x,y,z),dp;
def R = freeAlgebra(r, 5);
setring R;
isOrderingShiftInvariant(0); // should be 1
↪ 1
ring r2 = 0,(x,y,z),dp;
def R2 = freeAlgebra(r2, 5);
list RL = ringlist(R2);
RL[3][1][1] = "wp";
intvec weights = 1,1,1,1,1,1,1,1,2,3,1,1,1,1,1,1;
RL[3][1][2] = weights;
attrib(RL,"isLetterplaceRing",3);
attrib(RL,"maxExp",1);
def Rw = setLetterplaceAttributes(ring(RL),5,3);
setring Rw;
/* printlevel = voice + 1; */
isOrderingShiftInvariant(0);
↪ 0
isOrderingShiftInvariant(1);
↪ 0
```

#### 7.10.4.8 makeLetterplaceRing

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `makeLetterplaceRing(d [,h]);`  $d$  an integer,  $h$  an optional integer (deprecated, use `freeAlgebra` instead)

**Return:** ring

**Purpose:** creates a ring with the ordering, used in letterplace computations

**Note:**  $h = -1$  (default) : the ordering of the current ring will be used  $h = 0$  : Dp ordering will be used

$h = 2$  : weights 1 used for all the variables, a tie breaker is a list of block of original ring

$h = 1$  : the pure homogeneous letterplace block ordering (applicable in the situation of homogeneous input ideals) will be used.

**Example:**

```
LIB "freegb.lib";
ring r = 0,(x,y,z),Dp;
def A = makeLetterplaceRing(2); // same as makeLetterplaceRing(2,0)
setring A; A;
↪ // coefficients: QQ
↪ // number of vars : 6
↪ // block 1 : ordering Dp
```

```

↳ // : names x y z x y z
↳ // block 2 : ordering C
↳ // letterplace ring (block size 3, nngen count 0)
lpVarBlockSize(A);
↳ 3
lpDegBound(A); // degree bound
↳ 2
setring r; def B = makeLetterplaceRing(2,1); // to compare:
setring B; B;
↳ // coefficients: QQ
↳ // number of vars : 6
↳ // block 1 : ordering Dp
↳ // : names x y z
↳ // block 2 : ordering Dp
↳ // : names x y z
↳ // block 3 : ordering C
↳ // letterplace ring (block size 3, nngen count 0)
lpVarBlockSize(B);
↳ 3
lpDegBound(B); // degree bound
↳ 2
setring r; def C = makeLetterplaceRing(2,2); // to compare:
setring C; C;
↳ // coefficients: QQ
↳ // number of vars : 6
↳ // block 1 : ordering a
↳ // : names x y z x y z
↳ // : weights 1 1 1 1 1 1
↳ // block 2 : ordering Dp
↳ // : names x y z
↳ // block 3 : ordering Dp
↳ // : names x y z
↳ // block 4 : ordering C
↳ // letterplace ring (block size 3, nngen count 0)
lpDegBound(C);
↳ 2
lpDegBound(C); // degree bound
↳ 2

```

#### 7.10.4.9 letplaceGBasis

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `letplaceGBasis(I)`; I an ideal/module

**Return:** ideal/module

**Assume:** basering is a Letterplace ring, input consists of Letterplace polynomials

**Purpose:** compute the two-sided Groebner basis of I via Letterplace algorithm (legacy routine)

**Note:** the degree bound for this computation is read off the letterplace structure of basering

**Example:**

```

LIB "freegb.lib";
ring r = 0, (x,y,z), Dp;

```

```

int degree_bound = 5;
def R = freeAlgebra(r, 5);
setring R;
ideal I = -x*y-7*y*y+3*x*x, x*y*x-y*x*y;
ideal J = letplaceGBasis(I);
J;
↳ J[1]=3*x*x-x*y-7*y*y
↳ J[2]=22*x*y*y-3*y*x*y-21*y*y*x+7*y*y*y
↳ J[3]=3*x*y*x-22*x*y*y+21*y*y*x-7*y*y*y
↳ J[4]=22803*y*y*y*x+19307*y*y*y*y
↳ J[5]=1933*y*y*x*y+2751*y*y*y*x+161*y*y*y*y
↳ J[6]=y*y*y*y*y

```

#### 7.10.4.10 lieBracket

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `lieBracket(a,b[N])`; a,b letterplace polynomials, N an optional integer

**Return:** `poly`

**Assume:** `basering` has a letterplace ring structure

**Purpose:** compute the Lie bracket  $[a,b] = ab - ba$  between letterplace polynomials

**Note:** if  $N > 1$  is specified, then the left normed bracket  $[a, \dots [a,b]]$  is computed.

**Example:**

```

LIB "freegb.lib";
ring r = 0,(x,y),dp;
ring R = freeAlgebra(r, 4);
poly a = x*y; poly b = y;
lieBracket(a,b);
↳ -y*x*y+x*y*y
lieBracket(x,y,2);
↳ y*x*x-2*x*y*x+x*x*y

```

#### 7.10.4.11 setLetterplaceAttributes

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `setLetterplaceAttributes(R, d, b)`; R a ring, b,d integers

**Return:** ring with special attributes set

**Purpose:** sets attributes for a letterplace ring:  
`'isLetterplaceRing' = 'IV' = b`, `'uptodeg' = d`, where `'uptodeg'` stands for the degree bound,  
`'IV'` for the number of variables in the block 0.

**Note:** Activate the resulting ring by using `setring`

**Example:**

```

LIB "freegb.lib";
ring r = 0,(x(1),y(1),x(2),y(2),x(3),y(3),x(4),y(4)),dp;
def R = setLetterplaceAttributes(r, 4, 2); setring R;
lpVarBlockSize(R);
↳ 2
lieBracket(x(1),y(1),2);
↳ y(1)*x(2)*x(3)-2*x(1)*y(2)*x(3)+x(1)*x(2)*y(3)

```

### 7.10.4.12 testLift

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `testLift(M,T)`; module M, matrix T

**Return:** module

**Purpose:** assembles the result of the lift procedure

**Assume:** T is the lift matrix of a submodule of M

**Note:** the inverse of the lift procedure

**Example:**

```
LIB "freegb.lib";
LIB "freegb.lib";
⇒ // ** redefining tstfreegb (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining tstfreegb (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining setLetterplaceAttributes (LIB "freegb.lib"); ./examples/\
testLift.sing:2
⇒ // ** redefining setLetterplaceAttributes (LIB "freegb.lib"); ./examples/\
testLift.sing:2
⇒ // ** redefining lst2str (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining mod2str (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining vct2str (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining isVar (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining letplaceGBasis (LIB "freegb.lib"); ./examples/testLift.s\
ing:2
⇒ // ** redefining letplaceGBasis (LIB "freegb.lib"); ./examples/testLift.s\
ing:2
⇒ // ** redefining lieBracket (LIB "freegb.lib"); ./examples/testLift.sing:\
2
⇒ // ** redefining lieBracket (LIB "freegb.lib"); ./examples/testLift.sing:\
2
⇒ // ** redefining lpPrint (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining lpPrint (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining freeGBasis (LIB "freegb.lib"); ./examples/testLift.sing:\
2
⇒ // ** redefining freeGBasis (LIB "freegb.lib"); ./examples/testLift.sing:\
2
⇒ // ** redefining crs (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining polylen (LIB "freegb.lib"); ./examples/testLift.sing:2
⇒ // ** redefining lpDegBound (LIB "freegb.lib"); ./examples/testLift.sing:\
2
⇒ // ** redefining lpDegBound (LIB "freegb.lib"); ./examples/testLift.sing:\
2
⇒ // ** redefining lpVarBlockSize (LIB "freegb.lib"); ./examples/testLift.s\
ing:2
⇒ // ** redefining lpVarBlockSize (LIB "freegb.lib"); ./examples/testLift.s\
ing:2
⇒ // ** redefining isFreeAlgebra (LIB "freegb.lib"); ./examples/testLift.si\
ng:2
⇒ // ** redefining isFreeAlgebra (LIB "freegb.lib"); ./examples/testLift.si\
ng:2
⇒ // ** redefining lpNcgenCount (LIB "freegb.lib"); ./examples/testLift.sin\
```

```

g:2
⇒ // ** redefining lpNcgenCount (LIB "freegb.lib");) ./examples/testLift.sin\
g:2
⇒ // ** redefining makeLetterplaceRing (LIB "freegb.lib");) ./examples/testL\
ift.sing:2
⇒ // ** redefining makeLetterplaceRing (LIB "freegb.lib");) ./examples/testL\
ift.sing:2
⇒ // ** redefining makeLetterplaceRing1 (LIB "freegb.lib");) ./examples/test\
Lift.sing:2
⇒ // ** redefining makeLetterplaceRing2 (LIB "freegb.lib");) ./examples/test\
Lift.sing:2
⇒ // ** redefining makeLetterplaceRing4 (LIB "freegb.lib");) ./examples/test\
Lift.sing:2
⇒ // ** redefining makeLetterplaceRing3 (LIB "freegb.lib");) ./examples/test\
Lift.sing:2
⇒ // ** redefining freegbold (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining stringpoly2lplace (LIB "freegb.lib");) ./examples/testLif\
t.sing:2
⇒ // ** redefining addplaces (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining sent2lplace (LIB "freegb.lib");) ./examples/testLift.sing\
:2
⇒ // ** redefining testnumber (LIB "freegb.lib");) ./examples/testLift.sing:\
2
⇒ // ** redefining str2lplace (LIB "freegb.lib");) ./examples/testLift.sing:\
2
⇒ // ** redefining strpower2rep (LIB "freegb.lib");) ./examples/testLift.sin\
g:2
⇒ // ** redefining shiftPoly (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining lastBlock (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining test_shift (LIB "freegb.lib");) ./examples/testLift.sing:\
2
⇒ // ** redefining lp2lstr (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining lp2lstr (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining strList2poly (LIB "freegb.lib");) ./examples/testLift.sin\
g:2
⇒ // ** redefining file2lplace (LIB "freegb.lib");) ./examples/testLift.sing\
:2
⇒ // ** redefining lpPower (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining lpNF (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining lpNF (LIB "freegb.lib");) ./examples/testLift.sing:2
⇒ // ** redefining lpDivision (LIB "freegb.lib");) ./examples/testLift.sing:\
2
⇒ // ** redefining lpDivision (LIB "freegb.lib");) ./examples/testLift.sing:\
2
⇒ // ** redefining lpGBPres2Poly (LIB "freegb.lib");) ./examples/testLift.si\
ng:2
⇒ // ** redefining lpGBPres2Poly (LIB "freegb.lib");) ./examples/testLift.si\
ng:2
⇒ // ** redefining getExpVecs (LIB "freegb.lib");) ./examples/testLift.sing:\
2
⇒ // ** redefining delSupZero (LIB "freegb.lib");) ./examples/testLift.sing:\
2
⇒ // ** redefining delSupZeroList (LIB "freegb.lib");) ./examples/testLift.s\

```

```

ing:2
↳ // ** redefining makeDVec (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining makeDVecL (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining makeDVecI (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining dShiftDiv (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining lpNormalForm1 (LIB "freegb.lib"); ./examples/testLift.si\
ng:2
↳ // ** redefining lpNormalForm2 (LIB "freegb.lib"); ./examples/testLift.si\
ng:2
↳ // ** redefining isOrderingShiftInvariant (LIB "freegb.lib"); ./examples/\
testLift.sing:2
↳ // ** redefining isOrderingShiftInvariant (LIB "freegb.lib"); ./examples/\
testLift.sing:2
↳ // ** redefining lpMonomialsWithHoles (LIB "freegb.lib"); ./examples/test\
Lift.sing:2
↳ // ** redefining getlpCoeffs (LIB "freegb.lib"); ./examples/testLift.sing\
:2
↳ // ** redefining lpReduce (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining entryViolation (LIB "freegb.lib"); ./examples/testLift.s\
ing:2
↳ // ** redefining checkAssumptionsLPIV (LIB "freegb.lib"); ./examples/test\
Lift.sing:2
↳ // ** redefining checkAssumptions (LIB "freegb.lib"); ./examples/testLift\
.sing:2
↳ // ** redefining checkLPRing (LIB "freegb.lib"); ./examples/testLift.sing\
:2
↳ // ** redefining checkAssumptionIdeal (LIB "freegb.lib"); ./examples/test\
Lift.sing:2
↳ // ** redefining checkAssumptionPoly (LIB "freegb.lib"); ./examples/testL\
ift.sing:2
↳ // ** redefining isContainedInVp (LIB "freegb.lib"); ./examples/testLift.\
sing:2
↳ // ** redefining extractLinearPart (LIB "freegb.lib"); ./examples/testLif\
t.sing:2
↳ // ** redefining isLinearVector (LIB "freegb.lib"); ./examples/testLift.s\
ing:2
↳ // ** redefining lpAssumeViolation (LIB "freegb.lib"); ./examples/testLif\
t.sing:2
↳ // ** redefining skip0 (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining ivL2lpI (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining ivL2lpI (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining iv2lp (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining iv2lp (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining iv2lpList (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining iv2lpList (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining iv2lpMat (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining iv2lpMat (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining lpId2ivLi (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining lpId2ivLi (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining lp2iv (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining lp2iv (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining lp2ivId (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining lp2ivId (LIB "freegb.lib"); ./examples/testLift.sing:2

```

```

↳ // ** redefining testLift (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining testLift (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining testSyz (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining testSyz (LIB "freegb.lib"); ./examples/testLift.sing:2
↳ // ** redefining mod_init (LIB "freegb.lib"); ./examples/testLift.sing:2
ring r = 0, (x,y), (c,Dp);
ring R = freeAlgebra(r, 7, 2);
ideal I = std(x*y*x + 1);
print(matrix(I)); // finite two-sided Groebner basis
↳ x*y-y*x,y*x*x+1
ideal SI = x*I[1]*y + y*x*I[2], I[1]*y*x + I[2]*y;
matrix T = lift(I, SI); // T is the lifting matrix of SI wrt I
print(T); //
↳ y*ncgen(1)*x*x+x*ncgen(1)*y,y*x*ncgen(1)+y*ncgen(1)*x+ncgen(1)*y*x,
↳ y*ncgen(2)*x, y*ncgen(2)
print(matrix(SI)); // the original generators of SI as a matrix
↳ y*x*y*x*x+x*x*y*y-x*y*x*y+y*x,x*y*y*x+y*x*x*y-y*x*y*x+y
print(matrix(testLift(I,T))); // and the result of testLift
↳ y*x*y*x*x+x*x*y*y-x*y*x*y+y*x,x*y*y*x+y*x*x*y-y*x*y*x+y

```

#### 7.10.4.13 testSyz

Procedure from library `freegb.lib` (see Section 7.10.4 [`freegb.lib`], page 652).

**Usage:** `testSyz(M,S)`; module M, S

**Return:** module

**Purpose:** tests the result of the syz procedure

**Assume:** S is the syzygy bimodule of M

**Example:**

```

LIB "freegb.lib";
LIB "freegb.lib";
↳ // ** redefining tstfreegb (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining tstfreegb (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining setLetterplaceAttributes (LIB "freegb.lib"); ./examples/\
testSyz.sing:2
↳ // ** redefining setLetterplaceAttributes (LIB "freegb.lib"); ./examples/\
testSyz.sing:2
↳ // ** redefining lst2str (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining mod2str (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining vct2str (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining isVar (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining letplaceGBasis (LIB "freegb.lib"); ./examples/testSyz.si\
ng:2
↳ // ** redefining letplaceGBasis (LIB "freegb.lib"); ./examples/testSyz.si\
ng:2
↳ // ** redefining lieBracket (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lieBracket (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpPrint (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpPrint (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining freeGBasis (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining freeGBasis (LIB "freegb.lib"); ./examples/testSyz.sing:2

```



```

↳ // ** redefining crs (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining polylen (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpDegBound (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpDegBound (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpVarBlockSize (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpVarBlockSize (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining isFreeAlgebra (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining isFreeAlgebra (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpNcgenCount (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpNcgenCount (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeLetterplaceRing (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeLetterplaceRing (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeLetterplaceRing1 (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeLetterplaceRing2 (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeLetterplaceRing4 (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeLetterplaceRing3 (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining freegbold (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining stringpoly2lplace (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining addplaces (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining sent2lplace (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining testnumber (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining str2lplace (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining strpower2rep (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining shiftPoly (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lastBlock (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining test_shift (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lp2lstr (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lp2lstr (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining strList2poly (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining file2lplace (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpPower (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpNF (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpNF (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpDivision (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpDivision (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpGBPres2Poly (LIB "freegb.lib"); ./examples/testSyz.sing:2

```

```

g:2
↳ // ** redefining lpGBPres2Poly (LIB "freegb.lib"); ./examples/testSyz.sin\
g:2
↳ // ** redefining getExpVecs (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining delSupZero (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining delSupZeroList (LIB "freegb.lib"); ./examples/testSyz.si\
ng:2
↳ // ** redefining makeDVec (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeDVecL (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining makeDVecI (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining dShiftDiv (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpNormalForm1 (LIB "freegb.lib"); ./examples/testSyz.sin\
g:2
↳ // ** redefining lpNormalForm2 (LIB "freegb.lib"); ./examples/testSyz.sin\
g:2
↳ // ** redefining isOrderingShiftInvariant (LIB "freegb.lib"); ./examples/\
testSyz.sing:2
↳ // ** redefining isOrderingShiftInvariant (LIB "freegb.lib"); ./examples/\
testSyz.sing:2
↳ // ** redefining lpMonomialsWithHoles (LIB "freegb.lib"); ./examples/test\
Syz.sing:2
↳ // ** redefining getlpCoeffs (LIB "freegb.lib"); ./examples/testSyz.sing:\
2
↳ // ** redefining lpReduce (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining entryViolation (LIB "freegb.lib"); ./examples/testSyz.si\
ng:2
↳ // ** redefining checkAssumptionsLPIV (LIB "freegb.lib"); ./examples/test\
Syz.sing:2
↳ // ** redefining checkAssumptions (LIB "freegb.lib"); ./examples/testSyz.\
sing:2
↳ // ** redefining checkLPRing (LIB "freegb.lib"); ./examples/testSyz.sing:\
2
↳ // ** redefining checkAssumptionIdeal (LIB "freegb.lib"); ./examples/test\
Syz.sing:2
↳ // ** redefining checkAssumptionPoly (LIB "freegb.lib"); ./examples/testS\
yz.sing:2
↳ // ** redefining isContainedInVp (LIB "freegb.lib"); ./examples/testSyz.s\
ing:2
↳ // ** redefining extractLinearPart (LIB "freegb.lib"); ./examples/testSyz\
.sing:2
↳ // ** redefining isLinearVector (LIB "freegb.lib"); ./examples/testSyz.si\
ng:2
↳ // ** redefining lpAssumeViolation (LIB "freegb.lib"); ./examples/testSyz\
.sing:2
↳ // ** redefining skip0 (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining ivL2lpI (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining ivL2lpI (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining iv2lp (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining iv2lp (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining iv2lpList (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining iv2lpList (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining iv2lpMat (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining iv2lpMat (LIB "freegb.lib"); ./examples/testSyz.sing:2

```

```

↳ // ** redefining lpId2ivLi (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lpId2ivLi (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lp2iv (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lp2iv (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lp2ivId (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining lp2ivId (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining testLift (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining testLift (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining testSyz (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining testSyz (LIB "freegb.lib"); ./examples/testSyz.sing:2
↳ // ** redefining mod_init (LIB "freegb.lib"); ./examples/testSyz.sing:2
ring r = 0, (x,y), (c,Dp);
ring R = freeAlgebra(r, 7, 2);
ideal I = twostd(x*y*x + 1);
print(matrix(I));
↳ x*y-y*x,y*x*x+1
module S = syz(I);
print(S);
↳ ncgen(1)*x*x,S[1,2],S[1,3],S[1,4],S[1,5],
↳ S[2,1], S[2,2],S[2,3],S[2,4],S[2,5]
testSyz(I,S); // returns zero
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0
↳ _[4]=0
↳ _[5]=0

```

### 7.10.5 ncHilb\_lib

**Library:** ncHilb.lib

**Purpose:** Computation of graded and multi-graded Hilbert series of non-commutative algebras (Letterplace).

**Author:** Sharwan K. Tiwari shrawant@gmail.com  
 Roberto La Scala  
 Viktor Levandovskyy (adaptation to the new Letterplace release)

**References:**

La Scala R.: Monomial right ideals and the Hilbert series of non-commutative modules, Journal of Symbolic Computation (2016).

La Scala R., Tiwari Sharwan K.: Multigraded Hilbert Series of noncommutative modules, <https://arxiv.org/abs/1705.01083>.

**Procedures:**

#### 7.10.5.1 nchilb

Procedure from library ncHilb.lib (see Section 7.10.5 [ncHilb\_lib], page 665).

**Usage:** nchilb(I, d[, L]), list I, int d, optional list L

**Purpose:** compute Hilbert series of a non-commutative algebra

**Assume:**

**Note:**  $d$  is an integer for the degree bound (maximal total degree of polynomials of the generating set of the input ideal),  
 $\#[]=1$ , computation for non-finitely generated regular ideals,  $\#[]=2$ , computation of multi-graded Hilbert series,  
 $\#[]=tdeg$ , for obtaining the truncated Hilbert series up to the total degree  $tdeg-1$  ( $tdeg$  should be  $> 2$ ), and  $\#[]=string(p)$ , to print the details about the orbit and system of equations. Let the orbit is  $O_I = \{T_{\{w_1\}}(I), \dots, T_{\{w_r\}}(I)\}$  ( $w_i \in W$ ), where we assume that if  $T_{\{w_i\}}(I) = T_{\{w'_i\}}(I)$  for some  $w'_i \in W$ , then  $\deg(w_i) \leq \deg(w'_i)$ .  
Then, it prints words description of orbit:  $w_1, \dots, w_r$ . It also prints the maximal degree and the cardinality of  $\sum_j R(w_i, b_j)$  corresponding to each  $w_i$ , where  $\{b_j\}$  is a basis of  $I$ .  
Moreover, it also prints the linear system (for the information about adjacency matrix) and its solving time.

**Note :** A Groebner basis of two-sided ideal of the input should be given in a special form. This form is a list of modules, where each generator of every module represents a monomial times a coefficient in the free associative algebra. The first entry, in each generator, represents a coefficient and every next entry is a variable.

Ex: module  $p1=[1,y,z],[-1,z,y]$ , represents the poly  $y*z-z*y$ ; module  $p2=[1,x,z,x],[-1,z,x,z]$ , represents the poly  $x*z*x-z*x*z$  for more details about the input, see examples.

**Example:**

```
LIB "ncHilb.lib";
ring r=0,(X,Y,Z),dp;
module p1 =[1,Y,Z]; //represents the poly Y*Z
module p2 =[1,Y,Z,X]; //represents the poly Y*Z*X
module p3 =[1,Y,Z,Z,X,Z];
module p4 =[1,Y,Z,Z,Z,X,Z];
module p5 =[1,Y,Z,Z,Z,Z,X,Z];
module p6 =[1,Y,Z,Z,Z,Z,Z,X,Z];
module p7 =[1,Y,Z,Z,Z,Z,Z,Z,X,Z];
module p8 =[1,Y,Z,Z,Z,Z,Z,Z,Z,X,Z];
list l1=list(p1,p2,p3,p4,p5,p6,p7,p8);
nchilb(l1,10);
↳
↳ maximal length of words = 2
↳
↳ length of the Orbit = 3
↳
↳
↳ Hilbert series:
↳ 1/(t2-3t+1)
ring r2=0,(x,y,z),dp;
module p1=[1,y,z],[-1,z,y]; //y*z-z*y
module p2=[1,x,z,x],[-1,z,x,z]; // x*z*x-z*x*z
module p3=[1,x,z,z,x,z],[-1,z,x,z,z,x]; // x*z^2*x*z-z*x*z^2*x
module p4=[1,x,z,z,z,x,z],[-1,z,x,z,z,x,x]; // x*z^3*x*z-z*x*z^2*x^2
list l2=list(p1,p2,p3,p4);
nchilb(l2,6,1); //third argument '1' is for non-finitely generated case
↳
↳ maximal length of words = 3
↳
```

```

↳ length of the Orbit = 7
↳
↳
↳ Hilbert series:
↳ (t3+t2+1)/(2t5-2t4-t3+2t2-3t+1)
ring r3=0,(a,b),dp;
module p1=[1,a,a,a];
module p2=[1,a,b,b];
module p3=[1,a,a,b];
list l3=list(p1,p2,p3);
nchilb(l3,5,2); //third argument '2' is to compute multi-graded HS
↳
↳ maximal length of words = 3
↳
↳ length of the Orbit = 5
↳
↳
↳ Hilbert series:
↳ (t1^2+t1+1)/(t1*t2^2-t1*t2-t2+1)
ring r4=0,(x,y,z),dp;
module p1=[1,x,z,y,z,x,z];
module p2=[1,x,z,x];
module p3=[1,x,z,y,z,z,x,z];
module p4=[1,y,z];
module p5=[1,x,z,z,x,z];
list l4=list(p1,p2,p3,p4,p5);
nchilb(l4,7,"p"); //third argument "p" is to print the details
↳
↳ maximal length of words = 3
↳
↳ length of the Orbit = 6
↳ words description of the Orbit:
↳ 1 x y x*z y*z x*z*z
↳
↳ maximal degree, #(sum_j R(w,w_j))
↳ NULL
↳ 6, 4
↳ 1, 1
↳ 5, 4
↳ 0, 1
↳ 2, 1
↳
↳ linear system:
↳ H(1) = (t)*H(2) + (t)*H(3) + (t)*H(1) + 1
↳ H(2) = (t)*H(2) + (t)*H(3) + (t)*H(4) + 1
↳ H(3) = (t)*H(2) + (t)*H(3) + (t)*H(5) + 1
↳ H(4) = (t)*H(5) + (t)*H(3) + (t)*H(6) + 1
↳ H(5) = (t)*H(5) + (t)*H(5) + (t)*H(5) + 0
↳ H(6) = (t)*H(3) + (t)*H(3) + (t)*H(1) + 1
↳ where H(1) represents the series corresp. to input ideal
↳ and i^th summand in the rhs of an eqn. is according
↳ to the right colon map corresp. to the i^th variable
↳

```

```

↳
↳ Hilbert series:
↳ (t3+t2+1)/(2t5-2t4-t3+2t2-3t+1)
// of the orbit and system

```

### 7.10.5.2 rcolon

Procedure from library `ncHilb.lib` (see Section 7.10.5 [`ncHilb.lib`], page 665).

**Usage:** `rcolon(list of relations, a monomial, an integer);`  
`L` is a list of modules (each module represents a monomial), `w` is a monomial  
`d` is an integer for the degree bound (maximal total degree of monomials of the generating set of the input monomial ideal),

**Note :** A two-sided monomial ideal and a monomial `w` for the input should be given in a special form. This form is a list of modules, where the generator of every module represents a monomial times a coefficient in the free associative algebra. The first entry, in each generator, represents a coefficient, that is 1, and every next entry is a variable.

Ex: module `p1=[1,y,z]`, represents the monomial  $y^*z$ ;

module `p2=[1,x,z,x]`, represents the monomial  $x^*z^*x$

for more details about the input, see examples.

**Example:**

```

LIB "ncHilb.lib";
ring r=0, (X,Y,Z), dp;
module w = [1,Y];
module p1 = [1,Y,Z];
module p2 = [1,Y,Z,X];
module p3 = [1,Y,Z,Z,X,Z];
module p4 = [1,Y,Z,Z,Z,X,Z];
module p5 = [1,Y,Z,Z,Z,Z,X,Z];
module p6 = [1,Y,Z,Z,Z,Z,Z,X,Z];
module p7 = [1,Y,Z,Z,Z,Z,Z,Z,X,Z];
module p8 = [1,Y,Z,Z,Z,Z,Z,Z,Z,X,Z];
list l1=list(p1,p2,p3,p4,p5,p6,p7,p8);
rcolon(l1,w,10);
↳ J[1]=Z
↳ + generators of the given ideal;

```

### 7.10.6 ncrat\_lib

Status: experimental

**Library:** `ncrat.lib`

**Purpose:** Framework for working with non-commutative rational functions

**Author:** Ricardo Schnur, email: `ricardo.schnur@math.uni-sb.de`

**Support:** This project has been funded by the SFB-TRR 195  
'Symbolic Tools in Mathematics and their Application'.

**Overview:** This library provides a framework for working with non-commutative rational functions (or rather, expressions) and their linearized representations

**References:**

T. Mai: On the analytic theory of non-commutative distributions in free probability. Universitaet des Saarlandes, Dissertation, 2017

**Note:** an almost self-explaining introduction to the possibilities of the framework can be achieved by running the example for the procedure `ncrepGetRegularMinimal`.

**Procedures:****7.10.6.1 ncInit**

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncInit(vars);`  
list vars containing strings

**Return:** datatypes `ncrat` and `ncrep` (and `token`, `tokenstream`, but they are not meant for users), sets ring as 'NCRING' with `nc` variables from list `l`

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
NCRING;
↪ // coefficients: QQ[I]/(I^2+1)
↪ // number of vars : 3
↪ // block 1 : ordering dp
↪ // : names x y z
↪ // block 2 : ordering C
```

**7.10.6.2 ncVarsGet**

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncVarsGet();`

**Returns:** `nc` variables that are in use

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncVarsGet();
↪ x,y,z
```

**7.10.6.3 ncVarsAdd**

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncVarsAdd(vars);`  
list vars contains variables

**Returns:** sets list elements as `nc` variables

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncVarsAdd("z");
```

```

↳ x,y,z
ncVarsAdd(list("a", "b", "c"));
↳ // ** killing the basering for level 0
ncVarsGet();
↳ x,y,z,a,b,c

```

#### 7.10.6.4 ncratDefine

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrat f = ncratDefine(s, l);`  
string `s` contains kind, list `l` contains expressions

**Return:** `ncrat` with kind `s` and expressions `l`

**Note:** assignment operator `'='` for `ncrat` is overloaded with this procedure, hence  
`ncrat f = s, l;`  
yields the same result as  
`ncrat f = ncratDefine(s, l);`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
number n = 5;
ncrat f = ncratDefine("Const", list(n));
typeof(f);
↳ ncrat
f.kind;
↳ Const
f.expr;
↳ [1]:
↳ 5
f;
↳ 5
↳
ncrat g = "Const", list(n);
g;
↳ 5
↳

```

#### 7.10.6.5 ncratAdd

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrat h = ncratAdd(f, g);`  
`f, g` both of type `ncrat`

**Return:** `h = f + g`

**Note:** operator `'+'` for `ncrat` is overloaded with this procedure, hence  
`ncrat h = f + g;`  
yields the same result as  
`ncrat h = ncratAdd(f, g);`



**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
print(f);
↪ 2*x*y
ncrat g = ncratFromString("z");
print(g);
↪ z
ncrat h1, h2;
h1 = ncratAdd(f, g);
print(h1);
↪ 2*x*y+z
h2 = f + g;
print(h2);
↪ 2*x*y+z

```

**7.10.6.6 ncratSubtract**

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     `ncrat h = ncratSubtract(f, g);`  
               `f, g` both of type `ncrat`

**Return:**    `h = f - g`

**Note:**     operator `'-'` for `ncrat` is overloaded  
               with this procedure, hence  
               `ncrat h = f - g;`  
               yields the same result as  
               `ncrat h = ncratSubtract(f, g);`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
print(f);
↪ 2*x*y
ncrat g = ncratFromString("z");
print(g);
↪ z
ncrat h1, h2;
h1 = ncratSubtract(f, g);
print(h1);
↪ 2*x*y-z
h2 = f - g;
print(h2);
↪ 2*x*y-z

```

**7.10.6.7 ncratMultiply**

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     `ncrat h = ncratMultiply(f, g);`  
               `f, g` both of type `ncrat`

**Return:**  $h = f * g$

**Note:** operator `'*'` for `ncrat` is overloaded with this procedure, hence  
`ncrat h = f * g;`  
yields the same result as  
`ncrat h = ncratMultiply(f, g);`

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
print(f);
↳ 2*x*y
ncrat g = ncratFromString("z");
print(g);
↳ z
ncrat h1, h2;
h1 = ncratMultiply(f, g);
print(h1);
↳ 2*x*y*z
h2 = f * g;
print(h2);
↳ 2*x*y*z
```

### 7.10.6.8 ncratInvert

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrat h = ncratInvert(f);`  
`f` of type `ncrat`

**Return:**  $h = \text{inv}(f)$

**Note:** `ncrat h = f^-1;`  
yields the same result as  
`ncrat h = ncratInvert(f);`

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
print(f);
↳ 2*x*y
ncrat h1, h2;
h1 = ncratInvert(f);
print(h1);
↳ inv(2*x*y)
h2 = f ^ -1;
print(h2);
↳ inv(2*x*y)
```

### 7.10.6.9 ncratSPrint

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     string s = ncratSPrint(f);  
              f of type ncrat

**Return:**    prints f to string

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
string s = ncratSPrint(f);
print(s);
↪ 2*x*y
```

#### 7.10.6.10 ncratPrint

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     ncratPrint(f);  
              f of type ncrat

**Return:**    prints f

**Note:**     print(f);  
              yields the same result as  
              ncratPrint(f);

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
ncratPrint(f);
↪ 2*x*y
print(f);
↪ 2*x*y
```

#### 7.10.6.11 ncratFromString

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     ncrat f = ncratFromString(s);  
              s of type string

**Return:**    read string s into ncrat f

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
print(f);
↪ 2*x*y
```

#### 7.10.6.12 ncratFromPoly

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     ncrat f = ncratFromPoly(p);  
              p of type poly

**Return:** convert poly to ncrat

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
poly p = 2 * x * y;
ncrat f = ncratFromPoly(p);
print(f);
↪ 2*x*y
```

### 7.10.6.13 ncratPower

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** ncrat h = ncratPower(f, n);  
f of type ncrat, n integer

**Return:** h = f<sup>n</sup>

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
ncrat h = ncratPower(f, 3);
print(h);
↪ 2*x*y*2*x*y*2*x*y
```

### 7.10.6.14 ncratEvaluateAt

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** matrix M = ncratEvaluateAt(f, vars, point);

**Return:** Evaluate the ncrat f by substituting in the matrices contained in point for the respective variables contained in var, that is, calculate f(point).

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("x+y");
matrix A[2][2] = 1, 2, 3, 4;
matrix B[2][2] = 5, 6, 7, 8;
matrix M = ncratEvaluateAt(f, list(x, y), list(A, B));
print(M);
↪ 6, 8,
↪ 10,12
```

### 7.10.6.15 ncrepGet

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** ncrep q = ncrepGet(f);  
f of type ncrat

**Return:** q = (u, Q, v) linear representation of f

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
ncrep q = ncrepGet(f);
print(q);
↳ lvec=
↳ 0,0,0,1
↳
↳ mat=
↳ 0, 0, 1/2*x,-1/2,
↳ 0, 1, -1/2, 0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0
↳
↳ rvec=
↳ 0,
↳ 0,
↳ 0,
↳ 1

```

**7.10.6.16 ncrepAdd**

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrep s = ncrepAdd(q, r);`  
`q, r` both of type `ncrep`

**Return:** representation `s` of  $h = f + g$   
if `q, r` are representations of `f, g`

**Note:** operator '+' for `ncrep` is overloaded  
with this procedure, hence  
`ncrep s = q + r;`  
yields the same result as  
`ncrep s = ncrepAdd(q, r);`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("x");
ncrat g = ncratFromString("y");
ncrep q = ncrepGet(f);
ncrep r = ncrepGet(g);
ncrep s1, s2;
s1 = ncrepAdd(q, r);
print(s1);
↳ lvec=
↳ 0,1,0,1
↳
↳ mat=
↳ x, -1,0, 0,
↳ -1,0, 0, 0,
↳ 0, 0, y, -1,

```

```

↳ 0, 0, -1,0
↳
↳ rvec=
↳ 0,
↳ 1,
↳ 0,
↳ 1
s2 = q + r;
print(s2);
↳ lvec=
↳ 0,1,0,1
↳
↳ mat=
↳ x, -1,0, 0,
↳ -1,0, 0, 0,
↳ 0, 0, y, -1,
↳ 0, 0, -1,0
↳
↳ rvec=
↳ 0,
↳ 1,
↳ 0,
↳ 1

```

### 7.10.6.17 ncrepSubtract

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrep s = ncrepSubtract(q, r);`  
`q, r` both of type `ncrep`

**Return:** representation `s` of  $h = f - g$   
if `q, r` are representations of `f, g`

**Note:** operator `'-'` for `ncrep` is overloaded  
with this procedure, hence  
`ncrep s = q - r;`  
yields the same result as  
`ncrep s = ncrepSubtract(q, r);`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("x");
ncrat g = ncratFromString("y");
ncrep q = ncrepGet(f);
ncrep r = ncrepGet(g);
ncrep s1, s2;
s1 = ncrepSubtract(q, r);
print(s1);
↳ lvec=
↳ 0,1,0,1
↳
↳ mat=
↳ x, -1,0, 0,

```

```

↳ -1,0, 0, 0,
↳ 0, 0, -y,1,
↳ 0, 0, 1, 0
↳
↳ rvec=
↳ 0,
↳ 1,
↳ 0,
↳ 1
s2 = q - r;
print(s2);
↳ lvec=
↳ 0,1,0,1
↳
↳ mat=
↳ x, -1,0, 0,
↳ -1,0, 0, 0,
↳ 0, 0, -y,1,
↳ 0, 0, 1, 0
↳
↳ rvec=
↳ 0,
↳ 1,
↳ 0,
↳ 1

```

### 7.10.6.18 ncrepMultiply

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrep s = ncrepMultiply(q, r);`  
`q, r` both of type `ncrep`

**Return:** representation `s` of  $h = f * g$   
if `q, r` are representations of `f, g`

**Note:** operator `'*'` for `ncrep` is overloaded  
with this procedure, hence  
`ncrep s = q * r;`  
yields the same result as  
`ncrep s = ncrepMultiply(q, r);`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("x");
ncrat g = ncratFromString("y");
ncrep q = ncrepGet(f);
ncrep r = ncrepGet(g);
ncrep s1, s2;
s1 = ncrepMultiply(q, r);
print(s1);
↳ lvec=
↳ 0,0,0,1
↳

```

```

↳ mat=
↳ 0, 0, x, -1,
↳ 0, 1, -1,0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0
↳
↳ rvec=
↳ 0,
↳ 0,
↳ 0,
↳ 1
s2 = q * r;
print(s2);
↳ lvec=
↳ 0,0,0,1
↳
↳ mat=
↳ 0, 0, x, -1,
↳ 0, 1, -1,0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0
↳
↳ rvec=
↳ 0,
↳ 0,
↳ 0,
↳ 1

```

### 7.10.6.19 ncrepInvert

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     `ncrep s = ncrepInvert(q);`  
               `q` of type `ncrep`

**Return:**    representation of  $h = \text{inv}(f)$   
               if `q` is a representation of `f`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
ncrep q = ncrepGet(f);
ncrep s = ncrepInvert(q);
print(s);
↳ lvec=
↳ 1,0,0,0,0
↳
↳ mat=
↳ 0,0, 0, 0, 1,
↳ 0,0, 0, -1/2*x,1/2,
↳ 0,0, -1,1/2, 0,
↳ 0,-y,1, 0, 0,
↳ 1,1, 0, 0, 0

```



```

↳
↳ rvec=
↳ 1,
↳ 0,
↳ 0,
↳ 0,
↳ 0

```

### 7.10.6.20 ncrepPrint

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrepPrint(q);`  
`q` of type `ncrep`

**Return:** prints `q`

**Note:** `print(q);`  
yields the same result as  
`ncrepPrint(q);`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
ncrep q = ncrepGet(f);
ncrepPrint(q);
↳ lvec=
↳ 0,0,0,1
↳
↳ mat=
↳ 0, 0, 1/2*x,-1/2,
↳ 0, 1, -1/2, 0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0
↳
↳ rvec=
↳ 0,
↳ 0,
↳ 0,
↳ 1
print(q);
↳ lvec=
↳ 0,0,0,1
↳
↳ mat=
↳ 0, 0, 1/2*x,-1/2,
↳ 0, 1, -1/2, 0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0
↳
↳ rvec=
↳ 0,
↳ 0,
↳ 0,

```

↪ 1

### 7.10.6.21 ncrepDim

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     `ncrepDim(q);`  
              `q` of type `ncrep`

**Return:**    dimension of `q`;  
              returns 0 if `q` represents the zero-function

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("2*x*y");
ncrep q = ncrepGet(f);
print(q);
↪ lvec=
↪ 0,0,0,1
↪
↪ mat=
↪ 0, 0, 1/2*x,-1/2,
↪ 0, 1, -1/2, 0,
↪ y, -1,0, 0,
↪ -1,0, 0, 0
↪
↪ rvec=
↪ 0,
↪ 0,
↪ 0,
↪ 1
ncrepDim(q);
↪ 4
```

### 7.10.6.22 ncrepSubstitute

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:**     `ncrep s = ncrepSubstitute(q, l);`  
              `q` of type `ncrep`, `vars = (x1, ..., xg)`,  
              `points = (A1, ... , Ag)` with `Ai` matrices of the  
              same dimension and `xi` of type `poly` are `nc` variables

**Return:**    substitutes in `Ai` for `xi` in `q`

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("x+y");
ncrep q = ncrepGet(f);
matrix A[2][2] = 1, 2, 3, 4;
matrix B[2][2] = 5, 6, 7, 8;
ncrep s = ncrepSubstitute(q, list(x, y), list(A, B));
print(q);
```

```

↳ lvec=
↳ 0,1,0,1
↳
↳ mat=
↳ x, -1,0, 0,
↳ -1,0, 0, 0,
↳ 0, 0, y, -1,
↳ 0, 0, -1,0
↳
↳ rvec=
↳ 0,
↳ 1,
↳ 0,
↳ 1
print(s);
↳ lvec=
↳ 0,0,1,0,0,0,1,0,
↳ 0,0,0,1,0,0,0,1
↳
↳ mat=
↳ 1, 2, -1,0, 0, 0, 0, 0,
↳ 3, 4, 0, -1,0, 0, 0, 0,
↳ -1,0, 0, 0, 0, 0, 0, 0,
↳ 0, -1,0, 0, 0, 0, 0, 0,
↳ 0, 0, 0, 0, 5, 6, -1,0,
↳ 0, 0, 0, 0, 7, 8, 0, -1,
↳ 0, 0, 0, 0, -1,0, 0, 0,
↳ 0, 0, 0, 0, 0, -1,0, 0
↳
↳ rvec=
↳ 0,0,
↳ 0,0,
↳ 1,0,
↳ 0,1,
↳ 0,0,
↳ 0,0,
↳ 1,0,
↳ 0,1

```

### 7.10.6.23 ncrepEvaluate

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** matrix M = ncrepEvaluate(q);

**Return:** for  $q=(u, Q, v)$  calculate  $-u*Q^{(-1)}*v$

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y", "z"));
ncrat f = ncratFromString("x+y");
ncrep q = ncrepGet(f);
matrix A[2][2] = 1, 2, 3, 4;
matrix B[2][2] = 5, 6, 7, 8;

```

```

ncrep s = ncrepSubstitute(q, list(x, y), list(A, B));
matrix M = ncrepEvaluate(s);
print(M);
↳ 6, 8,
↳ 10,12

```

### 7.10.6.24 ncrepEvaluateAt

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** matrix M = ncrepEvaluateAt(q, vars, point);

**Return:** For  $q=(u, Q, v)$  calculate  $-u*Q(\text{point})^{(-1)}*v$ , that is to say, evaluate the `ncrat` represented by `q` at point (scalar or matrix point).

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("x+y");
ncrep q = ncrepGet(f);
matrix A[2][2] = 1, 2, 3, 4;
matrix B[2][2] = 5, 6, 7, 8;
matrix M = ncrepEvaluateAt(q, list(x, y), list(A, B));
print(M);
↳ 6, 8,
↳ 10,12

```

### 7.10.6.25 ncrepIsDefinedDim

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** list l = ncrepIsDefinedDim(q, N, vars, n, maxcoeff);

**Return:** list(k, list vars, list(A1, ..., Ak)), where:  
 If  $k = N$  then there are matrices  $A_1, \dots, A_k$  of size  $N$  such that  $q$  is defined at  $A = (A_1, \dots, A_k)$ , i.e.,  $q.\text{mat}$  is invertible at  $A$ .  
 If  $k = 0$  then no such point was found.

**Note:** Test whether  $q.\text{mat}$  is invertible via evaluation at random matrix points with integer coefficients in  $[-\text{maxcoeff}, \text{maxcoeff}]$ . Stops after  $n$  tries. Use square matrices of dimension  $N$ . The list `vars` contains the `nc` variables which occur in `q`.

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("inv(x*y-y*x)");
ncrep q = ncrepGet(f);
ncrepIsDefinedDim(q, 1, list(x, y), 10, 100);
↳ [1]:
↳ 0
↳ [2]:
↳ [1]:

```

```

↳ x
↳ [2]:
↳ y
↳ [3]:
↳ empty list
ncrepIsDefinedDim(q, 2, list(x, y), 10, 100);
↳ [1]:
↳ 2
↳ [2]:
↳ [1]:
↳ x
↳ [2]:
↳ y
↳ [3]:
↳ [1]:
↳ _[1,1]=-55
↳ _[1,2]=-24
↳ _[2,1]=39
↳ _[2,2]=-17
↳ [2]:
↳ _[1,1]=36
↳ _[1,2]=-58
↳ _[2,1]=-13
↳ _[2,2]=-55

```

### 7.10.6.26 ncrepIsDefined

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `list l = ncrepIsDefined(q, vars, n, maxcoeff);`

**Return:** `list(dim, list vars, list(A1, ..., Ak))`, where:  
 If `dim > 0` then there are matrices  $A_1, \dots, A_k$  of size `dim` such that  $q$  is defined at  $A = (A_1, \dots, A_k)$ , i.e.,  $q.mat$  is invertible at  $A$ .  
 If `dim = 0` then no such point was found.

**Note:** Test whether  $q.mat$  is invertible via evaluation at random matrix points with integer coefficients in `[-maxcoeff, maxcoeff]`. Stops after `n` tries. Use `ixi-matrix` in  $i$ -th try. The list `vars` contains the `nc` variables which occur in  $q$ .

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("inv(x*y-y*x)");
ncrep q = ncrepGet(f);
ncrepIsDefined(q, list(x, y), 5, 10);
↳ [1]:
↳ 2
↳ [2]:
↳ [1]:
↳ x

```

```

↳ [2]:
↳ y
↳ [3]:
↳ [1]:
↳ _[1,1]=0
↳ _[1,2]=-9
↳ _[2,1]=-2
↳ _[2,2]=7
↳ [2]:
↳ _[1,1]=8
↳ _[1,2]=-9
↳ _[2,1]=-4
↳ _[2,2]=-2
ncrat g = ncratFromString("inv(x-x)");
ncrep r = ncrepGet(g);
ncrepIsDefined(r, list(x), 5, 10);
↳ [1]:
↳ 0
↳ [2]:
↳ [1]:
↳ x
↳ [3]:
↳ empty list

```

### 7.10.6.27 ncrepIsRegular

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `list l = ncrepIsRegular(q, vars, n, maxcoeff);`

**Return:** `list(k, list vars, list(a1, ..., ak))`, where:

If  $k = 1$  then there are scalars ( $1 \times 1$ -matrices)  $a_1, \dots, a_k$  such that  $q$  is defined at  $a = (a_1, \dots, a_k)$ , i.e.,  $q.mat$  is invertible at  $a$ .  
If  $k = 0$  then no such point was found.

**Note:** Test whether  $q.mat$  is invertible via evaluation at random integers in  $[-maxcoeff, maxcoeff]$ . Stops after  $n$  tries. The list `vars` contains the  $nc$  variables which occur in  $q$ .

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("inv(x*y-y*x)");
ncrep q = ncrepGet(f);
ncrepIsRegular(q, list(x, y), 10, 100);
↳ [1]:
↳ 0
↳ [2]:
↳ [1]:
↳ x
↳ [2]:
↳ y

```

```

↳ [3]:
↳ empty list
ncrat g = ncratFromString("inv(1+x*y-y*x)");
ncrep r = ncrepGet(g);
ncrepIsRegular(r, list(x, y), 10, 100);
↳ [1]:
↳ 1
↳ [2]:
↳ [1]:
↳ x
↳ [2]:
↳ y
↳ [3]:
↳ [1]:
↳ _[1,1]=-55
↳ [2]:
↳ _[1,1]=-24

```

### 7.10.6.28 ncrepRegularZeroMinimize

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrep s = ncrepRegularZeroMinimize(q, l);`

**Return:** `ncrep s` representing the same rational function as `ncrep q`, where `s` is of minimal size

**Assumption:**

`q` is regular at zero, i.e.,  
if one substitutes in 0 for all `nc` variables in `q` then `q.mat` has to be invertible

**Note:** `list l = list(x1, ..., xn)` has to consist exactly of the `nc` variables occurring in `q`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("inv(1+x*y-y*x)");
ncrep q = ncrepGet(f);
ncrepDim(q);
↳ 11
ncrep s = ncrepRegularZeroMinimize(q, list(x, y));
ncrepDim(s);
↳ 3
s;
↳ lvec=
↳ 0,1,0
↳
↳ mat=
↳ 1, y, 0,
↳ -x,1, -y,
↳ 0, -x,1
↳
↳ rvec=
↳ 0,

```

```

↳ -1,
↳ 0
↳

```

### 7.10.6.29 ncregRegularMinimize

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrep s = ncregRegularMinimize(q, vars, point);`

**Return:** `ncrep s` representing the same rational function as `ncrep q`, where `s` is of minimal size

**Assumption:**

`q` is regular at scalar point `a`, i.e.,  
if one substitutes in `ai` for all `nc` variables `xi` in `q` then `q.mat` has to be invertible

**Note:** `list vars = list(x1, ..., xn)` has to consist exactly of the `nc` variables occurring in `q` and `list point = list(a1, ..., an)` consists of scalars

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("inv(x*y)");
ncrep q = ncrepGet(f);
ncrepDim(q);
↳ 5
ncrep s = ncregRegularMinimize(q, list(x, y), list(1, 1));
ncrepDim(s);
↳ 2
s;
↳ lvec=
↳ -1,0
↳
↳ mat=
↳ -y,x+1,
↳ 0, -x
↳
↳ rvec=
↳ 1,
↳ -1
↳

```

### 7.10.6.30 ncrepGetRegularZeroMinimal

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrep q = ncrepGetRegularZeroMinimal(f, vars);`

**Return:** `q` is a representation of `f` with minimal dimension

**Assumption:**

`f` is regular at zero, i.e.,  
`f(0)` has to be defined



**Note:** `list vars = list(x1, ..., xn)` has to consist exactly of the nc variables occurring in `f`

**Example:**

```
LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("inv(1+x*y-y*x)");
list vars = list(x, y);
ncrep q = ncrepGetRegularZeroMinimal(f, vars);
q;
↳ lvec=
↳ 0,1,0
↳
↳ mat=
↳ 1, y, 0,
↳ -x,1, -y,
↳ 0, -x,1
↳
↳ rvec=
↳ 0,
↳ -1,
↳ 0
↳
```

### 7.10.6.31 ncrepGetRegularMinimal

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `ncrep q = ncrepGetRegularMinimal(f, vars, point);`

**Return:** `q` is a representation of `f` with minimal dimension

**Assumption:**

`f` is regular at point, i.e.,  
`f(point)` has to be defined

**Note:** `list vars = list(x1, ..., xn)` has to consist exactly of the nc variables occurring in `f` and `list point = (p1, ..., pn)` of scalars such that `f(point)` is defined

**Example:**

```
LIB "ncrat.lib";
// We want to prove the Hua's identity, telling that for two
// invertible elements x,y from a division ring, one has
// inv(x+x*inv(y)*x)+inv(x+y) = inv(x)
// where inv(t) stands for the two-sided inverse of t
ncInit(list("x", "y"));
ncrat f = ncratFromString("inv(x+x*inv(y)*x)+inv(x+y)-inv(x)");
print(f);
↳ inv(x+x*inv(y)*x)+inv(x+y)-inv(x)
ncrep r = ncrepGet(f);
ncrepDim(r);
↳ 18
```

```

ncrep s = ncrepGetRegularMinimal(f, list(x, y), list(1, 1));
ncrepDim(s);
↳ 0
print(s);
↳ lvec=
↳ 0
↳
↳ mat=
↳ 1
↳
↳ rvec=
↳ 0
// since s represents the zero element, Hua's identity holds.

```

### 7.10.6.32 ncrepPencilGet

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** `list pencil = ncrepPencilGet(r, vars);`

**Return:** `pencil = list(vars, matrices)`,  
 where `vars = list(1, x1, ..., xg)` are the variables  
 occurring in `r` and `matrices = (Q0, ..., Qg)` is a list of matrices such that  
 $r.mat = Q_0 * x_0 + \dots + Q_g * x_g$   
 with  $x_0 = 1$

**Note:** `list vars = list(x1, ..., xn)` has to consist  
 exactly of the nc variables occurring in `f`

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("x*y");
ncrep r = ncrepGet(f);
print(r.mat);
↳ 0, 0, x, -1,
↳ 0, 1, -1,0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0
list l = ncrepPencilGet(r, list(x, y));
print(l[1]);
↳ [1]:
↳ 1
↳ [2]:
↳ x
↳ [3]:
↳ y
print(l[2][1]);
↳ 0, 0, 0, -1,
↳ 0, 1, -1,0,
↳ 0, -1,0, 0,
↳ -1,0, 0, 0
print(l[2][2]);
↳ 0,0,1,0,
↳ 0,0,0,0,

```

```

↳ 0,0,0,0,
↳ 0,0,0,0
print(l[2][3]);
↳ 0,0,0,0,
↳ 0,0,0,0,
↳ 1,0,0,0,
↳ 0,0,0,0

```

### 7.10.6.33 ncrepPencilCombine

Procedure from library `ncrat.lib` (see Section 7.10.6 [`ncrat.lib`], page 668).

**Usage:** matrix  $Q = \text{ncrepPencilCombine}(\text{pencil});$

**Return:** matrix  $Q = Q_0 \cdot x_0 + \dots + Q_g \cdot x_g,$   
 where  $\text{vars} = \text{list}(x_0, \dots, x_g)$  consists of polynomials and matrices  $= (Q_0, \dots, Q_g)$  is a list of matrices

**Example:**

```

LIB "ncrat.lib";
ncInit(list("x", "y"));
ncrat f = ncratFromString("x*y");
ncrep r = ncrepGet(f);
print(r.mat);
↳ 0, 0, x, -1,
↳ 0, 1, -1,0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0
list l = ncrepPencilGet(r, list(x, y));
matrix Q = ncrepPencilCombine(l);
print(Q);
↳ 0, 0, x, -1,
↳ 0, 1, -1,0,
↳ y, -1,0, 0,
↳ -1,0, 0, 0

```

## 7.11 Release Notes (letterplace)

### NEWS in SINGULAR:LETTERPLACE 4-2-0

#### News for SINGULAR:LETTERPLACE version 4-2-0

New functions:

added support for free bimodules of a fixed rank (Section 7.8.1 [`freeAlgebra (letterplace)`], page 619, Section 7.8.4 [`ncgen`], page 621)

several types of monomial orderings become available, among them three types of elimination orderings

`twostd` (Section 7.8.9 [`twostd (letterplace)`], page 624), `reduce` (Section 7.8.5 [`reduce (letterplace)`], page 622) and other functions support subbimodules

`syz` (Section 7.8.8 [`syz` (letterplace)], page 623), `lift` (Section 7.8.2 [`lift` (letterplace)], page 620), `liftstd` (Section 7.8.3 [`liftstd` (letterplace)], page 621) implemented

`bracket` (Section 7.3.2 [`bracket`], page 330) and `maxideal` (Section 5.1.88 [`maxideal`], page 216) work in Letterplace

the options `redSB`, `redTail` are effective for computations related to Groebner bases

the options `prot`, `mem` are effective for the whole LETTERPLACE subsystem

New libraries:

`fpaprops.lib`: Algorithms for properties of quotient algebras (Section 7.10.3 [`fpaprops.lib`], page 649)

`ncHilb.lib`: Hilbert functions for non-commutative algebras (Section 7.10.5 [`ncHilb.lib`], page 665)

Changed libraries:

`fpadim.lib`: Vector space dimension, basis and Hilbert series for finitely presented algebras (Section 7.10.1 [`fpadim.lib`], page 629), numerous enhancements, partially implemented in the kernel

`freegb.lib`: Main initialization and convenience tools (Section 7.10.4 [`freegb.lib`], page 652)

Changes in the kernel/build system:

SINGULAR:LETTERPLACE is available as the dynamical module

adaptions/functions for `Singular.jl` (<https://github.com/oscar-system/Singular.jl>)

## News for SINGULAR:LETTERPLACE version 4-1-2

New libraries:

`fpalgebras.lib`: Generation of various algebras in the letterplace case (Section 7.10.2 [`fpalgebras.lib`], page 634)

`ncrat.lib`: Manipulating non-commutative rational functions (Section 7.10.6 [`ncrat.lib`], page 668)

Changed/updated libraries:

`freegb.lib`: `lpDivision`, `lpPrint` (Section 7.10.4 [`freegb.lib`], page 652)

`fpadim.lib` (Section 7.10.1 [`fpadim.lib`], page 629)

`ncfactor.lib` (Section 7.5.12 [`ncfactor.lib`], page 480) is available for Letterplace rings

Changes in the kernel/build system:

code for free algebras (letterplace rings) rewritten (using now the standard `+, -, *, ^, std, ...`) (Section 7.7 [LETTERPLACE], page 613)

new command `rightstd` (Section 7.8.6 [`rightstd` (letterplace)], page 623)

extended `twostd` to LETTERPLACE (Section 7.8.9 [`twostd` (letterplace)], page 624, Section 7.3.29 [`twostd` (plural)], page 358)

## Appendix A Examples

### A.1 Programming

#### A.1.1 Basic programming

We show in the example below the following:

- define the ring  $R$  of characteristic 32003, variables  $x, y, z$ , monomial ordering  $dp$  (implementing  $F_{32003}[x, y, z]$ )
- list the information about  $R$  by typing its name
- check the order of the variables
- define the integers  $a, b, c, t$
- define a polynomial  $f$  (depending on  $a, b, c, t$ ) and display it
- define the jacobian ideal  $i$  of  $f$
- compute a Groebner basis of  $i$
- compute the dimension of the algebraic set defined by  $i$  (requires the computation of a Groebner basis)
- create and display a string in order to comment the result (text between quotes " "; is a 'string')
- load a library (see Section D.4.26 [primdec.lib], page 828)
- compute a primary decomposition for  $i$  and assign the result to a list  $L$  (which is a list of lists of ideals)
- display the number of primary components and the first primary and prime components (entries of the list  $L[1]$ )
- implement the localization of  $F_{32003}[x, y, z]$  at the homogeneous maximal ideal (generated by  $x, y, z$ ) by defining a ring with local monomial ordering ( $ds$  in place of  $dp$ )
- map  $i$  to this ring (see Section 5.1.59 [imap], page 195) - we may use the same name  $i$ , since ideals are ring dependent data
- compute the local dimension of the algebraic set defined by  $i$  at the origin (= dimension of the ideal generated by  $i$  in the localization)
- compute the local dimension of the algebraic set defined by  $i$  at the point  $(-2000, -6961, -7944)$  (by applying a linear coordinate transformation)

For a more basic introduction to programming in SINGULAR, we refer to Section 2.3 [Getting started], page 6.

```

ring R = 32003, (x,y,z), dp;
R;
↳ // coefficients: ZZ/32003
↳ // number of vars : 3
↳ // block 1 : ordering dp
↳ // : names x y z
↳ // block 2 : ordering C
x > y;
↳ 1
y > z;
↳ 1

```

```

int a,b,c,t = 1,2,-1,4;
poly f = a*x3+b*xy3-c*xz3+t*xy2z2;
f;
↳ 4xy2z2+2xy3+xz3+x3
ideal i = jacob(f); // Jacobian Ideal of f
ideal si = std(i); // compute Groebner basis
int dimi = dim(si);
string s = "The dimension of V(i) is "+string(dimi)+".";
s;
↳ The dimension of V(i) is 1.
LIB "primdec.lib"; // load library primdec.lib
list L = primdecGTZ(i);
size(L); // number of prime components
↳ 6
L[1][1]; // first primary component
↳ _[1]=2y2z2+y3-16001z3
↳ _[2]=x
L[1][2]; // corresponding prime component
↳ _[1]=2y2z2+y3-16001z3
↳ _[2]=x
ring Rloc = 32003,(x,y,z),ds; // ds = local monomial ordering
ideal i = imap(R,i);
dim(std(i));
↳ 1
map phi = R, x-2000, y-6961, z-7944;
dim(std(phi(i)));
↳ 0

```

### A.1.2 Writing procedures and libraries

The user may add their own commands to the commands already available in SINGULAR by writing SINGULAR procedures. There are basically two kinds of procedures:

- procedures written in the SINGULAR programming language (which are usually collected in SINGULAR libraries).
- procedures written in C/C++ (collected in dynamic modules).

At this point, we restrict ourselves to describing the first kind of (library) procedures, which are sufficient for most applications. The syntax and general structure of a library (procedure) is described in Section 3.7 [Procedures], page 50, and Section 3.8 [Libraries], page 54.

The probably most efficient way of writing a new library is to use one of the official SINGULAR libraries, say `ring.lib` as a sample. On a Unix-like operating system, type `LIB "ring.lib"`; to get information on where the libraries are stored on your disk.

SINGULAR provides several commands and tools, which may be useful when writing a procedure, for instance, to have a look at intermediate results (see Section 3.9 [Debugging tools], page 67).

If such a library should be contributed to SINGULAR some formal requirements are needed:

the library header must explain the purpose of the library and (for non-trivial algorithm) a pointer to the algorithm (text book, article, etc.)

all global procedures must have a help string and an example which shows its usage.

it is strongly recommend also to provide test scripts which test the functionality: one should test the essential functionality of the library/command in a relatively short time (say, in no more than 30s), other tests should check the functionality of the library/command in detail so

that, if possible, all relevant cases/results are tested. Nevertheless, such a test should not run longer than, say, 10 minutes.

We give short examples of procedures to demonstrate the following:

- Write procedures which return an integer (ring independent), see also Section A.4.1 [Milnor and Tjurina number], page 726. (Here we restrict ourselves to the main body of the procedures).
  - The procedure `milnorNumber` must be called with one parameter, a polynomial. The name `g` is local to the procedure and is killed automatically when leaving the procedure. `milnorNumber` returns the Milnor number (and displays a comment).
  - The procedure `tjurinaNumber` has no specified number of parameters. Here, the parameters are referred to by `#[1]` for the 1st, `#[2]` for the 2nd parameter, etc. `tjurinaNumber` returns the Tjurina number (and displays a comment).
  - the procedure `milnor_tjurina` which returns a list consisting of two integers, the Milnor and the Tjurina number.
- Write a procedure which creates a new ring and returns data dependent on this new ring (two numbers) and an int. In this example, we also show how to write a help text for the procedure (which is optional, but recommended).

```

proc milnorNumber (poly g)
{
 "Milnor number:";
 return(vdim(std(jacob(g))));
}

proc tjurinaNumber
{
 "Tjurina number:";
 return(vdim(std(jacob(#[1])+#[1])));
}

proc milnor_tjurina (poly f)
{
 ideal j=jacob(f);
 list L=vdim(std(j)),vdim(std(j+f));
 return(L);
}

proc real_sols (number b, number c)
"USAGE: real_sols (b,c); b,c number
ASSUME: active basering has characteristic 0
RETURN: list: first entry is an integer (the number of different real
solutions). If this number is non-negative, the list has as second
entry a ring in which the list SOL of real solutions of x^2+bx+c=0
is stored (as floating point number, precision 30 digits).
NOTE: This procedure calls laguerre_solve from solve.lib.
"
{
 def oldring = basering; // assign name to the ring active when
 // calling the procedure

 number disc = b^2-4*c;
 if (disc>0) { int n_of_sols = 2; }
 if (disc==0) { int n_of_sols = 1; }
 string s = nameof(var(1)); // name of first ring variable

```

```

if (disc>=0) {
 execute("ring rinC =(complex,30),("+s+"),lp;");
 if (not(defined(laguerre_solve))) { LIB "solve.lib"; }
 poly f = x2+imap(olddring,b)*x+imap(olddring,c);
 // f is a local ring-dependent variable
 list SOL = laguerre_solve(f,30);
 export SOL; // make SOL a global ring-dependent variable
 // such variables are still accessible when the
 // ring is among the return values of the proc

 setring olddring;
 return(list(n_of_sols,rinC));
}
else {
 return(list(0));
}
}

//
// We now apply the procedures which are defined by the
// lines of code above:
//
ring r = 0,(x,y),ds;
poly f = x7+y7+(x-y)^2*x2y2;

milnorNumber(f);
↳ Milnor number:
↳ 28
tjurinaNumber(f);
↳ Tjurina number:
↳ 24
milnor_tjurina(f); // a list containing Milnor and Tjurina number
↳ [1]:
↳ 28
↳ [2]:
↳ 24

def L=real_sols(2,1);
L[1]; // number of real solutions of x^2+2x+1
↳ 1
def R1=L[2];
setring R1;
listvar(R1); // only global ring-dependent objects are still alive
↳ // R1 [0] *ring
↳ // SOL [0] list, size: 2
SOL; // the real solutions
↳ [1]:
↳ -1
↳ [2]:
↳ -1

setring r;
L=real_sols(1,1);
L[1]; // number of reals solutions of x^2+x+1

```



```

↳ 0

setring r;
L=real_sols(1,-5);
L[1]; // number of reals solutions of x^2+x-5
↳ 2
def R3=L[2];
setring R3; SOL; // the real solutions
↳ [1]:
↳ -2.79128784747792000329402359686
↳ [2]:
↳ 1.79128784747792000329402359686

```

Writing a dynamic module is not as simple as writing a library procedure, since it does not only require some knowledge of C/C++, but also about the way the SINGULAR kernel works. See also Section A.1.9 [Dynamic modules], page 701.

### A.1.3 Rings associated to monomial orderings

In SINGULAR we may implement localizations of the polynomial ring by choosing an appropriate monomial ordering (when defining the ring by the `ring` command). We refer to Section B.2 [Monomial orderings], page 760 for a thorough discussion of the monomial orderings available in SINGULAR.

At this point, we restrict ourselves to describing the relation between a monomial ordering and the ring (as mathematical object) which is implemented by the ordering. This is most easily done by describing the set of units: if  $>$  is a monomial ordering then precisely those elements which have leading monomial 1 are considered as units (in all computations performed with respect to this ordering).

In mathematical terms: choosing a monomial ordering  $>$  implements the localization of the polynomial ring with respect to the multiplicatively closed set of polynomials with leading monomial 1.

That is, choosing  $>$  implements the ring

$$K[x]_{>} := \left\{ \frac{f}{u} \mid f, u \in K[x], LM(u) = 1 \right\}.$$

If  $>$  is global (that is, 1 is the smallest monomial), the implemented ring is just the polynomial ring. If  $>$  is local (that is, if 1 is the largest monomial), the implemented ring is the localization of the polynomial ring w.r.t. the homogeneous maximal ideal. For a mixed ordering, we obtain "something in between these two rings":

```

ring R = 0,(x,y,z),dp; // polynomial ring (global ordering)
poly f = y4z3+2x2y2z2+4z4+5y2+1;
f; // display f in a degrevlex-ordered way
↳ y4z3+2x2y2z2+4z4+5y2+1
short=0; // avoid short notation
f;
↳ y^4*z^3+2*x^2*y^2*z^2+4*z^4+5*y^2+1
short=1;
leadmonom(f); // leading monomial
↳ y4z3

ring r = 0,(x,y,z),ds; // local ring (local ordering)

```

```

poly f = fetch(R,f);
f; // terms of f sorted by degree
↪ 1+5y2+4z4+2x2y2z2+y4z3
leadmonom(f); // leading monomial
↪ 1

// Now we implement more "advanced" examples of rings:
//
// 1) (K[y]_<y>)[x]
//
int n,m=2,3;
ring A1 = 0,(x(1..n),y(1..m)),(dp(n),ds(m));
poly f = x(1)*x(2)^2+1+y(1)^10+x(1)*y(2)^5+y(3);
leadmonom(f);
↪ x(1)*x(2)^2
leadmonom(1+y(1)); // unit
↪ 1
leadmonom(1+x(1)); // no unit
↪ x(1)

//
// 2) some ring in between (K[x]_<x>)[y] and K[x,y]_<x>
//
ring A2 = 0,(x(1..n),y(1..m)),(ds(n),dp(m));
leadmonom(1+x(1)); // unit
↪ 1
leadmonom(1+x(1)*y(1)); // unit
↪ 1
leadmonom(1+y(1)); // no unit
↪ y(1)

//
// 3) K[x,y]_<x>
//
ring A4 = (0,y(1..m)),(x(1..n)),ds;
leadmonom(1+y(1)); // in ground field
↪ 1
leadmonom(1+x(1)*y(1)); // unit
↪ 1
leadmonom(1+x(1)); // unit
↪ 1

```

Note, that even if we implicitly compute over the localization of the polynomial ring, most computations are explicitly performed with polynomial data only. In particular,  $1/(1-x)$ ; does not return a power series expansion or a fraction but 0 (division with remainder in polynomial ring). See Section 5.1.26 [division], page 172 for division with remainder in the localization and [invunit], page 866 for a procedure returning a truncated power series expansion of the inverse of a unit.

#### A.1.4 Long coefficients

The following innocent example produces in its standard basis extremely long coefficients in char 0 for the lexicographical ordering. But a very small deformation does not (the undeformed example

is degenerated with respect to the Newton boundary). This example demonstrates that it might be wise, for complicated examples, to do the calculation first in positive char (e.g., 32003). It has been shown, that in complicated examples, more than 95 percent of the time needed for a standard basis computation is used in the computation of the coefficients (in char 0). The representation of long integers with real is demonstrated.

```

timer = 1; // activate the timer
ring R0 = 0,(x,y),lp;
poly f = x5+y11+xy9+x3y9;
ideal i = jacob(f);
ideal i1 = i,i[1]*i[2]; // undeformed ideal
ideal i2 = i,i[1]*i[2]+1/1000000*x5y8; // deformation of i1
i1; i2;
⇒ i1[1]=5x4+3x2y9+y9
⇒ i1[2]=9x3y8+9xy8+11y10
⇒ i1[3]=45x7y8+27x5y17+45x5y8+55x4y10+36x3y17+33x2y19+9xy17+11y19
⇒ i2[1]=5x4+3x2y9+y9
⇒ i2[2]=9x3y8+9xy8+11y10
⇒ i2[3]=45x7y8+27x5y17+4500001/1000000x5y8+55x4y10+36x3y17+33x2y19+9xy17+1\
1y19
ideal j = std(i1);
j;
⇒ j[1]=264627y39+26244y35-1323135y30-131220y26+1715175y21+164025y17+1830125\
y16
⇒ j[2]=12103947791971846719838321886393392913750065060875xy8-28639152114168\
3198701331939250003266767738632875y38-31954402206909026926764622877573565\
78554430672591y37+57436621420822663849721381265738895282846320y36+1657764\
214948799497573918210031067353932439400y35+213018481589308191195677223898\
98682697001205500y34+1822194158663066565585991976961565719648069806148y33\
-4701709279892816135156972313196394005220175y32-1351872269688192267600786\
97600850686824231975y31-3873063305929810816961516976025038053001141375y30\
+1325886675843874047990382005421144061861290080000y29+1597720195476063141\
9467945895542406089526966887310y28-26270181336309092660633348002625330426\
7126525y27-7586082690893335269027136248944859544727953125y26-867853074106\
49464602285843351672148965395945625y25-5545808143273594102173252331151835\
700278863924745y24+19075563013460437364679153779038394895638325y23+548562\
322715501761058348996776922561074021125y22+157465452677648386073957464715\
68100780933983125y21-1414279129721176222978654235817359505555191156250y20\
-20711190069445893615213399650035715378169943423125y19+272942733337472665\
573418092977905322984009750y18+789065115845334505801847294677413365720955\
3750y17+63554897038491686787729656061044724651089803125y16-22099251729923\
906699732244761028266074350255961625y14+147937139679655904353579489722585\
91339027857296625y10
⇒ j[3]=5x4+3x2y9+y9
// Compute average coefficient length (=51) by
// - converting j[2] to a string in order to compute the number
// of characters
// - divide this by the number of monomials:
size(string(j[2])) div size(j[2]);
⇒ 51
vdim(j);
⇒ 63
// For a better representation normalize the long coefficients

```

```

// of the polynomial j[2] and map it to real:
poly p=(1/12103947791971846719838321886393392913750065060875)*j[2];
ring R1=real,(x,y),lp;
short=0; // force the long output format
poly p=imap(R0,p);
p;
↳ x*y^8-(2.366e-02)*y^38-(2.640e-01)*y^37+(4.745e-06)*y^36+(1.370e-04)*y^35\
+(1.760e-03)*y^34+(1.505e-01)*y^33-(3.884e-07)*y^32-(1.117e-05)*y^31-(3.2\
00e-04)*y^30+(1.095e-01)*y^29+(1.320e+00)*y^28-(2.170e-05)*y^27-(6.267e-0\
4)*y^26-(7.170e-03)*y^25-(4.582e-01)*y^24+(1.576e-06)*y^23+(4.532e-05)*y^\
22+(1.301e-03)*y^21-(1.168e-01)*y^20-(1.711e+00)*y^19+(2.255e-05)*y^18+(6\
.519e-04)*y^17+(5.251e-03)*y^16-(1.826e+00)*y^14+(1.222e+00)*y^10
// Compute a standard basis for the deformed ideal:
setring R0; // return to the original ring R0
j = std(i2);
j;
↳ j[1]=y16
↳ j[2]=65610xy8+17393508y27+7223337y23+545292y19+6442040y18-119790y14+80190\
y10
↳ j[3]=5x4+3x2y9+y9
vdim(j);
↳ 40

```

### A.1.5 Parameters

Let us deform the ideal in Section A.1.4 [Long coefficients], page 696 by introducing a parameter  $t$  and compute over the ground field  $Q(t)$ . We compute the dimension at the generic point, i.e.,  $\dim_{Q(t)} Q(t)[x, y]/j$ . (This gives the same result as for the deformed ideal above. Hence, the above small deformation was "generic".)

For almost all  $a \in Q$  this is the same as  $\dim_Q Q[x, y]/j_0$ , where  $j_0 = j|_{t=a}$ .

```

ring Rt = (0,t),(x,y),lp;
Rt;
↳ // coefficients: QQ(t)
↳ // number of vars : 2
↳ // block 1 : ordering lp
↳ // : names x y
↳ // block 2 : ordering C
poly f = x5+y11+xy9+x3y9;
ideal i = jacob(f);
ideal j = i,i[1]*i[2]+t*x5y8; // deformed ideal, parameter t
vdim(std(j));
↳ 40
ring R=0,(x,y),lp;
ideal i=imap(Rt,i);
int a=random(1,30000);
ideal j=i,i[1]*i[2]+a*x5y8; // deformed ideal, fixed integer a
vdim(std(j));
↳ 40

```

### A.1.6 Formatting output

We show how to insert the result of a computation inside a text by using strings. First we compute the powers of 2 and comment the result with some text. Then we do the same and give the output a nice format by computing and adding appropriate space.

```
// The powers of 2:
int n;
for (n = 2; n <= 128; n = n * 2)
{"n = " + string (n);}
↳ n = 2
↳ n = 4
↳ n = 8
↳ n = 16
↳ n = 32
↳ n = 64
↳ n = 128
// The powers of 2 in a nice format
int j;
string space = "";
for (n = 2; n <= 128; n = n * 2)
{
 space = "";
 for (j = 1; j <= 5 - size (string (n)); j = j+1)
 { space = space + " "; }
 "n =" + space + string (n);
}
↳ n = 2
↳ n = 4
↳ n = 8
↳ n = 16
↳ n = 32
↳ n = 64
↳ n = 128
```

### A.1.7 Cyclic roots

We write a procedure returning a string that enables us to create automatically the ideal of cyclic roots over the basering with  $n$  variables. The procedure assumes that the variables consist of a single letter each (hence no indexed variables are allowed; the procedure `cyclic` in `polylib.lib` does not have this restriction). Then we compute a standard basis of this ideal and some numerical information. (This ideal is used as a classical benchmark for standard basis computations).

```
// We call the procedure 'cyclic':
proc cyclic (int n)
{
 string vs = varstr(basing)+varstr(basing);
 int c=find(vs,",");
 while (c!=0)
 {
 vs=vs[1,c-1]+vs[c+1,size(vs)];
 c=find(vs,",");
 }
 string t,s;
 int i,j;
```

```

 for (j=1; j<=n-1; j=j+1)
 {
 t="";
 for (i=1; i <=n; i=i+1)
 {
 t = t + vs[i,j] + "+";
 }
 t = t[1,size(t)-1] + ","+newline;
 s=s+t;
 }
 s=s+vs[1,n]+"-1";
 return (s);
}

ring r=0,(a,b,c,d,e),lp; // basering, char 0, lex ordering
string sc=cyclic(nvars(basing));
sc; // the string of the ideal
↳ a+b+c+d+e,
↳ ab+bc+cd+de+ea,
↳ abc+bcd+cde+dea+eab,
↳ abcd+bcde+cdea+deab+eabc,
↳ abcde-1
execute("ideal i="+sc+"); // this defines the ideal of cyclic roots
i;
↳ i[1]=a+b+c+d+e
↳ i[2]=ab+bc+cd+de+ea
↳ i[3]=abc+bcd+cde+dea+eab
↳ i[4]=abcd+bcde+cdea+deab+eabc
↳ i[5]=abcde-1
timer=1;
ideal j=std(i);
↳ //used time: 7.5 sec
size(j); // number of elements in the std basis
↳ 11
degree(j);
↳ // codimension = 5
↳ // dimension = 0
↳ // degree = 70

```

### A.1.8 Parallelization with ssi links

In this example, we demonstrate how ssi links can be used to parallelize computations.

To compute a standard basis for a zero-dimensional ideal in the lexicographical ordering, one of the two powerful routines `stdhilb` (see [stdhilb], page 785) and `stdfglm` (see [stdfglm], page 785) should be used. However, in general one cannot predict which one of the two commands is faster. This very much depends on the (input) example. Therefore, we use ssi links to let both commands work on the problem independently and in parallel, so that the one which finishes first delivers the result.

The example we use is the so-called "omndi example". See *Tim Wichmann; Der FGLM-Algorithmus: verallgemeinert und implementiert in Singular; Diplomarbeit Fachbereich Mathematik, Universitaet Kaiserslautern; 1997* for more details.

```
ring r=0,(a,b,c,u,v,w,x,y,z),lp;
```

```

ideal i=a+c+v+2x-1, ab+cu+2vw+2xy+2xz-2/3, ab2+cu2+2vw2+2xy2+2xz2-2/5,
ab3+cu3+2vw3+2xy3+2xz3-2/7, ab4+cu4+2vw4+2xy4+2xz4-2/9, vw2+2xyz-1/9,
vw4+2xy2z2-1/25, vw3+xyz2+xy2z-1/15, vw4+xyz3+xy3z-1/21;

link l_hilb,l_fgml = "ssi:fork","ssi:fork"; // 1.

open(l_fgml); open(l_hilb);

write(l_hilb, quote(stdhilb(i))); // 2.
write(l_fgml, quote(stdfgml(eval(i))));

list L=list(l_hilb,l_fgml); // 3.
int l_index=waitfirst(L);

if (l_index==1)
{
 "stdhilb won !!!!"; size(read(L[1]));
 close(L[1]); close(L[2]);
}
else
{
 "stdfgml won !!!!"; size(read(L[2]));
 close(L[1]); close(L[2]);
}
⇒ stdfgml won !!!!
⇒ 9

```

Some explanatory remarks are in order:

1. Instead of using links of the type `ssi:fork`, we alternatively could use `ssi:tcp` links such that the two "competing" SINGULAR processes run on different machines. This has the advantage of "true" parallel computing since no resource sharing is involved (as it usually is with forked processes).
2. Notice how quoting is used in order to prevent local evaluation (i.e., local computation of results). Since we "forked" the two competing processes, the identifier `i` is defined and has identical values in both child processes. Therefore, the innermost `eval` can be omitted (as is done for the `l_hilb` link), and only the identifier `i` needs to be communicated to the children. However, when `ssi:tcp` links are used, the inner evaluation must be applied so that actual values, and not the identifiers are communicated (as is done for the `l_fgml` link in our example).
3. We wait until one of the two children finished the computation. The main process sleeps (i.e., suspends its execution) in the intermediate time.
4. The child which has won delivers the result and is terminated with the usual `close` command. The other child which is still computing needs to be terminated by an explicit (i.e., system) kill command if running on a different computer. For `ssi:fork` a `close` is sufficient.

### A.1.9 Dynamic modules

The purpose of the following example is to illustrate the use of dynamic modules. Giving an example on how to write a dynamic module is beyond the scope of this manual. A technical reference is given at <https://www.singular.uni-kl.de/Manual/modules.pdf>.

In this example, we use a dynamic module, residing in the file `kstd.so`, which allows ignoring all but the first `j` entries of vectors when forming the pairs in the standard basis computation.

```

ring r=0,(x,y),dp;
module mo=[x^2-y^2,1,0,0],[xy+y^2,0,1,0],[y^2,0,0,1];
print(mo);

// load dynamic module - at the same time creating package Kstd
// procedures will be available in the packages Top and Kstd
LIB("kstd.so");
listvar(package);

// set the number of components to be considered to 1
module mostd=kstd(mo,1); // calling procedure in Top
 // obviously computation ignored pairs with leading
 // term in the second entry
print(mostd);

// now consider 2 components
module mostd2=Kstd::kstd(mo,2); // calling procedure in Kstd
 // this time the previously unconsidered pair was
 // treated too
print(mostd2);

```

## A.2 Computing Groebner and Standard Bases

### A.2.1 groebner and std

The basic version of Buchberger's algorithm leaves a lot of freedom in carrying out the computational process. Considerable improvements are obtained by implementing criteria for reducing the number of S-polynomials to be actually considered (e.g., by applying the product criterion or the chain criterion). We refer to Cox, Little, and O'Shea [1997], Chapter 2 for more details and references on these criteria and on further strategies for improving the performance of Buchberger's algorithm (see also Greuel, Pfister [2002]).

SINGULAR's implementation of Buchberger's algorithm is available via the `std` command ('std' referring to standard basis). The computation of reduced Groebner and standard bases may be forced by setting `option(redSB)` (see Section 5.1.110 [option], page 230).

However, depending on the monomial ordering of the active basering, it may be advisable to use the `groebner` command instead. This command is provided by the SINGULAR library `standard.lib` which is automatically loaded when starting a SINGULAR session. Depending on some heuristics, `groebner` either refers to the `std` command (e.g., for rings with ordering `dp`), or to one of the algorithms described in the sections Section A.2.2 [Groebner basis conversion], page 704, Section A.2.3 [slim Groebner bases], page 706. For information on the heuristics behind `groebner`, see the library file `standard.lib` (see also Section 2.3.3 [Procedures and libraries], page 10).

We apply the commands `std` and `groebner` to compute a lexicographic Groebner basis for the ideal of cyclic roots over the basering with 6 variables (see Section A.1.7 [Cyclic roots], page 699). We set `option(prot)` to make SINGULAR display some information on the performed computations (see Section 5.1.110 [option], page 230 for an interpretation of the displayed symbols). For long running computations, it is always recommended to set this option.

```

LIB "polylib.lib";
ring r=32003,(a,b,c,d,e,f),lp;
ideal I=cyclic(6);

```



```

option(prot);
int t=timer;
system("--ticks-per-sec", 100); // give time in 1/100 sec
ideal sI=std(I);
↳ [1048575:2]1(5)s2(4)s3(3)s4s(4)s5(6)s(9)s(11)s(14)s(17)-s6s(19)s(21)s(24)\
s(27)s(30)s(33)s(35)s(38)s(41)ss(42)-s----s7(41)s(43)s(46)s(48)s(51)s(54)\
s(56)s(59)s(62)s(63)s(65)s(66)s(68)s(70)s(73)s(75)s(78)---ss(81)-----\
--s(73)-----8-s(66)s(69)s(72)s(75)s(77)s(80)s(81)s(83)s(85)s(88)s(91)s\
(93)s(96)s(99)s(102)s(105)s(107)s(110)s(113)----- (100)-----\
s(101)s(108)s(110)----- (100)-----9-s(94)s(97)s(99)s(84)s(74)s(77)\
s(80)---ss(83)s(86)s(73)s(76)s10(78)s(81)s(82)s(84)s(86)s(89)s(92)s(94)s(\
97)s(100)s(103)s(82)s(84)s(86)s(89)s(92)s(95)s11(98)s(87)s(90)s(93)s(95)s\
(98)s(101)s(104)----(100)---12-s(99)s(90)s(93)s(92)-----s(86)-----\
---13-s(74)s(77)s(79)s(82)s(85)s(88)-----14-s(64)s(67)ss(70)\
s(73)s(77)s(81)-----15-s(57)s(65)s(68)ss(71)-----\
-----s(57)----16-s(55)ss(56)-----17-s(34)s(32)-----\
-18-s(26)s(28)s-----19-s(25)s(28)s(31)-----20-s(27)s(30)s(35)-----21-s\
(23)s(26)-----22-s(22)-----23-s(15)24-s(17)-s(19)--25-s(18)s(19)s26-s(2\
1)-----27-s(11)28-s(13)--29-s(12)-30--s--31-s(11)---32-s33(7)s(10)---\
34-s-35----36-s37(6)s38s39s40---42-s43(5)s44s45--48-s49s50s51---54-s55(4)\
--67-86-
↳ product criterion:664 chain criterion:2844
timer-t; // used time (in 1/100 secs)
↳ 10
size(sI);
↳ 17
t=timer;
sI=groebner(I);
↳ compute hilbert series with std in ring (ZZ/32003),(a,b,c,d,e,f,@),(dp(7)\
,C)
↳ weights used for hilbert series: 1,1,1,1,1,1
↳ [65535:2]1(5)s2(4)s3(3)s4ss5(4)s(5)s(7)-s6(8)s(9)s(11)s(13)s(16)s(18)s(21\
)--s7(22)s(23)s(24)s(27)s(29)s(31)s(32)s(35)-s(37)s(40)s(42)s(44)s(45)--s\
(46)s(48)----8-s(44)s(47)s(50)s(52)s(55)s(57)s(59)s(61)-s(63)----s(62)--\
--s(61)s(64)-s(66)-----s(58)-----9-s(53)s(56)s(59)s(62)s(65)s(68)\
s(71)s(74)s(77)s(80)s(83)s(86)s(90)s(95)s(102)s(108)----- (100)-----\
-----s(81)---10-s(83)s(88)s(90)s(94)s(99)s(104)s(109)s(114)-s(11\
6)s(121)s(126)s(128)s(132)----- (100)-----\
----11-s(87)-----12-s(50)-----13-s(4\
4)s(47)s(51)s(55)-----14-s(45)s(48)s(51)s(55)s(58)s(61)s(64)s(67)\
s(70)-----15-s(52)s(55)s(58)s(61)s(64)s(67)s(70)s(73)s(76)\
s(79)s(82)-----16-----\
-----17-
↳ product criterion:284 chain criterion:4184
↳ std with hilb in (ZZ/32003),(a,b,c,d,e,f,@),(lp(6),dp(1),C)
↳ [65535:2]1(98)s2(97)s3(96)s4s(97)-s5(98)s(101)s(103)s(106)s(109)---s6(107\
)s(109)s(111)s(114)s(117)s(120)s(123)s(125)s(128)s(131)ss(132)-s-----\
-s7(125)s(127)s(130)s(132)s(135)s(138)s(140)s(143)s(146)s(147)s(149)s(150\
)s(152)s(154)s(157)s(159)s(162)---ss(165)-----shhhhhhhhhhhhhhhhhhh\
hhh8(134)s(136)s(139)s(142)s(145)s(147)s(150)s(151)s(153)s(155)s(158)s(16\
1)s(163)s(166)s(169)s(172)s(175)s(177)s(180)s(183)-----\
-s(171)s(178)shhhhhhhhhhhhhhhhhhhhhhhhhhhhhhhhhhhhh9(147)s(150)s(153)s(155)s(\
181)s(184)s(187)s(190)s(203)s(208)s(213)s(217)s(218)s(220)s(222)s(225)---\

```





```

↳ 0
 size(j2); // size (no. of polys) in computed GB
↳ 5
 // usual Groebner basis computation for lex ordering
 t=timer;
 ideal j0 =std(i);
↳ [1048575:2]1(4)s2(3)s3(2)s4s(3)s5(5)s(4)s6(6)s(7)s(9)s(8)sss7(10)s(11)s(1\
0)s(11)s(13)s8(12)s(13)s(15)s.s(14).s.9.s(16)s(17)s(19).....10.s(20).s\
(21)ss..11.s(23)s(25).ss(27)...s(28)s(26)...12.s(25)sss(23)sss.....s(22\
)...13.s(23)sssssss(21)s(22)sssss(21)ss..14.ss(22)s.s.sssss(21)s(22)sss.\
s...15.ssss(21)s(22)sssssssss(21)s(22)sss16.ssssssss(21)s(22)sssssssss\
17ss(21)s(22)sssssssss(21)sss(22)ss(21)ss18(22)s(21)s(22)s.s.....\
..19.ssss(21)ss(22)sssssssss(21)s(22)s20.ssssssssss(21)s.....21.s(22\
)sssssssssssssss(21)s(22)ssss22sssssssssssss(21)s(22)sssssss23sssssssssss(\
21)s(22)sssssssss24sssssssssssss(21)s(22)sssssssss25sssssssssss(21)s(22)ssssss\
sss26sssssssssss(21)s(20)sssssssss27.ssssssssss.....s28.ssssss.....\
...29.sssssssssssssssssss30ssssssssssssssssss31.sssssssssssssssss32.s\
ssssssssssssssssss33ssssssssssssssssss34ssssssssssssssssss35ssssssss\
ssssssssssss36ssssssssssssssssss37ssssssssssssssssss38ssssssssssssss\
ssss39ssssssssssssssssss40ssssssssssssssssss41ssss-----42-\
s(4)--43-s44s45s46s47s48s49s50s51s52s53s54s55s56s
↳ product criterion:1395 chain criterion:904
 option(noprot);
 timer-t;
↳ 0

```

### A.2.3 slim Groebner bases

The command `slimgb` calls an implementation of an algorithm to compute Groebner bases which is designed for keeping the polynomials slim (short with small coefficients) during a Groebner basis computation. It provides, in particular, a fast algorithm for computing Groebner bases over function fields or over the rational numbers, but also in several other cases. The algorithm which is still under development was developed in the diploma thesis of Michael Brickenstein. It has been published as [https://www.singular.uni-kl.de/reports/35/paper\\_35\\_full.ps.gz](https://www.singular.uni-kl.de/reports/35/paper_35_full.ps.gz).

In the example below (Groebner basis with respect to degree reverse lexicographic ordering over function field) `slimgb` is much faster than the `std` command.

```

ring r=(32003,u1, u2, u3, u4),(x1, x2, x3, x4, x5, x6, x7),dp;
timer=1;
ideal i=
-x4*u3+x5*u2,
x1*u3+2*x2*u1-2*x2*u2-2*x3*u3-u1*u4+u2*u4,
-2*x1*x5+4*x4*x6+4*x5*x7+x1*u3-2*x4*u1-2*x4*u4-2*x6*u2-2*x7*u3+u1*u2+u2*u4,
-x1*x5+x1*x7-x4*u1+x4*u2-x4*u4+x5*u3+x6*u1-x6*u2+x6*u4-x7*u3,
-x1*x4+x1*u1-x5*u1+x5*u4,
-2*x1*x3+x1*u3-2*x2*u4+u1*u4+u2*u4,
x1^2*u3+x1*u1*u2-x1*u2^2-x1*u3^2-u1*u3*u4+u3*u4^2;
i=slimgb(i);

```

For detailed information and limitations see Section 5.1.143 [`slimgb`], page 260.

## A.3 Commutative Algebra

### A.3.1 Saturation

For any two ideals  $i, j$  in the basering  $R$  let

$$\text{sat}(i, j) = \{x \in R \mid \exists n \text{ s.t. } x \cdot (j^n) \subseteq i\} = \bigcup_{n=1}^{\infty} i : j^n$$

denote the saturation of  $i$  with respect to  $j$ . This defines, geometrically, the closure of the complement of  $V(j)$  in  $V(i)$  (where  $V(i)$  denotes the variety defined by  $i$ ).

The saturation is computed by the procedure `sat` in `elim.lib` by computing iterated ideal quotients with the maximal ideal. `sat` returns a list of two elements: the saturated ideal and the number of iterations.

We apply saturation to show that a variety has no singular points outside the origin (see also Section A.4.2 [Critical points], page 728). We choose  $m$  to be the homogeneous maximal ideal (note that `maxideal(n)` denotes the  $n$ -th power of the maximal ideal). Then  $V(i)$  has no singular point outside the origin if and only if  $\text{sat}(j + (f), m)$  is the whole ring, that is, generated by 1.

```
LIB "elim.lib"; // loading library elim.lib
ring r2 = 32003, (x,y,z), dp;
poly f = x^11+y^5+z^(3*3)+x^(3+2)*y^(3-1)+x^(3-1)*y^(3-1)*z3+
 x^(3-2)*y^3*(y^2)^2;
ideal j=jacob(f);
sat(j+f,maxideal(1));
⇒ [1]:
⇒ _[1]=1
⇒ [2]:
⇒ 17
 // list the variables defined so far:
 listvar();
⇒ // r2 [0] *ring
⇒ // j [0] ideal, 3 generator(s)
⇒ // f [0] poly
```

### A.3.2 Finite fields

We define a variety in the  $n$ -space of codimension 2 defined by polynomials of degree  $d$  with generic coefficients over the prime field  $Z/p$  and look for zeros on the torus. First over the prime field and then in the finite extension field with  $p^k$  elements. In general there will be many more solutions in the second case. (Since the SINGULAR language is interpreted, the evaluation of many `for`-loops is not very fast):

```
int p=3; int n=3; int d=5; int k=2;
ring rp = p, (x(1..n)), dp;
int s = size(maxideal(d));
s;
⇒ 21
 // create a dense homogeneous ideal m, all generators of degree d, with
 // generic (random) coefficients:
 ideal m = maxideal(d)*random(p,s,n-2);
 m;
⇒ m[1]=x(1)^3*x(2)^2-x(1)*x(2)^4+x(1)^4*x(3)-x(1)^3*x(2)*x(3)+x(1)*x(2)^3*x\
 (3)+x(2)^4*x(3)+x(2)^3*x(3)^2+x(1)*x(2)*x(3)^3+x(1)*x(3)^4-x(3)^5
 // look for zeros on the torus by checking all points (with no component 0)
```

```

// of the affine n-space over the field with p elements :
ideal mt;
int i(1..n); // initialize integers i(1),...,i(n)
int l;
s=0;
for (i(1)=1;i(1)<p;i(1)=i(1)+1)
{
 for (i(2)=1;i(2)<p;i(2)=i(2)+1)
 {
 for (i(3)=1;i(3)<p;i(3)=i(3)+1)
 {
 mt=m;
 for (l=1;l<=n;l=l+1)
 {
 mt=subst(mt,x(l),i(l));
 }
 if (size(mt)==0)
 {
 "solution:",i(1..n);
 s=s+1;
 }
 }
 }
}
}

```

$\mapsto$  solution: 1 1 2  
 $\mapsto$  solution: 1 2 1  
 $\mapsto$  solution: 1 2 2  
 $\mapsto$  solution: 2 1 1  
 $\mapsto$  solution: 2 1 2  
 $\mapsto$  solution: 2 2 1

```

"//",s,"solutions over GF("+string(p)+")";

```

$\mapsto$  // 6 solutions over GF(3)  
// Now go to the field with  $p^3$  elements:  
// As long as there is no map from  $\mathbb{Z}/p$  to the field with  $p^3$  elements  
// implemented, use the following trick: convert the ideal to be mapped  
// to the new ring to a string and then execute this string in the  
// new ring  
string ms="ideal m="+string(m)+";";
ms;

$\mapsto$  ideal m= $x(1)^3x(2)^2-x(1)x(2)^4+x(1)^4x(3)-x(1)^3x(2)x(3)+x(1)x(2)^3x(3)+x(2)^4x(3)+x(2)^3x(3)^2+x(1)x(2)x(3)^3+x(1)x(3)^4-x(3)^5$ ;  
// define a ring rpk with  $p^k$  elements, call the primitive element z. Hence  
// 'solution exponent: 0 1 5' means that  $(z^0, z^1, z^5)$  is a solution  
ring rpk=( $p^k, z$ ), (x(1..n)), dp;  
rpk;

$\mapsto$  // coefficients: ZZ/9[z]  
 $\mapsto$  // minpoly :  $1*z^2+2*z^1+2*z^0$   
 $\mapsto$  // number of vars : 3  
 $\mapsto$  // block 1 : ordering dp  
 $\mapsto$  // : names x(1) x(2) x(3)  
 $\mapsto$  // block 2 : ordering C  
execute(ms);  
s=0;



**WARNING:** In the case of a local or a mixed ordering, elimination needs special care.  $f$  may be considered as a map of germs  $f : (k^r, 0) \rightarrow (k^n, 0)$ , but even if this map germ is finite, we are in general not able to compute the image germ because for this we would need an implementation of the Weierstrass preparation theorem. What we can compute, and what `eliminate` actually does, is the following: let  $V(J)$  be the zero-set of  $J$  in  $k^r \times (k^n, 0)$ , then the closure of the image of  $V(J)$  under the projection

$$\text{pr} : k^r \times (k^n, 0) \rightarrow (k^n, 0)$$

can be computed. (Note that this germ contains also those components of  $V(J)$  which meet the fiber of  $\text{pr}$  outside the origin.) This is achieved by an ordering with the block of  $t$ -variables having a global ordering (and preceding the  $x$ -variables) and the  $x$ -variables having a local ordering.

In any case, if the input is weighted homogeneous (=quasihomogeneous), the weights given to the variables should be chosen accordingly. SINGULAR offers a function `weight` which proposes, given an ideal or module, integer weights for the variables, such that the ideal, resp. module, is as homogeneous as possible with respect to these weights. The function finds correct weights, if the input is weighted homogeneous (but is rather slow for many variables). In order to check, whether the input is quasihomogeneous, use the function `qhweight`, which returns an `intvec` of correct weights if the input is quasihomogeneous and an `intvec` of zeros otherwise.

Let us give three examples:

1. First we compute the equations of the simple space curve 'T[7]' consisting of two tangential cusps given in parametric form.
2. We compute weights for the equations such that the equations are quasihomogeneous w.r.t. these weights.
3. Then we compute the tangent developable of the rational normal curve in  $P^4$ .

```
// 1. Compute equations of curve given in parametric form:
// Two transversal cusps in (k^3,0):
ring r1 = 0,(t,x,y,z),ls;
ideal i1 = x-t^2,y-t^3,z; // parametrization of the first branch
ideal i2 = y-t^2,z-t^3,x; // parametrization of the second branch
ideal j1 = eliminate(i1,t);
j1; // equations of the first branch
=> j1[1]=z
=> j1[2]=y^2-x^3
ideal j2 = eliminate(i2,t);
j2; // equations of the second branch
=> j2[1]=z^2-y^3
=> j2[2]=x
// Now map to a ring with only x,y,z as variables and compute the
// intersection of j1 and j2 there:
ring r2 = 0,(x,y,z),ds;
ideal j1= imap(r1,j1); // imap is a convenient ringmap for
ideal j2= imap(r1,j2); // inclusions and projections of rings
ideal i = intersect(j1,j2);
i; // equations of both branches
=> i[1]=z^2-y^3+x^3y
=> i[2]=xz
=> i[3]=xy^2-x^4
//
// 2. Compute the weights:
intvec v= qhweight(i); // compute weights
v;
```



```

↳ 4,6,9
//
// 3. Compute the tangent developable
// The tangent developable of a projective variety given parametrically
// by $F=(f_1,\dots,f_n) : P^r \rightarrow P^n$ is the union of all tangent spaces
// of the image. The tangent space at a smooth point $F(t_1,\dots,t_r)$
// is given as the image of the tangent space at (t_1,\dots,t_r) under
// the tangent map (affine coordinates)
// $T(t_1,\dots,t_r): (y_1,\dots,y_r) \rightarrow \text{jacob}(f)*\text{transpose}((y_1,\dots,y_r))$
// where $\text{jacob}(f)$ denotes the jacobian matrix of f with respect to the
// t 's evaluated at the point (t_1,\dots,t_r) .
// Hence we have to create the graph of this map and then to eliminate
// the t 's and y 's.
// The rational normal curve in P^4 is given as the image of
// $F(s,t) = (s^4,s^3t,s^2t^2,st^3,t^4)$
// each component being homogeneous of degree 4.
ring P = 0,(s,t,x,y,a,b,c,d,e),dp;
ideal M = maxideal(1);
ideal F = M[1..2]; // take the 1st two generators of M
F=F^4;
// simplify(...,2); deletes 0-columns
matrix jac = simplify(jacob(F),2);
ideal T = x,y;
ideal J = jac*transpose(T);
ideal H = M[5..9];
ideal i = matrix(H)-matrix(J); // this is tricky: difference between two
// ideals is not defined, but between two
// matrices. By type conversion
// the ideals are converted to matrices,
// subtracted and afterwards converted
// to an ideal. Note that '+' is defined
// and adds (concatenates) two ideals

i;
↳ i[1]=-4s3x+a
↳ i[2]=-3s2tx-s3y+b
↳ i[3]=-2st2x-2s2ty+c
↳ i[4]=-t3x-3st2y+d
↳ i[5]=-4t3y+e
// Now we define a ring with product ordering and weights 4
// for the variables a,...,e.
// Then we map i from P to P1 and eliminate s,t,x,y from i.
ring P1 = 0,(s,t,x,y,a,b,c,d,e),(dp(4),wp(4,4,4,4,4));
ideal i = fetch(P,i);
ideal j= eliminate(i,stxy); // equations of tangent developable
j;
↳ j[1]=3c2-4bd+ae
↳ j[2]=2bcd-3ad2-3b2e+4ace
↳ j[3]=8b2d2-9acd2-9b2ce+14abde-4a2e2
// We can use the product ordering to eliminate s,t,x,y from i
// by a std-basis computation.
// We need proc 'nselect' from elim.lib.
LIB "elim.lib";
j = std(i); // compute a std basis j

```

```

 j = nselect(j,1..4); // select generators from j not
 j; // containing variable 1,...,4
 ↪ j[1]=3c2-4bd+ae
 ↪ j[2]=2bcd-3ad2-3b2e+4ace
 ↪ j[3]=8b2d2-9acd2-9b2ce+12ac2e-2abde

```

### A.3.4 Free resolution

In SINGULAR a free resolution of a module or ideal has its own type: **resolution**. It is a structure that stores all information related to free resolutions. This allows partial computations of resolutions via the command **res**. After applying **res**, only a pre-format of the resolution is computed which allows to determine invariants like Betti-numbers or homological dimension. To see the differentials of the complex, a resolution must be converted into the type list which yields a list of modules: the  $k$ -th module in this list is the first syzygy-module (module of relations) of the  $(k-1)$ st module. There are the following commands to compute a resolution:

|               |                                                                                                                                                                                                          |
|---------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <b>res</b>    | [res], page 785<br>computes a free resolution of an ideal or module using a heuristically chosen method. This is the preferred method to compute free resolutions of ideals or modules.                  |
| <b>fres</b>   | Section 5.1.48 [fres], page 186<br>improved version of Section 5.1.147 [sres], page 264, computes a free resolution of an ideal or module using Schreyer's method. The input has to be a standard basis. |
| <b>lres</b>   | Section 5.1.83 [lres], page 212<br>computes a free resolution of an ideal or module with LaScala's method. The input needs to be homogeneous.                                                            |
| <b>mres</b>   | Section 5.1.98 [mres], page 222<br>computes a minimal free resolution of an ideal or module with the syzygy method.                                                                                      |
| <b>sres</b>   | Section 5.1.147 [sres], page 264<br>computes a free resolution of an ideal or module with Schreyer's method. The input has to be a standard basis.                                                       |
| <b>nres</b>   | Section 5.1.105 [nres], page 228<br>computes a free resolution of an ideal or module with the standard basis method.                                                                                     |
| <b>minres</b> | Section 5.1.93 [minres], page 220<br>minimizes a free resolution of an ideal or module.                                                                                                                  |
| <b>syz</b>    | Section 5.1.154 [syz], page 275<br>computes the first syzygy module.                                                                                                                                     |

**res(i,r)**, **lres(i,r)**, **sres(i,r)**, **mres(i,r)**, **nres(i,r)** compute the first  $r$  modules of the resolution of  $i$ , resp. the full resolution if  $r=0$  and the basering is not a quing. See the manual for a precise description of these commands.

Note: The command **betti** does not require a minimal resolution for the minimal Betti numbers.

Now let us take a look at an example which uses resolutions: The Hilbert-Burch theorem says that the ideal  $i$  of a reduced curve in  $K^3$  has a free resolution of length 2 and that  $i$  is given by the  $2 \times 2$  minors of the 2nd matrix in the resolution. We test this for two transversal cusps in  $K^3$ . Afterwards we compute the resolution of the ideal  $j$  of the tangent developable of the rational normal curve in  $P^4$  from above. Finally we demonstrate the use of the type **resolution** in connection with the **lres** command.

```

// Two transversal cusps in (k^3,0):
ring r2 =0,(x,y,z),ds;
ideal i =z2-1y3+x3y,xz,-1xy2+x4,x3z;
resolution rs=mres(i,0); // computes a minimal resolution
rs; // the standard representation of complexes
↳ 1 3 2
↳ r2 <-- r2 <-- r2
↳
↳ 0 1 2
↳
 list resi=rs; // conversion to a list
 print(resi[1]); // the 1st module is i minimized
↳ xz,
↳ z2-y3+x3y,
↳ xy2-x4
 print(resi[2]); // the 1st syzygy module of i
↳ -z,-y2+x3,
↳ x, 0,
↳ y, z
 resi[3]; // the 2nd syzygy module of i
↳ _[1]=0
 ideal j=minor(resi[2],2);
 reduce(j,std(i)); // check whether j is contained in i
↳ _[1]=0
↳ _[2]=0
↳ _[3]=0
 size(reduce(i,std(j))); // check whether i is contained in j
↳ 0
// size(<ideal>) counts the non-zero generators
// -----
// The tangent developable of the rational normal curve in P^4:
ring P = 0,(a,b,c,d,e),dp;
ideal j= 3c2-4bd+ae, -2bcd+3ad2+3b2e-4ace,
 8b2d2-9acd2-9b2ce+9ac2e+2abde-1a2e2;
resolution rs=mres(j,0);
rs;
↳ 1 2 1
↳ P <-- P <-- P
↳
↳ 0 1 2
↳
 list L=rs;
 print(L[2]);
↳ 2bcd-3ad2-3b2e+4ace,
↳ -3c2+4bd-ae
// create an intmat with graded Betti numbers
intmat B=betti(rs);
// this gives a nice output of Betti numbers
print(B,"betti");
↳ 0 1 2
↳ -----
↳ 0: 1 - -
↳ 1: - 1 -

```

```

↳ 2: - 1 -
↳ 3: - - 1
↳ -----
↳ total: 1 2 1
↳
 // the user has access to all Betti numbers
 // the 2-nd column of B:
 B[1..4,2];
↳ 0 1 1 0
 ring cyc5=32003,(a,b,c,d,e,h),dp;
 ideal i=
 a+b+c+d+e,
 ab+bc+cd+de+ea,
 abc+bcd+cde+dea+eab,
 abcd+bcde+cdea+deab+eabc,
 h5-abcde;
 resolution rs=lres(i,0); //computes the resolution according LaScala
 rs; //the shape of the minimal resolution
↳ 1 5 10 10 5 1
↳ cyc5 <-- cyc5 <-- cyc5 <-- cyc5 <-- cyc5 <-- cyc5
↳
↳ 0 1 2 3 4 5
↳
 print(betti(rs),"betti"); //shows the Betti-numbers of cyclic 5
↳ 0 1 2 3 4 5
↳ -----
↳ 0: 1 1 - - - -
↳ 1: - 1 1 - - -
↳ 2: - 1 1 - - -
↳ 3: - 1 2 1 - -
↳ 4: - 1 2 1 - -
↳ 5: - - 2 2 - -
↳ 6: - - 1 2 1 -
↳ 7: - - 1 2 1 -
↳ 8: - - - 1 1 -
↳ 9: - - - 1 1 -
↳ 10: - - - - 1 1
↳ -----
↳ total: 1 5 10 10 5 1
↳
 dim(rs); //the homological dimension
↳ 4
 size(list(rs)); //gets the full (non-reduced) resolution
↳ 6
 minres(rs); //minimizes the resolution
↳ 1 5 10 10 5 1
↳ cyc5 <-- cyc5 <-- cyc5 <-- cyc5 <-- cyc5
↳
↳ 0 1 2 3 4 5
↳
 size(list(rs)); //gets the minimized resolution
↳ 6

```

### A.3.5 Handling graded modules

How to deal with graded modules in SINGULAR is best explained by looking at an example:

```

ring R = 0, (w,x,y,z), dp;
module I = [-x,0,-z2,0,y2z], [0,-x,-yz,0,y3], [-w,0,0,yz,-z3],
 [0,-w,0,y2,-yz2], [0,-1,-w,0,xz], [0,-w,0,y2,-yz2],
 [x2,-y2,-wy2+xz2];

print(I);
↳ -x, 0, -w, 0, 0, 0, x2,
↳ 0, -x, 0, -w, -1,-w, -y2,
↳ -z2,-yz,0, 0, -w,0, -wy2+xz2,
↳ 0, 0, yz, y2, 0, y2, 0,
↳ y2z,y3, -z3,-yz2,xz,-yz2,0

// (1) Check on degrees:
// =====
attrib(I,"isHomog"); // attribute not set => empty output
↳
homog(I);
↳ 1
attrib(I,"isHomog");
↳ 2,2,1,1,0

print(betti(I,0),"betti"); // read degrees from Betti diagram
↳ 0 1
↳ -----
↳ 0: 1 -
↳ 1: 2 1
↳ 2: 2 5
↳ 3: - 1
↳ -----
↳ total: 5 7
↳

// (2) Shift degrees:
// =====
def J=I;
intvec DV = 0,0,-1,-1,-2;
attrib(J,"isHomog",DV); // assign new weight vector
attrib(J,"isHomog");
↳ 0,0,-1,-1,-2
print(betti(J,0),"betti");
↳ 0 1
↳ -----
↳ -2: 1 -
↳ -1: 2 1
↳ 0: 2 5
↳ 1: - 1
↳ -----
↳ total: 5 7
↳

intmat bettiI=betti(I,0); // degree corresponding to first non-zero row

```

```

// of Betti diagram is accessible via
// attribute "rowShift"

attrib(bettiI);
↳ attr:rowShift, type int
intmat bettiJ=betti(J,0);
attrib(bettiJ);
↳ attr:rowShift, type int

// (3) Graded free resolutions:
// =====
resolution resJ = mres(J,0);
attrib(resJ);
↳ attr:isHomog, type intvec
print(betti(resJ),"betti");
↳ 0 1 2
↳ -----
↳ -2: 1 - -
↳ -1: 2 - -
↳ 0: 1 4 -
↳ 1: - - 1
↳ -----
↳ total: 4 4 1
↳
attrib(betti(resJ));
↳ attr:rowShift, type int

```

A check on degrees ((1), by using the `homog` command) shows that this is a graded matrix. The `homog` command assigns an admissible weight vector (here: 2,2,1,1,0) to the module `I` which is accessible via the attribute `"isHomog"`. Thus, we may think of `I` as a graded submodule of the graded free  $R$ -module

$$F = R(-2)^2 \oplus R(-1)^2 \oplus R.$$

We may also read the degrees from the Betti diagram as shown above. The degree on the left of the first nonzero row of the Betti diagram is accessible via the attribute `"rowShift"` of the betti matrix (which is of type `intmat`):

(2) We may shift degrees by assigning another admissible degree vector. Note that SINGULAR does not check whether the assigned degree vector really is admissible. Moreover, note that all assigned attributes are lost under a type conversion (e.g. from `module` to `matrix`).

(3) These considerations may be applied when computing data from free resolutions (see Section A.3.6 [Computation of Ext], page 716).

### A.3.6 Computation of Ext

We start by showing how to calculate the  $n$ -th Ext group of an ideal. The ingredients to do this are by the definition of Ext the following: calculate a (minimal) resolution at least up to length  $n$ , apply the Hom functor, and calculate the  $n$ -th homology group, that is, form the quotient  $\ker/\text{im}$  in the resolution sequence.

The Hom functor is given simply by transposing (hence dualizing) the module or the corresponding matrix with the command `transpose`. The image of the  $(n-1)$ -st map is generated by the columns of the corresponding matrix. To calculate the kernel apply the command `syz` at the  $(n-1)$ -st transposed entry of the resolution. Finally, the quotient is obtained by the command `modulo`, which gives for two modules  $A = \ker$ ,  $B = \text{Im}$  the module of relations of

$$A/(A \cap B)$$

in the usual way. As we have a chain complex, this is obviously the same as  $\ker/\text{Im}$ .

We collect these statements in the following short procedure:

```
proc ext(int n, ideal I)
{
 resolution rs = mres(I,n+1);
 module tAn = transpose(rs[n+1]);
 module tAn_1 = transpose(rs[n]);
 module ext_n = modulo(syz(tAn),tAn_1);
 return(ext_n);
}
```

Now consider the following example:

```
ring r5 = 32003,(a,b,c,d,e),dp;
ideal I = a2b2+ab2c+b2cd, a2c2+ac2d+c2de,a2d2+ad2e+bd2e,a2e2+abe2+bce2;
print(ext(2,I));
↳ 1,0,0,0,0,0,0,
↳ 0,1,0,0,0,0,0,
↳ 0,0,1,0,0,0,0,
↳ 0,0,0,1,0,0,0,
↳ 0,0,0,0,1,0,0,
↳ 0,0,0,0,0,1,0,
↳ 0,0,0,0,0,0,1,0,
↳ 0,0,0,0,0,0,0,1
ext(3,I); // too big to be displayed here
```

The library `homolog.lib` contains several procedures for computing Ext-modules and related modules, which are much more general and sophisticated than the above one. They are used in the following example:

If  $M$  is a module, then  $\text{Ext}^1(M, M)$ , resp.  $\text{Ext}^2(M, M)$ , are the modules of infinitesimal deformations, respectively of obstructions, of  $M$  (like T1 and T2 for a singularity). Similar to the treatment of singularities, the semiuniversal deformation of  $M$  can be computed (if  $\text{Ext}^1$  is finite dimensional) with the help of  $\text{Ext}^1$ ,  $\text{Ext}^2$  and the cup product. There is an extra procedure for  $\text{Ext}^k(R/J, R)$  if  $J$  is an ideal in  $R$ , since this is faster than the general Ext.

We compute

- the infinitesimal deformations ( $= \text{Ext}^1(K, K)$ ) and obstructions ( $= \text{Ext}^2(K, K)$ ) of the residue field  $K = R/m$  of an ordinary cusp,  $R = K[x, y]_m/(x^2 - y^3)$ ,  $m = (x, y)$ . To compute  $\text{Ext}^1(m, m)$  we have to apply `Ext(1, syz(m), syz(m))` with `syz(m)` the first syzygy module of  $m$ , which is isomorphic to  $\text{Ext}^2(K, K)$ .
- $\text{Ext}^k(R/i, R)$  for some ideal  $i$  and with an extra option.

```
LIB "homolog.lib";
ring R=0,(x,y),ds;
ideal i=x2-y3;
qring q = std(i); // defines the quotient ring k[x,y]_m/(x2-y3)
ideal m = maxideal(1);
module T1K = Ext(1,m,m); // computes Ext^1(R/m,R/m)
↳ // dimension of Ext^1: 0
↳ // vdim of Ext^1: 2
↳
print(T1K);
↳ 0,x,0,y,
↳ x,0,y,0
printlevel=2; // gives more explanation
module T2K=Ext(2,m,m); // computes Ext^2(R/m,R/m)
```

```

⇒ // Computing Ext^2 (help Ext; gives an explanation):
⇒ // Let 0←--coker(M)←--F0←--F1←--F2←--... be a resolution of coker(M),
⇒ // and 0←--coker(N)←--G0←--G1 a presentation of coker(N),
⇒ // then Hom(F2,G0)→Hom(F3,G0) is given by:
⇒ y2,x,
⇒ x, y
⇒ // and Hom(F1,G0) + Hom(F2,G1)→Hom(F2,G0) is given by:
⇒ -y,x, x,0,y,0,
⇒ x, -y2,0,x,0,y
⇒
⇒ // dimension of Ext^2: 0
⇒ // vdim of Ext^2: 2
⇒
 print(std(T2K));
⇒ 0,x,0,y,
⇒ x,0,y,0
 printlevel=0;
 module E = Ext(1,syz(m),syz(m));
⇒ // dimension of Ext^1: 0
⇒ // vdim of Ext^1: 2
⇒
 print(std(E));
⇒ x, 0,-y2,x,0,y,
⇒ -y,0,x, 0,y,0,
⇒ 0, 1,0, 0,0,0,
⇒ 1, 0,0, 0,0,0
 //The matrices which we have just computed are presentation matrices
 //of the modules T2K and E. Hence we may ignore those columns
 //containing 1 as an entry and see that T2K and E are isomorphic
 //as expected, but differently presented.
 //-----
 ring S=0,(x,y,z),dp;
 ideal i = x2y,y2z,z3x;
 module E = Ext_R(2,i);
⇒ // dimension of Ext^2: 1
⇒
 print(E);
⇒ 0,y,0,z2,
⇒ z,0,0,-x,
⇒ 0,0,x,-y
 // if a 3-rd argument of type int is given,
 // a list of Ext^k(R/i,R), a SB of Ext^k(R/i,R) and a vector space basis
 // is returned:
 list LE = Ext_R(3,i,0);
⇒ // dimension of Ext^3: 0
⇒ // vdim of Ext^3: 2
⇒
 LE;
⇒ [1]:
⇒ _[1]=y*gen(1)
⇒ _[2]=x*gen(1)
⇒ _[3]=z2*gen(1)
⇒ [2]:

```



```

↳ _[1]=y*gen(1)
↳ _[2]=x*gen(1)
↳ _[3]=z2*gen(1)
↳ [3]:
↳ _[1,1]=z
↳ _[1,2]=1
 print(LE[2]);
↳ y,x,z2
 print(kbase(LE[2]));
↳ z,1

```

### A.3.7 Depth

We compute the depth of the module of Kaehler differentials  $D_k(R)$  of the variety defined by the  $(m+1)$ -minors of a generic symmetric  $(n \times n)$ -matrix. We do this by computing the resolution over the polynomial ring. Then, by the Auslander-Buchsbaum formula, the depth is equal to the number of variables minus the length of a minimal resolution. This example was suggested by U. Vetter in order to check whether his bound  $\text{depth}(D_k(R)) \geq m(m+1)/2 + m - 1$  could be improved.

```

LIB "matrix.lib"; LIB "sing.lib";
int n = 4;
int m = 3;
int N = n*(n+1) div 2; // will become number of variables
ring R = 32003,x(1..N),dp;
matrix X = symmat(n); // proc from matrix.lib
 // creates the symmetric generic nxn matrix

print(X);
↳ x(1),x(2),x(3),x(4),
↳ x(2),x(5),x(6),x(7),
↳ x(3),x(6),x(8),x(9),
↳ x(4),x(7),x(9),x(10)
ideal J = minor(X,m);
J=std(J);
// Kaehler differentials D_k(R)
// of R=k[x1..xn]/J:
module D = J*freemodule(N)+transpose(jacob(J));
ncols(D);
↳ 110
nrows(D);
↳ 10
//
// Note: D is a submodule with 110 generators of a free module
// of rank 10 over a polynomial ring in 10 variables.
// Compute a full resolution of D with sres.
// This takes about 17 sec on a Mac PB 520c and 2 sec an a HP 735
int time = timer;
module sD = std(D);
list Dres = sres(sD,0); // the full resolution
timer-time; // time used for std + sres
↳ 0
intmat B = betti(Dres);
print(B,"betti");
↳ 0 1 2 3 4 5 6
↳ -----

```

```

↳ 0: 10 - - - - - -
↳ 1: - 10 - - - - -
↳ 2: - 84 144 60 - - -
↳ 3: - - 35 80 60 16 1
↳ -----
↳ total: 10 94 179 140 60 16 1
↳
↳ N-ncols(B)+1; // the desired depth
↳ 4

```

### A.3.8 Factorization

The factorization of polynomials is implemented in the C++ libraries `Factory` (written mainly by Ruediger Stobbe) and `libfac` (written by Michael Messollen) which are part of the SINGULAR system. For the factorization of univariate polynomials these libraries make use of the library NTL written by Victor Shoup.

```

ring r = 0, (x,y), dp;
poly f = 9x16-18x13y2-9x12y3+9x10y4-18x11y2+36x8y4
 +18x7y5-18x5y6+9x6y4-18x3y6-9x2y7+9y8;
// = 9 * (x5-1y2)^2 * (x6-2x3y2-1x2y3+y4)
factorize(f);
↳ [1]:
↳ _[1]=9
↳ _[2]=x6-2x3y2-x2y3+y4
↳ _[3]=-x5+y2
↳ [2]:
↳ 1,1,2
// returns factors and multiplicities,
// first factor is a constant.
poly g = (y4+x8)*(x2+y2);
factorize(g);
↳ [1]:
↳ _[1]=1
↳ _[2]=x2+y2
↳ _[3]=x8+y4
↳ [2]:
↳ 1,1,1
// The same in characteristic 2:
ring s = 2, (x,y), dp;
poly g = (y4+x8)*(x2+y2);
factorize(g);
↳ [1]:
↳ _[1]=1
↳ _[2]=x+y
↳ _[3]=x2+y
↳ [2]:
↳ 1,2,4
// factorization over algebraic extension fields
ring rext = (0,i), (x,y), dp;
minpoly = i2+1;
poly g = (y4+x8)*(x2+y2);
factorize(g);
↳ [1]:

```

```

↳ _[1]=1
↳ _[2]=x+(i)*y
↳ _[3]=x+(-i)*y
↳ _[4]=x4+(i)*y2
↳ _[5]=x4+(-i)*y2
↳ [2]:
↳ 1,1,1,1,1

```

### A.3.9 Primary decomposition

There are two algorithms implemented in SINGULAR which provide primary decomposition: `primdecGTZ`, based on Gianni/Trager/Zacharias (written by Gerhard Pfister) and `primdecSY`, based on Shimoyama/Yokoyama (written by Wolfram Decker and Hans Schoenemann).

The result of `primdecGTZ` and `primdecSY` is returned as a list of pairs of ideals, where the second ideal is the prime ideal and the first ideal the corresponding primary ideal.

```

LIB "primdec.lib";
ring r = 0,(a,b,c,d,e,f),dp;
ideal i= f3, ef2, e2f, bcf-adf, de+cf, be+af, e3;
primdecGTZ(i);
↳ [1]:
↳ [1]:
↳ _[1]=f
↳ _[2]=e
↳ [2]:
↳ _[1]=f
↳ _[2]=e
↳ [2]:
↳ [1]:
↳ _[1]=f3
↳ _[2]=ef2
↳ _[3]=e2f
↳ _[4]=e3
↳ _[5]=de+cf
↳ _[6]=be+af
↳ _[7]=-bc+ad
↳ [2]:
↳ _[1]=f
↳ _[2]=e
↳ _[3]=-bc+ad
// We consider now the ideal J of the base space of the
// miniversal deformation of the cone over the rational
// normal curve computed in section *8* and compute
// its primary decomposition.
ring R = 0,(A,B,C,D),dp;
ideal J = CD, BD+D2, AD;
primdecGTZ(J);
↳ [1]:
↳ [1]:
↳ _[1]=D
↳ [2]:
↳ _[1]=D
↳ [2]:
↳ [1]:

```

```

↳ _[1]=C
↳ _[2]=B+D
↳ _[3]=A
↳ [2]:
↳ _[1]=C
↳ _[2]=B+D
↳ _[3]=A
// We see that there are two components which are both
// prime, even linear subspaces, one 3-dimensional,
// the other 1-dimensional.
// (This is Pinkhams example and was the first known
// surface singularity with two components of
// different dimensions)
//
// Let us now produce an embedded component in the last
// example, compute the minimal associated primes and
// the radical. We use the Characteristic set methods
// from primdec.lib.
J = intersect(J,maxideal(3));
// The following shows that the maximal ideal defines an embedded
// (prime) component.
primdecSY(J);
↳ [1]:
↳ [1]:
↳ _[1]=D
↳ [2]:
↳ _[1]=D
↳ [2]:
↳ [1]:
↳ _[1]=C
↳ _[2]=B+D
↳ _[3]=A
↳ [2]:
↳ _[1]=C
↳ _[2]=B+D
↳ _[3]=A
↳ [3]:
↳ [1]:
↳ _[1]=D2
↳ _[2]=C2
↳ _[3]=B2
↳ _[4]=AB
↳ _[5]=A2
↳ _[6]=BCD
↳ _[7]=ACD
↳ [2]:
↳ _[1]=D
↳ _[2]=C
↳ _[3]=B
↳ _[4]=A
minAssChar(J);
↳ [1]:
↳ _[1]=C

```

```

↳ _[2]=B+D
↳ _[3]=A
↳ [2]:
↳ _[1]=D
 radical(J);
↳ _[1]=CD
↳ _[2]=BD+D2
↳ _[3]=AD

```

### A.3.10 Normalization

The normalization will be computed for a reduced ring  $R/I$ . The result is a list of rings; ideals are always called `norid` in the rings of this list. The normalization of  $R/I$  is the product of the factor rings of the rings in the list divided out by the ideals `norid`.

```

LIB "normal.lib";
// ----- first example: rational quadruple point -----
ring R=32003,(x,y,z),wp(3,5,15);
ideal I=z*(y3-x5)+x10;
list pr=normal(I);
↳
↳ // 'normal' created a list, say nor, of two elements.
↳ // To see the list type
↳ nor;
↳
↳ // * nor[1] is a list of 1 ring(s).
↳ // To access the i-th ring nor[1][i], give it a name, say Ri, and type
↳ def R1 = nor[1][1]; setring R1; norid; normap;
↳ // For the other rings type first (if R is the name of your base ring)
↳ setring R;
↳ // and then continue as for R1.
↳ // Ri/norid is the affine algebra of the normalization of R/P_i where
↳ // P_i is the i-th component of a decomposition of the input ideal id
↳ // and normap the normalization map from R to Ri/norid.
↳
↳ // * nor[2] is a list of 1 ideal(s). Let ci be the last generator
↳ // of the ideal nor[2][i]. Then the integral closure of R/P_i is
↳ // generated as R-submodule of the total ring of fractions by
↳ // 1/ci * nor[2][i].
↳ def S=pr[1][1];
↳ setring S;
↳ norid;
↳ norid[1]=T(2)*x+y*z
↳ norid[2]=T(1)*x^2-T(2)*y
↳ norid[3]=-T(1)*y+x^7-x^2*z
↳ norid[4]=T(1)*y^2*z+T(2)*x^8-T(2)*x^3*z
↳ norid[5]=T(1)^2+T(2)*z+x^4*y*z
↳ norid[6]=T(1)*T(2)+x^6*z-x*z^2
↳ norid[7]=T(2)^2+T(1)*x*z
↳ norid[8]=x^10-x^5*z+y^3*z
↳ // ----- second example: union of straight lines -----
↳ ring R1=0,(x,y,z),dp;
↳ ideal I=(x-y)*(x-z)*(y-z);
↳ list qr=normal(I);

```

```

↳
↳ // 'normal' created a list, say nor, of two elements.
↳ // To see the list type
↳ nor;
↳
↳ // * nor[1] is a list of 2 ring(s).
↳ // To access the i-th ring nor[1][i], give it a name, say Ri, and type
↳ def R1 = nor[1][1]; setring R1; norid; normap;
↳ // For the other rings type first (if R is the name of your base ring)
↳ setring R;
↳ // and then continue as for R1.
↳ // Ri/norid is the affine algebra of the normalization of R/P_i where
↳ // P_i is the i-th component of a decomposition of the input ideal id
↳ // and normap the normalization map from R to Ri/norid.
↳
↳ // * nor[2] is a list of 2 ideal(s). Let ci be the last generator
↳ // of the ideal nor[2][i]. Then the integral closure of R/P_i is
↳ // generated as R-submodule of the total ring of fractions by
↳ // 1/ci * nor[2][i].
↳ def S1=qr[1][1]; def S2=qr[1][2];
↳ setring S1; norid;
↳ norid[1]=-T(1)*y+T(1)*z+x-z
↳ norid[2]=T(1)*x-T(1)*y
↳ norid[3]=T(1)^2-T(1)
↳ norid[4]=x^2-x*y-x*z+y*z
↳ setring S2; norid;
↳ norid[1]=y-z

```

### A.3.11 Kernel of module homomorphisms

Let  $A, B$  be two matrices of size  $m \times r$  and  $m \times s$  over the ring  $R$  and consider the corresponding maps

$$R^r \xrightarrow{A} R^m \xleftarrow{B} R^s.$$

We want to compute the kernel of the map  $R^r \xrightarrow{A} R^m \longrightarrow R^m/\text{Im}(B)$ . This can be done using the `modulo` command:

$$\text{modulo}(A, B) = \ker(R^r \xrightarrow{A} R^m/\text{Im}(B)).$$

More precisely, the output of `modulo(A,B)` is a module such that the given generating vectors span the kernel on the right-hand side.

```

 ring r=0,(x,y,z),(c,dp);
 matrix A[2][2]=x,y,z,1;
 matrix B[2][2]=x2,y2,z2,xz;
 print(B);
↳ x2,y2,
↳ z2,xz
 def C=modulo(A,B);
 print(C); // matrix of generators for the kernel
↳ yz2-x2, xyz-y2, x2z-xy, x3-y2z,
↳ x2z-xz2, -x2z+y2z, xyz-yz2, 0
 print(A*matrix(C)); // should be in Im(B)
↳ x2yz-x3, y3z-xy2, x3z+xy2z-y2z2-x2y, x4-xy2z,
↳ yz3-xz2, xyz2-x2z, x2z2-yz2, x3z-y2z2

```

### A.3.12 Algebraic dependence

Let  $g, f_1, \dots, f_r \in K[x_1, \dots, x_n]$ . We want to check whether

1.  $f_1, \dots, f_r$  are algebraically dependent.

Let  $I = \langle Y_1 - f_1, \dots, Y_r - f_r \rangle \subseteq K[x_1, \dots, x_n, Y_1, \dots, Y_r]$ . Then  $I \cap K[Y_1, \dots, Y_r]$  are the algebraic relations between  $f_1, \dots, f_r$ .

2.  $g \in K[f_1, \dots, f_r]$ .

$g \in K[f_1, \dots, f_r]$  if and only if the normal form of  $g$  with respect to  $I$  and a block ordering with respect to  $X = (x_1, \dots, x_n)$  and  $Y = (Y_1, \dots, Y_r)$  with  $X > Y$  is in  $K[Y]$ .

Both questions can be answered using the following procedure. If the second argument is zero, it checks for algebraic dependence and returns the ideal of relations between the generators of the given ideal. Otherwise it checks for subring membership and returns the normal form of the second argument with respect to the ideal  $I$ .

```

proc algebraicDep(ideal J, poly g)
{
 def R=basing; // give a name to the basering
 int n=size(J);
 int k=nvars(R);
 int i;
 intvec v;

 // construction of the new ring:

 // construct a weight vector
 v[n+k]=0; // gives a zero vector of length n+k
 for(i=1;i<=k;i++)
 {
 v[i]=1;
 }
 string orde="(a("+string(v)+"),dp)";
 string ri="ring Rhelp="+charstr(R)+"",
 ("+"varstr(R)+"",Y(1.."+"string(n)+"")),"+orde;
 // ring definition as a string
 execute(ri); // execution of the string

 // construction of the new ideal I=(J[1]-Y(1),...,J[n]-Y(n))
 ideal I=imap(R,J);
 for(i=1;i<=n;i++)
 {
 I[i]=I[i]-var(k+i);
 }
 poly g=imap(R,g);
 if(g==0)
 {
 // construction of the ideal of relations by elimination
 poly el=var(1);
 for(i=2;i<=k;i++)
 {
 el=el*var(i);
 }
 ideal KK=eliminate(I,el);
 }
}

```

```

 keepring(Rhelp);
 return(KK);
 }
 // reduction of g with respect to I
 ideal KK=reduce(g,std(I));
 keepring(Rhelp);
 return(KK);
}

// applications of the procedure
ring r=0,(x,y,z),dp;
ideal i=xz,yz;
algebraicDep(i,0);
⇨ _[1]=0
// Note: after call of algebraicDep(), the basering is Rhelp.
setring r; kill Rhelp;
ideal j=xy+z2,z2+y2,x2y2-2xy3+y4;
algebraicDep(j,0);
⇨ _[1]=Y(1)^2-2*Y(1)*Y(2)+Y(2)^2-Y(3)
setring r; kill Rhelp;
poly g=y2z2-xz;
algebraicDep(i,g);
⇨ _[1]=Y(2)^2-Y(1)
// this shows that g is contained in i.
setring r; kill Rhelp;
algebraicDep(j,g);
⇨ _[1]=-z^4+z^2*Y(2)-x*z
// this shows that g is contained in j.

```

## A.4 Singularity Theory

### A.4.1 Milnor and Tjurina number

The Milnor number, resp. the Tjurina number, of a power series  $f$  in  $K[[x_1, \dots, x_n]]$  is

$$\text{milnor}(f) = \dim_K(K[[x_1, \dots, x_n]]/\text{jacob}(f)),$$

respectively

$$\text{tjurina}(f) = \dim_K(K[[x_1, \dots, x_n]]/((f) + \text{jacob}(f)))$$

where  $\text{jacob}(f)$  is the ideal generated by the partials of  $f$ .  $\text{tjurina}(f)$  is finite, if and only if  $f$  has an isolated singularity. The same holds for  $\text{milnor}(f)$  if  $K$  has characteristic 0. SINGULAR displays -1 if the dimension is infinite.

SINGULAR cannot compute with infinite power series. But it can work in  $\text{Loc}_{(x)}K[x_1, \dots, x_n]$ , the localization of  $K[x_1, \dots, x_n]$  at the maximal ideal  $(x_1, \dots, x_n)$ . To do this, one has to define a ring with a local monomial ordering such as ds, Ds, ls, ws, Ws (the second letter 's' referring to power 's'eries), or an appropriate matrix ordering. See Section B.2 [Monomial orderings], page 760 for a menu of possible orderings.

For theoretical reasons, the vector space dimension computed over the localization ring coincides with the Milnor (resp. Tjurina) number as defined above (in the power series ring).

We show in the example below the following:



- set option `prot` to have a short protocol during standard basis computation
- define the ring `r1` of characteristic 32003 with variables `x,y,z`, monomial ordering `ds`, series ring (i.e.,  $K[x,y,z]$  localized at  $(x,y,z)$ )
- list the information about `r1` by typing its name
- define the integers `a,b,c,t`
- define a polynomial `f` (depending on `a,b,c,t`) and display it
- define the jacobian ideal `i` of `f`
- compute a standard basis of `i`
- compute the Milnor number (=250) with `vdim` and create and display a string in order to comment the result (text between quotes " "; is a 'string')
- compute a standard basis of `i+(f)`
- compute the Tjurina number (=195) with `vdim`
- then compute the Milnor number (=248) and the Tjurina number (=195) for `t=1`
- reset the option to `noprot`

See also Section D.6.20 [sing\_lib], page 868 for the library commands for the computation of the Milnor and Tjurina number.

```

option(prot);
ring r1 = 32003, (x,y,z), ds;
r1;
↳ // coefficients: ZZ/32003
↳ // number of vars : 3
↳ // block 1 : ordering ds
↳ // : names x y z
↳ // block 2 : ordering C
int a,b,c,t=11,5,3,0;
poly f = x^a+y^b+z^(3*c)+x^(c+2)*y^(c-1)+x^(c-1)*y^(c-1)*z3+
 x^(c-2)*y^c*(y^2+t*x)^2;

f;
↳ y5+x5y2+x2y2z3+xy7+z9+x11
ideal i=jacob(f);
i;
↳ i[1]=5x4y2+2xy2z3+y7+11x10
↳ i[2]=5y4+2x5y+2x2yz3+7xy6
↳ i[3]=3x2y2z2+9z8
ideal j=std(i);
↳ 7(2)s8s10s11s12s(3)s13(4)s(5)s14(6)s(7)15--.s(6)-16-.s(5)17.s(7)s--s18(6\
).--19-.sH(24)20(3)...21....22....23.--24-
↳ product criterion:10 chain criterion:69
"The Milnor number of f(11,5,3) for t=0 is", vdim(j);
↳ The Milnor number of f(11,5,3) for t=0 is 250
j=i+f; // override j
j=std(j);
↳ 7(3)s8(2)s10s11(3)ss12(4)s(5)s13(6)s(8)s14(9).s(10).15--sH(23)(8)...16...\
...17.....sH(21)(9)sH(20)16(10).17.....18.....19..----.sH(19)
↳ product criterion:10 chain criterion:53
vdim(j); // compute the Tjurina number for t=0
↳ 195
t=1;
f=x^a+y^b+z^(3*c)+x^(c+2)*y^(c-1)+x^(c-1)*y^(c-1)*z3

```

```

+x^(c-2)*y^c*(y^2+t*x)^2;
ideal i1=jacob(f);
ideal j1=std(i1);
↳ 7(2)s8s10s11s12s13(3)ss(4)s14(5)s(6)s15(7).....s(8)16.s...s(9)..17.....\
.....s18(10).....s(11)..-.19.....sH(24)(10).....20.....21.....\
..22.....23.....24.---\
-----.25.26
↳ product criterion:11 chain criterion:83
 "The Milnor number of f(11,5,3) for t=1:",vdim(j1);
↳ The Milnor number of f(11,5,3) for t=1: 248
 vdim(std(j1+f)); // compute the Tjurina number for t=1
↳ 7(16)s8(15)s10s11ss(16)-12.s-s13s(17)s(18)s(19)-s(18).-14-s(17)-s(16)ss(1\
7)s15(18)..-s...--.16....-.....s(16).sH(23)s(18)...17.....18.....\
...sH(20)17(17).....18.....19..---.....-.-.-----20.\
-----...s17(9).....18.....19.-.....20.-.....21.....s\
H(19)16(5).....18.....19.-----
↳ product criterion:15 chain criterion:174
↳ 195
option(noprot);

```

## A.4.2 Critical points

The same computation which computes the Milnor, resp. the Tjurina, number, but with ordering `dp` instead of `ds` (i.e., in  $K[x_1, \dots, x_n]$  instead of  $\text{Loc}_{(x)}K[x_1, \dots, x_n]$ ) gives:

- the number of critical points of  $\mathbf{f}$  in the affine space (counted with multiplicities)
- the number of singular points of  $\mathbf{f}$  on the affine hypersurface  $\mathbf{f}=0$  (counted with multiplicities).

We start with the ring `r1` from section Section A.4.1 [Milnor and Tjurina number], page 726 and its elements.

The following will be implemented below:

- reset the protocol option and activate the timer
- define the ring `r2` of characteristic 32003 with variables `x,y,z` and monomial ordering `dp` (= `degrevlex`) (i.e., the polynomial ring =  $K[x,y,z]$ ).
- Note that polynomials, ideals, matrices (of polys), vectors, modules belong to a ring, hence we have to define `f` and `jacob(f)` again in `r2`. Since these objects are local to a ring, we may use the same names. Instead of defining `f` again we map it from ring `r1` to `r2` by using the `imap` command (`imap` is a convenient way to map variables from some ring identically to variables with the same name in the basering, even if the ground field is different. Compare with `fetch` which works for almost identical rings, e.g., if the rings differ only by the ordering or by the names of the variables and which may be used to rename variables). Integers and strings, however, do not belong to any ring. Once defined they are globally known.
- The result of the computation here (together with the previous one in Section A.4.1 [Milnor and Tjurina number], page 726) shows that (for  $t=0$ )  $\dim_K(\text{Loc}_{(x,y,z)}K[x,y,z]/\text{jacob}(f)) = 250$  (previously computed) while  $\dim_K(K[x,y,z]/\text{jacob}(f)) = 536$ . Hence  $\mathbf{f}$  has 286 critical points, counted with multiplicity, outside the origin. Moreover, since  $\dim_K(\text{Loc}_{(x,y,z)}K[x,y,z]/(\text{jacob}(f) + (f))) = 195 = \dim_K(K[x,y,z]/(\text{jacob}(f) + (f)))$ , the affine surface  $\mathbf{f}=0$  is smooth outside the origin.

```

ring r1 = 32003, (x,y,z), ds;
int a,b,c,t=11,5,3,0;
poly f = x^a+y^b+z^(3*c)+x^(c+2)*y^(c-1)+x^(c-1)*y^(c-1)*z3+
 x^(c-2)*y^c*(y^2+t*x)^2;

```

```

option(noprot);
timer=1;
ring r2 = 32003,(x,y,z),dp;
poly f=imap(r1,f);
ideal j=jacob(f);
vdim(std(j));
↳ 536
vdim(std(j+f));
↳ 195
timer=0; // reset timer

```

### A.4.3 Polar curves

The polar curve of a hypersurface given by a polynomial  $f \in k[x_1, \dots, x_n, t]$  with respect to  $t$  (we may consider  $f = 0$  as a family of hypersurfaces parametrized by  $t$ ) is defined as the Zariski closure of  $V(\partial f/\partial x_1, \dots, \partial f/\partial x_n) \setminus V(f)$  if this happens to be a curve. Some authors consider  $V(\partial f/\partial x_1, \dots, \partial f/\partial x_n)$  itself as polar curve.

We may consider projective hypersurfaces (in  $P^n$ ), affine hypersurfaces (in  $k^n$ ) or germs of hypersurfaces (in  $(k^n, 0)$ ), getting in this way projective, affine or local polar curves.

Now let us compute this for a family of curves. We need the library `elim.lib` for saturation and `sing.lib` for the singular locus.

```

LIB "elim.lib";
LIB "sing.lib";
// Affine polar curve:
ring R = 0,(x,z,t),dp; // global ordering dp
poly f = z5+xz3+x2-tz6;
dim_slocus(f); // dimension of singular locus
↳ 1
ideal j = diff(f,x),diff(f,z);
dim(std(j)); // dim V(j)
↳ 1
dim(std(j+ideal(f))); // V(j,f) also 1-dimensional
↳ 1
// j defines a curve, but to get the polar curve we must remove the
// branches contained in f=0 (they exist since dim V(j,f) = 1). This
// gives the polar curve set theoretically. But for the structure we
// may take either j:f or j:f^k for k sufficiently large. The first is
// just the ideal quotient, the second the iterated ideal quotient
// or saturation. In our case both coincide.
ideal q = quotient(j,ideal(f)); // ideal quotient
ideal qsat = sat(j,f)[1]; // saturation, proc from elim.lib
ideal sq = std(q);
dim(sq);
↳ 1
// 1-dimensional, hence q defines the affine polar curve
//
// to check that q and qsat are the same, we show both inclusions, i.e.,
// both reductions must give the 0-ideal
size(reduce(qsat,sq));
↳ 0
size(reduce(q,std(qsat)));
↳ 0

```

```

qsat;
⇒ qsat[1]=12zt+3z-10
⇒ qsat[2]=5z2+12xt+3x
⇒ qsat[3]=144xt2+72xt+9x+50z
// We see that the affine polar curve does not pass through the origin,
// hence we expect the local polar "curve" to be empty
// -----
// Local polar curve:
ring r = 0,(x,z,t),ds; // local ordering ds
poly f = z5+xz3+x2-tz6;
ideal j = diff(f,x),diff(f,z);
dim(std(j)); // V(j) 1-dimensional
⇒ 1
dim(std(j+ideal(f))); // V(j,f) also 1-dimensional
⇒ 1
ideal q = quotient(j,ideal(f)); // ideal quotient
q;
⇒ q[1]=1
// The local polar "curve" is empty, i.e., V(j) is contained in V(f)
// -----
// Projective polar curve: (we need "sing.lib" and "elim.lib")
ring P = 0,(x,z,t,y),dp; // global ordering dp
poly f = z5+xz3y2+x2y4-tz6;
// but consider t as parameter
dim_slocus(f); // projective 1-dimensional singular locus
⇒ 2
ideal j = diff(f,x),diff(f,z);
dim(std(j)); // V(j), projective 1-dimensional
⇒ 2
dim(std(j+ideal(f))); // V(j,f) also projective 1-dimensional
⇒ 2
ideal q = quotient(j,ideal(f));
ideal qsat = sat(j,f)[1]; // saturation, proc from elim.lib
dim(std(qsat));
⇒ 2
// projective 1-dimensional, hence q and/or qsat define the projective
// polar curve. In this case, q and qsat are not the same, we needed
// 2 quotients.
// Let us check both reductions:
size(reduce(qsat,std(q)));
⇒ 4
size(reduce(q,std(qsat)));
⇒ 0
// Hence q is contained in qsat but not conversely
q;
⇒ q[1]=12zty+3zy-10y2
⇒ q[2]=60z2t-36xty-9xy-50zy
⇒ q[3]=12xty2+5z2y+3xy2
⇒ q[4]=z3y+2xy3
qsat;
⇒ qsat[1]=12zt+3z-10y
⇒ qsat[2]=12xty+5z2+3xy
⇒ qsat[3]=144xt2+72xt+9x+50z

```

```

↳ qsat[4]=z3+2xy2
 //
 // Now consider again the affine polar curve,
 // homogenize it with respect to y (deg t=0) and compare:
 // affine polar curve:
 ideal qa = 12zt+3z-10,5z2+12xt+3x,-144xt2-72xt-9x-50z;
 // homogenized:
 ideal qh = 12zt+3z-10y,5z2+12xyt+3xy,-144xt2-72xt-9x-50z;
 size(reduce(qh,std(qsat)));
↳ 0
 size(reduce(qsat,std(qh)));
↳ 0
 // both ideals coincide

```

#### A.4.4 T1 and T2

$T^1$ , resp.  $T^2$ , of an ideal  $j$  usually denote the modules of infinitesimal deformations, resp. of obstructions. In SINGULAR there are procedures `T_1` and `T_2` in `sing.lib` such that `T_1(j)` and `T_2(j)` compute a standard basis of a presentation of these modules. If  $T^1, T^2$  are finite dimensional  $K$ -vector spaces (e.g., for isolated singularities), a basis can be computed by applying `kbase(T_1(j))`; resp. `kbase(T_2(j))`; the dimensions by applying `vdim`. For a complete intersection  $j$  the procedure `Tjurina` also computes  $T^1$ , but faster ( $T^2 = 0$  in this case). For a non complete intersection, it is faster to use the procedure `T_12` instead of `T_1` and `T_2`. Type `help T_1`; (or `help T_2`; or `help T_12`;) to obtain more detailed information about these procedures.

We give three examples, the first being a hypersurface, the second a complete intersection, the third not a complete intersection:

- load `sing.lib`
- check whether the ideal  $j$  is a complete intersection. It is, if number of variables = dimension + minimal number of generators
- compute the Tjurina number
- compute a vector space basis (`kbase`) of  $T^1$
- compute the Hilbert function of  $T^1$
- create a polynomial encoding the Hilbert series
- compute the dimension of  $T^2$

```

LIB "sing.lib";
ring R=32003,(x,y,z),ds;
// -----
// hypersurface case (from series T[p,q,r]):
int p,q,r = 3,3,4;
poly f = x^p+y^q+z^r+xyz;
tjurina(f);
↳ 8
 // Tjurina number = 8
 kbase(Tjurina(f));
↳ // Tjurina number = 8
↳ _[1]=z3
↳ _[2]=z2
↳ _[3]=yz
↳ _[4]=xz
↳ _[5]=z

```

```

↳ _[6]=y
↳ _[7]=x
↳ _[8]=1
 // -----
 // complete intersection case (from series P[k,1]):
 int k,l =3,2;
 ideal j=xy,x^k+y^l+z2;
 dim(std(j)); // Krull dimension
↳ 1
 size(minbase(j)); // minimal number of generators
↳ 2
 tjurina(j); // Tjurina number
↳ 6
 module T=Tjurina(j);
↳ // Tjurina number = 6
 kbase(T); // a sparse output of the k-basis of T_1
↳ _[1]=z*gen(1)
↳ _[2]=gen(1)
↳ _[3]=y*gen(2)
↳ _[4]=x2*gen(2)
↳ _[5]=x*gen(2)
↳ _[6]=gen(2)
 print(kbase(T)); // columns of matrix are a k-basis of T_1
↳ z,1,0,0, 0,0,
↳ 0,0,y,x2,x,1
 // -----
 // general case (cone over rational normal curve of degree 4):
 ring r1=0,(x,y,z,u,v),ds;
 matrix m[2][4]=x,y,z,u,y,z,u,v;
 ideal i=minor(m,2); // 2x2 minors of matrix m
 module M=T_1(i); // a presentation matrix of T_1
↳ // dim T_1 = 4
 vdim(M); // Tjurina number
↳ 4
 hilb(M); // display of both Hilbert series
↳ // 4 t^0
↳ // -20 t^1
↳ // 40 t^2
↳ // -40 t^3
↳ // 20 t^4
↳ // -4 t^5
↳
↳ // 4 t^0
↳ // dimension (local) = 0
↳ // multiplicity = 4
 intvec v1=hilb(M,1); // first Hilbert series as intvec
 intvec v2=hilb(M,2); // second Hilbert series as intvec
 v1;
↳ 4,-20,40,-40,20,-4,0
 v2;
↳ 4,0
 v1[3]; // 3rd coefficient of the 1st Hilbert series
↳ 40

```

```

 module N=T_2(i);
 ⇨ // dim T_2 = 3

 // In some cases it might be useful to have a polynomial in some ring
 // encoding the Hilbert series. This polynomial can then be
 // differentiated, evaluated etc. It can be done as follows:
 ring H = 0,t,ls;
 poly h1;
 int ii;
 for (ii=1; ii<=size(v1); ii=ii+1)
 {
 h1=h1+v1[ii]*t^(ii-1);
 }
 h1; // 1st Hilbert series
 ⇨ 4-20t+40t2-40t3+20t4-4t5
 diff(h1,t); // differentiate h1
 ⇨ -20+80t-120t2+80t3-20t4
 subst(h1,t,1); // substitute t by 1
 ⇨ 0

 // The procedures T_1, T_2, T_12 may be called with two arguments and then
 // they return a list with more information (type help T_1; etc.)
 // e.g., T_12(i,<any>); returns a list with 9 nonempty objects where
 // _[1] = std basis of T_1-module, _[2] = std basis of T_2-module,
 // _[3]= vdim of T_1, _[4]= vdim of T_2
 setring r1; // make r1 again the basering
 list L = T_12(i,1);
 ⇨ // dim T_1 = 4
 ⇨ // dim T_2 = 3
 kbase(L[1]); // kbase of T_1
 ⇨ _[1]=1*gen(2)
 ⇨ _[2]=1*gen(3)
 ⇨ _[3]=1*gen(6)
 ⇨ _[4]=1*gen(7)
 kbase(L[2]); // kbase of T_2
 ⇨ _[1]=1*gen(6)
 ⇨ _[2]=1*gen(8)
 ⇨ _[3]=1*gen(9)
 L[3]; // vdim of T_1
 ⇨ 4
 L[4]; // vdim of T_2
 ⇨ 3

```

#### A.4.5 Deformations

- The libraries `sing.lib`, respectively `deform.lib`, contain procedures to compute total and base space of the miniversal (= semiuniversal) deformation of an isolated complete intersection singularity, respectively of an arbitrary isolated singularity.
- The procedure `deform` in `sing.lib` returns a matrix whose columns  $h_1, \dots, h_r$  represent all 1st order deformations. More precisely, if  $I \subset R$  is the ideal generated by  $f_1, \dots, f_s$ , then any infinitesimal deformation of  $R/I$  over  $K[\varepsilon]/(\varepsilon^2)$  is given by  $f + \varepsilon g$ , where  $f = (f_1, \dots, f_s)$ , and where  $g$  is a  $K$ -linear combination of the  $h_i$ .

- The procedure `versal` in `deform.lib` computes a formal miniversal deformation up to a certain order which can be prescribed by the user. For a complete intersection the 1st order part is already miniversal.
- The procedure `versal` extends the basering to a new ring with additional deformation parameters which contains the equations for the miniversal base space and the miniversal total space.
- There are default names for the objects created, but the user may also choose their own names.
- If the user sets `printlevel=2;` before running `versal`, some intermediate results are shown. This is useful since `versal` is already complicated and might run for some time on more complicated examples. (type `help versal;`)

We compute for the same examples as in the section Section A.4.4 [T1 and T2], page 731 the miniversal deformations:

```
LIB "deform.lib";
ring R=32003,(x,y,z),ds;
//-----
// hypersurface case (from series T[p,q,r]):
int p,q,r = 3,3,4;
poly f = x^p+y^q+z^r+xyz;
print(deform(f));
↳ z3,z2,yz,xz,z,y,x,1
// the miniversal deformation of f=0 is the projection from the
// miniversal total space to the miniversal base space:
// { (A,B,C,D,E,F,G,H,x,y,z) | x3+y3+xyz+z4+A+Bx+Cxz+Dy+Eyz+Fz+Gz2+Hz3 =0 }
// --> { (A,B,C,D,E,F,G,H) }
//-----
// complete intersection case (from series P[k,l]):
int k,l =3,2;
ideal j=xy,x^k+y^l+z2;
print(deform(j));
↳ 0,0, 0,0,z,1,
↳ y,x2,x,1,0,0
def L=versal(j); // using default names
↳ // smooth base space
↳ // ready: T_1 and T_2
↳
↳
↳ // 'versal' returned a list, say L, of four rings. In L[1] are stored:
↳ // as matrix Fs: Equations of total space of the miniversal deformation\
,
↳ // as matrix Js: Equations of miniversal base space,
↳ // as matrix Rs: syzygies of Fs mod Js.
↳ // To access these data, type
↳ def Px=L[1]; setring Px; print(Fs); print(Js); print(Rs);
↳
↳ // L[2] = L[1]/Fo extending Qo=Po/Fo,
↳ // L[3] = the embedding ring of the versal base space,
↳ // L[4] = L[1]/Js extending L[3]/Js.
↳
def Px=L[1]; setring Px;
show(Px); // show is a procedure from inout.lib
↳ // ring: (ZZ/32003),(A,B,C,D,E,F,x,y,z),(ds(6),ds(3),C);
```



```

⇒ // minpoly = 0
⇒ // objects belonging to this ring:
⇒ // Rs [0] matrix 2 x 1
⇒ // Fs [0] matrix 1 x 2
⇒ // Js [0] matrix 1 x 0
 listvar(matrix);
⇒ // Rs [0] matrix 2 x 1
⇒ // Fs [0] matrix 1 x 2
⇒ // Js [0] matrix 1 x 0
 // ___ Equations of miniversal base space ___:
 Js;
⇒
 // ___ Equations of miniversal total space ___:
 Fs;
⇒ Fs[1,1]=y2+z2+x3+Cy+Dx2+Ex+F
⇒ Fs[1,2]=xy+Az+B
 // the miniversal deformation of V(j) is the projection from the
 // miniversal total space to the miniversal base space:
 // { (A,B,C,D,E,F,x,y,z) | xy+F+Ez=0, y2+z2+x3+D+Cx+Bx2+Ay=0 }
 // --> { (A,B,C,D,E,F) }
 //-----
 // general case (cone over rational normal curve of degree 4):
 kill L;
 ring r1=0,(x,y,z,u,v),ds;
 matrix m[2][4]=x,y,z,u,y,z,u,v;
 ideal i=minor(m,2); // 2x2 minors of matrix m
 int time=timer;
 // Call parameters of the miniversal base A(1),A(2),...:
 def L=versal(i,0,"","A(");
⇒ // ready: T_1 and T_2
⇒ // start computation in degree 2.
⇒
⇒
⇒ // 'versal' returned a list, say L, of four rings. In L[1] are stored:
⇒ // as matrix Fs: Equations of total space of the miniversal deformation\
,
⇒ // as matrix Js: Equations of miniversal base space,
⇒ // as matrix Rs: syzygies of Fs mod Js.
⇒ // To access these data, type
⇒ def Px=L[1]; setring Px; print(Fs); print(Js); print(Rs);
⇒
⇒ // L[2] = L[1]/Fo extending Qo=Po/Fo,
⇒ // L[3] = the embedding ring of the versal base space,
⇒ // L[4] = L[1]/Js extending L[3]/Js.
⇒
 // used time:",timer-time,"sec"; // time of last command
⇒ // used time: 0 sec
 def Def_rPx=L[1]; setring Def_rPx;
 Fs;
⇒ Fs[1,1]=u2-z*v-A(2)*u+A(4)*v
⇒ Fs[1,2]=z*u-y*v-A(1)*u+A(4)*u
⇒ Fs[1,3]=y*u-x*v+A(3)*u+A(4)*z
⇒ Fs[1,4]=z2-y*u-A(1)*z+A(2)*y

```

```

↳ Fs[1,5]=y*z-x*u+A(2)*x+A(3)*z
↳ Fs[1,6]=y^2-x*z+A(1)*x+A(3)*y
 Js;
↳ Js[1,1]=A(2)*A(4)
↳ Js[1,2]=-A(1)*A(4)+A(4)^2
↳ Js[1,3]=A(3)*A(4)
 // the miniversal deformation of V(i) is the projection from the
 // miniversal total space to the miniversal base space:
 // { (A(1..4),x,y,z,u,v) |
 // -u^2+x*v+A(2)*u+A(4)*v=0, -z*u+y*v-A(1)*u+A(3)*u=0,
 // -y*u+x*v+A(3)*u+A(4)*z=0, z^2-y*u+A(1)*z+A(2)*y=0,
 // y*z-x*u+A(2)*x-A(3)*z=0, -y^2+x*z+A(1)*x+A(3)*y=0 }
 // --> { A(1..4) |
 // A(2)*A(4) = -A(3)*A(4) = -A(1)*A(4)+A(4)^2 = 0 }
 //-----

```

#### A.4.6 Invariants of plane curve singularities

The Puiseux pairs of an irreducible and reduced plane curve singularity are probably its most important invariants. They can be computed from its Hamburger-Noether expansion (which is the analogue of the Puiseux expansion in characteristic 0 for fields of arbitrary characteristic).

The library `hnoether.lib` (see Section D.6.15 [hnoether.lib], page 864) uses the algorithm of Antonio Campillo in "Algebroid curves in positive characteristic" SLN 813, 1980. This algorithm has the advantage that it needs least possible field extensions and, moreover, works in any characteristic. This fact can be used to compute the invariants over a field of finite characteristic, say 32003, which will most probably be the same as in characteristic 0.

We compute the Hamburger-Noether expansion of a plane curve singularity given by a polynomial  $f$  in two variables. This expansion is given by a matrix, and it allows us to compute a primitive parametrization (up to a given order) for the curve singularity defined by  $f$  and numerical invariants such as the

- characteristic exponents,
- Puiseux pairs (of a complex model),
- degree of the conductor,
- delta invariant,
- generators of the semigroup.

Besides commands for computing a parametrization and the invariants mentioned above, the library `hnoether.lib` provides commands for the computation of the Newton polygon of  $f$ , the square-free part of  $f$  and a procedure to convert one set of invariants to another.

```

LIB "hnoether.lib";
// ===== The irreducible case =====
ring s = 0,(x,y),ds;
poly f = y4-2x3y2-4x5y+x6-x7;
list hn = develop(f);
show(hn[1]); // Hamburger-Noether matrix
↳ // matrix, 3x3
↳ 0,x, 0,
↳ 0,1, x,
↳ 0,1/4,-1/2
 displayHNE(hn); // Hamburger-Noether development
↳ y = z(1)*x

```

```

⇒ x = z(1)^2+z(1)^2*z(2)
⇒ z(1) = 1/4*z(2)^2-1/2*z(2)^3 + (terms of degree >=4)
 setring s;
 displayInvariants(hn);
⇒ characteristic exponents : 4,6,7
⇒ generators of semigroup : 4,6,13
⇒ Puiseux pairs : (3,2)(7,2)
⇒ degree of the conductor : 16
⇒ delta invariant : 8
⇒ sequence of multiplicities: 4,2,2,1,1
 // invariants(hn); returns the invariants as list
 // partial parametrization of f: param takes the first variable
 // as infinite except the ring has more than 2 variables. Then
 // the 3rd variable is chosen.
 param(hn);
⇒ // ** Warning: result is exact up to order 5 in x and 7 in y !
⇒ _[1]=1/16x4-3/16x5+1/4x7
⇒ _[2]=1/64x6-5/64x7+3/32x8+1/16x9-1/8x10
 ring extring=0,(x,y,t),ds;
 poly f=x3+2xy2+y2;
 list hn=develop(f,-1);
 param(hn); // partial parametrization of f
⇒ // ** Warning: result is exact up to order 2 in x and 3 in y !
⇒ _[1]=-t2
⇒ _[2]=-t3
 list hn1=develop(f,6);
 param(hn1); // a better parametrization
⇒ // ** Warning: result is exact up to order 6 in x and 7 in y !
⇒ _[1]=-t2+2t4-4t6
⇒ _[2]=-t3+2t5-4t7
 // instead of recomputing you may extend the development:
 list hn2=extdevelop(hn,12);
 param(hn2); // a still better parametrization
⇒ // ** Warning: result is exact up to order 12 in x and 13 in y !
⇒ _[1]=-t2+2t4-4t6+8t8-16t10+32t12
⇒ _[2]=-t3+2t5-4t7+8t9-16t11+32t13
 //
 // ===== The reducible case =====
 ring r = 0,(x,y),dp;
 poly f=x11-2y2x8-y3x7-y2x6+y4x5+2y4x3+y5x2-y6;
 // = (x5-1y2) * (x6-2x3y2-1x2y3+y4)
 list L=hnexpansion(f);
⇒ // No change of ring necessary, return value is HN expansion.
 show(L[1][1]); // Hamburger-Noether matrix of 1st branch
⇒ // matrix, 3x3
⇒ 0,x,0,
⇒ 0,1,x,
⇒ 0,1,-1
 displayInvariants(L);
⇒ --- invariants of branch number 1 : ---
⇒ characteristic exponents : 4,6,7
⇒ generators of semigroup : 4,6,13
⇒ Puiseux pairs : (3,2)(7,2)

```

```

↳ degree of the conductor : 16
↳ delta invariant : 8
↳ sequence of multiplicities: 4,2,2,1,1
↳
↳ --- invariants of branch number 2 : ---
↳ characteristic exponents : 2,5
↳ generators of semigroup : 2,5
↳ Puiseux pairs : (5,2)
↳ degree of the conductor : 4
↳ delta invariant : 2
↳ sequence of multiplicities: 2,2,1,1
↳
↳ ----- contact numbers : -----
↳
↳ branch | 2
↳ -----+-----
↳ 1 | 2
↳
↳ ----- intersection multiplicities : -----
↳
↳ branch | 2
↳ -----+-----
↳ 1 | 12
↳
↳ ----- delta invariant of the curve : 22
 param(L[2]); // parametrization of 2nd branch
↳ _[1]=x2
↳ _[2]=x5

```

#### A.4.7 Branches of space curve singularities

In this example, the number of branches of a given quasihomogeneous isolated space curve singularity will be computed as an example of the pitfalls appearing in the use of primary decomposition. When dealing with singularities, two situations are possible in which the primary decomposition algorithm might not lead to a complete decomposition: first of all, one of the computed components could be globally irreducible, but analytically reducible (this is impossible for quasihomogeneous singularities) and, as a second possibility, a component might be irreducible over the rational numbers, but reducible over the complex numbers.

```

ring r=0,(x,y,z),ds;
ideal i=x^4-y*z^2,x*y-z^3,y^2-x^3*z; // the space curve singularity
qhweight(i);
↳ 1,2,1
// The given space curve singularity is quasihomogeneous. Hence we can pass
// to the polynomial ring.
ring rr=0,(x,y,z),dp;
ideal i=imap(r,i);
resolution ires=mres(i,0);
ires;
↳ 1 3 2
↳ rr <-- rr <-- rr
↳
↳ 0 1 2
↳

```

```

// From the structure of the resolution, we see that the Cohen-Macaulay
// type of the given singularity is 2
//
// Let us now look for the branches using the primdec library.
LIB "primdec.lib";
primdecSY(i);
⇨ [1]:
⇨ [1]:
⇨ _[1]=z3-xy
⇨ _[2]=x3+x2z+xz2+xy+yz
⇨ _[3]=x2z2+x2y+xyz+yz2+y2
⇨ [2]:
⇨ _[1]=z3-xy
⇨ _[2]=x3+x2z+xz2+xy+yz
⇨ _[3]=x2z2+x2y+xyz+yz2+y2
⇨ [2]:
⇨ [1]:
⇨ _[1]=x-z
⇨ _[2]=z2-y
⇨ [2]:
⇨ _[1]=x-z
⇨ _[2]=z2-y
 def li=_[1];
 ideal i2=li[2]; // call the first ideal i1
 // The curve seems to have 2 branches by what we computed using the
 // algorithm of Shimoyama-Yokoyama.
 // Now the same computation by the Gianni-Trager-Zacharias algorithm:
 primdecGTZ(i);
⇨ [1]:
⇨ [1]:
⇨ _[1]=-z2+y
⇨ _[2]=x-z
⇨ [2]:
⇨ _[1]=-z2+y
⇨ _[2]=x-z
⇨ [2]:
⇨ [1]:
⇨ _[1]=z8+yz6+y2z4+y3z2+y4
⇨ _[2]=xz5+z6+yz4+y2z2+y3
⇨ _[3]=-z3+xy
⇨ _[4]=x2z2+xz3+xyz+yz2+y2
⇨ _[5]=x3+x2z+xz2+xy+yz
⇨ [2]:
⇨ _[1]=z8+yz6+y2z4+y3z2+y4
⇨ _[2]=xz5+z6+yz4+y2z2+y3
⇨ _[3]=-z3+xy
⇨ _[4]=x2z2+xz3+xyz+yz2+y2
⇨ _[5]=x3+x2z+xz2+xy+yz
 // Having computed the primary decomposition in 2 different ways and
 // having obtained the same number of branches, we might expect that the
 // number of branches is really 2, but we can check this by formulae
 // for the invariants of space curve singularities:
 //

```

```

// mu = tau - t + 1 (for quasihomogeneous curve singularities)
// where mu denotes the Milnor number, tau the Tjurina number and
// t the Cohen-Macaulay type
//
// mu = 2 delta - r + 1
// where delta denotes the delta-Invariant and r the number of branches
//
// tau can be computed by using the corresponding procedure T1 from
// sing.lib.
setring r;
LIB "sing.lib";
T_1(i);
⇒ // dim T_1 = 13
⇒ _[1]=gen(6)+2z*gen(5)
⇒ _[2]=gen(4)+3x2*gen(2)
⇒ _[3]=gen(3)+gen(1)
⇒ _[4]=x*gen(5)-y*gen(2)-z*gen(1)
⇒ _[5]=x*gen(1)-z2*gen(2)
⇒ _[6]=y*gen(5)+3x2z*gen(2)
⇒ _[7]=y*gen(2)-z*gen(1)
⇒ _[8]=2y*gen(1)-z2*gen(5)
⇒ _[9]=z2*gen(5)
⇒ _[10]=z2*gen(1)
⇒ _[11]=x3*gen(2)
⇒ _[12]=x2z2*gen(2)
⇒ _[13]=xz3*gen(2)
⇒ _[14]=z4*gen(2)
setring rr;
// Hence tau is 13 and therefore mu is 12. But then it is impossible that
// the singularity has two branches, since mu is even and delta is an
// integer!
// So obviously, we did not decompose completely. Because the second branch
// is smooth, only the first ideal can be the one which can be decomposed
// further.
// Let us now consider the normalization of this first ideal i1.
LIB "normal.lib";
normal(i2);
⇒
⇒ // 'normal' created a list, say nor, of two elements.
⇒ // To see the list type
⇒ nor;
⇒
⇒ // * nor[1] is a list of 1 ring(s).
⇒ // To access the i-th ring nor[1][i], give it a name, say Ri, and type
⇒ def R1 = nor[1][1]; setring R1; norid; normap;
⇒ // For the other rings type first (if R is the name of your base ring)
⇒ setring R;
⇒ // and then continue as for R1.
⇒ // Ri/norid is the affine algebra of the normalization of R/P_i where
⇒ // P_i is the i-th component of a decomposition of the input ideal id
⇒ // and normap the normalization map from R to Ri/norid.
⇒
⇒ // * nor[2] is a list of 1 ideal(s). Let ci be the last generator

```

```

⇒ // of the ideal nor[2][i]. Then the integral closure of R/P_i is
⇒ // generated as R-submodule of the total ring of fractions by
⇒ // 1/ci * nor[2][i].
⇒ [1]:
⇒ [1]:
⇒ // coefficients: QQ
⇒ // number of vars : 6
⇒ // block 1 : ordering dp
⇒ // : names T(1) T(2) T(3)
⇒ // block 2 : ordering dp
⇒ // : names x y z
⇒ // block 3 : ordering C
⇒ [2]:
⇒ [1]:
⇒ _[1]=y
⇒ _[2]=xz
⇒ _[3]=x2
⇒ _[4]=z2
⇒ def rno=_[1][1];
⇒ setring rno;
⇒ norid;
⇒ norid[1]=-T(2)*z+x
⇒ norid[2]=T(1)*x-z
⇒ norid[3]=T(2)*x-T(3)*z
⇒ norid[4]=T(1)*z+T(2)*z+T(3)*x+T(3)*z+z
⇒ norid[5]=-T(2)*y+z^2
⇒ norid[6]=T(1)*z^2-y
⇒ norid[7]=T(2)*z^2-T(3)*y
⇒ norid[8]=T(1)*y+T(2)*y+T(3)*z^2+T(3)*y+y
⇒ norid[9]=T(1)^2+T(1)+T(2)+T(3)+1
⇒ norid[10]=T(1)*T(2)-1
⇒ norid[11]=T(2)^2-T(3)
⇒ norid[12]=T(1)*T(3)-T(2)
⇒ norid[13]=T(2)*T(3)+T(1)+T(2)+T(3)+1
⇒ norid[14]=T(3)^2-T(1)
⇒ norid[15]=z^3-x*y
⇒ norid[16]=x^3+x^2*z+x*z^2+x*y+y*z
⇒ norid[17]=x^2*z^2+x^2*y+x*y*z+y*z^2+y^2
⇒ // The ideal is generated by a polynomial in one variable of degree 4 which
⇒ // factors completely into 4 polynomials of type T(2)+a.
⇒ // From this, we know that the ring of the normalization is the direct sum of
⇒ // 4 polynomial rings in one variable.
⇒ // Hence our original curve has these 4 branches plus a smooth one
⇒ // which we already determined by primary decomposition.
⇒ // Our final result is therefore: 5 branches.

```

#### A.4.8 Classification of hypersurface singularities

Classification of isolated hypersurface singularities with respect to right equivalence is provided by the command `classify` of the library `classify.lib`. The classification is done by using the algorithm of Arnold. Before entering this algorithm, a first guess based on the Hilbert polynomial of the Milnor algebra is made.

```
LIB "classify.lib";
```

```

ring r=0,(x,y,z),ds;
poly p=singularity("E[6k+2]",2)[1];
p=p+z^2;
p;
↳ z2+x3+xy6+y8
// We received an E_14 singularity in normal form
// from the database of normal forms. Since only the residual
// part is saved in the database, we added z^2 to get an E_14
// of embedding dimension 3.
//
// Now we apply a coordinate change in order to deal with a
// singularity which is not in normal form:
map phi=r,x+y,y+z,x;
poly q=phi(p);
// Yes, q really looks ugly, now:
q;
↳ x2+x3+3x2y+3xy2+y3+xy6+y7+6xy5z+6y6z+15xy4z2+15y5z2+20xy3z3+20y4z3+15xy2z\
 4+15y3z4+6xyz5+6y2z5+xz6+yz6+y8+8y7z+28y6z2+56y5z3+70y4z4+56y3z5+28y2z6+8\
 yz7+z8
// Classification
classify(q);
↳ About the singularity :
↳ Milnor number(f) = 14
↳ Corank(f) = 2
↳ Determinacy <= 12
↳ Guessing type via Milnorcode: E[6k+2]=E[14]
↳
↳ Computing normal form ...
↳ I have to apply the splitting lemma. This will take some time....:-)
↳ Arnold step number 9
↳ The singularity
↳ x3-9/4x4+27/4x5-189/8x6+737/8x7+6x6y+15x5y2+20x4y3+15x3y4+6x2y5+xy6-24\
 089/64x8-x7y+11/2x6y2+26x5y3+95/2x4y4+47x3y5+53/2x2y6+8xy7+y8+104535/64x9\
 +27x8y+135/2x7y2+90x6y3+135/2x5y4+27x4y5+9/2x3y6-940383/128x10-405/4x9y-2\
 025/8x8y2-675/2x7y3-2025/8x6y4-405/4x5y5-135/8x4y6+4359015/128x11+1701/4x\
 10y+8505/8x9y2+2835/2x8y3+8505/8x7y4+1701/4x6y5+567/8x5y6-82812341/512x12\
 -15333/8x11y-76809/16x10y2-25735/4x9y3-78525/16x8y4-16893/8x7y5-8799/16x6\
 y6-198x5y7-495/4x4y8-55x3y9-33/2x2y10-3xy11-1/4y12
↳ is R-equivalent to E[14].
↳ Milnor number = 14
↳ modality = 1
↳ 2z2+x3+xy6+y8
// The library also provides routines to determine the corank of q
// and its residual part without going through the whole
// classification algorithm.
corank(q);
↳ 2
morsesplit(q);
↳ y3-9/4y4+27/4y5-189/8y6+737/8y7+6y6z+15y5z2+20y4z3+15y3z4+6y2z5+yz6-24089\
 /64y8-y7z+11/2y6z2+26y5z3+95/2y4z4+47y3z5+53/2y2z6+8yz7+z8+104535/64y9+27\
 y8z+135/2y7z2+90y6z3+135/2y5z4+27y4z5+9/2y3z6-940383/128y10-405/4y9z-2025\
 /8y8z2-675/2y7z3-2025/8y6z4-405/4y5z5-135/8y4z6+4359015/128y11+1701/4y10z\
 +8505/8y9z2+2835/2y8z3+8505/8y7z4+1701/4y6z5+567/8y5z6-82812341/512y12-15\

```



333/8y11z-76809/16y10z2-25735/4y9z3-78525/16y8z4-16893/8y7z5-8799/16y6z6-\  
198y5z7-495/4y4z8-55y3z9-33/2y2z10-3yz11-1/4z12

#### A.4.9 Resolution of singularities

Resolution of singularities and applications thereof are provided by the libraries `resolve.lib` and `reszeta.lib`; graphical output may be generated automatically by using external programs `surf` and `dot` respectively to which a specialized interface is provided by the library `resgraph.lib`. In this example, the basic functionality of the resolution of singularities package is illustrated by the computation of the intersection matrix and genera of the exceptional curves on a surface obtained from resolving the A6 surface singularity. A separate tutorial, which introduces the complete functionality of the package and explains the rather complicated data structures appearing in intermediate results, can be found at [https://www.singular.uni-kl.de/tutor\\_resol.pdf](https://www.singular.uni-kl.de/tutor_resol.pdf).

```
LIB"resolve.lib"; // load the resolution algorithm
LIB"reszeta.lib"; // load its application algorithms

ring R=0,(x,y,z),dp; // define the ring Q[x,y,z]
ideal I=x7+y2-z2; // an A6 surface singularity
list L=resolve(I); // compute the resolution
list iD=intersectionDiv(L); // compute intersection properties
iD; // show the output
↳ [1]:
↳ -2,0,1,0,0,0,
↳ 0,-2,0,1,0,0,
↳ 1,0,-2,0,1,0,
↳ 0,1,0,-2,0,1,
↳ 0,0,1,0,-2,1,
↳ 0,0,0,1,1,-2
↳ [2]:
↳ 0,0,0,0,0,0
↳ [3]:
↳ [1]:
↳ [1]:
↳ 2,1,1
↳ [2]:
↳ 4,1,1
↳ [2]:
↳ [1]:
↳ 2,1,2
↳ [2]:
↳ 4,1,2
↳ [3]:
↳ [1]:
↳ 4,2,1
↳ [2]:
↳ 6,2,1
↳ [4]:
↳ [1]:
↳ 4,2,2
↳ [2]:
↳ 6,2,2
↳ [5]:
↳ [1]:
```

```

↳ 6,3,1
↳ [2]:
↳ 7,3,1
↳ [6]:
↳ [1]:
↳ 6,3,2
↳ [2]:
↳ 7,3,2
↳ [4]:
↳ 1,1,1,1,1,1
// The output is a list whose first entry contains the intersection matrix
// of the exceptional divisors. The second entry is the list of genera
// of these divisors. The third and fourth entry contain the information
// how to find the corresponding divisors in the respective charts.

```

## A.5 Invariant Theory

### A.5.1 G<sub>a</sub>-Invariants

We work in characteristic 0 and use the Lie algebra generated by one vectorfield of the form  $\sum x_i \partial / \partial x_{i+1}$ .

```

LIB "ainvar.lib";
int n=5;
int i;
ring s=32003,(x(1..n)),wp(1,2,3,4,5);
// definition of the vectorfield m=sum m[i,1]*d/dx(i)
matrix m[n][1];
for (i=1;i<=n-1;i=i+1)
{
 m[i+1,1]=x(i);
}
// computation of the ring of invariants
ideal in=invariantRing(m,x(2),x(1),0);
in; //invariant ring is generated by 5 invariants
↳ in[1]=x(1)
↳ in[2]=x(2)^2-2*x(1)*x(3)
↳ in[3]=x(3)^2-2*x(2)*x(4)+2*x(1)*x(5)
↳ in[4]=x(2)^3-3*x(1)*x(2)*x(3)+3*x(1)^2*x(4)
↳ in[5]=x(3)^3-3*x(2)*x(3)*x(4)-15997*x(1)*x(4)^2+3*x(2)^2*x(5)-6*x(1)*x(3)\
*x(5)
ring q=32003,(x,y,z,u,v,w),dp;
matrix m[6][1];
m[2,1]=x;
m[3,1]=y;
m[5,1]=u;
m[6,1]=v;
// the vectorfield is: xd/dy+yd/dz+ud/dv+vd/dw
ideal in=invariantRing(m,y,x,0);
in; //invariant ring is generated by 6 invariants
↳ in[1]=x
↳ in[2]=u
↳ in[3]=v^2-2uw

```

```

↳ in[4]=zu-yv+xw
↳ in[5]=yu-xv
↳ in[6]=y2-2xz

```

### A.5.2 Invariants of a finite group

Two algorithms to compute the invariant ring are implemented in SINGULAR, `invariant_ring` and `invariant_ring_random`, both by Agnes E. Heydtmann ([agnes@math.uni-sb.de](mailto:agnes@math.uni-sb.de)).

Bases of homogeneous invariants are generated successively and those are chosen as primary invariants that lower the dimension of the ideal generated by the previously found invariants (see paper "Generating a Noetherian Normalization of the Invariant Ring of a Finite Group" by Decker, Heydtmann, Schreyer (J.Symb.Comput. 25, No.6, 727-731, 1998). In the non-modular case secondary invariants are calculated by finding a basis (in terms of monomials) of the basering modulo the primary invariants, mapping to invariants with the Reynolds operator and using those or their power products such that they are linearly independent modulo the primary invariants (see paper "Some Algorithms in Invariant Theory of Finite Groups" by Kemper and Steel (In: Proceedings of the Euroconference in Essen 1997, Birkhäuser Prog. Math. 173, 267-285, 1999)). In the modular case they are generated according to "Calculating Invariant Rings of Finite Groups over Arbitrary Fields" by Kemper (J.Symb.Comput. 21, No.3, 351-366, 1996).

We calculate now an example from Sturmfels: "Algorithms in Invariant Theory 2.3.7":

```

LIB "finvar.lib";
ring R=0,(x,y,z),dp;
matrix A[3][3]=0,1,0,-1,0,0,0,0,-1;
// the group G is generated by A in Gl(3,Q);
print(A);
↳ 0, 1,0,
↳ -1,0,0,
↳ 0, 0,-1
print(A*A*A*A); // the fourth power of A is 1
↳ 1,0,0,
↳ 0,1,0,
↳ 0,0,1
// Use the first method to compute the invariants of G:
matrix B(1..3);
B(1..3)=invariant_ring(A);
// SINGULAR returns 2 matrices, the first containing
// primary invariants and the second secondary
// invariants, i.e., module generators over a Noetherian
// normalization
// the third result are the irreducible secondary invariants
// if the Molien series was available
print(B(1));
↳ z2,x2+y2,x2y2
print(B(2));
↳ 1,xyz,x2z-y2z,x3y-xy3
print(B(3));
↳ xyz,x2z-y2z,x3y-xy3
// Use the second method,
// with random numbers between -1 and 1:
B(1..3)=invariant_ring_random(A,1);
print(B(1..3));
↳ z2,x2+y2,x4+y4-z4

```

```

↦ 1,xyz,x2z-y2z,x3y-xy3
↦ xyz,x2z-y2z,x3y-xy3

```

## A.6 Geometric Invariant Theory

### A.6.1 GIT-Fans

Dolgachev/Hu and Thaddeus assigned to an algebraic variety with the action of an algebraic group the GIT-fan, a polyhedral fan enumerating the GIT-quotients in the sense of Mumford. The case of the action of an algebraic torus  $H$  on an affine variety  $X$  has been treated by Berchtold/Hausen. Based on their construction, an algorithm to compute the GIT-fan in this setting has been proposed by Keicher. Note that this setting is essential for many applications, since the torus case can be used to investigate the GIT-variation of the action of a connected reductive group  $G$ . In many important examples,  $X$  is symmetric under the action of a finite group which either is known directly from its geometry or can be computed. A prominent instance is the Deligne-Mumford compactification  $M_{0,6}$  of the moduli space of 6-pointed stable curves of genus zero, which has a natural action of the symmetric group  $S_6$ . The library `gitfan.lib` implements an efficient algorithm for computing GIT-fans, which makes use of symmetries. We have applied this algorithm to determine the Mori chamber decomposition of the cone of movable divisor classes of  $M_{0,6}$ . Each cone is represented by a single integer. The algorithm relies on Groebner basis techniques, convex geometry and actions of finite symmetry groups. It demonstrates the strength of cross-boarder methods in computer algebra, and the efficiency of the algorithms implemented in all involved areas. The algorithm is also suitable for parallel computations.

As an example we address in the following the computation of the GIT-Fan of  $M_{0,5}$ .

We first compute the GIT-fan using the single line command provided by the library:

```

LIB "gitfan.lib";
setcores(4);
ring R = 0,T(1..10),wp(1,1,1,1,1,1,1,1,1,1);
ideal J =
 T(5)*T(10)-T(6)*T(9)+T(7)*T(8),
 T(1)*T(9)-T(2)*T(7)+T(4)*T(5),
 T(1)*T(8)-T(2)*T(6)+T(3)*T(5),
 T(1)*T(10)-T(3)*T(7)+T(4)*T(6),
 T(2)*T(10)-T(3)*T(9)+T(4)*T(8);
intmat Q[5][10] =
 1, 1, 1, 1, 0, 0, 0, 0, 0, 0,
 1, 0, 0, 0, 1, 1, 1, 0, 0, 0,
 0, 1, 1, 0, 0, 0, -1, 1, 0, 0,
 0, 1, 0, 1, 0, -1, 0, 0, 1, 0,
 0, 0, 1, 1, -1, 0, 0, 0, 0, 1;
fan GIT = GITfan(J,Q);
fVector(GIT);

```

The GIT-Fan can be computed using symmetries as follows:

```

LIB "gitfan.lib";
setcores(4);
ring R = 0,T(1..10),wp(1,1,1,1,1,1,1,1,1,1);
ideal J =
 T(5)*T(10)-T(6)*T(9)+T(7)*T(8),
 T(1)*T(9)-T(2)*T(7)+T(4)*T(5),
 T(1)*T(8)-T(2)*T(6)+T(3)*T(5),

```

```

T(1)*T(10)-T(3)*T(7)+T(4)*T(6),
T(2)*T(10)-T(3)*T(9)+T(4)*T(8);
intmat Q[5][10] =
 1, 1, 1, 1, 0, 0, 0, 0, 0, 0,
 1, 0, 0, 0, 1, 1, 1, 0, 0, 0,
 0, 1, 1, 0, 0, 0, -1, 1, 0, 0,
 0, 1, 0, 1, 0, -1, 0, 0, 1, 0,
 0, 0, 1, 1, -1, 0, 0, 0, 0, 1;
list simplexSymmetryGroup = G25Action();
fan GIT2 = GITfan(J,Q,simplexSymmetryGroup);
GIT2;

```

Although we provide a procedure to compute the orbit decomposition of the group action on the simplex of variables this is not fast in Singular. In the following we describe how to use GAP to obtain the orbit decomposition and then continue with this data in Singular. This is particularly useful for more complicated examples.

The file `orbits.gp` in the directory `doc` of the Singular source tree contains GAP code to do this computation. This result is provided in the file `doc/simplexOrbitRepresentativesG25.sing`.

The file `doc/simplexSymmetryGroupG25.sing` contains the symmetry group (which here is  $S_5$ ).

Moreover the file `doc/elementsInTermsOfGeneratorsG25.sing` contains a representation of the elements of the symmetry group in terms of generators.

```

LIB "gitfan.lib";
setcores(4);
ring R = 0,T(1..10),wp(1,1,1,1,1,1,1,1,1,1);
ideal J =
 T(5)*T(10)-T(6)*T(9)+T(7)*T(8),
 T(1)*T(9)-T(2)*T(7)+T(4)*T(5),
 T(1)*T(8)-T(2)*T(6)+T(3)*T(5),
 T(1)*T(10)-T(3)*T(7)+T(4)*T(6),
 T(2)*T(10)-T(3)*T(9)+T(4)*T(8);
intmat Q[5][10] =
 1, 1, 1, 1, 0, 0, 0, 0, 0, 0,
 1, 0, 0, 0, 1, 1, 1, 0, 0, 0,
 0, 1, 1, 0, 0, 0, -1, 1, 0, 0,
 0, 1, 0, 1, 0, -1, 0, 0, 1, 0,
 0, 0, 1, 1, -1, 0, 0, 0, 0, 1;
intmat Qt = transpose(Q);
<"doc/simplexOrbitRepresentativesG25.sing";
list afaceOrbitRepresentatives=afaces(J,simplexOrbitRepresentatives);
<"doc/simplexSymmetryGroupG25.sing";
list fullDimAfaceOrbitRepresentatives=fullDimImages(afaceOrbitRepresentatives,Q);
list afaceOrbits=computeAfaceOrbits(fullDimAfaceOrbitRepresentatives,simplexSymmetry);
apply(afaceOrbits,size);
list minAfaceOrbits = minimalAfaceOrbits(afaceOrbits);
apply(minAfaceOrbits,size);
list listOfOrbitConeOrbits = orbitConeOrbits(minAfaceOrbits,Q);
apply(listOfOrbitConeOrbits,size);
list listOfMinimalOrbitConeOrbits = minimalOrbitConeOrbits(listOfOrbitConeOrbits);
size(listOfMinimalOrbitConeOrbits);
cone mov = coneViaPoints(transpose(Q));
mov = canonicalizeCone(mov);
list OC = listOfOrbitConeOrbits;

```

```

<"doc/elementsInTermsOfGeneratorsG25.sing";
list Asigmagens = groupActionOnQImage(generatorsG,Q);
list actionOnOrbitconeIndicesForGenerators = groupActionOnHashes(Asigmagens,OC);
list actionOnOrbitconeIndices;
for (int i =1; i<=size(elementsInTermsOfGenerators);i++)
{
 actionOnOrbitconeIndices[i]=evaluateProduct(actionOnOrbitconeIndicesForGenerators,
}
list OClist = OC[1];
for (i =2;i<=size(OC);i++)
{
 OClist = OClist + OC[i];
}
list SigmaHashes = GITfanParallelSymmetric(OClist, Q, mov, actionOnOrbitconeIndices)
SigmaHashes;
fan Sigma = hashesToFan(SigmaHashes,OClist);

```

Note that the result is not the complete fan but only the fan generated by a minimal set of representatives of maximal cones for the group action (by the group generated by `Asigmagens`).

## A.7 Non-commutative Algebra

### A.7.1 Left and two-sided Groebner bases

For a set of polynomials (resp. vectors)  $S$  in a non-commutative  $G$ -algebra, `SINGULAR:PLURAL` provides two algorithms for computing Groebner bases.

The command `std` computes a left Groebner basis of a left module, generated by the set  $S$  (see Section 7.3.26 [`std (plural)`], page 355). The command `twostd (plural)` computes a two-sided Groebner basis (which is in particular also a left Groebner basis) of a two-sided ideal, generated by the set  $S$  (see Section 7.3.29 [`twostd (plural)`], page 358).

In the example below, we consider a particular set  $S$  in the algebra  $A := U(sl_2)$  with the degree reverse lexicographic ordering. We compute a left Groebner basis  $L$  of the left ideal generated by  $S$  and a two-sided Groebner basis  $T$  of the two-sided ideal generated by  $S$ .

Then, we read off the information on the vector space dimension of the factor modules  $A/L$  and  $A/T$  using the command `vdim` (see Section 7.3.30 [`vdim (plural)`], page 359).

Further on, we use the command `reduce` (see Section 7.3.23 [`reduce (plural)`], page 351) to compare the left ideals generated by  $L$  and  $T$ .

We set `option(redSB)` and `option(redTail)` to make `SINGULAR` compute completely reduced minimal bases of ideals (see Section 5.1.110 [`option`], page 230 and Section 7.4.2 [Groebner bases in  $G$ -algebras], page 361 for definitions and further details).

For long running computations, it is always recommended to set `option(prot)` to make `SINGULAR` display some information on the performed computations (see Section 5.1.110 [`option`], page 230 for an interpretation of the displayed symbols).

```

// ----- 1. setting up the algebra
ring R = 0,(e,f,h),dp;
matrix D[3][3];
D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
def A=nc_algebra(1,D); setring A;
// ----- equivalently, you may use the following:
// LIB "ncalg.lib";

```

```

// def A = makeUs12();
// setring A;
// ----- 2. defining the set S
ideal S = e^3, f^3, h^3 - 4*h;
option(redSB);
option(redTail);
option(prot); // let us activate the protocol
ideal L = std(S);
↳ 3(2)s
↳ s
↳ s
↳ 5s
↳ s
↳ (4)s
↳ 4(5)(4)s
↳ (6)(5)(4)s
↳ 3(7)4(5)(4)(3)s
↳ 3(4)(3)4(2)s
↳ (3)(2)s
↳ 3(5)(4)4(2)5
↳ (S:5)-----
↳ product criterion:7 chain criterion:12
L;
↳ L[1]=h3-4h
↳ L[2]=fh2-2fh
↳ L[3]=eh2+2eh
↳ L[4]=2efh-h2-2h
↳ L[5]=f3
↳ L[6]=e3
vdim(L); // the vector space dimension of the module A/L
↳ 15
option(noprot); // turn off the protocol
ideal T = twostd(S);
T;
↳ T[1]=h3-4h
↳ T[2]=fh2-2fh
↳ T[3]=eh2+2eh
↳ T[4]=f2h-2f2
↳ T[5]=2efh-h2-2h
↳ T[6]=e2h+2e2
↳ T[7]=f3
↳ T[8]=ef2-fh
↳ T[9]=e2f-eh-2e
↳ T[10]=e3
vdim(T); // the vector space dimension of the module A/T
↳ 10
print(matrix(reduce(L,T))); // reduce L with respect to T
↳ 0,0,0,0,0,0
// as we see, L is included in the left ideal generated by T
print(matrix(reduce(T,L))); // reduce T with respect to L
↳ 0,0,0,f2h-2f2,0,e2h+2e2,0,ef2-fh,e2f-eh-2e,0
// the non-zero elements belong to T only
ideal LT = twostd(L); // the two-sided Groebner basis of L

```

```

// LT and T coincide as left ideals:
size(reduce(LT,T));
↳ 0
size(reduce(T,LT));
↳ 0

```

### A.7.2 Right Groebner bases and syzygies

Most of the SINGULAR:PLURAL commands correspond to the *left-sided* computations, that is left Groebner bases, left syzygies, left resolutions and so on. However, the *right-sided* computations can be done, using the *left-sided* functionality and *opposite* algebras.

In the example below, we consider the algebra  $A := U(sl_2)$  and a set of generators  $I = \{e^2, f\}$ .

We will compute a left Groebner basis LI and a left syzygy module LS of a left ideal, generated by the set  $I$ .

Then, we define the opposite algebra Aop of A, set it as a basering, and create opposite objects of already computed ones.

Further on, we compute a right Groebner basis RI and a right syzygy module RS of a right ideal, generated by the set  $I$  in  $A$ .

```

// ----- setting up the algebra:
LIB "ncalg.lib";
def A = makeUs12();
setring A; A;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ // block 1 : ordering dp
↳ // : names e f h
↳ // block 2 : ordering C
↳ // noncommutative relations:
↳ // fe=ef-h
↳ // he=eh+2e
↳ // hf=fh-2f
// ----- equivalently, you may use
// ring AA = 0,(e,f,h),dp;
// matrix D[3][3];
// D[1,2]=-h; D[1,3]=2*e; D[2,3]=-2*f;
// def A=nc_algebra(1,D); setring A;
option(redSB);
option(redTail);
matrix T;
// --- define a generating set
ideal I = e2,f;
ideal LI = std(I); // the left Groebner basis of I
LI; // we see that I was not a Groebner basis
↳ LI[1]=f
↳ LI[2]=h2+h
↳ LI[3]=eh+e
↳ LI[4]=e2
module LS = syz(I); // the left syzygy module of I
print(LS);
↳ -ef-2h+6,-f3, -ef2-fh+4f, -e2f2-4efh+16ef-6h2+42h-72\
,

```



```

↳ e3, e2f2-6efh-6ef+6h2+18h+12,e3f-3e2h-6e2,e4f
 // check: LS is a left syzygy, if T=0:
T = transpose(LS)*transpose(I);
print(T);
↳ 0,
↳ 0,
↳ 0,
↳ 0
// --- let us define the opposite algebra of A
def Aop = opposite(A);
setring Aop; Aop; // see how Aop looks like
↳ // coefficients: QQ
↳ // number of vars : 3
↳ // block 1 : ordering a
↳ // : names H F E
↳ // : weights 1 1 1
↳ // block 2 : ordering ls
↳ // : names H F E
↳ // block 3 : ordering C
↳ // noncommutative relations:
↳ // FH=HF-2F
↳ // EH=HE+2E
↳ // EF=FE-H
// --- we "oppose" (transfer) objects from A to Aop
ideal Iop = oppose(A,I);
ideal RIop = std(Iop); // the left Groebner basis of Iop in Aop
module RSop = syz(Iop); // the left syzygy module of Iop in Aop
module LSop = oppose(A,LS);
module RLS = syz(transpose(LSop));
// RLS is the left syzygy of transposed LSop in Aop
// --- let us return to A and transfer (i.e. oppose)
// all the computed objects back
setring A;
ideal RI = oppose(Aop,RIop); // the right Groebner basis of I
RI; // it differs from the left Groebner basis LI
↳ RI[1]=f
↳ RI[2]=h2-h
↳ RI[3]=eh+e
↳ RI[4]=e2
module RS = oppose(Aop,RSop); // the right syzygy module of I
print(RS);
↳ -ef+3h+6,-f3, -ef2+3fh,-e2f2+4efh+4ef,
↳ e3, e2f2+2efh-6ef+2h2-10h+12,e3f, e4f
 // check: RS is a right syzygy, if T=0:
T = matrix(I)*RS;
T;
↳ T[1,1]=0
↳ T[1,2]=0
↳ T[1,3]=0
↳ T[1,4]=0
module RLS;
RLS = transpose(oppose(Aop,RLS));
// RLS is the right syzygy of a left syzygy of I

```

```
// it is I itself ?
print(RLS);
↳ e2,f
```

## A.8 Applications

### A.8.1 Solving systems of polynomial equations

Here we turn our attention to the probably most popular aspect of the solving problem: given a system of complex polynomial equations with only finitely many solutions, compute floating point approximations for these solutions. This is widely considered as a task for numerical analysis. However, due to rounding errors, purely numerical methods are often unstable in an unpredictable way.

Therefore, in many cases, it is worth investing more computing power to derive additional knowledge on the geometric structure of the set of solutions (not to mention the question of how to decide whether the set of solutions is finite or not). The symbolic-numerical approach to the solving problem combines numerical methods with a symbolic preprocessing.

Depending on whether we want to preserve the multiplicities of the solutions or not, possible goals for a symbolic preprocessing are

- to find another system of generators (for instance, a reduced Groebner basis) for the ideal  $I$  generated by the polynomial equations. Alternatively, find a system of polynomials defining an ideal which has the same radical as  $I$  (see Section A.2 [Computing Groebner and Standard Bases], page 702, resp. [radical], page 829).

In any case, the goal should be to find a system for which a numerical solution can be found more easily and in a more stable way. For systems with a large number of generators, the first step in a SINGULAR computation could be to reduce the number of generators by applying the `interred` command (see Section 5.1.64 [interred], page 199). Another goal might be

- to decompose the system into several smaller (or, at least, more accessible) systems of polynomial equations. Then, the set of solutions of the original system is obtained by taking the union of the sets of solutions of the new systems.

Such a decomposition can be obtained in several ways: for instance, by computing a triangular decomposition (see Section D.8.5 [triang\_lib], page 878) for the ideal  $I$ , or by applying the factorizing Buchberger algorithm (see Section 5.1.34 [facstd], page 177), or by computing a primary decomposition of  $I$  (see Section D.4.26 [primdec\_lib], page 828).

Moreover, the equational modelling of a problem frequently causes unwanted solutions, for instance, zero as a multiple solution. Not only for stability reasons, one is frequently interested to get rid of those. This can be done by computing the saturation of  $I$  with respect to an ideal having the excess components as set of solutions (see [sat], page 812).

The SINGULAR libraries `solve.lib` and `triang.lib` provide several commands for solving systems of polynomial equations (based on a symbolic-numerical approach via Groebner bases, resp. resultants). In the example below, we show some of these commands at work.

```
LIB "solve.lib";
ring r=0,x(1..5),dp;
poly f0= x(1)^3+x(2)^2+x(3)^2+x(4)^2-x(5)^2;
poly f1= x(2)^3+x(1)^2+x(3)^2+x(4)^2-x(5)^2;
poly f2=x(3)^3+x(1)^2+x(2)^2+x(4)^2-x(5)^2;
poly f3=x(4)^2+x(1)^2+x(2)^2+x(3)^2-x(5)^2;
poly f4=x(5)^2+x(1)^2+x(2)^2+x(3)^2;
```

```

ideal i=f0,f1,f2,f3,f4;
ideal si=std(i);
//
// dimension of a solution set (here: 0) can be read from a Groebner bases
// (with respect to any global monomial ordering)
dim(si);
↳ 0
//
// the number of complex solutions (counted with multiplicities) is:
vdim(si);
↳ 108
//
// The given system has a multiple solution at the origin. We use facstd
// to compute equations for the non-zero solutions:
option(redSB);
ideal maxI=maxideal(1);
ideal j=sat(si,maxI)[1]; // output is Groebner basis
vdim(j); // number of non-zero solutions (with mult's)
↳ 76
//
// We compute a triangular decomposition for the ideal I. This requires first
// the computation of a lexicographic Groebner basis (we use the FGLM
// conversion algorithm):
ring R=0,x(1..5),lp;
ideal j=fglm(r,j);
list L=triangMH(j);
size(L); // number of triangular components
↳ 7
L[1]; // the first component
↳ _[1]=x(5)^2+1
↳ _[2]=x(4)^2+2
↳ _[3]=x(3)-1
↳ _[4]=x(2)^2
↳ _[5]=x(1)^2
//
// We compute floating point approximations for the solutions (with 30 digits)
def S=triang_solve(L,30);
↳
↳ // 'triang_solve' created a ring, in which a list rlist of numbers (the
↳ // complex solutions) is stored.
↳ // To access the list of complex solutions, type (if the name R was assign\
ned
↳ // to the return value):
↳ setring R; rlist;
setring S;
size(rlist); // number of different non-zero solutions
↳ 28
rlist[1]; // the first solution
↳ [1]:
↳ 0
↳ [2]:
↳ 0
↳ [3]:

```

```

⇒ 1
⇒ [4]:
⇒ (-I*1.41421356237309504880168872421)
⇒ [5]:
⇒ -I
//
// Alternatively, we could have applied directly the solve command:
setring r;
def T=solve(i,30,1,"nodisplay"); // compute all solutions with mult's
⇒
⇒ // 'solve' created a ring, in which a list SOL of numbers (the complex so\
 lutions)
⇒ // is stored.
⇒ // To access the list of complex solutions, type (if the name R was assign\
 ed
⇒ // to the return value):
⇒ setring R; SOL;
setring T;
size(SOL); // number of different solutions
⇒ 4
SOL[1][1]; SOL[1][2]; // first solution and its multiplicity
⇒ [1]:
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ 1
⇒ [3]:
⇒ 1
⇒ [4]:
⇒ (i*2.44948974278317809819728407471)
⇒ [5]:
⇒ (i*1.73205080756887729352744634151)
⇒ [2]:
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ 1
⇒ [3]:
⇒ 1
⇒ [4]:
⇒ (-i*2.44948974278317809819728407471)
⇒ [5]:
⇒ (i*1.73205080756887729352744634151)
⇒ [3]:
⇒ [1]:
⇒ 1
⇒ [2]:
⇒ 1
⇒ [3]:
⇒ 1
⇒ [4]:
⇒ (i*2.44948974278317809819728407471)
⇒ [5]:

```

```

↳ (-i*1.73205080756887729352744634151)
↳ [4]:
↳ [1]:
↳ 1
↳ [2]:
↳ 1
↳ [3]:
↳ 1
↳ [4]:
↳ (-i*2.44948974278317809819728407471)
↳ [5]:
↳ (-i*1.73205080756887729352744634151)
↳ 1
SOL[size(SOL)]; // solutions of highest multiplicity
↳ [1]:
↳ [1]:
↳ [1]:
↳ 0
↳ [2]:
↳ 0
↳ [3]:
↳ 0
↳ [4]:
↳ 0
↳ [5]:
↳ 0
↳ [2]:
↳ 32
//
// Or, we could remove the multiplicities first, by computing the
// radical:
setring r;
ideal k=std(radical(i));
vdim(k); // number of different complex solutions
↳ 29
def T1=solve(k,30,"nodisplay"); // compute all solutions with mult's
↳
↳ // 'solve' created a ring, in which a list SOL of numbers (the complex so\
 lutions)
↳ // is stored.
↳ // To access the list of complex solutions, type (if the name R was assign\
 ed
↳ // to the return value):
↳ setring R; SOL;
setring T1;
size(SOL); // number of different solutions
↳ 29
SOL[1];
↳ [1]:
↳ 1
↳ [2]:
↳ 1
↳ [3]:

```

```

↳ 1
↳ [4]:
↳ (-i*2.44948974278317809819728407471)
↳ [5]:
↳ (-i*1.73205080756887729352744634151)

```

### A.8.2 AG codes

The library `brnoeth.lib` provides an implementation of the Brill-Noether algorithm for solving the Riemann-Roch problem and applications to Algebraic Geometry codes. The procedures can be applied to plane (singular) curves defined over a prime field of positive characteristic.

```

LIB "brnoeth.lib";
ring s=2,(x,y),lp; // characteristic 2
poly f=x3y+y3+x; // the Klein quartic
list KLEIN=Adj_div(f); // compute the conductor
↳ Computing affine singular points ...
↳ Computing all points at infinity ...
↳ Computing affine singular places ...
↳ Computing singular places at infinity ...
↳ Computing non-singular places at infinity ...
↳ Adjunction divisor computed successfully
↳
↳ The genus of the curve is 3
KLEIN=NSplaces(1..3,KLEIN); // computes places up to degree 3
↳ Computing non-singular affine places of degree 1 ...
↳ Computing non-singular affine places of degree 2 ...
↳ Computing non-singular affine places of degree 3 ...
KLEIN=extcurve(3,KLEIN); // construct Klein quartic over F_8
↳
↳ Total number of rational places : NrRatPl = 24
↳
KLEIN[3]; // display places (degree, number)
↳ [1]:
↳ 1,1
↳ [2]:
↳ 1,2
↳ [3]:
↳ 1,3
↳ [4]:
↳ 2,1
↳ [5]:
↳ 3,1
↳ [6]:
↳ 3,2
↳ [7]:
↳ 3,3
↳ [8]:
↳ 3,4
↳ [9]:
↳ 3,5
↳ [10]:
↳ 3,6
↳ [11]:

```

```

⇒ 3,7
// We define a divisor G of degree 14=6*1+4*2:
intvec G=6,0,0,4,0,0,0,0,0,0,0; // 6 * place #1 + 4 * place #4
// We compute an evaluation code which evaluates at all rational places
// outside the support of G (place #4 is not rational)
intvec D=2..24;
// in D, the number i refers to the i-th element of the list POINTS in
// the ring KLEIN[1][5].
def RR=KLEIN[1][5];
setring RR; POINTS[1]; // the place in the support of G (not in supp(D))
⇒ [1]:
⇒ 0
⇒ [2]:
⇒ 1
⇒ [3]:
⇒ 0
setring s;
def RR=KLEIN[1][4];
⇒ // ** redefining RR (def RR=KLEIN[1][4];) ./examples/AG_codes.sing:18
setring RR;
matrix C=AGcode_L(G,D,KLEIN); // generator matrix for the evaluation AG code
⇒ Forms of degree 5 :
⇒ 21
⇒
⇒ Vector basis successfully computed
⇒
nrows(C);
⇒ 12
ncols(C);
⇒ 23
//
// We can also compute a generator matrix for the residual AG code
matrix CO=AGcode_Omega(G,D,KLEIN);
⇒ Forms of degree 5 :
⇒ 21
⇒
⇒ Vector basis successfully computed
⇒
//
// Preparation for decoding:
// We need a divisor of degree at least 6 whose support is disjoint with the
// support of D:
intvec F=6; // F = 6*point #1
// in F, the i-th entry refers to the i-th element of the list POINTS in
// the ring KLEIN[1][5]
list K=prepSV(G,D,F,KLEIN);
⇒ Forms of degree 5 :
⇒ 21
⇒
⇒ Vector basis successfully computed
⇒
⇒ Forms of degree 4 :
⇒ 15

```

```

↳
↳ Vector basis successfully computed
↳
↳ Forms of degree 4 :
↳ 15
↳
↳ Vector basis successfully computed
↳
K[size(K)][1]; // error-correcting capacity
↳ 3
//
// Encoding and Decoding:
matrix word[1][11]; // a word of length 11 is encoded
word = 1,1,1,1,1,1,1,1,1,1,1;
def y=word*CO; // the code word (length: 23)
matrix disturb[1][23];
disturb[1,1]=1;
disturb[1,10]=a;
disturb[1,12]=1+a;
y=y+disturb; // disturb the code word (3 errors)
def yy=decodeSV(y,K); // error correction
yy-y; // display the error
↳ _[1,1]=1
↳ _[1,2]=0
↳ _[1,3]=0
↳ _[1,4]=0
↳ _[1,5]=0
↳ _[1,6]=0
↳ _[1,7]=0
↳ _[1,8]=0
↳ _[1,9]=0
↳ _[1,10]=(a)
↳ _[1,11]=0
↳ _[1,12]=(a+1)
↳ _[1,13]=0
↳ _[1,14]=0
↳ _[1,15]=0
↳ _[1,16]=0
↳ _[1,17]=0
↳ _[1,18]=0
↳ _[1,19]=0
↳ _[1,20]=0
↳ _[1,21]=0
↳ _[1,22]=0
↳ _[1,23]=0

```



## Appendix B Polynomial data

### B.1 Representation of mathematical objects

SINGULAR distinguishes between objects which do not belong to a ring and those which belong to a specific ring (see Section 3.3 [Rings and orderings], page 30). We comment only on the latter ones.

Internally all ring-dependent objects are polynomials or structures built from polynomials (and some additional information). Note that SINGULAR stores (and hence prints) a polynomial automatically w.r.t. the monomial ordering.

The definition of ideals and matrices, respectively, is straight forward: The user gives a list of polynomials which generate the ideal, resp. which are the entries of the matrix. (The number of rows and columns need to be provided when creating the matrix.)

A vector in SINGULAR is always an element of a free module over the basering. It is given as a list of polynomials in one of the following formats  $[f_1, \dots, f_n]$  or  $f_1 * gen(1) + \dots + f_n * gen(n)$ , where  $gen(i)$  denotes the  $i$ -th canonical generator of a free module (with 1 at index  $i$  and 0 everywhere else). Both forms are equivalent. A vector is internally represented in the second form with the  $gen(i)$  being "special" ring variables, ordered accordingly to the monomial ordering. Therefore, the form  $[f_1, \dots, f_n]$  serves as output only if the monomial ordering gives priority to the component, i.e., is of the form  $(c, \dots)$  (see Section B.2.5 [Module orderings], page 761). However, in any case the procedure `show` from the library `inout.lib` displays the bracket format.

A vector  $v = [f_1, \dots, f_n]$  should always be considered as a column vector in a free module of rank equal to `nrows(v)` where `nrows(v)` is equal to the maximal index  $r$  such that  $f_r \neq 0$ . This is due to the fact, that internally  $v$  is a polynomial in a sparse representation, i.e.,  $f_i * gen(i)$  is not stored if  $f_i = 0$  (for reasons of efficiency), hence the last 0-entries of  $v$  are lost. Only more complex structures are able to keep the rank.

A module  $M$  in SINGULAR is given by a list of vectors  $v_1, \dots, v_k$  which generate the module as a submodule of the free module of rank equal to `nrows(M)` which is the maximum of `nrows(v_i)`.

If one wants to create a module with a larger rank than given by its generators, one has to use the command `attrib(M, "rank", r)` (see Section 5.1.2 [attrib], page 154, Section 5.1.106 [nrows], page 228) or to define a matrix first, then converting it into a module. Modules in SINGULAR are almost the same as matrices, they may be considered as sparse representations of matrices. A module of a matrix is generated by the columns of the matrix and a matrix of a module has as columns the generators of the module. These conversions preserve the rank and the number of generators, resp. the number of rows and columns.

By the above remarks it might appear that SINGULAR is only able to handle submodules of a free module. However, this is not true. SINGULAR can compute with any finitely generated module over the basering  $R$ . Such a module, say  $N$ , is not represented by its generators but by its (generators and) relations. This means that  $N = R^n/M$  where  $n$  is the number of generators of  $N$  and  $M \subseteq R^n$  is the module of relations. In other words, defining a module  $M$  as a submodule of a free module  $R^n$  can also be considered as the definition of  $N = R^n/M$ .

Note that most functions, when applied to a module  $M$ , really deal with  $M$ . However, there are some functions which deal with  $N = R^n/M$  instead of  $M$ .

For example, `std(M)` computes a standard basis of  $M$  (and thus gives another representation of  $N$  as  $N = R^n/std(M)$ ). However, `dim(M)`, resp. `vdim(M)`, return  $\dim(R^n/M)$ , resp.  $\dim_k(R^n/M)$  (if  $M$  is given by a standard basis).

The function `syz(M)` returns the first syzygy module of  $M$ , i.e., the module of relations of the given generators of  $M$  which is equal to the second syzygy module of  $N$ . Refer to the description of each

function in Section 5.1 [Functions], page 154 to get information which module the function deals with.

The numbering in `res` and other commands for computing resolutions refers to a resolution of  $N = R^n/M$  (see [res], page 785; Section C.3 [Syzygies and resolutions], page 767).

It is possible to compute in any field which is a valid ground field in SINGULAR. For doing so, one has to define a ring with the desired ground field and at least one variable. The elements of the field are of type number, but may also be considered as polynomials (of degree 0). Large computations should be faster if the elements of the field are defined as numbers.

The above remarks do also apply to quotient rings. Polynomial data are stored internally in the same manner, the only difference is that this polynomial representation is in general not unique. `reduce(f, std(O))` computes a normal form of a polynomial `f` in a quotient ring (cf. Section 5.1.129 [reduce], page 246).

## B.2 Monomial orderings

### B.2.1 Introduction to orderings

SINGULAR offers a great variety of monomial orderings which provide an enormous functionality, if used diligently. However, this flexibility might also be confusing for the novice user. Therefore, we recommend to those not familiar with monomial orderings to generally use the ordering `dp` for computations in the polynomial ring  $K[x_1, \dots, x_n]$ , resp. `ds` for computations in the localization  $\text{Loc}_{(x)}K[x_1, \dots, x_n]$ .

For inhomogeneous input ideals, standard (resp. groebner) bases computations are generally faster with the orderings  $\text{Wp}(w_1, \dots, w_n)$  (resp.  $\text{Ws}(w_1, \dots, w_n)$ ) if the input is quasihomogenous w.r.t. the weights  $w_1, \dots, w_n$  of  $x_1, \dots, x_n$ .

If the output needs to be "triangular" (resp. "block-triangular"), the lexicographical ordering `lp` (resp. lexicographical block-orderings) need to be used. However, these orderings usually result in much less efficient computations.

### B.2.2 General definitions for orderings

A monomial ordering (term ordering) on  $K[x_1, \dots, x_n]$  is a total ordering  $<$  on the set of monomials (power products)  $\{x^\alpha \mid \alpha \in \mathbf{N}^n\}$  which is compatible with the natural semigroup structure, i.e.,  $x^\alpha < x^\beta$  implies  $x^\gamma x^\alpha < x^\gamma x^\beta$  for any  $\gamma \in \mathbf{N}^n$ . We do not require  $<$  to be a wellordering. See the literature cited in Section C.9 [References], page 783.

It is known that any monomial ordering can be represented by a matrix  $M$  in  $GL(n, R)$ , but, of course, only integer coefficients are of relevance in practice.

Global orderings are wellorderings (i.e.,  $1 < x_i$  for each variable  $x_i$ ), local orderings satisfy  $1 > x_i$  for each variable. If some variables are ordered globally and others locally we call it a mixed ordering. Local or mixed orderings are not wellorderings.

Let  $K$  be the ground field,  $x = (x_1, \dots, x_n)$  the variables and  $<$  a monomial ordering, then  $\text{Loc } K[x]$  denotes the localization of  $K[x]$  with respect to the multiplicatively closed set

$$\{1 + g \mid g = 0 \text{ or } g \in K[x] \setminus \{0\} \text{ and } L(g) < 1\}.$$

Here,  $L(g)$  denotes the leading monomial of  $g$ , i.e., the biggest monomial of  $g$  with respect to  $<$ . The result of any computation which uses standard basis computations has to be interpreted in  $\text{Loc } K[x]$ .

Note that the definition of a ring includes the definition of its monomial ordering (see Section 3.3 [Rings and orderings], page 30). SINGULAR offers the monomial orderings described in the following sections.

### B.2.3 Global orderings

For all these orderings, we have  $\text{Loc } K[x] = K[x]$

- lp: lexicographical ordering:  
 $x^\alpha < x^\beta \Leftrightarrow \exists 1 \leq i \leq n : \alpha_1 = \beta_1, \dots, \alpha_{i-1} = \beta_{i-1}, \alpha_i < \beta_i.$
- rp: reverse lexicographical ordering:  
 $x^\alpha < x^\beta \Leftrightarrow \exists 1 \leq i \leq n : \alpha_n = \beta_n, \dots, \alpha_{i+1} = \beta_{i+1}, \alpha_i < \beta_i.$
- dp: degree reverse lexicographical ordering:  
 let  $\deg(x^\alpha) = \alpha_1 + \dots + \alpha_n$ , then  $x^\alpha < x^\beta \Leftrightarrow \deg(x^\alpha) < \deg(x^\beta)$  or  
 $\deg(x^\alpha) = \deg(x^\beta)$  and  $\exists 1 \leq i \leq n : \alpha_n = \beta_n, \dots, \alpha_{i+1} = \beta_{i+1}, \alpha_i > \beta_i.$
- Dp: degree lexicographical ordering:  
 let  $\deg(x^\alpha) = \alpha_1 + \dots + \alpha_n$ , then  $x^\alpha < x^\beta \Leftrightarrow \deg(x^\alpha) < \deg(x^\beta)$  or  
 $\deg(x^\alpha) = \deg(x^\beta)$  and  $\exists 1 \leq i \leq n : \alpha_1 = \beta_1, \dots, \alpha_{i-1} = \beta_{i-1}, \alpha_i < \beta_i.$
- wp: weighted reverse lexicographical ordering:  
 let  $w_1, \dots, w_n$  be positive integers. Then  $\text{wp}(w_1, \dots, w_n)$  is defined as **dp** but with  
 $\deg(x^\alpha) = w_1\alpha_1 + \dots + w_n\alpha_n.$
- Wp: weighted lexicographical ordering:  
 let  $w_1, \dots, w_n$  be positive integers. Then  $\text{Wp}(w_1, \dots, w_n)$  is defined as **Dp** but with  
 $\deg(x^\alpha) = w_1\alpha_1 + \dots + w_n\alpha_n.$

### B.2.4 Local orderings

For **ls**, **ds**, **Ds** and, if the weights are positive integers, also for **ws** and **Ws**, we have  $\text{Loc } K[x] = K[x]_{(x)}$ , the localization of  $K[x]$  at the maximal ideal  $(x) = (x_1, \dots, x_n)$ .

- ls: negative lexicographical ordering:  
 $x^\alpha < x^\beta \Leftrightarrow \exists 1 \leq i \leq n : \alpha_1 = \beta_1, \dots, \alpha_{i-1} = \beta_{i-1}, \alpha_i > \beta_i.$
- ds: negative degree reverse lexicographical ordering:  
 let  $\deg(x^\alpha) = \alpha_1 + \dots + \alpha_n$ , then  $x^\alpha < x^\beta \Leftrightarrow \deg(x^\alpha) > \deg(x^\beta)$  or  
 $\deg(x^\alpha) = \deg(x^\beta)$  and  $\exists 1 \leq i \leq n : \alpha_n = \beta_n, \dots, \alpha_{i+1} = \beta_{i+1}, \alpha_i > \beta_i.$
- Ds: negative degree lexicographical ordering:  
 let  $\deg(x^\alpha) = \alpha_1 + \dots + \alpha_n$ , then  $x^\alpha < x^\beta \Leftrightarrow \deg(x^\alpha) > \deg(x^\beta)$  or  
 $\deg(x^\alpha) = \deg(x^\beta)$  and  $\exists 1 \leq i \leq n : \alpha_1 = \beta_1, \dots, \alpha_{i-1} = \beta_{i-1}, \alpha_i < \beta_i.$
- ws: (general) weighted reverse lexicographical ordering:  
 $\text{ws}(w_1, \dots, w_n)$ ,  $w_1$  a nonzero integer,  $w_2, \dots, w_n$  any integer (including 0), is defined as **ds** but with  $\deg(x^\alpha) = w_1\alpha_1 + \dots + w_n\alpha_n.$
- Ws: (general) weighted lexicographical ordering:  
 $\text{Ws}(w_1, \dots, w_n)$ ,  $w_1$  a nonzero integer,  $w_2, \dots, w_n$  any integer (including 0), is defined as **Ds** but with  $\deg(x^\alpha) = w_1\alpha_1 + \dots + w_n\alpha_n.$

### B.2.5 Module orderings

SINGULAR offers also orderings on the set of “monomials”  $\{x^a e_i \mid a \in N^n, 1 \leq i \leq r\}$  in  $\text{Loc } K[x]^r = \text{Loc } K[x]e_1 + \dots + \text{Loc } K[x]e_r$ , where  $e_1, \dots, e_r$  denote the canonical generators of  $\text{Loc } K[x]^r$ , the  $r$ -fold direct sum of  $\text{Loc } K[x]$ . (The function  $\text{gen}(i)$  yields  $e_i$ ).

We have two possibilities: either to give priority to the component of a vector in  $\text{Loc } K[x]^r$  or (which is the default in SINGULAR) to give priority to the coefficients. The orderings  $(<, c)$  and

$(\langle, C)$  give priority to the coefficients; whereas  $(c, \langle)$  and  $(C, \langle)$  give priority to the components. Let  $\langle$  be any of the monomial orderings of  $\text{Loc } K[x]$  as above.

$(\langle, C)$ :  $\langle_m = (\langle, C)$  denotes the module ordering (giving priority to the coefficients):  
 $x^\alpha e_i \langle_m x^\beta e_j \Leftrightarrow x^\alpha < x^\beta$  or  $(x^\alpha = x^\beta$  and  $i < j)$ .

**Example:**

```
ring r = 0, (x,y,z), ds;
// the same as ring r = 0, (x,y,z), (ds, C);
[x+y2,z3+xy];
↦ x*gen(1)+xy*gen(2)+y2*gen(1)+z3*gen(2)
[x,x,x];
↦ x*gen(3)+x*gen(2)+x*gen(1)
```

$(C, \langle)$ :  $\langle_m = (C, \langle)$  denotes the module ordering (giving priority to the component):  
 $x^\alpha e_i \langle_m x^\beta e_j \Leftrightarrow i < j$  or  $(i = j$  and  $x^\alpha < x^\beta)$ .

**Example:**

```
ring r = 0, (x,y,z), (C,lp);
[x+y2,z3+xy];
↦ xy*gen(2)+z3*gen(2)+x*gen(1)+y2*gen(1)
[x,x,x];
↦ x*gen(3)+x*gen(2)+x*gen(1)
```

$(\langle, c)$ :  $\langle_m = (\langle, c)$  denotes the module ordering (giving priority to the coefficients):  
 $x^\alpha e_i \langle_m x^\beta e_j \Leftrightarrow x^\alpha < x^\beta$  or  $(x^\alpha = x^\beta$  and  $i > j)$ .

**Example:**

```
ring r = 0, (x,y,z), (lp,c);
[x+y2,z3+xy];
↦ xy*gen(2)+x*gen(1)+y2*gen(1)+z3*gen(2)
[x,x,x];
↦ x*gen(1)+x*gen(2)+x*gen(3)
```

$(c, \langle)$ :  $\langle_m = (c, \langle)$  denotes the module ordering (giving priority to the component):  
 $x^\alpha e_i \langle_m x^\beta e_j \Leftrightarrow i > j$  or  $(i = j$  and  $x^\alpha < x^\beta)$ .

**Example:**

```
ring r = 0, (x,y,z), (c,lp);
[x+y2,z3+xy];
↦ [x+y2,xy+z3]
[x,x,x];
↦ [x,x,x]
```

The output of a vector  $v$  in  $K[x]^r$  with components  $v_1, \dots, v_r$  has the format  $v_1 * \text{gen}(1) + \dots + v_r * \text{gen}(r)$  (up to permutation) unless the ordering starts with  $c$ . In this case a vector is written as  $[v_1, \dots, v_r]$ . In all cases SINGULAR can read input in both formats.

## B.2.6 Matrix orderings

Let  $M$  be an invertible  $(n \times n)$ -matrix with integer coefficients and  $M_1, \dots, M_n$  the rows of  $M$ .

The  $M$ -ordering  $\langle$  is defined as follows:

$$x^a < x^b \Leftrightarrow \exists 1 \leq i \leq n : M_1 a = M_1 b, \dots, M_{i-1} a = M_{i-1} b \text{ and } M_i a < M_i b.$$

Thus,  $x^a < x^b$  if and only if  $Ma$  is smaller than  $Mb$  with respect to the lexicographical ordering.

The following matrices represent (for 3 variables) the global and local orderings defined above (note that the matrix is not uniquely determined by the ordering):

$$\begin{aligned}
 \text{lp: } & \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix} & \text{dp: } & \begin{pmatrix} 1 & 1 & 1 \\ 0 & 0 & -1 \\ 0 & -1 & 0 \end{pmatrix} & \text{Dp: } & \begin{pmatrix} 1 & 1 & 1 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \end{pmatrix} \\
 \text{wp(1,2,3): } & \begin{pmatrix} 1 & 2 & 3 \\ 0 & 0 & -1 \\ 0 & -1 & 0 \end{pmatrix} & \text{Wp(1,2,3): } & \begin{pmatrix} 1 & 2 & 3 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \end{pmatrix} \\
 \text{ls: } & \begin{pmatrix} -1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & -1 \end{pmatrix} & \text{ds: } & \begin{pmatrix} -1 & -1 & -1 \\ 0 & 0 & -1 \\ 0 & -1 & 0 \end{pmatrix} & \text{Ds: } & \begin{pmatrix} -1 & -1 & -1 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \end{pmatrix} \\
 \text{ws(1,2,3): } & \begin{pmatrix} -1 & -2 & -3 \\ 0 & 0 & -1 \\ 0 & -1 & 0 \end{pmatrix} & \text{Ws(1,2,3): } & \begin{pmatrix} -1 & -2 & -3 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \end{pmatrix}
 \end{aligned}$$

Product orderings (see next section) represented by a matrix:

$$\begin{aligned}
 (\text{dp}(3), \text{wp}(1,2,3)): & \begin{pmatrix} 1 & 1 & 1 & 0 & 0 & 0 \\ 0 & 0 & -1 & 0 & 0 & 0 \\ 0 & -1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 2 & 3 \\ 0 & 0 & 0 & 0 & 0 & -1 \\ 0 & 0 & 0 & 0 & -1 & 0 \end{pmatrix} \\
 (\text{Dp}(3), \text{ds}(3)): & \begin{pmatrix} 1 & 1 & 1 & 0 & 0 & 0 \\ 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & -1 & -1 & -1 \\ 0 & 0 & 0 & 0 & 0 & -1 \\ 0 & 0 & 0 & 0 & -1 & 0 \end{pmatrix}
 \end{aligned}$$

Orderings with extra weight vector (see below) represented by a matrix:

$$\begin{aligned}
 (\text{dp}(3), \text{a}(1,2,3), \text{dp}(3)): & \begin{pmatrix} 1 & 1 & 1 & 0 & 0 & 0 \\ 0 & 0 & -1 & 0 & 0 & 0 \\ 0 & -1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 2 & 3 \\ 0 & 0 & 0 & 1 & 1 & 1 \\ 0 & 0 & 0 & 0 & 0 & -1 \\ 0 & 0 & 0 & 0 & -1 & 0 \end{pmatrix} \\
 (\text{a}(1,2,3,4,5), \text{Dp}(3), \text{ds}(3)): & \begin{pmatrix} 1 & 2 & 3 & 4 & 5 & 0 \\ 1 & 1 & 1 & 0 & 0 & 0 \\ 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & -1 & -1 & -1 \\ 0 & 0 & 0 & 0 & 0 & -1 \\ 0 & 0 & 0 & 0 & -1 & 0 \end{pmatrix}
 \end{aligned}$$

**Example:**

$$\text{ring } r = 0, (x,y,z), M(1, 0, 0, \quad 0, 1, 0, \quad 0, 0, 1);$$

which may also be written as:

```

intmat m[3][3]=1, 0, 0, 0, 1, 0, 0, 0, 1;
m;
↳ 1,0,0,
↳ 0,1,0,
↳ 0,0,1

```

```

ring r = 0, (x,y,z), M(m);
r;
↳ // coefficients: QQ
↳ // number of vars : 3
↳ // block 1 : ordering M
↳ // : names x y z
↳ // : weights 1 0 0
↳ // : weights 0 1 0
↳ // : weights 0 0 1
↳ // block 2 : ordering C

```

If the ring has  $n$  variables and the matrix does not contain  $n \times n$  entries, an error message is given.

**WARNING:** SINGULAR does not check whether the matrix has full rank. In such a case some computations might not terminate, others may not give a sensible result.

Having these matrix orderings SINGULAR can compute standard bases for any monomial ordering which is compatible with the natural semigroup structure. In practice the global and local orderings together with block orderings should be sufficient in most cases. These orderings are faster than the corresponding matrix orderings, since evaluating a matrix product is time consuming.

## B.2.7 Product orderings

Let  $x = (x_1, \dots, x_n)$  and  $y = (y_1, \dots, y_m)$  be two ordered sets of variables,  $<_1$  a monomial ordering on  $K[x]$  and  $<_2$  a monomial ordering on  $K[y]$ . The product ordering (or block ordering)  $< := (<_1, <_2)$  on  $K[x, y]$  is the following:

$$x^a y^b < x^A y^B \Leftrightarrow x^a <_1 x^A \text{ or } (x^a = x^A \text{ and } y^b <_2 y^B).$$

Inductively one defines the product ordering of more than two monomial orderings.

In SINGULAR, any of the above global orderings, local orderings or matrix orderings may be combined (in an arbitrary manner and length) to a product ordering. E.g., `(lp(3), M(1, 2, 3, 1, 1, 1, 1, 0, 0), ds(4), ws(1,2,3))` defines: `lp` on the first 3 variables, the matrix ordering `M(1, 2, 3, 1, 1, 1, 1, 0, 0)` on the next 3 variables, `ds` on the next 4 variables and `ws(1,2,3)` on the last 3 variables.

## B.2.8 Extra weight vector

`a(w1, ..., wn)`,  $w_1, \dots, w_n$  any integers (including 0), defines  $\deg(x^\alpha) = w_1 \alpha_1 + \dots + w_n \alpha_n$  and

$$\deg(x^\alpha) < \deg(x^\beta) \Rightarrow x^\alpha < x^\beta,$$

$$\deg(x^\alpha) > \deg(x^\beta) \Rightarrow x^\alpha > x^\beta.$$

An extra weight vector does not define a monomial ordering by itself: it can only be used in combination with other orderings to insert an extra line of weights into the ordering matrix.

**Example:**

```

ring r = 0, (x,y,z), (a(1,2,3), wp(4,5,2));
ring s = 0, (x,y,z), (a(1,2,3), dp);
ring q = 0, (a,b,c,d), (lp(1), a(1,2,3), ds);

```

### B.2.9 Pseudo ordering L

`L(max_exponent)` is not an ordering but sets the maximal allowed exponent for polynomial in this ring. The default is 32767. The current value for a ring is reflected in the attribute "maxExp". This attribute is also set (and acknowledged) for the list constructed by `ringlist` and the construction of a ring from such a list.

## Appendix C Mathematical background

This chapter introduces some of the mathematical notions and definitions used throughout the manual. It is mostly a collection of the most prominent definitions and properties. For details, please, refer to articles or text books (see Section C.9 [References], page 783).

### C.1 Standard bases

#### Definition

Let  $R = \text{Loc}_{<}K[x]$  and let  $I$  be a submodule of  $R^r$ . Note that for  $r=1$  this means that  $I$  is an ideal in  $R$ . Denote by  $L(I)$  the submodule of  $R^r$  generated by the leading terms of elements of  $I$ , i.e. by  $\{L(f) \mid f \in I\}$ . Then  $f_1, \dots, f_s \in I$  is called a **standard basis** of  $I$  if  $L(f_1), \dots, L(f_s)$  generate  $L(I)$ .

A standard basis is **minimal** if  $\forall i : (f_1, \dots, f_{i-1}, f_{i+1}, \dots, f_s) \neq I$ .

A minimal standard basis is **completely reduced** if  $\forall i : \text{reduce}(f_i, (f_1, \dots, f_{i-1}, f_{i+1}, \dots, f_s)) = f_i$

#### Properties

normal form:

A function  $\text{NF} : R^r \times \{G \mid G \text{ a standard basis}\} \rightarrow R^r, (p, G) \mapsto \text{NF}(p|G)$ , is called a **normal form** if for any  $p \in R^r$  and any standard basis  $G$  the following holds: if  $\text{NF}(p|G) \neq 0$  then  $L(g)$  does not divide  $L(\text{NF}(p|G))$  for all  $g \in G$ . The function may also be applied to any generating set of an ideal: the result is then not uniquely defined.

$\text{NF}(p|G)$  is called a **normal form of  $p$  with respect to  $G$**

ideal membership:

For a standard basis  $G$  of  $I$  the following holds:  $f \in I$  if and only if  $\text{NF}(f, G) = 0$ .

Hilbert function:

Let  $I \subseteq K[x]^r$  be a homogeneous module, then the Hilbert function  $H_I$  of  $I$  (see below) and the Hilbert function  $H_{L(I)}$  of the leading module  $L(I)$  coincide, i.e.,  $H_I = H_{L(I)}$ .

### C.2 Hilbert function

Let  $M = \bigoplus_{i \in \mathbb{Z}} M_i$  be a graded module over  $K[x_1, \dots, x_n]$  with respect to weights  $(w_1, \dots, w_n)$ . The **Hilbert function** of  $M$ ,  $H_M$ , is defined (on the integers) by

$$H_M(k) := \dim_K M_k.$$

The **Hilbert-Poincare series** of  $M$  is the power series

$$\text{HP}_M(t) := \sum_{i=-\infty}^{\infty} H_M(i)t^i = \sum_{i=-\infty}^{\infty} \dim_K M_i \cdot t^i.$$

It turns out that  $\text{HP}_M(t)$  can be written in two useful ways for weights  $(1, \dots, 1)$ :

$$\text{HP}_M(t) = \frac{Q(t)}{(1-t)^n} = \frac{P(t)}{(1-t)^{\dim(M)}}$$

where  $Q(t)$  and  $P(t)$  are polynomials in  $\mathbb{Z}[t]$ .  $Q(t)$  is called the **first Hilbert series**, and  $P(t)$  the **second Hilbert series**. If  $P(t) = \sum_{k=0}^N a_k t^k$ , and  $d = \dim(M)$ , then  $H_M(s) = \sum_{k=0}^N a_k \binom{d+s-k-1}{d-1}$  (the



**Hilbert polynomial**) for  $s \geq N$ .

Generalizing this to quasihomogeneous modules we get

$$\text{HP}_M(t) = \frac{Q(t)}{\prod_{i=1}^n (1 - t^{w_i})}$$

where  $Q(t)$  is a polynomial in  $\mathbf{Z}[t]$ .  $Q(t)$  is called the **first (weighted) Hilbert series** of  $M$ .

### C.3 Syzygies and resolutions

#### Syzygies

Let  $R$  be a quotient of  $\text{Loc}_{<}K[\underline{x}]$  and let  $I = (g_1, \dots, g_s)$  be a submodule of  $R^r$ . Then the **module of syzygies** (or **1st syzygy module, module of relations**) of  $I$ ,  $\text{syz}(I)$ , is defined to be the kernel of the map  $R^s \rightarrow R^r$ ,  $\sum_{i=1}^s w_i e_i \mapsto \sum_{i=1}^s w_i g_i$ .

The  **$k$ -th syzygy module** is defined inductively to be the module of syzygies of the  $(k-1)$ -st syzygy module.

Note, that the syzygy modules of  $I$  depend on a choice of generators  $g_1, \dots, g_s$ . But one can show that they depend on  $I$  uniquely up to direct summands.

**Example:**

```
ring R= 0,(u,v,x,y,z),dp;
ideal i=ux, vx, uy, vy;
print(syz(i));
↳ -y,0, -v,0,
↳ 0, -y,u, 0,
↳ x, 0, 0, -v,
↳ 0, x, 0, u
```

#### Free resolutions

Let  $I = (g_1, \dots, g_s) \subseteq R^r$  and  $M = R^r/I$ . A **free resolution** of  $M$  is a long exact sequence

$$\dots \longrightarrow F_2 \xrightarrow{A_2} F_1 \xrightarrow{A_1} F_0 \longrightarrow M \longrightarrow 0,$$

where the columns of the matrix  $A_1$  generate  $I$ . Note that resolutions need not to be finite (i.e., of finite length). The Hilbert Syzygy Theorem states that for  $R = \text{Loc}_{<}K[\underline{x}]$  there exists a ("minimal") resolution of length not exceeding the number of variables.

**Example:**

```
ring R= 0,(u,v,x,y,z),dp;
ideal I = ux, vx, uy, vy;
resolution resI = mres(I,0); resI;
↳ 1 4 4 1
↳ R <-- R <-- R <-- R
↳
↳ 0 1 2 3
↳
// The matrix A_1 is given by
print(matrix(resI[1]));
```

```

↳ vy,uy,vx,ux
 // We see that the columns of A_1 generate I.
 // The matrix A_2 is given by
 print(matrix(resI[3]));
↳ u,
↳ -v,
↳ -x,
↳ y

```

### Betti numbers and regularity

Let  $R$  be a graded ring (e.g.,  $R = \text{Loc}_{<}K[x]$ ) and let  $I \subset R^r$  be a graded submodule. Let

$$R^r = \bigoplus_a R \cdot e_{a,0} \xleftarrow{A_1} \bigoplus_a R \cdot e_{a,1} \xleftarrow{\dots} \bigoplus_a R \cdot e_{a,n} \xleftarrow{\dots} 0$$

be a minimal free resolution of  $R^r/I$  considered with homogeneous maps of degree 0. Then the **graded Betti number**  $b_{i,j}$  of  $R^r/I$  is the minimal number of generators  $e_{a,j}$  in degree  $i + j$  of the  $j$ -th syzygy module of  $R^r/I$  (i.e., the  $(j - 1)$ -st syzygy module of  $I$ ). Note that, by definition, the 0-th syzygy module of  $R^r/I$  is  $R^r$  and the 1st syzygy module of  $R^r/I$  is  $I$ .

The **regularity** of  $I$  is the smallest integer  $s$  such that

$$\deg(e_{a,j}) \leq s + j - 1 \quad \text{for all } j.$$

**Example:**

```

ring R= 0,(u,v,x,y,z),dp;
ideal I = ux, vx, uy, vy;
resolution resI = mres(I,0); resI;
↳ 1 4 4 1
↳ R <-- R <-- R <-- R
↳
↳ 0 1 2 3
↳
 // the betti number:
 print(betti(resI), "betti");
↳ 0 1 2 3
↳ -----
↳ 0: 1 - - -
↳ 1: - 4 4 1
↳ -----
↳ total: 1 4 4 1
↳
 // the regularity:
 regularity(resI);
↳ 2

```

### C.4 Characteristic sets

Let  $<$  be the lexicographical ordering on  $R = K[x_1, \dots, x_n]$  with  $x_1 < \dots < x_n$ . For  $f \in R$  let  $\text{lvar}(f)$  (the leading variable of  $f$ ) be the largest variable in  $f$ , i.e., if  $f = a_s(x_1, \dots, x_{k-1})x_k^s + \dots + a_0(x_1, \dots, x_{k-1})$  for some  $k \leq n$  then  $\text{lvar}(f) = x_k$ .

Moreover, let  $\text{ini}(f) := a_s(x_1, \dots, x_{k-1})$ . The pseudoremainder  $r = \text{prem}(g, f)$  of  $g$  with respect to  $f$  is defined by the equality  $\text{ini}(f)^a \cdot g = qf + r$  with  $\deg_{\text{lvar}(f)}(r) < \deg_{\text{lvar}(f)}(f)$  and  $a$  minimal.

A set  $T = \{f_1, \dots, f_r\} \subset R$  is called triangular if  $\text{lvar}(f_1) < \dots < \text{lvar}(f_r)$ . Moreover, let  $U \subset T$ , then  $(T, U)$  is called a triangular system, if  $T$  is a triangular set such that  $\text{ini}(T)$  does not vanish on  $V(T) \setminus V(U)(=: V(T \setminus U))$ .

$T$  is called irreducible if for every  $i$  there are no  $d_i, f'_i, f''_i$  such that

$$\begin{aligned} \text{lvar}(d_i) < \text{lvar}(f_i) = \text{lvar}(f'_i) = \text{lvar}(f''_i), \\ 0 \notin \text{prem}(\{d_i, \text{ini}(f'_i), \text{ini}(f''_i)\}, \{f_1, \dots, f_{i-1}\}), \\ \text{prem}(d_i f_i - f'_i f''_i, \{f_1, \dots, f_{i-1}\}) = 0. \end{aligned}$$

Furthermore,  $(T, U)$  is called irreducible if  $T$  is irreducible.

The main result on triangular sets is the following: Let  $G = \{g_1, \dots, g_s\} \subset R$ , then there are irreducible triangular sets  $T_1, \dots, T_l$  such that  $V(G) = \bigcup_{i=1}^l (V(T_i) \setminus I_i)$  where  $I_i = \{\text{ini}(f) \mid f \in T_i\}$ . Such a set  $\{T_1, \dots, T_l\}$  is called an **irreducible characteristic series** of the ideal  $(G)$ .

**Example:**

```
ring R= 0, (x,y,z,u), dp;
ideal i=-3zu+y2-2x+2,
 -3x2u-4yz-6xz+2y2+3xy,
 -3z2u-xu+y2z+y;
print(char_series(i));
→ _[1,1], 3x2z-y2+2yz, 3x2u-3xy-2y2+2yu,
→ x, -y+2z, -2y2+3yu-4
```

### C.5 Gauss-Manin connection

Let  $f: (C^{n+1}, 0) \rightarrow (C, 0)$  be a complex isolated hypersurface singularity given by a polynomial with algebraic coefficients which we also denote by  $f$ . Let  $O = C[x_0, \dots, x_n]_{(x_0, \dots, x_n)}$  be the local ring at the origin and  $J_f$  the Jacobian ideal of  $f$ .

A **Milnor representative** of  $f$  defines a differentiable fibre bundle over the punctured disc with fibres of homotopy type of  $\mu$   $n$ -spheres. The  $n$ -th cohomology bundle is a flat vector bundle of dimension  $n$  and carries a natural flat connection with covariant derivative  $\partial_t$ . The **monodromy operator** is the action of a positively oriented generator of the fundamental group of the punctured disc on the Milnor fibre. Sections in the cohomology bundle of **moderate growth** at 0 form a regular  $D = C\{t\}[\partial_t]$ -module  $G$ , the **Gauss-Manin connection**.

By integrating along flat multivalued families of cycles, one can consider fibrewise global holomorphic differential forms as elements of  $G$ . This factors through an inclusion of the **Brieskorn lattice**  $H'' := \Omega_{C^{n+1}, 0}^{n+1}/df \wedge d\Omega_{C^{n+1}, 0}^{n-1}$  in  $G$ .

The  $D$ -module structure defines the **V-filtration**  $V$  on  $G$  by  $V^\alpha := \sum_{\beta \geq \alpha} C\{t\} \ker(t\partial_t - \beta)^{n+1}$ . The Brieskorn lattice defines the **Hodge filtration**  $F$  on  $G$  by  $F_k = \partial_t^k H''$  which comes from the **mixed Hodge structure** on the Milnor fibre. Note that  $F_{-1} = H'$ .

The induced V-filtration on the Brieskorn lattice determines the **singularity spectrum**  $Sp$  by  $Sp(\alpha) := \dim_C Gr_V^\alpha Gr_0^F G$ . The spectrum consists of  $\mu$  rational numbers  $\alpha_1, \dots, \alpha_\mu$  such that  $e^{2\pi i \alpha_1}, \dots, e^{2\pi i \alpha_\mu}$  are the eigenvalues of the monodromy. These **spectral numbers** lie in the open interval  $(-1, n)$ , symmetric about the midpoint  $(n - 1)/2$ .

The spectrum is constant under  $\mu$ -constant deformations and has the following semicontinuity property: The number of spectral numbers in an interval  $(a, a + 1]$  of all singularities of a small deformation of  $f$  is greater than or equal to that of  $f$  in this interval. For semiquasihomogeneous singularities, this also holds for intervals of the form  $(a, a + 1)$ .

Two given isolated singularities  $f$  and  $g$  determine two spectra and from these spectra we get an integer. This integer is the maximal positive integer  $k$  such that the semicontinuity holds for the spectrum of  $f$  and  $k$  times the spectrum of  $g$ . These numbers give bounds for the maximal number of isolated singularities of a specific type on a hypersurface  $X \subset P^n$  of degree  $d$ : such a hypersurface has a smooth hyperplane section, and the complement is a small deformation of a cone over this hyperplane section. The cone itself being a  $\mu$ -constant deformation of  $x_0^d + \dots + x_n^d = 0$ , the singularities are bounded by the spectrum of  $x_0^d + \dots + x_n^d$ .

Using the library `gmssing.lib` one can compute the **monodromy**, the V-filtration on  $H''/H'$ , and the spectrum.

Let us consider as an example  $f = x^5 + x^2y^2 + y^5$ . First, we compute a matrix  $M$  such that  $\exp(2\pi iM)$  is a monodromy matrix of  $f$  and the Jordan normal form of  $M$  :

```
LIB "mondromy.lib";
ring R=0,(x,y),ds;
poly f=x5+x2y2+y5;
matrix M=mondromyB(f);
print(M);
⇒ 11/10,0, 0, 0, 0, 0,-1/4,0, 0, 0, 0,
⇒ 0, 13/10,0, 0, 0, 0,0, 15/8,0, 0, 0,
⇒ 0, 0, 13/10,0, 0, 0,0, 0, 15/8,0, 0,
⇒ 0, 0, 0, 11/10,-1/4,0,0, 0, 0, 0, 0,
⇒ 0, 0, 0, 0, 9/10,0,0, 0, 0, 0, 0,
⇒ 0, 0, 0, 0, 0, 1,0, 0, 0, 0, 3/5,
⇒ 0, 0, 0, 0, 0, 0,9/10,0, 0, 0, 0,
⇒ 0, 0, 0, 0, 0, 0,0, 7/10,0, 0, 0,
⇒ 0, 0, 0, 0, 0, 0,0, 0, 7/10,0, 0,
⇒ 0, 0, 0, 0, 0, 0,0, 0, 0, 1, -2/5,
⇒ 0, 0, 0, 0, 0, 0,0, 0, 0, 5/8,0
```

Now, we compute the V-filtration on  $H''/H'$  and the spectrum:

```
LIB "gmssing.lib";
ring R=0,(x,y),ds;
poly f=x5+x2y2+y5;
list l=vfilt(f);
print(l[1]); // spectral numbers
⇒ -1/2,
⇒ -3/10,
⇒ -1/10,
⇒ 0,
⇒ 1/10,
⇒ 3/10,
⇒ 1/2
print(l[2]); // corresponding multiplicities
⇒ 1,
⇒ 2,
⇒ 2,
⇒ 1,
⇒ 2,
⇒ 2,
⇒ 1
print(l[3]); // vector space of i-th graded part
⇒ [1]:
⇒ _[1]=gen(11)
```

```

↳ [2] :
↳ _[1]=gen(10)
↳ _[2]=gen(6)
↳ [3] :
↳ _[1]=gen(9)
↳ _[2]=gen(4)
↳ [4] :
↳ _[1]=gen(5)
↳ [5] :
↳ _[1]=gen(3)
↳ _[2]=gen(8)
↳ [6] :
↳ _[1]=gen(2)
↳ _[2]=gen(7)
↳ [7] :
↳ _[1]=gen(1)
print(l[4]); // monomial vector space basis of H''/s*H''
↳ y5,
↳ y4,
↳ y3,
↳ y2,
↳ xy,
↳ y,
↳ x4,
↳ x3,
↳ x2,
↳ x,
↳ 1
print(l[5]); // standard basis of Jacobian ideal
↳ 2x2y+5y4,
↳ 5x5-5y5,
↳ 2xy2+5x4,
↳ 10y6+25x3y4

```

Here  $l[1]$  contains the spectral numbers,  $l[2]$  the corresponding multiplicities,  $l[3]$  a  $C$ -basis of the  $V$ -filtration on  $H''/H'$  in terms of the monomial basis of  $O/J_f \cong H''/H'$  in  $l[4]$  (separated by degree).

If the principal part of  $f$  is  $C$ -nondegenerate, one can compute the spectrum using the library `spectrum.lib`. In this case, the  $V$ -filtration on  $H''$  coincides with the Newton-filtration on  $H''$  which allows to compute the spectrum more efficiently.

Let us calculate one specific example, the maximal number of triple points of type  $\tilde{E}_6$  on a surface  $X \subset P^3$  of degree seven. This calculation can be done over the rationals. We choose a local ordering on  $Q[x, y, z]$ . Here we take the negative degree lexicographical ordering, in SINGULAR denoted by `ds`:

```

ring r=0, (x,y,z), ds;
LIB "spectrum.lib";
poly f=x^7+y^7+z^7;
list s1=spectrumnd(f);
s1;
↳ [1] :
↳ _[1]=-4/7
↳ _[2]=-3/7
↳ _[3]=-2/7

```

```

⇒ _[4]=-1/7
⇒ _[5]=0
⇒ _[6]=1/7
⇒ _[7]=2/7
⇒ _[8]=3/7
⇒ _[9]=4/7
⇒ _[10]=5/7
⇒ _[11]=6/7
⇒ _[12]=1
⇒ _[13]=8/7
⇒ _[14]=9/7
⇒ _[15]=10/7
⇒ _[16]=11/7
⇒ [2]:
⇒ 1,3,6,10,15,21,25,27,27,25,21,15,10,6,3,1

```

The command `spectrumnd(f)` computes the spectrum of  $f$  and returns a list with six entries: The Milnor number  $\mu(f)$ , the geometric genus  $p_g(f)$  and the number of different spectrum numbers. The other three entries are of type `intvec`. They contain the numerators, denominators and multiplicities of the spectrum numbers. So  $x^7 + y^7 + z^7 = 0$  has Milnor number 216 and geometrical genus 35. Its spectrum consists of the 16 different rationals

$\frac{3}{7}, \frac{4}{7}, \frac{5}{7}, \frac{6}{7}, \frac{1}{1}, \frac{8}{7}, \frac{9}{7}, \frac{10}{7}, \frac{11}{7}, \frac{12}{7}, \frac{13}{7}, \frac{2}{1}, \frac{15}{7}, \frac{16}{7}, \frac{17}{7}, \frac{18}{7}$

appearing with multiplicities

1,3,6,10,15,21,25,27,27,25,21,15,10,6,3,1.

The singularities of type  $\tilde{E}_6$  form a  $\mu$ -constant one parameter family given by  $x^3 + y^3 + z^3 + \lambda xyz = 0$ ,  $\lambda^3 \neq -27$ . Therefore they have all the same spectrum, which we compute for  $x^3 + y^3 + z^3$ .

```

poly g=x^3+y^3+z^3;
list s2=spectrumnd(g);
s2;
⇒ [1]:
⇒ 8
⇒ [2]:
⇒ 1
⇒ [3]:
⇒ 4
⇒ [4]:
⇒ 1,4,5,2
⇒ [5]:
⇒ 1,3,3,1
⇒ [6]:
⇒ 1,3,3,1

```

Evaluating semicontinuity is very easy:

```

semicont(s1,s2);
⇒ 18

```

This tells us that there are at most 18 singularities of type  $\tilde{E}_6$  on a septic in  $P^3$ . But  $x^7 + y^7 + z^7$  is semiquasihomogeneous (sqh), so we can also apply the stronger form of semicontinuity:

```

semicontsqh(s1,s2);
⇒ 17

```

So in fact a septic has at most 17 triple points of type  $\tilde{E}_6$ .

Note that `spectrumnd(f)` works only if  $f$  has a nondegenerate principal part. In fact `spectrumnd` will detect a degenerate principal part in many cases and print out an error message. However if it

is known in advance that  $f$  has nondegenerate principal part, then the spectrum may be computed much faster using `spectrumnd(f,1)`.

## C.6 Toric ideals and integer programming

### C.6.1 Toric ideals

Let  $A$  denote an  $m \times n$  matrix with integral coefficients. For  $u \in \mathbb{Z}^n$ , we define  $u^+, u^-$  to be the uniquely determined vectors with nonnegative coefficients and disjoint support (i.e.,  $u_i^+ = 0$  or  $u_i^- = 0$  for each component  $i$ ) such that  $u = u^+ - u^-$ . For  $u \geq 0$  component-wise, let  $x^u$  denote the monomial  $x_1^{u_1} \cdot \dots \cdot x_n^{u_n} \in K[x_1, \dots, x_n]$ .

The ideal

$$I_A := (x^{u^+} - x^{u^-} \mid u \in \ker(A) \cap \mathbb{Z}^n) \subset K[x_1, \dots, x_n]$$

is called a **toric ideal**.

The first problem in computing toric ideals is to find a finite generating set: Let  $v_1, \dots, v_r$  be a lattice basis of  $\ker(A) \cap \mathbb{Z}^n$  (i.e., a basis of the  $\mathbb{Z}$ -module). Then

$$I_A := I : (x_1 \cdot \dots \cdot x_n)^\infty$$

where

$$I = \langle x^{v_i^+} - x^{v_i^-} \mid i = 1, \dots, r \rangle$$

The required lattice basis can be computed using the LLL-algorithm (Section 5.1.153 [system], page 270, see see [[Coh93]], page 776). For the computation of the saturation, there are various possibilities described in the section Algorithms.

### C.6.2 Algorithms

The following algorithms are implemented in Section D.4.35 [toric.lib], page 834.

#### C.6.2.1 The algorithm of Conti and Traverso

The algorithm of Conti and Traverso (see [[CoTr91]], page 776) computes  $I_A$  via the extended matrix  $B = (I_m \mid A)$ , where  $I_m$  is the  $m \times m$  unity matrix. A lattice basis of  $B$  is given by the set of vectors  $(a^j, -e_j) \in \mathbb{Z}^{m+n}$ , where  $a^j$  is the  $j$ -th row of  $A$  and  $e_j$  the  $j$ -th coordinate vector. We look at the ideal in  $K[y_1, \dots, y_m, x_1, \dots, x_n]$  corresponding to these vectors, namely

$$I_1 = \langle y^{a_j^+} - x_j y^{a_j^-} \mid j = 1, \dots, n \rangle.$$

We introduce a further variable  $t$  and adjoin the binomial  $t \cdot y_1 \cdot \dots \cdot y_m - 1$  to the generating set of  $I_1$ , obtaining an ideal  $I_2$  in the polynomial ring  $K[t, y_1, \dots, y_m, x_1, \dots, x_n]$ .  $I_2$  is saturated w.r.t. all variables because all variables are invertible modulo  $I_2$ . Now  $I_A$  can be computed from  $I_2$  by eliminating the variables  $t, y_1, \dots, y_m$ .

Because of the big number of auxiliary variables needed to compute a toric ideal, this algorithm is rather slow in practice. However, it has a special importance in the application to integer programming (see Section C.6.4 [Integer programming], page 775).

### C.6.2.2 The algorithm of Pottier

The algorithm of Pottier (see [[Pot94]], page 776) starts by computing a lattice basis  $v_1, \dots, v_r$  for the integer kernel of  $A$  using the LLL-algorithm (Section 5.1.153 [system], page 270). The ideal corresponding to the lattice basis vectors

$$I_1 = \langle x^{v_i^+} - x^{v_i^-} \mid i = 1, \dots, r \rangle$$

is saturated – as in the algorithm of Conti and Traverso – by inversion of all variables: One adds an auxiliary variable  $t$  and the generator  $t \cdot x_1 \cdot \dots \cdot x_n - 1$  to obtain an ideal  $I_2$  in  $K[t, x_1, \dots, x_n]$  from which one computes  $I_A$  by elimination of  $t$ .

### C.6.2.3 The algorithm of Hosten and Sturmfels

The algorithm of Hosten and Sturmfels (see [[HoSt95]], page 776) allows to compute  $I_A$  without any auxiliary variables, provided that  $A$  contains a vector  $w$  with positive coefficients in its row space. This is a real restriction, i.e., the algorithm will not necessarily work in the general case.

A lattice basis  $v_1, \dots, v_r$  is again computed via the LLL-algorithm. The saturation step is performed in the following way: First note that  $w$  induces a positive grading w.r.t. which the ideal

$$I = \langle x^{v_i^+} - x^{v_i^-} \mid i = 1, \dots, r \rangle$$

is homogeneous corresponding to our lattice basis. We use the following lemma:

Let  $I$  be a homogeneous ideal w.r.t. the weighted reverse lexicographical ordering with weight vector  $w$  and variable order  $x_1 > x_2 > \dots > x_n$ . Let  $G$  denote a Groebner basis of  $I$  w.r.t. this ordering. Then a Groebner basis of  $(I : x_n^\infty)$  is obtained by dividing each element of  $G$  by the highest possible power of  $x_n$ .

From this fact, we can successively compute

$$I_A = I : (x_1 \cdot \dots \cdot x_n)^\infty = (((I : x_1^\infty) : x_2^\infty) : \dots : x_n^\infty);$$

in the  $i$ -th step we take  $x_i$  as the smallest variable and apply the lemma with  $x_i$  instead of  $x_n$ .

This procedure involves  $n$  Groebner basis computations. Actually, this number can be reduced to at most  $n/2$  (see [[HoSh98]], page 776), and each computation – except for the first one – proves to be simple and fast in practice.

### C.6.2.4 The algorithm of Di Biase and Urbanke

Like the algorithm of Hosten and Sturmfels, the algorithm of Di Biase and Urbanke (see [[DBUr95]], page 776) performs up to  $n/2$  Groebner basis computations. It needs no auxiliary variables, but a supplementary precondition; namely, the existence of a vector without zero components in the kernel of  $A$ .

The main idea comes from the following observation:

Let  $B$  be an integer matrix,  $u_1, \dots, u_r$  a lattice basis of the integer kernel of  $B$ . Assume that all components of  $u_1$  are positive. Then

$$I_B = \langle x^{u_i^+} - x^{u_i^-} \mid i = 1, \dots, r \rangle,$$

i.e., the ideal on the right is already saturated w.r.t. all variables.

The algorithm starts by finding a lattice basis  $v_1, \dots, v_r$  of the kernel of  $A$  such that  $v_1$  has no zero component. Let  $\{i_1, \dots, i_l\}$  be the set of indices  $i$  with  $v_{1,i} < 0$ . Multiplying the components



$i_1, \dots, i_l$  of  $v_1, \dots, v_r$  and the columns  $i_1, \dots, i_l$  of  $A$  by  $-1$  yields a matrix  $B$  and a lattice basis  $u_1, \dots, u_r$  of the kernel of  $B$  that fulfill the assumption of the observation above. It is then possible to compute a generating set of  $I_A$  by applying the following “variable flip” successively to  $i = i_1, \dots, i_l$ : Let  $>$  be an elimination ordering for  $x_i$ . Let  $A_i$  be the matrix obtained by multiplying the  $i$ -th column of  $A$  by  $-1$ . Let

$$\{x_i^{r_j} x^{a_j} - x^{b_j} | j \in J\}$$

be a Groebner basis of  $I_{A_i}$  w.r.t.  $>$  (where  $x_i$  is neither involved in  $x^{a_j}$  nor in  $x^{b_j}$ ). Then

$$\{x^{a_j} - x_i^{r_j} x^{b_j} | j \in J\}$$

is a generating set for  $I_A$ .

### C.6.2.5 The algorithm of Bigatti, La Scala and Robbiano

The algorithm of Bigatti, La Scala and Robbiano (see [[BLR98]], page 776) combines the ideas of the algorithms of Pottier and of Hosten and Sturmfels. The computations are performed on a graded ideal with one auxiliary variable  $u$  and one supplementary generator  $x_1 \cdot \dots \cdot x_n - u$  (instead of the generator  $t \cdot x_1 \cdot \dots \cdot x_n - 1$  in the algorithm of Pottier). The algorithm uses a quite unusual technique to get rid of the variable  $u$  again.

There is another algorithm of the authors which tries to parallelize the computations (but which is not implemented in this library).

### C.6.3 The Buchberger algorithm for toric ideals

Toric ideals have a very special structure that allows us to improve the Buchberger algorithm in many aspects: They are prime ideals and generated by binomials. Pottier used this fact to describe all operations of the Buchberger algorithm on the ideal generators in terms of vector additions and subtractions. Some other strategies like multiple reduction (see [[CoTr91]], page 776) or the use of bit vectors to represent the support of a monomial (see [[Big97]], page 776) may be applied to more general ideals, but show to be especially useful in the toric case.

### C.6.4 Integer programming

Let  $A$  be an  $m \times n$  matrix with integral coefficients,  $b \in \mathbb{Z}^m$  and  $c \in \mathbb{Z}^n$ . The problem

$$\min\{c^T x | x \in \mathbb{Z}^n, Ax = b, x \geq 0 \text{ component-wise}\}$$

is called an instance of the **integer programming problem** or **IP problem**.

The IP problem is very hard; namely, it is NP-complete.

For the following discussion let  $c \geq 0$  (component-wise). We consider  $c$  as a weight vector; because of its nonnegativity,  $c$  can be refined into a monomial ordering  $>_c$ . It turns out that we can solve such an IP instance with the help of toric ideals:

First we assume that an initial solution  $v$  (i.e.,  $v \in \mathbb{Z}^n, v \geq 0, Av = b$ ) is already known. We obtain the optimal solution  $v_0$  (i.e., with  $c^T v_0$  minimal) by the following procedure:

- (1) Compute the toric ideal  $I(A)$  using one of the algorithms in the previous section.
- (2) Compute the reduced Groebner basis  $G(c)$  of  $I(A)$  w.r.t.  $>_c$ .
- (3) Reduce  $x^v$  modulo  $G(c)$  using the Hironaka division algorithm. If the result of this reduction is  $x^w$ , then  $w$  is an optimal solution of the given instance.

If no initial solution is known, we are nevertheless able to solve the problem with similar techniques. For this purpose we replace our instance by an extended instance with the matrix used in the Conti-Traverso algorithm. Indeed, the Conti-Traverso algorithm offers the possibility to verify solvability of a given instance and to find an initial solution in the case of existence (but none of the other algorithms does!). Details can be found in see [[CoTr91]], page 776 and see [[The99]], page 776.

An implementation of the above algorithm and some examples can be found in Section D.4.12 [intprog\_lib], page 817.

In general, classical methods for solving IP instances like Branch-and-Bound methods seem to be faster than the methods using toric ideals. But the latter have one great advantage: If one wants to solve various instances that differ only by the vector  $b$ , one has to perform steps (1) and (2) above only once. As the running time of step (3) is very short, solving all the instances is not much harder than solving one single instance.

For a detailed discussion see see [[The99]], page 776.

### C.6.5 Relevant References

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## C.7 Non-commutative algebra

See Section 7.4 [Mathematical background (plural)], page 360, Section 7.9 [Mathematical background (letterplace)], page 625.

## C.8 Decoding codes with Groebner bases

This section introduces some of the mathematical notions, definitions, and results for solving decoding problems and finding the minimum distance of linear (and in particular cyclic) codes. The material presented here should assist the user in working with Section D.10.2 [decodegb\_lib], page 887. More details can be obtained from [[BP2008b]], page 782.

## C.8.1 Codes and the decoding problem

### Codes

- Let  $F_q$  be a field with  $q$  elements. A *linear code*  $C$  is a linear subspace of  $F_q^n$  endowed with the **Hamming metric**.
- **Hamming distance** between  $\mathbf{x}, \mathbf{y} \in F_q^n$  :  $d(x, y) = \#\{i | x_i \neq y_i\}$ . **Hamming weight** of  $\mathbf{x} \in F_q^n$  :  $wt(x) = \#\{i | x_i \neq 0\}$ .
- **Minimum distance** of the code  $C$  :  $d(C) := \min_{\mathbf{x}, \mathbf{y} \in C, \mathbf{x} \neq \mathbf{y}} (d(\mathbf{x}, \mathbf{y}))$ .
- The code  $C$  of dimension  $k$  and minimum distance  $d$  is denoted as  $[n, k, d]$ .
- A matrix  $G$  whose rows are the base vectors of  $C$  is the **generator matrix**.
- A matrix  $H$  with the property  $\mathbf{c} \in C \iff H\mathbf{c}^T = 0$  is the **check matrix**.

### Cyclic codes

The code  $C$  is **cyclic**, if for every codeword  $\mathbf{c} = (c_0, \dots, c_{n-1})$  in  $C$  its cyclic shift  $(c_{n-1}, c_0, \dots, c_{n-2})$  is again a codeword in  $C$ . When working with cyclic codes, vectors are usually presented as polynomials. So  $\mathbf{c}$  is represented by the polynomial  $c(x) = \sum_{i=0}^{n-1} c_i x^i$  with  $x^n = 1$ , more precisely  $c(x)$  is an element of the factor ring  $F_q[X]/\langle X^n - 1 \rangle$ . Cyclic codes over  $F_q$  of length  $n$  correspond one-to-one to ideals in this factor ring. We assume for cyclic codes that  $(q, n) = 1$ . Let  $F = F_{q^m}$  be the splitting field of  $X^n - 1$  over  $F_q$ . Then  $F$  has a **primitive  $n$ -th root of unity** which will be denoted by  $a$ . A cyclic code is uniquely given by a **defining set**  $S_C$  which is a subset of  $\mathbb{Z}_n$  such that

$$c(x) \in C \text{ if } c(a^i) = 0 \text{ for all } i \in S_C.$$

A cyclic code has several defining sets.

### Decoding problem

- **Complete decoding**: Given  $y \in F_q^n$  and a code  $C \subseteq F_q^n$ , so that  $y$  is at distance  $d(y, C)$  from the code, find  $c \in C$  :  $d(y, c) = d(y, C)$ .
- **Bounded up to half the minimum distance**: With the additional assumption  $d(\mathbf{y}, C) \leq (d(C) - 1)/2$ , a codeword with the above property is unique.

### Decoding via systems solving

One distinguishes between two concepts:

- **Generic decoding**: Solve some system  $S(C)$  and obtain some "closed" formulas  $CF$ . Evaluating these formulas at data specific to a received word  $\mathbf{r}$  should yield a solution to the decoding problem. For example for  $f \in CF$  :  $f(\text{syndrome}(\mathbf{r}), x) = \text{poly}(x)$ . The roots of  $\text{poly}(x) = 0$  yield error positions, see the section on the general error-locator polynomial.
- **Online decoding**: Solve some system  $S(C, \mathbf{r})$ . The solutions should solve the decoding problem.

### Computational effort

- Generic decoding. Here, preprocessing is very hard, whereas decoding is relatively simple (if the formulas are sparse).
- Online decoding. In this case, decoding is the hard part.

### C.8.2 Cooper philosophy

#### Computing syndromes in cyclic code case

Let  $C$  be an  $[n, k]$  cyclic code over  $F_q$ ;  $F$  is a splitting field with  $a$  being a primitive  $n$ -th root of unity. Let  $S_C = \{i_1, \dots, i_{n-k}\}$  be the complete defining set of  $C$ . Let  $\mathbf{r} = \mathbf{c} + \mathbf{e}$  be a received word with  $\mathbf{c} \in C$  and  $\mathbf{e}$  an error vector. Denote the corresponding polynomials in  $F_q[x]/\langle x^n - 1 \rangle$  by  $r(x)$ ,  $c(x)$  and  $e(x)$ , resp. Compute syndromes

$$s_{i_m} = r(a^{i_m}) = e(a^{i_m}) = \sum_{l=1}^t e_{j_l} (a^{i_m})^{j_l}, \quad 1 \leq m \leq n - k,$$

where  $t$  is the number of errors,  $j_1, \dots, j_t$  are the **error positions** and  $e_{j_1}, \dots, e_{j_t}$  are the **error values**. Define  $z_l = a^{j_l}$  and  $y_l = e_{j_l}$ . Then  $z_1, \dots, z_t$  are the error locations and  $y_1, \dots, y_t$  are the error values and the syndromes above become **generalized power sum functions**  $s_{i_m} = \sum_{l=1}^t y_l z_l^{i_m}$ ,  $1 \leq m \leq n - k$ .

#### CRHT-ideal

Replace the concrete values above by variables and add some natural restrictions. Introduce

- $f_u := \sum_{l=1}^e Y_l Z_l^{i_u} - X_u = 0, 1 \leq u \leq n - k$ ;
- $\epsilon_j := X_j^{q^m} - X_j = 0, 1 \leq j \leq n - k$ , since  $s_j \in F$ ;
- $\eta_i := Z_i^{n+1} - Z_i = 0, 1 \leq i \leq e$ , since  $a^{j_i}$  are either  $n$ -th roots of unity or zero;
- $\lambda_i := Y_i^{q-1} - 1 = 0, 1 \leq i \leq e$ , since  $y_l \in F_q \setminus \{0\}$ .

We obtain the following set of polynomials in the variables  $X = (X_1, \dots, X_{n-k})$ ,  $Z = (Z_1, \dots, Z_e)$  and  $Y = (Y_1, \dots, Y_e)$ :

$$F_C = \{f_j, \epsilon_j, \eta_i, \lambda_i : 1 \leq j \leq n - k, 1 \leq i \leq e\} \subset F_q[X, Z, Y].$$

The zero-dimensional ideal  $I_C$  generated by  $F_C$  is the **CRHT-syndrome ideal** associated to the code  $C$ , and the variety  $V(F_C)$  defined by  $F_C$  is the **CRHT-syndrome variety**, after Chen, Reed, Helleseht and Truong.

#### General error-locator polynomial

Adding some more polynomials to  $F_C$ , thus obtaining some  $F'_C$ , it is possible to prove the following **Theorem**:

Every cyclic code  $C$  possesses a **general error-locator polynomial**  $L_C$  from  $F_q[X_1, \dots, X_{n-k}, Z]$  that satisfies the following two properties:

- $L_C = Z^e + a_{t-1}Z^{e-1} + \dots + a_0$  with  $a_j \in F_q[X_1, \dots, X_{n-k}]$ ,  $0 \leq j \leq e - 1$ , where  $e$  is the error-correcting capacity;
- given a syndrome  $\mathbf{s} = (s_{i_1}, \dots, s_{i_{n-k}}) \in F^{n-k}$  corresponding to an error of weight  $t \leq e$  and error locations  $\{k_1, \dots, k_t\}$ , if we evaluate the  $X_u = s_{i_u}$  for all  $1 \leq u \leq n - k$ , then the roots of  $L_C(\mathbf{s}, Z)$  are exactly  $a^{k_1}, \dots, a^{k_t}$  and 0 of multiplicity  $e - t$ , in other words  $L_C(\mathbf{s}, Z) = Z^{e-t} \prod_{i=1}^t (Z - a^{k_i})$ .

The general error-locator polynomial actually is an element of the reduced Gröbner basis of  $\langle F'_C \rangle$ . Having this polynomial, decoding of the cyclic code  $C$  reduces to univariate factorization.

For an example see `sysCRHT` in Section D.10.2 [decodegb.lib], page 887. More on Cooper's philosophy and the general error-locator polynomial can be found in [[OS2005]], page 782.

## Finding the minimum distance

The method described above can be adapted to find the minimum distance of a code. More concretely, the following holds:

Let  $C$  be the binary  $[n, k, d]$  cyclic code with the defining set  $S_C = \{i_1, \dots, i_v\}$ . Let  $1 \leq w \leq n$  and let  $J_C(w)$  denote the system:

$$\begin{aligned} Z_1^{i_1} + \dots + Z_w^{i_1} &= 0, \\ &\vdots \\ Z_1^{i_v} + \dots + Z_w^{i_v} &= 0, \\ Z_1^n - 1 &= 0, \\ &\vdots \\ Z_w^n - 1 &= 0, \\ p(n, Z_i, Z_j) &= 0, 1 \leq i < j \leq w. \end{aligned}$$

Then the number of solutions of  $J_C(w)$  is equal to  $w!$  times the number of codewords of weight  $w$ . And for  $1 \leq w \leq d$ , either  $J_C(w)$  has no solutions, which is equivalent to  $w < d$ , or  $J_C(w)$  has some solutions, which is equivalent to  $w = d$ .

For an example see `sysCRHTMindist` in Section D.10.2 [decodegb.lib], page 887. More on finding the minimum distance with Groebner bases can be found in [[S2007]], page 782. See [[OS2005]], page 782, for the definition of the polynomial  $p$  above.

### C.8.3 Generalized Newton identities

The **error-locator polynomial** is defined by

$$\sigma(Z) = \prod_{l=1}^t (Z - z_l).$$

If this product is expanded,

$$\sigma(Z) = Z^t + \sigma_1 Z^{t-1} + \dots + \sigma_{t-1} Z + \sigma_t,$$

then the coefficients  $\sigma_i$  are the **elementary symmetric functions** in the error locations  $z_1, \dots, z_t$

$$\sigma_i = (-1)^i \sum_{1 \leq j_1 < j_2 < \dots < j_i \leq t} z_{j_1} z_{j_2} \dots z_{j_i}, \quad 1 \leq i \leq t.$$

### Generalized Newton identities

The syndromes  $s_i = r(a^i) = e(a^i)$  and the coefficients  $\sigma_i$  satisfy the following **generalized Newton identities**:

$$s_i + \sum_{j=1}^t \sigma_j s_{i-j} = 0, \quad \text{for all } i \in \mathbb{Z}_n.$$

## Decoding up to error-correcting capacity

We have  $s_{i+n} = s_i$ , for all  $i \in \mathbb{Z}_n$ , since  $s_{i+n} = r(a^{i+n}) = r(a^i)$ . Furthermore

$$s_i^q = (e(a^i))^q = e(a^{iq}) = s_{qi}, \text{ for all } i \in \mathbb{Z}_n,$$

and  $\sigma_i^{q^m} = \sigma_i$ , for all  $1 \leq i \leq t$ . Replace the syndromes by variables and obtain the following set of polynomials  $Newton_t$  in the variables  $S_1, \dots, S_n$  and  $\sigma_1, \dots, \sigma_t$ :

$$\begin{aligned} \sigma_i^{q^m} - \sigma_i, \quad \forall 1 \leq i \leq t, \\ S_{i+n} - S_i, \quad \forall i \in \mathbb{Z}_n, \\ S_i^q - S_{qi}, \quad \forall i \in \mathbb{Z}_n, \\ S_i + \sum_{j=1}^t \sigma_j S_{i-j}, \quad \forall i \in \mathbb{Z}_n, \\ S_i - s_i(r) \quad \forall i \in S_C. \end{aligned}$$

For an example see `sysNewton` in Section D.10.2 [`decodegb_lib`], page 887. More on this method and the method based on Waring function can be found in [[ABF2002]], page 782. See also [[ABF2008]], page 782.

### C.8.4 Fitzgerald-Lax method

#### Affine codes

Let  $I = \langle g_1, \dots, g_m \rangle \subseteq F_q[X_1, \dots, X_s]$  be an ideal. Define

$$I_q := I + \langle X_1^q - X_1, \dots, X_s^q - X_s \rangle.$$

So  $I_q$  is a zero-dimensional ideal. Define also  $V(I_q) = \{P_1, \dots, P_n\}$ . Every  $q$ -ary linear code  $C$  with parameters  $[n, k]$  can be seen as an **affine variety code**  $C(I, L)$ , that is, the image of a vector space  $L$  of the **evaluation map**

$$\begin{aligned} \phi : R &\rightarrow F_q^n \\ \bar{f} &\mapsto (f(P_1), \dots, f(P_n)), \end{aligned}$$

where  $R := F_q[U_1, \dots, U_s]/I_q$ ,  $L$  is a vector subspace of  $R$  and  $\bar{f}$  the coset of  $f$  in  $F_q[U_1, \dots, U_s]$  modulo  $I_q$ .

#### Decoding affine variety codes

Given a  $q$ -ary  $[n, k]$  code  $C$  with a generator matrix  $G = (g_{ij})$ :

1. choose  $s$ , such that  $q^s \geq n$ , and construct  $s$  distinct points  $P_1, \dots, P_s$  in  $F_q^s$ .
2. Construct a Gröbner basis  $\{g_1, \dots, g_m\}$  for an ideal  $I$  of polynomials from  $F_q[X_1, \dots, X_s]$  that vanish at the points  $P_1, \dots, P_s$ . Define  $\xi_i \in F_q[X_1, \dots, X_s]$  such that  $\xi_i(P_i) = 1, \xi_i(P_j) = 0, i \neq j$ .
3. Then  $f_i = \sum_{j=1}^n g_{ij} \xi_j$  span the space  $L$ , so that  $g_{ij} = f_i(P_j)$ .

In this way we obtain that the code  $C$  is the image of the evaluation above, thus  $C = C(I, L)$ . In the same way by considering a parity check matrix instead of a generator matrix we have that the dual code is also an affine variety code.

The method of decoding is a generalization of CRHT. One needs to add polynomials  $(g_l(X_{k_1}, \dots, X_{k_s}))_{l=1, \dots, m; k=1, \dots, t}$  for every error position. We also assume that field equations on  $X_{ij}$ 's are included among the polynomials above. Let  $C$  be a  $q$ -ary  $[n, k]$  linear code such that its dual is written as an affine variety code of the form  $C^\perp = C(I, L)$ . Let  $\mathbf{r} = \mathbf{c} + \mathbf{e}$  as usual and  $t \leq e$ . Then the syndromes are computed by  $s_i = \sum_{j=1}^n r_j f_i(P_j) = \sum_{j=1}^n e_j f_i(P_j)$  for  $i = 1, \dots, n - k$ .

Consider the ring  $F_q[X_{11}, \dots, X_{1s}, \dots, X_{t1}, \dots, X_{ts}, E_1, \dots, E_t]$ , where  $(X_{i1}, \dots, X_{is})$  correspond to the  $i$ -th error position and  $E_i$  to the  $i$ -th error value. Consider the ideal  $Id_C$  generated by

$$\begin{aligned} \sum_{j=1}^t E_j f_i(X_{j1}, \dots, X_{js}) - s_i, 1 \leq i \leq n - k, \\ g_l(X_{j1}, \dots, X_{js}), 1 \leq l \leq m, \\ E_k^{q-1} - 1. \end{aligned}$$

**Theorem:** Let  $G$  be the reduced Gröbner basis for  $Id_C$  with respect to an elimination order  $X_{11} < \dots < X_{1s} < E_1$ . Then we may solve for the error locations and values by applying elimination theory to the polynomials in  $G$ .

For an example see `sysFL` in Section D.10.2 [decodegb-lib], page 887. More on this method can be found in [[FL1998]], page 782.

### C.8.5 Decoding method based on quadratic equations

#### Preliminary definitions

Let  $\mathbf{b}_1, \dots, \mathbf{b}_n$  be a basis of  $F_q^n$  and let  $B$  be the  $n \times n$  matrix with  $\mathbf{b}_1, \dots, \mathbf{b}_n$  as rows. The **unknown syndrome**  $\mathbf{u}(B, \mathbf{e})$  of a word  $\mathbf{e}$  w.r.t  $B$  is the column vector  $\mathbf{u}(B, \mathbf{e}) = B\mathbf{e}^T$  with entries  $u_i(B, \mathbf{e}) = \mathbf{b}_i \cdot \mathbf{e}$  for  $i = 1, \dots, n$ .

For two vectors  $\mathbf{x}, \mathbf{y} \in F_q^n$  define  $\mathbf{x} * \mathbf{y} = (x_1 y_1, \dots, x_n y_n)$ . Then  $\mathbf{b}_i * \mathbf{b}_j$  is a linear combination of  $\mathbf{b}_1, \dots, \mathbf{b}_n$ , so there are constants  $\mu_l^{ij} \in F_q$  such that  $\mathbf{b}_i * \mathbf{b}_j = \sum_{l=1}^n \mu_l^{ij} \mathbf{b}_l$ . The elements  $\mu_l^{ij} \in F_q$  are the **structure constants** of the basis  $\mathbf{b}_1, \dots, \mathbf{b}_n$ .

Let  $B_s$  be the  $s \times n$  matrix with  $\mathbf{b}_1, \dots, \mathbf{b}_s$  as rows ( $B = B_n$ ). Then  $\mathbf{b}_1, \dots, \mathbf{b}_n$  is an **ordered MDS basis** and  $B$  an **MDS matrix** if all the  $s \times s$  submatrices of  $B_s$  have rank  $s$  for all  $s = 1, \dots, n$ .

#### Expressing known syndromes

Let  $C$  be an  $F_q$ -linear code with parameters  $[n, k, d]$ . W.l.o.g  $n \leq q$ .  $H$  is a check matrix of  $C$ . Let  $\mathbf{h}_1, \dots, \mathbf{h}_{n-k}$  be the rows of  $H$ . One can express  $\mathbf{h}_i = \sum_{j=1}^n a_{ij} \mathbf{b}_j$  with some  $a_{ij} \in F_q$ . In other words  $H = AB$  where  $A$  is the  $(n - k) \times n$  matrix with entries  $a_{ij}$ .

Let  $\mathbf{r} = \mathbf{c} + \mathbf{e}$  be a received word with  $\mathbf{c} \in C$  and  $\mathbf{e}$  an error vector. The syndromes of  $\mathbf{r}$  and  $\mathbf{e}$  w.r.t  $H$  are equal and known:

$$s_i(\mathbf{r}) := \mathbf{h}_i \cdot \mathbf{r} = \mathbf{h}_i \cdot \mathbf{e} = s_i(\mathbf{e}).$$

They can be expressed in the unknown syndromes of  $\mathbf{e}$  w.r.t  $B$ :

$$s_i(\mathbf{r}) = s_i(\mathbf{e}) = \sum_{j=1}^n a_{ij} u_j(\mathbf{e})$$

since  $\mathbf{h}_i = \sum_{j=1}^n a_{ij} \mathbf{b}_j$  and  $\mathbf{b}_j \cdot \mathbf{e} = u_j(\mathbf{e})$ .

## Contracting the system

Let  $B$  be an MDS matrix with structure constants  $\mu_l^{ij}$ . Define  $U_{ij}$  in the variables  $U_1, \dots, U_n$  by

$$U_{ij} = \sum_{l=1}^n \mu_l^{ij} U_l.$$

The ideal  $J(\mathbf{r})$  in  $F_q[U_1, \dots, U_n]$  is generated by

$$\sum_{l=1}^n a_{jl} U_l - s_j(\mathbf{r}) \text{ for } j = 1, \dots, n-k.$$

The ideal  $I(t, U, V)$  in  $F_q[U_1, \dots, U_n, V_1, \dots, V_t]$  is generated by

$$\sum_{j=1}^t U_{ij} V_j - U_{i,t+1} \text{ for } i = 1, \dots, n$$

Let  $J(t, \mathbf{r})$  be the ideal in  $F_q[U_1, \dots, U_n, V_1, \dots, V_t]$  generated by  $J(\mathbf{r})$  and  $I(t, U, V)$ .

## Main theorem

Let  $B$  be an MDS matrix with structure constants  $\mu_l^{ij}$ . Let  $H$  be a check matrix of the code  $C$  such that  $H = AB$  as above. Let  $\mathbf{r} = \mathbf{c} + \mathbf{e}$  be a received word with  $\mathbf{c} \in C$  the codeword sent and  $\mathbf{e}$  the error vector. Suppose that  $wt(\mathbf{e}) \neq 0$  and  $wt(\mathbf{e}) \leq \lfloor (d(C) - 1)/2 \rfloor$ . Let  $t$  be the smallest positive integer such that  $J(t, \mathbf{r})$  has a solution  $(\mathbf{u}, \mathbf{v})$  over the algebraic closure of  $F_q$ . Then

- $wt(\mathbf{e}) = t$  and the solution is unique and of multiplicity one satisfying  $\mathbf{u} = \mathbf{u}(\mathbf{e})$ .
- the reduced Gröbner basis  $G$  for the ideal  $J(t, \mathbf{r})$  w.r.t any monomial ordering is

$$\begin{aligned} U_i - u_i(\mathbf{e}), i = 1, \dots, n, \\ V_j - v_j, j = 1, \dots, t, \end{aligned}$$

where  $(\mathbf{u}(\mathbf{e}), \mathbf{v})$  is the unique solution.

For an example see `sysQE` in Section D.10.2 [`decodegb_lib`], page 887. More on this method can be found in [[BP2008a]], page 782.

### C.8.6 References for decoding with Groebner bases

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## C.9 References

The Centre for Computer Algebra Kaiserslautern publishes a series of preprints which are electronically available at <https://www.singular.uni-kl.de/reports>. Other sources to check are <http://symbolicnet.org/>, <http://www-sop.inria.fr/galaad/>,... and the following list of books.

For references on non-commutative algebras and algorithms, see Section 7.4.4 [References (plural)], page 364.

### Text books on computational algebraic geometry

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### Descriptions of algorithms

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## Appendix D SINGULAR libraries

SINGULAR comes with a set of standard libraries. Their content is described in the following subsections.

Use the LIB command (see Section 5.1.79 [LIB], page 208) for loading of single libraries, and the command LIB "all.lib"; for loading all libraries.

### Interpreter libraries:

See also Section 7.5 [PLURAL libraries], page 365 and Section 7.10 [LETTERPLACE libraries], page 629.

### D.1 standard\_lib

The library `standard.lib` provides extensions to the set of built-in commands and is automatically loaded during the start of SINGULAR, unless SINGULAR is started up with the `--no-stdlib` command line option (see Section 3.1.6 [Command line options], page 19).

**Library:** `standard.lib`

**Purpose:** Procedures which are always loaded at Start-up

#### Procedures:

```
stdfglm(ideal[,ord])
 standard basis of ideal via fglm [and ordering ord]

stdhilb(ideal[,h])
 Hilbert driven Groebner basis of ideal

groebner(ideal,...)
 standard basis using a heuristically chosen method

res(ideal/module,[i])
 free resolution of ideal or module

sprintf(fmt,...)
 returns formatted string

fprintf(link,fmt,..)
 writes formatted string to link

printf(fmt,...)
 displays formatted string

weightKB(stc,dd,vl)
 degree dd part of a kbase w.r.t. some weights

qslimb(i)
 computes a standard basis with slimb in a qring

par2varRing([i])
 create a ring making pars to vars, together with i

datetime()
 return date and time as a string

max(i_1,...,i_k)
 maximum of i_1, ..., i_k
```

```

min(i_1, ..., i_k)
 minimum of i_1, ..., i_k

create_ring(l1,l2,l3,l4)
 return ring(list(l1, l2, l3, l4))

```

### D.1.1 qslimgb

Procedure from library `standard.lib` (see Section D.1 [`standard.lib`], page 785).

**Usage:** `qslimgb(i)`;  $i$  ideal or module

**Return:** same type as input, a standard basis of  $i$  computed with `slimgb`

**Note:** Only as long as `slimgb` does not know `qgrings` `qslimgb` should be used in case the basering is (possibly) a quotient ring.  
The quotient ideal is added to the input and `slimgb` is applied.

**Example:**

```

ring R = (0,v),(x,y,z,u),dp;
qring Q = std(x2-y3);
ideal i = x+y2,xy+yz+zu+u*v,xyzu*v-1;
ideal j = qslimgb(i); j;
↳ j[1]=y-1
↳ j[2]=x+1
↳ j[3]=(v)*z+(v2)*u+(-v-1)
↳ j[4]=(-v2)*u2+(v+1)*u+1
module m = [x+y2,1,0], [1,1,x2+y2+xyz];
print(qslimgb(m));
↳ y2+x,x2+xy,1, 0, 0, -x, -xy-xz-x,
↳ 1, y, 1, y3-x2,0, y2-1, y2z-xy-x-z,
↳ 0, 0, xyz+x2+y2,0, y3-x2,x2y2+x3z+x2y,x3z2-x3y-2*x3-xy2

```

### D.1.2 par2varRing

Procedure from library `standard.lib` (see Section D.1 [`standard.lib`], page 785).

**Usage:** `par2varRing(l)`;  $l$  list of ideals/modules [default: $l$ =empty list]

**Return:** list, say  $L$ , with  $L[1]$  a ring where the parameters of the basering have been converted to an additional last block of variables, all of weight 1, and ordering `dp`.  
If a list  $l$  with  $l[i]$  an ideal/module is given, then  
 $l[i] + \text{minpoly} * \text{freemodule}(\text{nrows}(l[i]))$  is mapped to an ideal/module in  $L[1]$  with name  $l[i]$ .  
If the basering has no parameters then  $L[1]$  is the basering.

**Example:**

```

ring R = (0,x),(y,z,u,v),lp;
minpoly = x2+1;
ideal i = x3,x2+y+z+u+v,xyzuv-1; i;
↳ i[1]=(-x)
↳ i[2]=y+z+u+v-1
↳ i[3]=(x)*xyzuv-1
def P = par2varRing(i)[1]; P;
↳ // coefficients: QQ
↳ // number of vars : 5

```

```

↳ // block 1 : ordering lp
↳ // : names y z u v
↳ // block 2 : ordering dp
↳ // : names x
↳ // block 3 : ordering C
setring(P);
Id[1];
↳ _[1]=-x
↳ _[2]=y+z+u+v-1
↳ _[3]=yzuvx-1
↳ _[4]=x2+1
setring R;
module m = x3*[1,1,1], (xyzuv-1)*[1,0,1];
def Q = par2varRing(m)[1]; Q;
↳ // coefficients: QQ
↳ // number of vars : 5
↳ // block 1 : ordering lp
↳ // : names y z u v
↳ // block 2 : ordering dp
↳ // : names x
↳ // block 3 : ordering C
setring(Q);
print(Id[1]);
↳ -x,yzuvx-1,x2+1,0, 0,
↳ -x,0, 0, x2+1,0,
↳ -x,yzuvx-1,0, 0, x2+1

```

## D.2 General purpose

### D.2.1 all.lib

The library `all.lib` provides a convenient way to load all libraries of the SINGULAR distribution.

**Example:**

```

option(loadLib);
LIB "all.lib";
↳ // ** loaded all.lib (4.1.1.0,Jan_2018)
↳ // ** loaded ratgb.lib (4.1.2.0,Feb_2019)
↳ // ** loaded qmatrix.lib (4.1.2.0,Feb_2019)
↳ // ** loaded purityfiltration.lib (4.1.2.0,Feb_2019)
↳ // ** loaded perron.lib (4.1.2.0,Feb_2019)
↳ // ** loaded nctools.lib (4.1.2.0,Feb_2019)
↳ // ** loaded ncpream.lib (4.1.2.0,Feb_2019)
↳ // ** loaded dmodloc.lib (4.1.2.0,Feb_2019)
↳ // ** loaded ncModslimgb.lib (4.1.3.0,Apr_2020)
↳ // ** loaded resources.lib (4.1.2.0,Feb_2019)
↳ // ** loaded parallel.lib (4.1.2.0,Feb_2019)
↳ // ** loaded tasks.lib (4.1.2.0,Feb_2019)
↳ // ** loaded nchomolog.lib (4.1.2.0,Feb_2019)
↳ // ** loaded ncfactor.lib (4.1.2.0,Feb_2019)
↳ // ** loaded ncdecomp.lib (4.1.2.0,Feb_2019)
↳ // ** loaded ncalg.lib (4.1.2.0,Feb_2019)
↳ // ** loaded ncall.lib (4.1.2.0,Feb_2019)

```

```
↳ // ** loaded involut.lib (4.1.2.0, Feb_2019)
↳ // ** loaded gkdim.lib (4.1.2.0, Feb_2019)
↳ // ** loaded freegb.lib (4.1.2.0, Feb_2019)
↳ // ** loaded /home/hannes/test/Singular/MOD/freealgebra.so
↳ // ** loaded fpaprops.lib (4.1.2.0, Feb_2019)
↳ // ** loaded fpalgebras.lib (4.1.2.0, Feb_2019)
↳ // ** loaded fpadim.lib (4.1.2.0, Feb_2019)
↳ // ** loaded dmodvar.lib (4.1.2.0, Feb_2019)
↳ // ** loaded dmodapp.lib (4.1.3.0, Mar_2020)
↳ // ** loaded dmod.lib (4.1.2.0, Feb_2019)
↳ // ** loaded central.lib (4.1.2.0, Feb_2019)
↳ // ** loaded bfun.lib (4.1.2.0, Feb_2019)
↳ // ** loaded bimodules.lib (4.1.2.0, Feb_2019)
↳ // ** loaded zeroset.lib (4.1.2.0, Feb_2019)
↳ // ** loaded weierstr.lib (4.1.2.0, Feb_2019)
↳ // ** loaded triang.lib (4.1.2.0, Feb_2019)
↳ // ** loaded toric.lib (4.1.2.0, Feb_2019)
↳ // ** loaded teachstd.lib (4.1.2.0, Feb_2019)
↳ // ** loaded surfex.lib (4.1.2.0, Feb_2019)
↳ // ** loaded surfacesignature.lib (4.1.2.0, Feb_2019)
↳ // ** loaded surf_jupyter.lib (4.1.2.0, Feb_2019)
↳ // ** loaded surf.lib (4.1.2.0, Feb_2019)
↳ // ** loaded stratify.lib (4.1.2.0, Feb_2019)
↳ // ** loaded stanleyreisner.lib (4.1.2.0, Feb_2019)
↳ // ** loaded /home/hannes/test/Singular/MOD/cohomo.so
↳ // ** loaded spectrum.lib (4.1.2.0, Feb_2019)
↳ // ** loaded spcurve.lib (4.1.2.0, Feb_2019)
↳ // ** loaded solve.lib (4.1.2.0, Feb_2019)
↳ // ** loaded signcond.lib (4.1.2.0, Feb_2019)
↳ // ** loaded sing4ti2.lib (4.1.2.0, Feb_2019)
↳ // ** loaded sing.lib (4.1.2.0, Feb_2019)
↳ // ** loaded sheafcoh.lib (4.1.2.0, Feb_2019)
↳ // ** loaded sagbi.lib (4.1.2.0, Feb_2019)
↳ // ** loaded rootsur.lib (4.1.2.0, Feb_2019)
↳ // ** loaded rootsmr.lib (4.1.2.0, Feb_2019)
↳ // ** loaded rinvar.lib (4.1.2.0, Feb_2019)
↳ // ** loaded ringgb.lib (4.1.2.0, Feb_2019)
↳ // ** loaded ring.lib (4.1.2.0, Feb_2019)
↳ // ** loaded reszeta.lib (4.1.2.0, Feb_2019)
↳ // ** loaded resolve.lib (4.1.2.0, Feb_2019)
↳ // ** loaded resjung.lib (4.1.2.0, Feb_2019)
↳ // ** loaded resgraph.lib (4.1.2.0, Feb_2019)
↳ // ** loaded resbinomial.lib (4.1.2.0, Feb_2019)
↳ // ** loaded reesclos.lib (4.1.2.0, Feb_2019)
↳ // ** loaded locnormal.lib (4.1.2.0, Feb_2019)
↳ // ** loaded redcgs.lib (4.1.2.0, Feb_2019)
↳ // ** loaded realrad.lib (4.1.2.0, Feb_2019)
↳ // ** loaded realclassify.lib (4.1.2.0, Feb_2019)
↳ // ** loaded rootisolation.lib (4.1.2.0, Feb_2019)
↳ // ** loaded /home/hannes/test/Singular/MOD/interval.so
↳ // ** loaded classify2.lib (4.1.2.0, Feb_2019)
↳ // ** loaded gfa.lib.so
↳ // ** loaded polyclass.lib (4.1.2.0, Feb_2019)
```

```
↳ // ** loaded random.lib (4.1.2.0, Feb_2019)
↳ // ** loaded qhmoduli.lib (4.1.2.0, Feb_2019)
↳ // ** loaded primitiv.lib (4.1.2.0, Feb_2019)
↳ // ** loaded primdecint.lib (4.1.2.0, Feb_2019)
↳ // ** loaded primdec.lib (4.1.2.0, Feb_2019)
↳ // ** loaded presolve.lib (4.1.2.0, Feb_2019)
↳ // ** loaded polylib.lib (4.1.2.0, Feb_2019)
↳ // ** loaded pointid.lib (4.1.2.0, Feb_2019)
↳ // ** loaded phindex.lib (4.1.2.0, Feb_2019)
↳ // ** loaded pfd.lib (4.1.3.2, Aug_2020)
↳ // ** loaded paraplanecurves.lib (4.1.2.0, Feb_2019)
↳ // ** loaded ntsolve.lib (4.1.2.0, Feb_2019)
↳ // ** loaded normaliz.lib (4.1.2.0, Feb_2019)
↳ // ** loaded normal.lib (4.1.2.0, Feb_2019)
↳ // ** loaded curveInv.lib (4.1.2.0, Feb_2019)
↳ // ** loaded noether.lib (4.1.2.0, Feb_2019)
↳ // ** loaded nfmodsyz.lib (4.1.2.0, Feb_2019)
↳ // ** loaded nfmodstd.lib (4.1.2.0, Feb_2019)
↳ // ** loaded mregular.lib (4.1.2.0, Feb_2019)
↳ // ** loaded mprimdec.lib (4.1.2.0, Feb_2019)
↳ // ** loaded monomialideal.lib (4.1.2.0, Feb_2019)
↳ // ** loaded mondromy.lib (4.1.2.0, Feb_2019)
↳ // ** loaded modstd.lib (4.1.2.0, Feb_2019)
↳ // ** loaded modular.lib (4.1.2.0, Feb_2019)
↳ // ** loaded modnormal.lib (4.1.2.0, Feb_2019)
↳ // ** loaded moddiq.lib (4.1.2.0, Feb_2020)
↳ // ** loaded matrix.lib (4.1.2.0, Feb_2019)
↳ // ** loaded makedbm.lib (4.1.2.0, Feb_2019)
↳ // ** loaded linalg.lib (4.1.2.0, Feb_2019)
↳ // ** loaded latex.lib (4.1.2.0, Feb_2019)
↳ // ** loaded kskernel.lib (4.1.2.0, Feb_2019)
↳ // ** loaded jacobson.lib (4.1.2.0, Feb_2019)
↳ // ** loaded intprog.lib (4.1.2.0, Feb_2019)
↳ // ** loaded inout.lib (4.1.2.0, Feb_2019)
↳ // ** loaded integralbasis.lib (4.1.2.0, Feb_2019)
↳ // ** loaded hyperel.lib (4.1.2.0, Feb_2019)
↳ // ** loaded homolog.lib (4.1.2.0, Feb_2019)
↳ // ** loaded hnoether.lib (4.1.2.0, Feb_2019)
↳ // ** loaded grwalk.lib (4.1.2.0, Feb_2019)
↳ // ** loaded groups.lib (4.1.2.0, Feb_2019)
↳ // ** loaded grobcov.lib (4.1.3, January_2020)
↳ // ** loaded graphics.lib (4.1.2.0, Feb_2019)
↳ // ** loaded gmssing.lib (4.1.2.0, Feb_2019)
↳ // ** loaded gmspoly.lib (4.1.2.0, Feb_2019)
↳ // ** loaded general.lib (4.1.2.0, Feb_2019)
↳ // ** loaded finvar.lib (4.1.2.0, Feb_2019)
↳ // ** loaded equising.lib (4.1.2.0, Feb_2019)
↳ // ** loaded elim.lib (4.1.2.0, Feb_2019)
↳ // ** loaded deform.lib (4.1.2.0, Feb_2019)
↳ // ** loaded decodegb.lib (4.1.2.0, Feb_2019)
↳ // ** loaded curvepar.lib (4.1.2.0, Feb_2019)
↳ // ** loaded crypto.lib (4.1.2.0, Feb_2019)
↳ // ** loaded control.lib (4.1.2.0, Feb_2019)
```

```

↳ // ** loaded compregb.lib (4.1.2.0, Feb_2019)
↳ // ** loaded classify.lib (4.1.2.0, Feb_2019)
↳ // ** loaded cisimplicial.lib (4.1.2.0, Feb_2019)
↳ // ** loaded brnoeth.lib (4.1.2.0, Feb_2019)
↳ // ** loaded atkins.lib (4.1.2.0, Feb_2019)
↳ // ** loaded assprimeszerodim.lib (4.1.2.0, Feb_2019)
↳ // ** loaded arcpoint.lib (4.1.2.0, Feb_2019)
↳ // ** loaded algebra.lib (4.1.2.0, Feb_2019)
↳ // ** loaded alexpoly.lib (4.1.2.0, Feb_2019)
↳ // ** loaded aksaka.lib (4.1.2.0, Feb_2019)
↳ // ** loaded ainvar.lib (4.1.2.0, Feb_2019)
↳ // ** loaded absfact.lib (4.1.2.0, Feb_2019)

```

## D.2.2 compregb.lib

**Library:** compregb.lib

**Purpose:** experimental implementation for comprehensive Groebner systems

**Author:** Akira Suzuki (<http://kurt.scitec.kobe-u.ac.jp/~sakira/CGBusingGB/>) (<sakira@kobe-u.ac.jp>)

**Overview:** see "A Simple Algorithm to compute Comprehensive Groebner Bases using Groebner Bases" by Akira Suzuki and Yosuke Sato for details.

**Procedures:**

```

cgs(polys, vars, pars, R1, R2)
 comprehensive Groebner systems

base2str(G)
 pretty print of the result G

```

## D.2.3 general.lib

**Library:** general.lib

**Purpose:** Elementary Computations of General Type

**Procedures:**

```

A_Z("a", n)
 string a,b,... of n comma separated letters

A_Z_L("a", n)
 list of strings a,b,... of n letters

ASCII([n,m])
 string of printable ASCII characters (number n to m)

absValue(c)
 absolute value of c

binomial(n,m[,.../.])
 n choose m (type int), [type bigint]

deleteSublist(iv,l)
 delete entries given by iv from list l

```



```

factorial(n[,.../...])
 n factorial (=n!) (type int), [type bigint]
fibonacci(n)
 nth Fibonacci number
kmemory([n[,v]])
 active [allocated] memory in kilobyte
killall()
 kill all user-defined variables
number_e(n)
 compute exp(1) up to n decimal digits
number_pi(n)
 compute pi (area of unit circle) up to n digits
primes(n,m)
 intvec of primes p, n<=p<=m
product(.../...[,v])
 multiply components of vector/ideal/...[indices v]
sort(ideal/module)
 sort generators according to monomial ordering
sum(vector/id/...[,v])
 add components of vector/ideal/...[with indices v]
watchdog(i,cmd)
 only wait for result of command cmd for i seconds
primecoeffs(J[,q])
 primefactors <= min(p,32003) of coeffs of J
timeStd(i,d)
 std(i) if the standard basis computation finished after d-1 seconds and i
 otherwise
timeFactorize(p,d)
 factorize(p) if the factorization finished after d-1 seconds otherwise f is
 considered to be irreducible
factorH(p)
 changes variables to become the last variable the principal one in the mul-
 tivariate factorization and factorizes then the polynomial

```

## D.2.4 grobcov\_lib

**Library:** grobcov.lib

**Purpose:**

"Groebner Cover for parametric ideals.", Comprehensive Groebner Systems, Groebner Cover, Canonical Forms, Parametric Polynomial Systems, Automatic Deduction of Geometric Theorems, Dynamic Geometry, Loci, Envelope, Constructible sets. See: A. Montes A, M. Wibmer, "Groebner Bases for Polynomial Systems with parameters", Journal of Symbolic Computation 45 (2010) 1391-1425. (<https://www.mat.upc.edu/en/people/antonio.montes/>).

**Important:**

Recently published book:

A. Montes. "The Groebner Cover":

Springer, Algorithms and Computation in Mathematics 27 (2019) ISSN 1431-1550

ISBN 978-3-030-03903-5

ISBN 978-3-030-03904-2 (e-Book)

Springer Nature Switzerland AG 2018

<https://www.springer.com/gp/book/9783030039035>

The book can also be used as a user manual of all the routines included in this library.

It defines and proves all the theoretic results used in the library, and shows examples of all the routines. There are many previous papers related to the subject, and the book actualices all the contents.

**Authors:** Antonio Montes (Universitat Politecnica de Catalunya), Hans Schoenemann (Technische Universitaet Kaiserslautern).

**Overview:** In 2010, the library was designed to contain Montes-Wibmer's algorithm for computing the Canonical Groebner Cover of a parametric ideal. The central routine is `grobcov`. Given a parametric ideal, `grobcov` outputs its Canonical Groebner Cover, consisting of a set of triplets of (lpp, basis, segment). The basis (after normalization) is the reduced Groebner basis for each point of the segment. The segments are disjoint, locally closed and correspond to constant lpp (leading power product) of the basis, and are represented in canonical representation. The segments cover the whole parameter space. The output is canonical, it only depends on the given parametric ideal and the monomial order, because the segments have different lpp of the homogenized system. This is much more than a simple Comprehensive Groebner System. The algorithm `grobcov` allows options to solve partially the problem when the whole automatic algorithm does not finish in reasonable time. Its existence was proved for the first time by Michael Wibmer "Groebner bases for families of affine or projective schemes", JSC, 42,803-834 (2007). `grobcov` uses a first algorithm `cgsdr` that outputs a disjoint reduced Comprehensive Groebner System with constant lpp. For this purpose, in this library, the implemented algorithm is Kapur-Sun-Wang algorithm, because it is actually the most efficient algorithm known for this purpose. D. Kapur, Y. Sun, and D.K. Wang "A New Algorithm for Computing Comprehensive Groebner Systems". Proceedings of ISSAC'2010, ACM Press, (2010), 29-36.

The library has evolved to include new applications of the Groebner Cover, and new theoretical developments have been done.

A routine locus has been included to compute loci of points, and determining the taxonomy of the components.

Additional routines to transform the output to string (locusdg, locusto) are also included and used in the Dynamic Geometry software GeoGebra. They were described in:

M.A. Abanades, F. Botana, A. Montes, T. Recio:  
"An Algebraic Taxonomy for Locus Computation in Dynamic Geometry".

Computer-Aided Design 56 (2014) 22-33.

Routines for determining the generalized envelope of a family of hypersurfaces (envelop, AssocTanToEnv, FamElementsToEnvCompPoints) are also included.

It also includes procedures for Automatic Deduction of Geometric Theorems (ADGT).

The actual version also includes a routine (ConsLevels) for computing the canonical form of a constructible set, given as a union of locally closed sets. It determines the canonical levels of a constructible set. It is described in:

J.M. Brunat, A. Montes, "Computing the canonical representation of constructible sets".  
Math. Comput. Sci. (2016) 19: 165-178.

A complementary routine Levels transforms the output of ConsLevels into the proper locally closed sets

forming the levels of the constructible.

Another complementary routine Grob1Levels has been included to select the locally closed sets of the segments of the grobcov that correspond to basis different from 1, add them together and return the canonical form of this constructible set.

More recently (2019) given two locally closed sets in canonical form the new routine DifConsLCSets determines a set of locally closed sets equivalent to the difference them. The description of the

routine is submitted to the Journal of Symbolic Computation. This routine can be also used internally by ADGT

with the option "neg", 1. With this option DifConsLCSets is used for the negative hypothesis and thesis in ADGT.

This version was finished on 26/5/2020,

**Notations:** Before calling any routine of the library grobcov, the user must define the ideal  $\mathbb{Q}[a][x]$ , and all the input polynomials and ideals defined on it.

Internally the routines define and use also other ideals:  $\mathbb{Q}[a]$ ,  $\mathbb{Q}[x,a]$  and so on.

**Procedures:**

grobcov(F)

Is the basic routine giving the canonical Groebner Cover of the parametric ideal F. This routine accepts many options, that allow to obtain results even when the canonical computation does not finish in reasonable time.

- cgsdr(F)** Is the procedure for obtaining a first disjoint, reduced Comprehensive Groebner System that is used in `grobcov`, but can also be used independently if only a CGS is required. It is a more efficient routine than `buildtree` (the own routine of 2010 that is no more available). Now, Kapur-Sun-Wang (KSW) algorithm is used.
- pdivi(f,F)**  
Performs a pseudodivision of a parametric polynomial by a parametric ideal.
- pnormalf(f,E,N)**  
Reduces a parametric polynomial  $f$  over  $V(E) \setminus V(N)$ .  $E$  is the null ideal and  $N$  the non-null ideal over the parameters.
- Crep(N,M)**  
Computes the canonical C-representation of  $V(N) \setminus V(M)$ . It can be called in  $\mathbb{Q}[a]$  or in  $\mathbb{Q}[a][x]$ , but the ideals  $N, M$  can only contain parameters of  $\mathbb{Q}[a]$ .
- Prep(N,M)**  
Computes the canonical P-representation of  $V(N) \setminus V(M)$ . It can be called in  $\mathbb{Q}[a]$  or in  $\mathbb{Q}[a][x]$ , but the ideals  $N, M$  can only contain parameters of  $\mathbb{Q}[a]$ .
- PtoCrep(L)**  
Starting from the canonical Prep of a locally closed set computes its Crep.
- extendpoly(f,p,q)**  
Given the generic representation  $f$  of an I-regular function  $F$  defined by  $\text{poly } f$  on  $V(p) \setminus V(q)$  it returns its full representation.
- extendGC(GC)**  
When the `grobcov` of an ideal has been computed with the default option ("ext",0) and the explicit option ("rep",2) (which is not the default), then one can call `extendGC(GC)` (and options) to obtain the full representation of the bases. With the default option ("ext",0) only the generic representation of the bases is computed, and one can obtain the full representation using `extendGC`.
- locus(G)** Special routine for determining the geometrical locus of points verifying given conditions. Given a parametric ideal  $J$  in  $\mathbb{Q}[x_1, \dots, x_n][u_1, \dots, u_m]$  with parameters  $(x_1, \dots, x_n)$  and variables  $(u_1, \dots, u_m)$ , representing the system determining the  $n$ -dimensional locus with tracer point  $(x_1, \dots, x_n)$  verifying certain properties, one can apply `locus` to the system  $F$ , for obtaining the locus. `locus` provides all the components of the locus and determines their taxonomy, that can be: "Normal", "Special", "Accumulation", "Degenerate". The mover are the  $u$ -variables. The user can eventually restrict them to a subset of them for geometrical reasons but this can change the true taxonomy.
- locusdg(G)**  
Is a special routine that determines the "Relevant" components of the locus in dynamic geometry. It is to be called to the output of `locus` and selects from it the "Normal", and "Accumulation" components.
- envelop(F,C)**

Special routine for determining the envelop of a family of hyper-surfaces  $F$  in  $Q[x_1, \dots, x_n][t_1, \dots, t_m]$  depending on an ideal of constraints  $C$  in  $Q[t_1, \dots, t_m]$ . It computes the locus of the envelop, and determines the different components as well as its taxonomy: "Normal", "Special", "Accumulation", "Degenerate". (See help for locus).

`locusto(L)`

Transforms the output of `locus`, `locusdg`, `envelop` into a string that can be read from different computational systems.

`stdlocus(F)`

Simple procedure to determine the components of the locus, alternative to `locus` that uses only standard GB computation. Cannot determine the taxonomy of the irreducible components.

`AssocTanToEnv(F, C, E)`

Having computed an envelop component  $E$  of a family of hyper-surfaces  $F$ , with constraints  $C$ , it returns the parameter values of the associated tangent hyper-surface of the family passing at one point of the envelop component  $E$ .

`FamElemsAtEnvCompPoints(F, C, E)`

Having computed an envelop component  $E$  of a family of hyper-surfaces  $F$ , with constraints  $C$ , it returns the parameter values of all the hyper-surfaces of the family passing at one point of the envelop component  $E$ .

`discrim(f, x)`

Determines the factorized discriminant of a degree 2 polynomial in the variable  $x$ . The polynomial can be defined on any ring where  $x$  is a variable. The polynomial  $f$  can depend on parameters and variables.

`WLemma(F, A)`

Given an ideal  $F$  in  $Q[a][x]$  and an ideal  $A$  in  $Q[a]$ , it returns the list  $(lpp, B, S)$  where  $B$  is the reduced Groebner basis of the specialized  $F$  over the segment  $S$ , subset of  $V(A)$  with top  $A$ , determined by Wibmer's Lemma.  $S$  is determined in  $P$ -representation (or optionally in  $C$ -representation). The basis is given by  $I$ -regular functions.

`WLCGS(F)` Given a parametric ideal  $F$  in  $Q[a][x]$  determines a CGS in full-representation using `WLemma`

`intersectpar(L)`

Auxiliary routine. Given a list of ideals defined on  $K[a][x]$  it determines the intersection of all of them in  $K[x, a]$

`ADGT(H, T, H1, T1)`

Given 4 ideals  $H, T, H1, T1$  in  $Q[a][x]$ , corresponding to a problem of Automatic Deduction of Geometric Theorems, it determines the supplementary conditions over the parameters for the Proposition  $(H \text{ and not } H1) \Rightarrow (T \text{ and not } T1)$  to be a Theorem. If  $H1=1$  then  $H1$  is not considered, and analogously for  $T1$ .

`ConsLevels(A)`

Given a list of locally closed sets, constructs the canonical representation of the levels of  $A$  and its complement.

`Levels(L)`

Transforms the output of `ConsLevels` into the proper `Levels` of the constructible set.

`Grob1Levels(G)`

From the output of `grobcov`, `Grob1Levels` selects the segments of `G` with basis different from 1 (having solutions), and determines the levels of the constructible set formed by them.

`DifConsLCSets(A,B)`

given the canonical forms of the constructible sets `A` and `B`,  $A=[a_1,a_2,\dots,a_k]$ ,  $B=[b_1,b_2,\dots,b_j]$ , `DifConsLCSets` returns a list of locally closed sets of the set `A` minus `B`, that can be transformed into the canonical form of `A` minus `B` applying `ConsLevels`.

See also: Section D.2.2 [`compregb_lib`], page 790.

## D.2.5 `inout_lib`

**Library:** `inout.lib`

**Purpose:** Printing and Manipulating In- and Output

**Procedures:**

`allprint(list)`

print list if `ALLprint` is defined, with pause if `>0`

`lprint(poly/...[,n])`

display poly/... fitting to pagewidth [size n]

`pmat(matrix[,n])`

print form-matrix [first n chars of each column]

`rMacaulay(string)`

read `Macaulay_1` output and return its Singular format

`show(any)`

display any object in a compact format

`showrecursive(id,p)`

display id recursively with respect to variables in p

`split(string,n)`

split given string into lines of length n

`tab(n)` string of n space tabs

`pause([prompt])`

stop the computation until user input

## D.2.6 `modular_lib`

**Library:** `modular.lib`

**Purpose:** An abstraction layer for modular techniques

**Author:** Andreas Steenpass, e-mail: [steenpass@mathematik.uni-kl.de](mailto:steenpass@mathematik.uni-kl.de)

**Overview:** This library is an abstraction layer for modular techniques which are well-known to speed up many computations and to be easy parallelizable.

The basic idea is to execute some computation modulo several primes and then to lift the result back to characteristic zero via the farey rational map and chinese remaindering. It is thus possible to overcome the often problematic coefficient swell and to run the modular computations in parallel.

In Singular, modular techniques have been quite successfully employed for several applications. A first implementation was done for Groebner bases in Singular's Section D.4.16 [modstd\_lib], page 818, a pioneering work by Stefan Steidel. Since the algorithm is basically the same for all applications, this library aims at preventing library authors from writing the same code over and over again by providing an appropriate abstraction layer. It also offers one-line commands for ordinary Singular users who want to take advantage of modular techniques for their own calculations. Thus modular techniques can be regarded as a parallel skeleton of their own.

The terminology (such as 'pTest' and 'finalTest') follows Singular's Section D.4.16 [modstd\_lib], page 818 and [1].

**References:**

[1] Nazeran Idrees, Gerhard Pfister, Stefan Steidel: Parallelization of Modular Algorithms. Journal of Symbolic Computation 46, 672-684 (2011). <http://arxiv.org/abs/1005.5663>

**Procedures:**

```
modular(...)
```

execute a command modulo several primes and lift the result back to characteristic zero

See also: Section D.4.3 [assprimeszerodim\_lib], page 810; Section 4.9 [link], page 94; Section D.4.16 [modstd\_lib], page 818; Section D.2.7 [parallel\_lib], page 797; Section D.2.13 [tasks\_lib], page 804.

## D.2.7 parallel\_lib

**Library:** parallel.lib

**Purpose:** An abstraction layer for parallel skeletons

**Author:** Andreas Steenpass, e-mail: [steenpass@mathematik.uni-kl.de](mailto:steenpass@mathematik.uni-kl.de)

**Overview:** This library provides implementations of several parallel 'skeletons' (i.e. ways in which parallel tasks rely upon and interact with each other). It is based on the library tasks.lib and aims at both ordinary Singular users as well as authors of Singular libraries.

**Procedures:**

```
parallelWaitN()
```

execute several jobs in parallel and wait for N of them to finish

```
parallelWaitFirst()
```

execute several jobs in parallel and wait for the first to finish

```
parallelWaitAll()
```

execute several jobs in parallel and wait for all of them to finish

```
parallelTestAND()
```

run several tests in parallel and determine if they all succeed

```
parallelTestOR()
```

run several tests in parallel and determine if any of them succeeds

See also: Section D.4.15 [modnormal\_lib], page 818; Section D.4.16 [modstd\_lib], page 818; Section D.2.11 [resources\_lib], page 802; Section D.2.13 [tasks\_lib], page 804.

## D.2.8 polylib\_lib

**Library:** polylib.lib

**Purpose:** Procedures for Manipulating Polys, Ideals, Modules

**Authors:** O. Bachmann, G.-M. Greuel, A. Fruehbis

**Procedures:**

`cyclic(int)`  
ideal of cyclic n-roots

`elemSymmId(int)`  
ideal of elementary symmetric polynomials

`katsura([i])`  
katsura [i] ideal

`freerank(poly/...)`  
rank of coker(input) if coker is free else -1

`is_zero(poly/...)`  
int, =1 resp. =0 if coker(input) is 0 resp. not

`lcm(ideal)`  
lcm of given generators of ideal

`maxcoef(poly/...)`  
maximal length of coefficient occurring in poly/...

`maxdeg(poly/...)`  
int/intmat = degree/s of terms of maximal order

`maxdeg1(poly/...)`  
int = [weighted] maximal degree of input

`mindeg(poly/...)`  
int/intmat = degree/s of terms of minimal order

`mindeg1(poly/...)`  
int = [weighted] minimal degree of input

`normalize(poly/...)`  
normalize poly/... such that leading coefficient is 1

`rad_con(p,I)`  
check radical containment of polynomial p in ideal I

`content(f)`  
content of polynomial/vector f

`mod2id(M,iv)`  
conversion of a module M to an ideal

`id2mod(i,iv)`  
conversion inverse to mod2id



```

substitute(I,...)
 substitute in I variables by polynomials

subrInterred(i1,i2,iv)
 interred w.r.t. a subset of variables

newtonDiag(f)
 Newton diagram of a polynomial

hilbPoly(I)
 Hilbert polynomial of basering/I

```

### D.2.9 redcgs\_lib

**Library:** redcgs.lib

**Purpose:** Reduced Comprehensive Groebner Systems.

**Overview:** Comprehensive Groebner Systems. Canonical Forms.

The library contains Monte's algorithms to compute disjoint, reduced Comprehensive Groebner Systems (CGS). A CGS is a set of pairs of (segment,basis). The segments  $S_i$  are subsets of the parameter space, and the bases  $B_i$  are sets of polynomials specializing to Groebner bases of the specialized ideal for every point in  $S_i$ .

The purpose of the routines in this library is to obtain CGS with better properties, namely disjoint segments forming a partition of the parameter space and reduced bases. Reduced bases are sets of polynomials that specialize to the reduced Groebner basis of the specialized ideal preserving the leading power products (lpp). The lpp characterize the type of solution in each segment.

A further objective is to summarize as much as possible the segments with the same lpp into a single segment, and if possible to obtain a final result that is canonical, i.e. independent of the algorithm and only attached to the given ideal.

There are three fundamental routines in the library: mrcgs, rcgs and crcgs. mrcgs (Minimal Reduced CGS) is an algorithm that packs so much as it is able to do (using algorithms adhoc) the segments with the same lpp, obtaining the minimal number of segments. The hypothesis is that the result is also canonical, but for the moment there is no proof of the uniqueness of this minimal packing. Moreover, the segments that are obtained are not locally closed, i.e. there are not difference of two varieties.

On the other side, Michael Wibmer has proved that for homogeneous ideals, all the segments with reduced bases having the same lpp admit a unique basis specializing well. For this purpose it is necessary to extend the description of the elements of the bases to functions, forming sheaves of polynomials instead of simple polynomials, so that the polynomials in a sheaf either preserve the lpp of the corresponding polynomial of the specialized Groebner basis (and then it specializes well) or it specializes to 0. Moreover, in a sheaf, for every point in the corresponding segment, at least one of the polynomials specializes well. And moreover Wibmer's Theorem ensures that the packed segments are locally closed, that is can be described as the difference of two varieties.

Using Wibmer's Theorem we proved that an affine ideal can be homogenized, than discussed by mrcgs and finally de-homogenized. The bases so obtained can be reduced and specialize well in the segment. If the theoretic objective is reached, and all the segments of the homogenized ideal have been packed, locally closed segments will be obtained.

If we only homogenize the given basis of the ideal, then we cannot ensure the canonicity of the partition obtained, because there are many different bases of the given ideal that can be homogenized, and the homogenized ideals are not identical. This corresponds to the algorithm `rcgs` and is recommended as the most practical routine. It provides locally closed segments and is usually faster than `mrcgs` and `crcgs`. But the given partition is not always canonical.

Finally it is possible to homogenize the whole affine ideal, and then the packing algorithm will provide canonical segments by dehomogenizing. This corresponds to `crcgs` routine. It provides the best description of the segments and bases. In contrast `crcgs` algorithm is usually much more time consuming and it will not always finish in a reasonable time. Moreover it will contain more segments than `mrcgs` and possibly also more than `rcgs`.

But the actual algorithms in the library to pack segments have some lacks. They are not theoretically always able to pack the segments that we know that can be packed. Nevertheless, thanks to Wibmer's Theorem, the algorithms `rcgs` and `crcgs` are able to detect if the objective has not been reached, and if so, to give a Warning. The warning does not invalidate the output, but it only recognizes that the theoretical objective is not completely reached by the actual computing methods and that some segments that can be packed have not been packed with a single basis.

The routine `buildtree` is the first algorithm used in all the previous methods providing a first disjoint CGS, and can be used if none of the three fundamental algorithms of the library finishes in a reasonable time.

There are also routines to visualize better the output of the previous algorithms: `finalcases` can be applied to the list provided by `buildtree` to obtain the CGS. The list provided by `buildtree` contains the whole discussion, and `finalcases` extracts the CGS. The output of `buildtree` can also be transformed into a file using `buildtreeToMaple` routine that can be read in Maple. Using Monte's `dpgb` library in Maple the output can be plotted (with the routine `tplot`). To plot the output of `mrcgs`, `rcgs` or `crcgs` in Maple, the library also provides the routine `cantreeToMaple`. The file written using it and read in Maple can then be plotted with the command `plotcantree` and printed with `printcantree` from the Monte's `dpgb` library in Maple. The output of `mrcgs`, `rcgs` and `crcgs` is given in form of tree using prime ideals in a canonical form that is described in the papers. Nevertheless this canonical form is somewhat uncomfortable to be interpreted. When the segments are all locally closed (and this is always the case for `rcgs` and `crcgs`) the routine `cantodiffcgs` transforms the output into a simpler form having only one list element for each segment and providing the two varieties whose difference represent the segment also in a canonical form.

**Authors:** Antonio Montes , Hans Schoenemann.

**Overview:** see "Minimal Reduced Comprehensive Groebner Systems" by Antonio Montes. (<http://www-ma2.upc.edu/~montes/>).

**Notations:** All given and determined polynomials and ideals are in the basering  $K[a][x]$ ; (a=parameters, x=variables)  
 After defining the ring and calling `setglobalrings()`; the rings  
 $@R (K[a][x])$ ,  
 $@P (K[a])$ ,  
 $@RP (K[x,a])$  are defined globally  
 They are used internally and can also be used by the user.  
 The fundamental routines are: `buildtree`, `mrcgs`, `rcgs` and `crcgs`

**Procedures:**

- setglobalrings()**  
It is called by the fundamental routines of the library: (buildtree, mrcgs, rcgs, crcgs). After calling it, the rings  $\mathbb{C}R$ ,  $\mathbb{C}P$  and  $\mathbb{C}RP$  are defined globally.
- memberpos(f, J)**  
Returns the list of two integers: the value 0 or 1 depending on if  $f$  belongs to  $J$  or not, and the position in  $J$  (0 if it does not belong).
- subset(F, G)**  
If all elements of  $F$  belong to the ideal  $G$  it returns 1, and 0 otherwise.
- pdivi2(f, F)**  
Pseudodivision of a polynomial  $f$  by an ideal  $F$  in  $\mathbb{C}R$ . Returns a list  $(r, q, m)$  such that  $m \cdot f = r + \text{sum}(q \cdot G)$ .
- facvar(ideal J)**  
Returns all the free-square factors of the elements of ideal  $J$  (non repeated). Integer factors are ignored, even 0 is ignored. It can be called from ideal  $\mathbb{C}R$ , but the given ideal  $J$  must only contain polynomials in the parameters.
- redspec(N, W)**  
Given null and non-null conditions depending only on the parameters it returns a red-specification.
- pnormalform(f, N, W)**  
Reduces the polynomial  $f$  w.r.t. to the null condition ideal  $N$  and the non-null condition ideal  $W$  (both depending on the parameters).
- buildtree(F)**  
Returns a list  $T$  describing a first reduced CGS of the ideal  $F$  in  $K[a][x]$ .
- buildtreetoMaple(T)**  
Writes into a file the output of buildtree in Maple readable form.
- finalcases(T)**  
From the output of buildtree it provides the list of its terminal vertices. That list represents the dichotomic, reduced CGS obtained by buildtree.
- mrcgs(F)** Returns a list  $T$  describing the Minimal Reduced CGS of the ideal  $F$  of  $K[a][x]$
- rcgs(F)** Returns a list  $T$  describing the Reduced CGS of the ideal  $F$  of  $K[a][x]$  obtained by direct homogenizing and de-homogenizing the basis of the given ideal.
- crcgs(F)** Returns a list  $T$  describing the Canonical Reduced CGS of the ideal  $F$  of  $K[a][x]$  obtained by homogenizing and de-homogenizing the initial ideal. cantreetoMaple)(M); Writes into a file the output of mrcgs, rcgs or crcgs in Maple readable form.
- cantodiffcgs(list L)**  
From the output of rcgs or crcgs (or even of mrcgs when it is possible) it returns a simpler list where the segments are given as difference of varieties.

See also: Section D.2.2 [compregb\_lib], page 790.

### D.2.10 random\_lib

**Library:** random.lib

**Purpose:** Creating Random and Sparse Matrices, Ideals, Polys

**Procedures:**

`genericid(i[,p,b])`  
generic sparse linear combinations of generators of  $i$

`randomid(id,[k,b])`  
random linear combinations of generators of  $id$

`randommat(n,m[,id,b])`  
 $n \times m$  matrix of random linear combinations of  $id$

`sparseid(k,u[,o,p,b])`  
ideal of  $k$  random sparse poly's of degree  $d$  [ $u \leq d \leq o$ ]

`sparsematrix(n,m,o[,.])`  
 $n \times m$  sparse matrix of polynomials of degree  $\leq o$

`sparsemat(n,m[,p,b])`  
 $n \times m$  sparse integer matrix with random coefficients

`sparsepoly(u[,o,p,b])`  
random sparse polynomial with terms of degree in  $[u,o]$

`sparsetriag(n,m[,.])`  
 $n \times m$  sparse lower-triag intmat with random coefficients

`sparseHomogIdeal(k,u[,.])`  
ideal with  $k$  sparse homogeneous generators of degree in  $[u,o]$

`triagmatrix(n,m,o[,.])`  
 $n \times m$  sparse lower-triag matrix of poly's of degree  $\leq o$

`randomLast(b)`  
random transformation of the last variable

`randomBinomial(k,u,..)`  
binomial ideal,  $k$  random generators of degree  $\geq u$

### D.2.11 resources\_lib

**Library:** resources.lib

**Purpose:** Tools to manage the computational resources

**Author:** Andreas Steenpass, e-mail: steenpass@mathematik.uni-kl.de

**Overview:** The purpose of this library is to manage the computational resources of a Singular session. The library `tasks.lib` and any library build upon `tasks.lib` respect these settings, i.e. they will not use more computational resources than provided via `resources.lib`.

The provided procedures and their implementation are currently quite simple. The library can be extended later on to support, e.g., distributed computations on several servers.

**Procedures:**

`addcores()`  
 add an integer to the number of available processor cores  
`setcores()`  
 set the number of available processor cores  
`getcores()`  
 get the number of available processor cores  
`semaphore()`  
 initialize a new semaphore

See also: Section D.2.7 [`parallel.lib`], page 797; Section D.2.13 [`tasks.lib`], page 804.

## D.2.12 `ring.lib`

**Library:** `ring.lib`

**Purpose:** Manipulating Rings and Maps

**Authors:** Singular team

**Procedures:**

`changechar(c[,r])`  
 make a copy of basering [ring r] with new char c  
`changeord(o[,r])`  
 make a copy of basering [ring r] with new ord o  
`changevar(v[,r])`  
 make a copy of basering [ring r] with new vars v  
`defring("R",c,n,v,o)`  
 define a ring R in specified char c, n vars v, ord o  
`defrings(n[,p])`  
 define ring  $S_n$  in n vars, char 32003 [p], ord ds  
`defringp(n[,p])`  
 define ring  $P_n$  in n vars, char 32003 [p], ord dp  
`extendring("R",n,v,o)`  
 extend given ring by n vars v, ord o and name it R  
`fetchall(R[,str])`  
 fetch all objects of ring R to basering  
`imapall(R[,str])`  
 imap all objects of ring R to basering  
`mapall(R,i[,str])`  
 map all objects of ring R via ideal i to basering  
`ord_test(R)`  
 test whether ordering of R is global, local or mixed  
`ringtensor(s,t,...)`  
 create ring, tensor product of rings s,t,...  
`ringweights(r)`  
 intvec of weights of ring variables of ring r

`preimageLoc(R,phi,Q)`  
 computes preimage for non-global orderings

`rootofUnity(n)`  
 the minimal polynomial for the n-th primitive root of unity (parameters in square brackets [] are optional)

`optionIsSet(opt)`  
 check if as a string given option is set or not. `hasFieldCoefficient` check if the coefficient ring is considered a field `hasGFCoefficient` check if the coefficient ring is  $\text{GF}(p,k)$  `hasZpCoefficient` check if the coefficient ring is  $\mathbb{Z}\mathbb{Z}/p$  `hasZp_aCoefficient` check if the coefficient ring is an elag. ext. of  $\mathbb{Z}\mathbb{Z}/p$  `hasQQCoefficient` check if the coefficient ring is  $\mathbb{Q}\mathbb{Q}$

`hasNumericCoeffs(rng)`  
 check for use of floating point numbers

`hasCommutativeVars(rng)`  
 non-commutative or commutative polynomial ring

`hasGlobalOrdering(rng)`  
 global versus mixed/local monomial ordering

`hasMixedOrdering()`  
 mixed versus global/local ordering

`hasAlgExtensionCoefficient(r)`  
 coefficients are an algebraic extension

`hasTransExtensionCoefficient(r)`  
 coefficients are rational functions

`isQuotientRing(rng)`  
 ring is a quotient ring

`isSubModule(I,J)`  
 check if I is in J as submodule

`changeordTo(r,o)`  
 change the ordering of a ring to a simple one

`addvarsTo(r,vars,i)`  
 add variables to a ring

`addNvarsTo(r,N,name,i)`  
 add N variables to a ring

### D.2.13 tasks.lib

**Library:** tasks.lib

**Purpose:** A parallel framework based on tasks

**Author:** Andreas Steenpass, e-mail: steenpass@mathematik.uni-kl.de

**Overview:** This library provides a parallel framework based on tasks. It introduces a new Singular type `task`; an object of this type is a command (given by a string) applied to a list of arguments. Tasks can be computed in parallel via the procedures in this library and they can even be started recursively, i.e. from within other tasks.

tasks.lib respects the limits for computational resources defined in Section D.2.11 [resources.lib], page 802, i.e., all tasks within the same Singular session will not use more computational resources than provided via resources.lib, even if tasks are started recursively.

The Singular library Section D.2.7 [parallel.lib], page 797 provides implementations of several parallel 'skeletons' based on tasks.lib.

**Procedures:**

```
createTask()
 create a task

killTask()
 kill a task

copyTask()
 copy a task

compareTasks()
 compare two tasks

printTask()
 print a task

startTasks()
 start tasks

stopTask()
 stop a task

waitTasks()
 wait for a certain number of tasks

waitAllTasks()
 wait for all tasks

pollTask()
 poll a task

getCommand()
 get the command of a task

getArguments()
 get the arguments of a task

getResult()
 get the result of a task

getState()
 get the state of a task
```

See also: Section D.2.7 [parallel.lib], page 797; Section D.2.11 [resources.lib], page 802.

### D.3 Linear algebra

### D.3.1 matrix\_lib

**Library:** matrix.lib

**Purpose:** Elementary Matrix Operations

**Procedures:**

`compress(A)`  
matrix, zero columns from A deleted

`concat(A1,A2,...)`  
matrix, concatenation of matrices A1,A2,...

`diag(p,n)`  
matrix, nxn diagonal matrix with entries poly p

`dsum(A1,A2,...)`  
matrix, direct sum of matrices A1,A2,...

`flatten(A)`  
ideal, generated by entries of matrix A

`genericmat(n,m[,id])`  
generic nxm matrix [entries from id]

`is_complex(c)`  
1 if list c is a complex, 0 if not

`outer(A,B)`  
matrix, outer product of matrices A and B

`power(A,n)`  
matrix/intmat, n-th power of matrix/intmat A

`skewmat(n[,id])`  
generic skew-symmetric nxn matrix [entries from id]

`submat(A,r,c)`  
submatrix of A with rows/cols specified by intvec r/c

`symmat(n[,id])`  
generic symmetric nxn matrix [entries from id]

`unitmat(n)`  
unit square matrix of size n

`gauss_col(A)`  
transform a matrix into col-reduced Gauss normal form

`gauss_row(A)`  
transform a matrix into row-reduced Gauss normal form

`addcol(A,c1,p,c2)`  
add  $p \cdot$  (c1-th col) to c2-th column of matrix A, p poly

`addrow(A,r1,p,r2)`  
add  $p \cdot$  (r1-th row) to r2-th row of matrix A, p poly

`multcol(A,c,p)`  
multiply c-th column of A with poly p



`multrow(A,r,p)`  
 multiply r-th row of A with poly p  
`permcop(A,i,j)`  
 permute i-th and j-th columns  
`permrow(A,i,j)`  
 permute i-th and j-th rows  
`rowred(A[,any])`  
 reduction of matrix A with elementary row-operations  
`colred(A[,any])`  
 reduction of matrix A with elementary col-operations  
`linear_relations(E)`  
 find linear relations between homogeneous vectors  
`rm_unitrow(A)`  
 remove unit rows and associated columns of A  
`rm_unitcol(A)`  
 remove unit columns and associated rows of A  
`headStand(A)`  
 $A[n-i+1,m-j+1]:=A[i,j]$   
`symmetricBasis(n,k[,s])`  
 basis of k-th symmetric power of n-dim v.space  
`exteriorBasis(n,k[,s])`  
 basis of k-th exterior power of n-dim v.space  
`symmetricPower(A,k)`  
 k-th symmetric power of a module/matrix A  
`exteriorPower(A,k)`  
 k-th exterior power of a module/matrix A

### D.3.2 linalg\_lib

**Library:** linalg.lib

**Purpose:** Algorithmic Linear Algebra

**Authors:** Ivor Saynisch (ivs@math.tu-cottbus.de)  
 Mathias Schulze (mschulze@mathematik.uni-kl.de)

**Procedures:**

`inverse(A)`  
 matrix, the inverse of A  
`inverse_B(A)`  
 list(matrix Inv,poly p), $Inv*A=p*En$  ( using busadj(A) )  
`inverse_L(A)`  
 list(matrix Inv,poly p), $Inv*A=p*En$  ( using lift )  
`sym_gauss(A)`  
 symmetric gaussian algorithm

`orthogonalize(A)`  
 Gram-Schmidt orthogonalization

`diag_test(A)`  
 test whether A can be diagonalized

`busadj(A)`  
 coefficients of  $\text{Adj}(E^*t-A)$  and coefficients of  $\det(E^*t-A)$

`charpoly(A,v)`  
 characteristic polynomial of A ( using `busadj(A)` )

`adjoint(A)`  
 adjoint of A ( using `busadj(A)` )

`det_B(A)` determinant of A ( using `busadj(A)` )

`gaussred(A)`  
 gaussian reduction:  $P^*A=U^*S$ , S a row reduced form of A

`gaussred_pivot(A)`  
 gaussian reduction:  $P^*A=U^*S$ , uses row pivoting

`gauss_nf(A)`  
 gaussian normal form of A

`mat_rk(A)`  
 rank of constant matrix A

`U_D_O(A)`  $P^*A=U^*D^*O$ , P,D,U,O=permutation,diag,lower-,upper-triang

`pos_def(A,i)`  
 test symmetric matrix for positive definiteness

`hessenberg(M)`  
 Hessenberg form of M

`eigenvals(M)`  
 eigenvalues with multiplicities of M

`minipoly(M)`  
 minimal polynomial of M

`spnf(sp)` normal form of spectrum sp

`spprint(sp)`  
 print spectrum sp

`jordan(M)`  
 Jordan data of M

`jordanbasis(M)`  
 Jordan basis and weight filtration of M

`jordanmatrix(jd)`  
 Jordan matrix with Jordan data jd

`jordannf(M)`  
 Jordan normal form of M

## D.4 Commutative algebra

### D.4.1 `absfact.lib`

**Library:** `absfact.lib`

**Purpose:** Absolute factorization for characteristic 0

**Authors:** Wolfram Decker, `decker@math.uni-sb.de`  
 Gregoire Lecerf, `lecerf@math.uvsq.fr`  
 Gerhard Pfister, `pfister@mathematik.uni-kl.de` Martin Lee, `mlee@mathematik.uni-kl.de`

**Overview:** A library for computing the absolute factorization of multivariate polynomials  $f$  with coefficients in a field  $K$  of characteristic zero. Using Trager's idea, the implemented algorithm computes an absolutely irreducible factor by factorizing over some finite extension field  $L$  (which is chosen such that  $V(f)$  has a smooth point with coordinates in  $L$ ). Then a minimal extension field is determined making use of the Rothstein-Trager partial fraction decomposition algorithm. `absFactorizeBCG` uses the algorithm of Bertone, Cheze and Galligo for bivariate polynomials and similar ideas as above to reduce to this case.

**References:**

G. Cheze, G. Lecerf: Lifting and recombination techniques for absolute factorization. *Journal of Complexity*, 23(3):380-420, 2007. C. Bertone, G. Cheze, and A. Galligo: Modular las vegas algorithms for polynomial absolute factorization. *J. Symb. Comput.*, 45(12):1280-1295, December 2010

**Procedures:**

`absFactorize()`  
 absolute factorization of poly

`absFactorizeBCG()`  
 absolute factorization of poly

See also: Section 5.1.36 [`factorize`], page 178.

### D.4.2 `algebra.lib`

**Library:** `algebra.lib`

**Purpose:** Compute with Algebras and Algebra Maps

**Authors:** Gert-Martin Greuel, `greuel@mathematik.uni-kl.de`,  
 Agnes Eileen Heydtmann, `agnes@math.uni-sb.de`,  
 Gerhard Pfister, `pfister@mathematik.uni-kl.de`

**Procedures:**

`algebra_containment()`  
 query of algebra containment

`module_containment()`  
 query of module containment over a subalgebra

`inSubring(p,I)`  
 test whether polynomial  $p$  is in subring generated by  $I$

`algDependent(I)`  
 computes algebraic relations between generators of  $I$   
`alg_kernel(phi)`  
 computes the kernel of the ringmap  $\phi$   
`is_injective(phi)`  
 test for injectivity of ringmap  $\phi$   
`is_surjective(phi)`  
 test for surjectivity of ringmap  $\phi$   
`is_bijective(phi)`  
 test for bijectivity of ring map  $\phi$   
`noetherNormal(id)`  
 noether normalization of ideal  $id$   
`mapIsFinite(R,phi,I)`  
 query for finiteness of map  $\phi:R \rightarrow \text{basering}/I$   
`finitenessTest(i,z)`  
 find variables which occur as pure power in  $\text{lead}(i)$   
`nonZeroEntry(id)`  
 list describing non-zero entries of an identifier

### D.4.3 `assprimeszerodim_lib`

**Library:** `assprimeszerodim.lib`

**Purpose:** associated primes of a zero-dimensional ideal

**Authors:** N. Idrees nazeranjawwad@gmail.com  
 G. Pfister pfister@mathematik.uni-kl.de  
 A. Steenpass steenpass@mathematik.uni-kl.de  
 S. Steidel steidel@mathematik.uni-kl.de

**Overview:** A library for computing the associated primes and the radical of a zero-dimensional ideal in the polynomial ring over the rational numbers,  $\mathbb{Q}[x_1, \dots, x_n]$ , using modular computations.

**Procedures:**

`zeroRadical(I)`  
 computes the radical of  $I$   
`assPrimes(I)`  
 computes the associated primes of  $I$

See also: Section D.4.26 [`primdec_lib`], page 828.

### D.4.4 `cisimplicial_lib`

**Library:** `cisimplicial.lib`

**Purpose:** . Determines if the toric ideal of a simplicial toric variety is a complete intersection

**Authors:** I.Bermejo, ibermejo@ull.es  
 I.Garcia-Marco, iggarcia@ull.es

**Overview:** A library for determining if a simplicial toric ideal is a complete intersection with NO NEED of computing explicitly a system of generators of such ideal. The procedures are based on two papers: I. Bermejo, I. Garcia-Marco and J.J. Salazar-Gonzalez: 'An algorithm for checking whether the toric ideal of an affine monomial curve is a complete intersection', J. Symbolic Computation 42 (2007) pags: 971–991 and I. Bermejo and I. Garcia-Marco: 'Complete intersections in simplicial toric varieties', Preprint (2010)

**Procedures:**

`minMult(a,b)`  
 computes the minimum multiple of  $a$  that belongs to the semigroup generated by  $b$

`belongSemigroup(v,A[,n])`  
 checks whether  $A \cdot x = v$  has a nonnegative integral solution

`oneDimBelongSemigroup(n,v[,m])`  
 checks whether  $v \cdot x = n$  has a nonnegative integral solution

`cardGroup(A)`  
 computes the cardinal of  $Z^m / ZA$

`isCI(A)` checks whether  $I(A)$  is a complete intersection

### D.4.5 curveInv\_lib

**Library:** curveInv.lib

**Purpose:** A library for computing invariants of curves

**Author:** Peter Chini, chini@rhrk.uni-kl.de

**Overview:** This library provides a collection of procedures for computing invariants of curve singularities. Invariants that can be computed are: - the delta invariant  
 - the multiplicity of the conductor: the length of  $\text{Normalization}(R)/C$ , where  $C$  denotes the conductor  
 - the Deligne number  
 - the colength of derivations along the normalization - the length of  $\text{Der}(\text{Normalization}(R/I)) / R/I$

In addition, it is possible to compute the conductor of a ring  $S = R/I$ , where  $R$  is a (localized) polynomial ring.

**Theory:** Computing the Deligne number of curve singularities and an algorithmic framework for differential algebras in SINGULAR;  
 Chapter 5 - Master's Thesis of Peter Chini - August 2015

**Procedures:**

`curveDeltaInv(ideal)`  
 computes the delta invariant of  $R/I$  for a given ideal  $I$

`curveConductorMult(ideal)`  
 returns the multiplicity of the conductor of  $R/I$

`curveDeligneNumber(ideal)`  
 computes the Deligne number of  $R/I$

`curveColengthDerivations(ideal)`  
 returns the colength of derivations, the length of  $\text{Der}(\text{Normalization}(R/I))/\text{Der}(R/I)$

### D.4.6 elim\_lib

**Library:** elim.lib

**Purpose:** Elimination, Saturation and Blowing up

**Procedures:**

```

blowup0(j[,s1,s2])
 create presentation of blowup ring of ideal j

elimRing(p)
 create ring with block ordering for eliminating vars in p

elim(id,..)
 variables .. eliminated from id (ideal/module)

elim1(id,p)
 variables .. eliminated from id (different algorithm)

elim2(id,..)
 variables .. eliminated from id (different algorithm)

nselect(id,v)
 select generators not containing variables given by v

sat(id,j)
 saturated quotient of ideal/module id by ideal j

select(id,v)
 select generators containing all variables given by v

select1(id,v)
 select generators containing one variable given by v

```

### D.4.7 ellipticcovers\_lib

**Library:** ellipticCovers.lib

**Purpose:** Gromov-Witten numbers of elliptic curves

**Authors:** J. Boehm, boehm @ mathematik.uni-kl.de  
 A. Buchholz, buchholz @ math.uni-sb.de  
 H. Markwig hannah @ math.uni-sb.de

**Overview:** We implement a formula for computing the number of covers of elliptic curves. It has been obtained by proving mirror symmetry for arbitrary genus by tropical methods in [BBM]. A Feynman graph of genus  $g$  is a trivalent, connected graph of genus  $g$  (with  $2g-2$  vertices and  $3g-3$  edges). The branch type  $b=(b_1,\dots,b_{(3g-3)})$  of a stable map is the multiplicity of the edge  $i$  over a fixed base point.

Given a Feynman graph  $G$  and a branch type  $b$ , we obtain the number  $N_-(G,b)$  of stable maps of branch type  $b$  from a genus  $g$  curve of topological type  $G$  to the elliptic curve by computing a path integral

over a rational function. The path integral is computed as a residue.

The sum of  $N_-(G,b)$  over all branch types  $b$  of sum  $d$  gives  $N_-(G,d)*|\text{Aut}(G)|$ , with the Gromov-Witten invariant  $N_-(G,d)$  of degree  $d$  stable maps from a genus  $g$  curve of topological type  $G$  to the elliptic curve.

The sum of  $N_-(G,d)$  over all such graphs gives the usual Gromov-Witten invariant  $N_-(g,d)$  of degree  $d$  stable maps from a genus  $g$  curve to the elliptic curve.

The key function computing the numbers  $N_-(G,b)$  and  $N_-(G,d)$  is `gromovWitten`.

**References:**

[BBM] J. Boehm, A. Buchholz, H. Markwig: Tropical mirror symmetry for elliptic curves, arXiv:1309.5893 (2013).

**Types:** graph

**Procedures:**

`makeGraph(list, list)`  
generate a graph from a list of vertices and a list of edges

`printGraph(graph)`  
print procedure for graphs

`propagator(list, int)`  
propagator factor of degree  $d$  in the quotient of two variables, or propagator for fixed graph and branch type

`computeConstant(number, number)`  
constant coefficient in the Laurent series expansion of a rational function in a given variable

`evalutateIntegral(number, list)`  
path integral for a given propagator and ordered sequence of variables

`gromovWitten(number)`  
sum of path integrals for a given propagator over all orderings of the variables, or Gromov Witten invariant for a given graph and a fixed branch type, or list of Gromov Witten invariants for a given graph and all branch types

`computeGromovWitten(graph, int, int)`  
compute the Gromov Witten invariants for a given graph and some branch types  
`generatingFunction (graph, int)` multivariate generating function for the Gromov Witten invariants of a graph up to fixed degree

`partitions(int, int)`  
partitions of an integer into a fixed number of summands

`permute(list)`  
all permutations of a list

`lsum(list)`  
sum of the elements of a list

### D.4.8 fmodstd\_lib

**Library:** fmodstd.lib

**Purpose:** Groebner bases of ideals in polynomial rings over rational function fields

**Authors:** D.K. Boku boku@mathematik.uni-kl.de  
W. Decker decker@mathematik.uni-kl.de  
C. Fieker fieker@mathematik.uni-kl.de  
A. Steenpass steenpass@mathematik.uni-kl.de

**Overview:** A library for computing a Groebner basis of an ideal in a polynomial ring over an algebraic function field  $Q(T) := Q(t_1, \dots, t_m)$  using modular methods and sparse multivariate rational interpolation, where the  $t_i$  are transcendental over  $Q$ . The idea is as follows: Given an ideal  $I$  in  $Q(T)[X]$ , we map  $I$  to  $J$  via the map sending  $T$  to  $Tz := (t_1z + s_1, \dots, t_mz + s_m)$  for a suitable point  $s$  in  $Q^m \setminus \{(0, \dots, 0)\}$  and for some extra variable  $z$  so that  $J$  is an ideal in  $Q(Tz)[X]$ . For a suitable point  $b$  in  $Z^m \setminus \{(0, \dots, 0)\}$ , we map  $J$  to  $K$  via the map sending  $(T, z)$  to  $(b, z)$ , where  $b := (b_1, \dots, b_m)$  (usually the  $b_i$ 's are distinct primes), so that  $K$  is an ideal in  $Q(z)[X]$ . For such a rational point  $b$ , we compute a Groebner basis  $G_b$  of  $K$  using modular algorithms [1], where prime numbers are replaced by maximal ideals of the form  $\langle z - z_i \rangle$ , and univariate rational interpolation [2,7]. Note that since  $Q[z]/\langle z - z_i \rangle = Q$  we also use (if required) modular algorithms [1] over  $Q$ . The procedure is repeated for many rational points  $b$  until their number is sufficiently large to recover the correct coefficients in  $Q(T)$ . Once we have these points, we obtain a set of polynomials  $G$  by applying the sparse multivariate rational interpolation algorithm from [4] coefficient-wise to the list of Groebner bases  $G_b$  in  $Q(z)[X]$ , where this algorithm makes use of the following algorithms: univariate polynomial interpolation [2], univariate rational function reconstruction [7], and multivariate polynomial interpolation [3]. The last algorithm uses the well-known Berlekamp/Massey algorithm [5] and its early termination version [6]. The set  $G$  is then a Groebner basis of  $I$  with high probability.

#### References:

- [1] E. A. Arnold: Modular algorithms for computing Groebner bases. *J. Symb. Comput.* 35, 403-419 (2003).
- [2] R. L. Burden and J. D. Faires: *Numerical analysis*. 9th ed. (1993).
- [3] M. Ben-Or and P. Tiwari: A deterministic algorithm for sparse multivariate polynomial interpolation. *Proc. of the 20th Annual ACM Symposium on Theory of Computing*, 301-309 (1988).
- [4] A. Cuyt and W.-s. Lee: Sparse interpolation of multivariate rational functions. *Theor. Comput. Sci.* 412, 1445-1456 (2011).
- [5] E. Kaltofen and W.-s. Lee: Early termination in sparse interpolation algorithms. *J. Symb. Comput.* 36, 365-400 (2003).
- [6] E. Kaltofen, W.-s. Lee and A. A. Lobo: Early termination in Ben-Or/Tiwari sparse interpolation and a hybrid of Zippel's algorithm. *Proc. ISSAC (ISSAC '00)*, 192-201 (2000).
- [7] K. Sara and M. Monagan: Fast Rational Function Reconstruction. *Proc. ISSAC (ISSAC '06)*, 184-190 (2006).

#### Procedures:

```

fareypoly(g,f)
 univariate rational function reconstruction

polyInterpolation(l,m)
 univariate polynomial interpolation

modrationalInterpolation(l,m)
 modular univariate rational interpolation

BerlekampMassey(L,i)
 Berlekamp/Massey algorithm

modberlekampMassey(L,i)
 modular Berlekamp/Massey algorithm

```



`sparseInterpolation(f,L,n)`  
 sparse multivariate polynomial interpolation

`ffmodStd(I)`  
 Groebner bases over algebraic function fields using modular methods and sparse multivariate rational interpolation

### D.4.9 grwalk\_lib

**Library:** grwalk.lib

**Purpose:** Groebner Walk Conversion Algorithms

**Author:** I Made Sulandra

**Procedures:**

`fwalk(ideal[,intvec])`  
 standard basis of ideal via fractalwalk alg

`twalk(ideal[,intvec])`  
 standard basis of ideal via Tran's alg

`awalk1(ideal[,intvec])`  
 standard basis of ideal via the first alt. alg

`awalk2(ideal[,intvec])`  
 standard basis of ideal via the second alt. alg

`pwalk(ideal[,intvec])`  
 standard basis of ideal via perturbation walk alg

`gwalk(ideal[,intvec])`  
 standard basis of ideal via groebnerwalk alg

See also: Section D.15.18 [rwalk.lib], page 940; Section D.15.21 [swalk.lib], page 942.

### D.4.10 homolog\_lib

**Library:** homolog.lib

**Purpose:** Procedures for Homological Algebra

**Authors:** Gert-Martin Greuel, greuel@mathematik.uni-kl.de,  
 Bernd Martin, martin@math.tu-cottbus.de  
 Christoph Lossen, lossen@mathematik.uni-kl.de

**Procedures:**

`canonMap(id)`  
 the kernel and the cokernel of the canonical map

`cup(M)` `cup: Ext1(M',M') x Ext1() -> Ext2()`

`cupproduct(M,N,P,p,q)`  
`cup: Extp(M',N') x Extq(N',P') -> Extp+q(M',P')`

`depth(I,M)`  
 depth(I,M'), I ideal, M module, M'=coker(M)

`Ext_R(k,M)`  
 Ext<sup>k</sup>(M',R), M module, R basering, M'=coker(M)

`Ext(k,M,N)`  
 $\text{Ext}^k(M',N')$ ,  $M,N$  modules,  $M'=\text{coker}(M)$ ,  $N'=\text{coker}(N)$

`fitting(M,n)`  
 $n$ -th Fitting ideal of  $M'=\text{coker}(M)$ ,  $M$  module,  $n$  int

`flatteningStrat(M)`  
 Flattening stratification of  $M'=\text{coker}(M)$ ,  $M$  module

`Hom(M,N)`  $\text{Hom}(M',N')$ ,  $M,N$  modules,  $M'=\text{coker}(M)$ ,  $N'=\text{coker}(N)$

`homology(A,B,M,N)`  
 $\ker(B)/\text{im}(A)$ , homology of complex  $R^k \rightarrow M' \rightarrow B \rightarrow N'$

`isCM(M)` test if  $\text{coker}(M)$  is Cohen-Macaulay,  $M$  module

`isFlat(M)`  
 test if  $\text{coker}(M)$  is flat,  $M$  module

`isLocallyFree(M,r)`  
 test if  $\text{coker}(M)$  is locally free of constant rank  $r$

`isReg(I,M)`  
 test if  $I$  is  $\text{coker}(M)$ -sequence,  $I$  ideal,  $M$  module

`hom_kernel(A,M,N)`  
 $\ker(M' \rightarrow N')$   $M,N$  modules,  $A$  matrix

`kohom(A,k)`  
 $\text{Hom}(R^k, A)$ ,  $A$  matrix over basering  $R$

`kontrahom(A,k)`  
 $\text{Hom}(A, R^k)$ ,  $A$  matrix over basering  $R$

`KoszulHomology(I,M,n)`  
 $n$ -th Koszul homology  $H_n(I, \text{coker}(M))$ ,  $I$ =ideal

`tensorMod(M,N)`  
 Tensor product of modules  $M'=\text{coker}(M)$ ,  $N'=\text{coker}(N)$

`Tor(k,M,N)`  
 $\text{Tor}_k(M',N')$ ,  $M,N$  modules,  $M'=\text{coker}(M)$ ,  $N'=\text{coker}(N)$

#### D.4.11 integralbasis\_lib

**Library:** integralbasis.lib

**Purpose:** Integral basis in algebraic function fields

**Authors:** J. Boehm, j.boehm at mx.uni-saarland.de  
 W. Decker, decker at mathematik.uni-kl.de  
 S. Laplagne, slaplagn at dm.uba.ar  
 F. Seelisch, seelisch at mathematik.uni-kl.de

**Overview:** Given an irreducible polynomial  $f$  in two variables defining a plane curve, this library implements an algorithm to compute an integral basis of the integral closure of the affine coordinate ring in the algebraic function field via normalization. The user can choose whether the algorithm will do the computation globally or (this is the default) compute in the localization at each component of the singular locus and put everything together.

**Procedures:**

`integralBasis(f, intVar)`  
Integral basis of an algebraic function field

**D.4.12 intprog\_lib****Library:** intprog.lib**Purpose:** Integer Programming with Groebner Basis Methods**Author:** Christine Theis, email: ctheis@math.uni-sb.de**Procedures:**

`solve_IP(..)`  
procedures for solving Integer Programming problems

**D.4.13 locnormal\_lib****Library:** locnormal.lib**Purpose:** Normalization of affine domains using local methods

**Authors:** J. Boehm boehm@mathematik.uni-kl.de  
W. Decker decker@mathematik.uni-kl.de  
S. Laplagne slaplagn@dm.uba.ar  
G. Pfister pfister@mathematik.uni-kl.de  
S. Steidel steidel@mathematik.uni-kl.de  
A. Steenpass steenpass@mathematik.uni-kl.de

**Overview:** Suppose  $A$  is an affine domain over a perfect field. This library implements a local-to-global strategy for finding the normalization of  $A$ . Following [1], the idea is to stratify the singular locus of  $A$ , apply the normalization algorithm given in [2] locally at each stratum, and put the local results together. This approach is inherently parallel. Furthermore we allow for the optional modular computation of the local results as provided by modnormal.lib. See again [1] for details.

**References:**

- [1] Janko Boehm, Wolfram Decker, Santiago Laplagne, Gerhard Pfister, Stefan Steidel, Andreas Steenpass: Parallel algorithms for normalization, <http://arxiv.org/abs/1110.4299>, 2011.
- [2] Gert-Martin Greuel, Santiago Laplagne, Frank Seelisch: Normalization of Rings, *Journal of Symbolic Computation* 9 (2010), p. 887-901

**Procedures:**

`locNormal(I, [...])`  
normalization of  $R/I$  using local methods

See also: Section D.4.15 [modnormal.lib], page 818; Section D.4.23 [normal.lib], page 824.

**D.4.14 moddiq\_lib****Library:** moddiq.lib**Purpose:** Double ideal quotient using modular methods

**Authors:** Y. Ishihara yishihara@rikkyo.ac.jp

**Overview:** A library for computing ideal quotient and saturation in the polynomial ring over the rational numbers using modular methods.

**References:**

M. Noro, K. Yokoyama: Usage of Modular Techniques for Efficient Computation of Ideal Operations. *Math.Comput.Sci.* 12: 1, 1-32. (2017).

**Procedures:**

`modQuotient(I, J)`  
                                   standard basis of (I:J) using modular methods

`modSat(I, J)`  
                                   standard basis of (I:J<sup>infty</sup>) using modular methods

### D.4.15 `modnormal.lib`

**Library:** `modnormal.lib`

**Purpose:** Normalization of affine domains using modular methods

**Authors:** J. Boehm boehm@mathematik.uni-kl.de  
 W. Decker decker@mathematik.uni-kl.de  
 S. Laplagne slaplagn@dm.uba.ar  
 G. Pfister pfister@mathematik.uni-kl.de  
 A. Steenpass steenpass@mathematik.uni-kl.de  
 S. Steidel steidel@mathematik.uni-kl.de

**Overview:** Suppose  $A$  is an affine domain over a perfect field.

This library implements a modular strategy for finding the normalization of  $A$ . Following [1], the idea is to apply the normalization algorithm given in [2] over finite fields and lift the results via Chinese remaindering and rational reconstruction as described in [3]. This approach is inherently parallel.

The strategy is available both as a randomized and as a verified algorithm.

**References:**

- [1] Janko Boehm, Wolfram Decker, Santiago Laplagne, Gerhard Pfister, Stefan Steidel, Andreas Steenpass: Parallel algorithms for normalization, preprint, 2011.
- [2] Gert-Martin Greuel, Santiago Laplagne, Frank Seelisch: Normalization of Rings, *Journal of Symbolic Computation* 9 (2010), p. 887-901
- [3] Janko Boehm, Wolfram Decker, Claus Fieker, Gerhard Pfister: The use of Bad Primes in Rational Reconstruction, preprint, 2012.

**Procedures:**

`modNormal(I)`  
                                   normalization of  $R/I$  using modular methods

See also: Section D.4.13 [`locnormal.lib`], page 817; Section D.4.23 [`normal.lib`], page 824.

### D.4.16 `modstd.lib`

**Library:** `modstd.lib`

**Purpose:** Groebner bases of ideals/modules using modular methods

**Authors:** A. Hashemi Amir.Hashemi@lip6.fr  
 G. Pfister pfister@mathematik.uni-kl.de  
 H. Schoenemann hannes@mathematik.uni-kl.de  
 A. Steenpass steenpass@mathematik.uni-kl.de  
 S. Steidel steidel@mathematik.uni-kl.de

**Overview:** A library for computing Groebner bases of ideals/modules in the polynomial ring over the rational numbers using modular methods.

**References:**

E. A. Arnold: Modular algorithms for computing Groebner bases. *J. Symb. Comp.* 35, 403-419 (2003).  
 N. Idrees, G. Pfister, S. Steidel: Parallelization of Modular Algorithms. *J. Symb. Comp.* 46, 672-684 (2011).

**Procedures:**

`modStd(I)`  
                   standard basis of I using modular methods  
`modSyz(I)`  
                   syzygy module of I using modular methods  
`modIntersect(I,J)`  
                   intersection of I and J using modular methods

#### D.4.17 monomialideal\_lib

**Library:** monomialideal.lib

**Purpose:** Primary and irreducible decompositions of monomial ideals

**Authors:** I.Bermejo, ibermejo@ull.es  
 E.Garcia-Llorente, evgarcia@ull.es  
 Ph.Gimenez, pgimenez@agt.uva.es

**Overview:** A library for computing a primary and the irreducible decompositions of a monomial ideal using several methods.

In this library we also take advantage of the fact that the ideal is monomial to make some computations that are Grobner free in this case (radical, intersection, quotient...).

**Procedures:**

`isMonomial(id)`  
                   checks whether an ideal id is monomial  
`minbaseMon(id)`  
                   computes the minimal monomial generating set of a monomial ideal id  
`gcdMon(f,g)`  
                   computes the gcd of two monomials f, g  
`lcmMon(f,g)`  
                   computes the lcm of two monomials f, g  
`membershipMon(f,id)`  
                   checks whether a polynomial f belongs to a monomial ideal id  
`intersectMon(id1,id2)`  
                   intersection of monomial ideals id1 and id2

```

quotientMon(id1,id2)
 quotient ideal id1:id2

radicalMon(id)
 computes the radical of a monomial ideal id

isprimeMon(id)
 checks whether a monomial ideal id is prime

isprimaryMon(id)
 checks whether a monomial ideal id is primary

isirreducibleMon(id)
 checks whether a monomial ideal id is irreducible

isartinianMon(id)
 checks whether a monomial ideal id is artinian

isgenericMon(id)
 checks whether a monomial ideal id is generic

dimMon(id)
 dimension of a monomial ideal id

irreddecMon(id,..)
 computes the irreducible decomposition of a monomial ideal id

primdecMon(id,..)
 computes a minimal primary decomposition of a monomial ideal id

```

#### D.4.18 mprimdec\_lib

**Library:** mprimdec.lib

**Purpose:** procedures for primary decomposition of modules

**Authors:** Alexander Dreyer, dreyer@mathematik.uni-kl.de; adreyer@web.de

**Overview:** Algorithms for primary decomposition for modules based on the algorithms of Gianni, Trager and Zacharias and Shimoyama and Yokoyama (generalization of the latter suggested by Hans-Gert Graebe, Leipzig ) using elements of primdec.lib

**Remark:** These procedures are implemented to be used in characteristic 0. They also work in positive characteristic  $\gg 0$ . In small characteristic and for algebraic extensions, the procedures via Gianni, Trager, Zacharias may not terminate.

**Procedures:**

```

separator(l)
 computes a list of separators of prime ideals

PrimdecA(N[,i])
 (not necessarily minimal) primary decomposition via Shi-
 moyama/Yokoyama (suggested by Graebe)

PrimdecB(N,p)
 (not necessarily minimal) primary decomposition for pseudo-primary ideals

```

`modDec(N[, i])`  
 minimal primary decomposition via Shimoyama/Yokoyama (suggested by Graebe)

`zeroMod(N[, check])`  
 minimal zero-dimensional primary decomposition via Gianni, Trager and Zacharias

`GTZmod(N[, check])`  
 minimal primary decomposition via Gianni, Trager and Zacharias

`de1var(N[, check[, ann]])`  
 primary decomposition for one variable

`annil(N)` the annihilator of  $M/N$  in the basering

`splitting(N[, check[, ann]])`  
 splitting to simpler modules

`primTest(i[, p])`  
 tests whether  $i$  is prime or homogeneous

`preComp(N, check[, ann])`  
 enhanced Version of splitting

`indSet(i)`  
 lists with varstrings of (in)dependent variables

`GTZopt(N[, check[, ann]])`  
 a faster version of GTZmod

`zeroOpt(N[, check[, ann]])`  
 a faster version of zeroMod

#### D.4.19 mregular\_lib

**Library:** mregular.lib

**Purpose:** Castelnuovo-Mumford regularity of homogeneous ideals

**Authors:** I.Bermejo, ibermejo@ull.es  
 Ph.Gimenez, pgimenez@agt.uva.es  
 G.-M.Greuel, greuel@mathematik.uni-kl.de

**Overview:** A library for computing the Castelnuovo-Mumford regularity of a homogeneous ideal that DOES NOT require the computation of a minimal graded free resolution of the ideal.

It also determines `depth(basering/ideal)` and `satiety(ideal)`. The procedures are based on 3 papers by Isabel Bermejo and Philippe Gimenez: 'On Castelnuovo-Mumford regularity of projective curves' Proc.Amer.Math.Soc. 128(5) (2000), 'Computing the Castelnuovo-Mumford regularity of some subschemes of  $P_n$  using quotients of monomial ideals', Proceedings of MEGA-2000, J. Pure Appl. Algebra 164 (2001), and 'Saturation and Castelnuovo-Mumford regularity', Preprint (2004).

**Procedures:**

`regIdeal(id, [, e])`  
 regularity of homogeneous ideal `id`

`depthIdeal(id, [, e])`  
depth of  $S/\text{id}$  with  $S=\text{basering}$ ,  $\text{id}$  homogeneous ideal

`satiety(id, [, e])`  
saturation index of homogeneous ideal  $\text{id}$

`regMonCurve(li)`  
regularity of projective monomial curve defined by  $\text{li}$

`NoetherPosition(id)`  
Noether normalization of ideal  $\text{id}$

`is_NP(id)`  
checks whether variables are in Noether position

`is_nested(id)`  
checks whether monomial ideal  $\text{id}$  is of nested type

#### D.4.20 `nfmodstd_lib`

**Library:** `nfmodstd.lib`

**Purpose:** Groebner bases of ideals in polynomial rings over algebraic number fields

**Authors:** D.K. Boku `boku@mathematik.uni-kl.de`  
W. Decker `decker@mathematik.uni-kl.de`  
C. Fieker `fieker@mathematik.uni-kl.de`

**Overview:** A library for computing the Groebner basis of an ideal in the polynomial ring over an algebraic number field  $Q(t)$  using the modular methods, where  $t$  is algebraic over the field of rational numbers  $Q$ . For the case  $Q(t) = Q$ , the procedure is inspired by Arnold [1]. This idea is then extended to the case  $t$  not in  $Q$  using factorization as follows:

Let  $f$  be the minimal polynomial of  $t$ .

For  $I, I'$  ideals in  $Q(t)[X]$ ,  $Q[X, t]/\langle f \rangle$  respectively, we map  $I$  to  $I'$  via the map sending  $t$  to  $t + \langle f \rangle$ . We first choose a prime  $p$  such that  $f$  has at least two factors in characteristic  $p$  and add each factor  $f_i$  to  $I'$  to obtain the ideal  $J'_i = I' + \langle f_i \rangle$ . We then compute a standard basis  $G'_i$  of  $J'_i$  for each  $i$  and combine the  $G'_i$  to  $G_p$  (a standard basis of  $I'_p$ ) using chinese remaindering for polynomials. The procedure is repeated for many primes  $p$ , where we compute the  $G_p$  in parallel until the number of primes is sufficiently large to recover the correct standard basis  $G'$  of  $I'$ . Finally, by mapping  $G'$  back to  $Q(t)[X]$ , a standard basis  $G$  of  $I$  is obtained.

The procedure also works if the input is a module. For this, we consider the rings  $A = Q(t)[X]$  and  $A' = (Q[t]/\langle f \rangle)[X]$ . For submodules  $I, I'$  in  $A^m, A'^m$ , respectively, we map  $I$  to  $I'$  via the map sending  $t$  to  $t + \langle f \rangle$ . As above, we first choose a prime  $p$  such that  $f$  has at least two factors in characteristic  $p$ . For each factor  $f_{\{i,p\}}$  of  $f_p := (f \bmod p)$ , we set  $I'_{\{i,p\}} := (I'_p \bmod f_{\{i,p\}})$ . We then compute a standard basis  $G'_i$  of  $I'_{\{i,p\}}$  over  $F_p[t]/\langle f_{\{i,p\}} \rangle$  for each  $i$  and combine the  $G'_i$  to  $G_p$  (a standard basis of  $I'_p$ ) using chinese remaindering for polynomials. The procedure is repeated for many primes  $p$  as described above and we finally obtain a standard basis of  $I$ .

**References:**

[1] E. A. Arnold: Modular algorithms for computing Groebner bases. *J. Symb. Comp.* 35, 403-419 (2003).

**Procedures:**

`chinrempoly(l, m)`  
chinese remaindering for polynomials



`nfmodStd(I)`

standard basis of  $I$  over algebraic number field using modular methods

#### D.4.21 `nfmodsyz_lib`

**Library:** `nfmodsyz.lib`

**Purpose:** Syzygy modules of submodules of free modules over algebraic number fields

**Authors:** D.K. Boku `boku@mathematik.uni-kl.de`  
 W. Decker `decker@mathematik.uni-kl.de`  
 C. Fieker `fieker@mathematik.uni-kl.de`

**Overview:** A library for computing the syzygy module of a given submodule  $I$  in a polynomial ring over an algebraic number field  $\mathbb{Q}(t)$ , where  $t$  is an algebraic number, using modular methods. For the case  $\mathbb{Q}(t)=\mathbb{Q}$ , that is, where  $t$  is an element of  $\mathbb{Q}$ , we compute, following [1], the syzygy module of  $I$  as follows: For a submodule  $I$  of  $A^m$  with  $A = \mathbb{Q}[X]$ , we first choose a sufficiently large set of primes  $P$  and compute the reduced Groebner basis of the syzygy module of  $I_p$ , for each  $p$  in  $P$ , in parallel. We then use the Chinese remainder algorithm and rational reconstruction to obtain the syzygy module of  $I$  over  $\mathbb{Q}$ . For the case where  $t$  is not in  $\mathbb{Q}$ , we compute, following [2], the syzygy module of  $I$  as follows:

Let  $f$  be the minimal polynomial of  $t$ . For a submodule  $I$  in  $A^m$  with  $A = \mathbb{Q}(t)[X]$ , we map  $I$  to a submodule  $I'$  in  $A^m$  with  $A = (\mathbb{Q}[t]/\langle f \rangle)[X]$  via the map sending  $t$  to  $t + \langle f \rangle$ . We first choose a prime  $p$  such that  $f$  has at least two factors in characteristic  $p$ . For each factor  $f_{i,p}$  of  $f_p := (f \bmod p)$ , we set  $I'_{i,p} := (I'_p \bmod f_{i,p})$ . We then compute the reduced Groebner bases  $G'_i$  of the syzygy modules of  $I'_{i,p}$  over  $F_p[t]/\langle f_{i,p} \rangle$  and combine the  $G'_i$  to  $G_p$  (the syzygy module of  $I'_p$ ) using chinese remaindering for polynomials. As described in [2], the procedure is repeated for many primes  $p$ , where we compute the  $G_p$  in parallel until the number of primes is sufficiently large to recover the correct generating set for the syzygy module  $G'$  of  $I'$  which is, considered over  $\mathbb{Q}(t)$ , also a generating set for the syzygy module of  $I$ .

**References:**

- [1] E. A. Arnold: Modular algorithms for computing Groebner bases. *J. Symb. Comp.* 35, 403-419 (2003).
- [2] D. Boku, W. Decker, C. Fieker, and A. Steenpass. Groebner bases over algebraic number fields. In: *Proceedings of the 2015 International Workshop on Parallel Symb. Comp. PASCO'15*, pages 16-24 (2015).

**Procedures:**

`nfmodSyz(I)`

syzygy module of  $I$  over algebraic number field using modular methods

#### D.4.22 `noether_lib`

**Library:** `noether.lib`

**Purpose:** Noether normalization of an ideal (not necessary homogeneous)

**Authors:** A. Hashemi, `Amir.Hashemi@lip6.fr`

**Overview:** A library for computing the Noether normalization of an ideal that DOES NOT require the computation of the dimension of the ideal. It checks whether an ideal is in Noether position. A modular version of these algorithms is also provided.

The procedures are based on a paper of Amir Hashemi 'Efficient Algorithms for Computing Noether Normalization' (presented in ASCM 2007)

This library computes also Castelnuovo-Mumford regularity and satiety of an ideal. A modular version of these algorithms is also provided. The procedures are based on a paper of Amir Hashemi 'Computation of Castelnuovo-Mumford regularity and satiety' (preprint 2008)

**Procedures:**

`NPos_test(id)`  
checks whether monomial ideal `id` is in Noether position

`modNpos_test(id)`  
the same as above using modular methods

`NPos(id)` Noether normalization of ideal `id`

`modNPos(id)`  
Noether normalization of ideal `id` by modular methods

`nsatiety(id)`  
Satiety of ideal `id`

`modsatiety(id)`  
Satiety of ideal `id` by modular methods

`regCM(id)`  
Castelnuovo-Mumford regularity of ideal `id`

`modregCM(id)`  
Castelnuovo-Mumford regularity of ideal `id` by modular methods

### D.4.23 `normal_lib`

**Library:** `normal.lib`

**Purpose:** Normalization of Affine Rings

**Authors:** G.-M. Greuel, greuel@mathematik.uni-kl.de,  
S. Laplagne, slaplagn@dm.uba.ar,  
G. Pfister, pfister@mathematik.uni-kl.de  
Peter Chini, chini@rhrk.uni-kl.de (normalConductor)

**Procedures:**

`normal(I, [...])`  
normalization of an affine ring

`normalP(I, [...])`  
normalization of an affine ring in positive characteristic

`normalC(I, [...])`  
normalization of an affine ring through a chain of rings

`HomJJ(L)` presentation of  $\text{End}_R(J)$  as affine ring, `J` an ideal

`genus(I)` computes the geometric genus of a projective curve

`primeClosure(L)`  
integral closure of  $R/p$ , `p` a prime ideal

`closureFrac(L)`  
 writes a poly in integral closure as element of  $\text{Quot}(R/p)$   
`iMult(L)` intersection multiplicity of the ideals of the list L  
`deltaLoc(f,S)`  
 sum of delta invariants at conjugated singular points  
`locAtZero(I)`  
 checks whether the zero set of I is located at 0  
`norTest(I,nor)`  
 checks the output of normal, normalP, normalC  
`getSmallest(J)`  
 computes the polynomial of smallest degree of J  
`getOneVar(J, vari)`  
 computes a polynomial of J in the variable vari  
`changeDenominator(U1, c1, c2, I)`  
 computes ideal U2 such that  $1/c1*U1=1/c2*U2$   
`normalConductor(ideal)`  
 computation of the conductor as ideal in the basering

See also: Section D.4.13 [`locnormal.lib`], page 817; Section D.4.15 [`modnormal.lib`], page 818.

#### D.4.24 `normaliz.lib`

**Library:** `normaliz.lib`

**Purpose:** Provides an interface for the use of Normaliz 2.11 or newer within SINGULAR.

**Authors:** Winfried Bruns, Winfried.Bruns@Uni-Osnabrueck.de  
 Christof Soeger, Christof.Soeger@Uni-Osnabrueck.de

**Overview:** The library `normaliz.lib` provides an interface for the use of Normaliz 2.11 or newer within SINGULAR. The exchange of data is via files. In addition to the top level functions that aim at objects of type `ideal` or `ring`, several other auxiliary functions allow the user to apply Normaliz to data of type `intmat`. Therefore SINGULAR can be used as a comfortable environment for the work with Normaliz.

Please see the `Normaliz.pdf` (included in the Normaliz distribution) for a more extensive documentation of Normaliz.

Normaliz allows the use of a grading. In the Singular functions that access Normaliz the parameter `grading` is an `intvec` that assigns a (not necessarily positive) degree to every variable of the ambient polynomial ring. But it must give positive degrees to the generators given to function.

Singular and Normaliz exchange data via files. These files are automatically created and erased behind the scenes. As long as one wants to use only the ring-theoretic functions there is no need for file management.

Note that the numerical invariants computed by Normaliz can be accessed in this "automatic file mode".

However, if Singular is used as a frontend for Normaliz or the user wants to inspect data not automatically returned to Singular, then an explicit filename and a path can be specified for the exchange of data. Moreover, the library provides functions for access to these files. Deletion of the files is left to the user.

Use of this library requires the program Normaliz to be installed. You can download it from <http://www.mathematik.uni-osnabrueck.de/normaliz/>. Please make sure that the executables are in the search path or use `setNmzExecPath` ([`setNmzExecPath`], page 826).

### Procedures:

```

intclToricRing(ideal I)
 computes the integral closure of the toric ring generated by the leading
 monomials of the elements of I in the basering

normalToricRing(ideal I)
 computes the normalization of the toric ring generated by the leading
 monomials of the elements of I

normalToricRingFromBinomials(ideal I)
 computes the normalization of the polynomial ring modulo the unique
 minimal binomial prime ideal of the binomial ideal I

ehrhartRing(ideal I)
 considers the exponent vectors of the elements of I as points of a lattice
 polytope and computes the integral closure of the polytopal algebra

intclMonIdeal(ideal I)
 Computes the integral closure of the Rees algebra of the ideal generated
 by the leading monomials of the elements of I

torusInvariants(intmat T)
 computes the ring of invariants of a torus action

finiteDiagInvariants(intmat C)
 computes the ring of invariants of a finite abelian group acting diagonally
 on a polynomial ring

diagInvariants(intmat C)
 computes the ring of invariants of a diagonalizable group

intersectionValRings(intmat V)
 computes the intersection of the polynomial ring with the valuation rings
 of monomial valuations

intersectionValRingIdeals(intmat V)
 computes ideals of monomial valuations

showNuminvs()
 prints the numerical invariants found by Normaliz

exportNuminvs()
 exports the numerical invariants found by Normaliz

setNmzOption(string s, int onoff)
 sets the option s to onoff

showNmzOptions()
 prints the enabled options to the standard output

normaliz(intmat sgr, int nmz_mode)
 applies Normaliz

setNmzExecPath(string nmz_exec_path_name)
 sets the path to the Normaliz executable

```

```

writeNmzData(intmat sgr, int n_mode)
 creates an input file for Normaliz

readNmzData(string nmz_suffix)
 reads the Normaliz output file with the specified suffix

setNmzFilename(string nmz_filename_name)
 sets the filename for the exchange of data

setNmzDataPath(string nmz_data_path_name)
 sets the directory for the exchange of data

writeNmzPaths()
 writes the path names into two files

startNmz()
 retrieves the path names written by writeNmzPaths

rmNmzFiles()
 removes the files created for and by Normaliz

mons2intmat(ideal I)
 returns the intmat whose rows represent the leading exponents of the elements of I

intmat2mons(intmat expo_vecs)
 returns the ideal generated by the monomials which have the rows of expo_vecs as exponent vector

binomials2intmat(ideal I)
 returns the intmat whose rows represent the exponents of the elements of the binomial ideal I

```

#### D.4.25 pointid\_lib

**Library:** pointid.lib

**Purpose:** Procedures for computing a factorized lex GB of the vanishing ideal of a set of points via the Axis-of-Evil Theorem (M.G. Marinari, T. Mora)

**Author:** Stefan Steidel, steidel@mathematik.uni-kl.de

**Overview:** The algorithm of Cerlienco-Mureddu [Marinari M.G., Mora T., A remark on a remark by Macaulay or Enhancing Lazard Structural Theorem. Bull. of the Iranian Math. Soc., 29 (2003), 103-145] associates to each ordered set of points  $A := \{a_1, \dots, a_s\}$  in  $K^n$ ,  $a_i := (a_{i1}, \dots, a_{in})$

- a set of monomials  $N$  and

- a bijection  $\phi: A \rightarrow N$ .

Here  $I(A) := \{f \in K[x(1), \dots, x(n)] \mid f(a_i) = 0, \text{ for all } 1 \leq i \leq s\}$  denotes the vanishing ideal of  $A$  and  $N = \text{Mon}(x(1), \dots, x(n)) \setminus \{LM(f) \mid f \in I(A)\}$  is the set of monomials which do not lie in the leading ideal of  $I(A)$  (w.r.t. the lexicographical ordering with  $x(n) > \dots > x(1)$ ).  $N$  is also called the set of non-monomials of  $I(A)$ . NOTE:  $\#A = \#N$  and  $N$  is a monomial basis of  $K[x(1..n)]/I(A)$ . In particular, this allows to deduce the set of corner-monomials, i.e. the minimal basis  $M := \{m_1, \dots, m_r\}$ ,  $m_1 < \dots < m_r$ , of its associated monomial ideal  $M(I(A))$ , such that

$M(I(A)) = \{k * m_i \mid k \in \text{Mon}(x(1), \dots, x(n)), m_i \in M\}$ ,

and (by interpolation) the unique reduced lexicographical Groebner basis  $G :=$

$\{f_1, \dots, f_r\}$  such that  $LM(f_i) = m_i$  for each  $i$ , that is,  $I(A) = \langle G \rangle$ . Moreover, a variation of this algorithm allows to deduce a canonical linear factorization of each element of such a Groebner basis in the sense of the Axis-of-Evil Theorem by M.G. Marinari and T. Mora. More precisely, a combinatorial algorithm and interpolation allow to deduce polynomials

$$y_{m,d} = x(m) - g_{m,d}(x(1), \dots, x(m-1)),$$

$i=1, \dots, r; m=1, \dots, n; d$  in a finite index-set  $F$ , satisfying

$$f_i = (\text{product of } y_{m,d}) \text{ modulo } (f_1, \dots, f_{i-1})$$

where the product runs over all  $m=1, \dots, n$ ; and all  $d$  in  $F$ .

**Procedures:**

`nonMonomials(id)`  
non-monomials of the vanishing ideal  $id$  of a set of points

`cornerMonomials(N)`  
corner-monomials of the set of non-monomials  $N$

`facGBIdeal(id)`  
GB  $G$  of  $id$  and linear factors of each element of  $G$

**D.4.26 primdec\_lib****Library:** primdec.lib**Purpose:** Primary Decomposition and Radical of Ideals

**Authors:** Gerhard Pfister, pfister@mathematik.uni-kl.de (GTZ)  
Wolfram Decker, decker@math.uni-sb.de (SY)  
Hans Schoenemann, hannes@mathematik.uni-kl.de (SY)  
Santiago Laplagne, slaplagn@dm.uba.ar (GTZ)

**Overview:** Algorithms for primary decomposition based on the ideas of Gianni, Trager and Zacharias (implementation by Gerhard Pfister), respectively based on the ideas of Shimoyama and Yokoyama (implementation by Wolfram Decker and Hans Schoenemann). The procedures are implemented to be used in characteristic 0. They also work in positive characteristic  $\gg 0$ . In small characteristic and for algebraic extensions, primdecGTZ may not terminate. Algorithms for the computation of the radical based on the ideas of Krick, Logar, Laplagne and Kemper (implementation by Gerhard Pfister and Santiago Laplagne). They work in any characteristic. Baserings must have a global ordering and no quotient ideal. Exceptions: primdecGTZ, absPrimdecGTZ, minAssGTZ, primdecSY, minAssChar, radical accept non-global ordering.

**Procedures:**

`Ann(M)` annihilator of  $R^n/M$ ,  $R$ =basing,  $M$  in  $R^n$

`primdecGTZ(I)`  
complete primary decomposition via Gianni,Trager,Zacharias

`primdecGTZE(I)`  
complete primary decomposition via Gianni,Trager,Zacharias. Returns empty list for the unit ideal

`primdecSY(I...)`  
 complete primary decomposition via Shimoyama-Yokoyama

`primdecSYE(I,...)`  
 complete primary decomposition via Shimoyama-Yokoyama. Returns empty list for the unit ideal

`minAssGTZ(I)`  
 the minimal associated primes via Gianni,Trager,Zacharias (with modifications by Laplagne)

`minAssGTZE(I)`  
 the minimal associated primes via Gianni,Trager,Zacharias. Returns empty list for unit ideal

`minAssChar(I...)`  
 the minimal associated primes using characteristic sets

`minAssCharE(I...)`  
 the minimal associated primes using characteristic sets. Returns empty list for unit ideal

`testPrimary(L,k)`  
 tests the result of the primary decomposition

`testPrimaryE(L,k)`  
 tests the result of the primary decomposition. Handles also empty list L.

`radical(I)`  
 computes the radical of I via Krick/Logar (with modifications by Laplagne) and Kemper

`radicalEHV(I)`  
 computes the radical of I via Eisenbud,Huneke,Vasconcelos

`equiRadical(I)`  
 the radical of the equidimensional part of the ideal I

`prepareAss(I)`  
 list of radicals of the equidimensional components of I

`equidim(I)`  
 weak equidimensional decomposition of I

`equidimMax(I)`  
 equidimensional locus of I

`equidimMaxEHV(I)`  
 equidimensional locus of I via Eisenbud,Huneke,Vasconcelos

`zerodec(I)`  
 zerodimensional decomposition via Monico

`absPrimdecGTZ(I)`  
 the absolute prime components of I

`absPrimdecGTZE(I)`  
 the absolute prime components of I. Assumes I is not unit ideal.

`sep(f,k)` the separabel part of f as polynomial in  $\mathbb{F}_p(t_1, \dots, t_m)$

See also: Section D.4.27 [primdecint.lib], page 830.

### D.4.27 `primdecint_lib`

**Library:** `primdecint.lib`

**Purpose:** primary decomposition of an ideal in the polynomial ring over the integers

**Authors:** G. Pfister `pfister@mathematik.uni-kl.de`  
A. Sadiq `afshanatiq@gmail.com`  
S. Steidel `steidel@mathematik.uni-kl.de`

**Overview:** A library for computing the primary decomposition of an ideal in the polynomial ring over the integers,  $\mathbb{Z}[x_1, \dots, x_n]$ .

The first procedure `'primdecZ'` can be used in parallel.

Reference: Pfister, Sadiq, Steidel, "An Algorithm for primary decomposition in polynomial rings over the integers", `arXiv:1008.2074`

**Procedures:**

```

primdecZ(I)
 compute the primary decomposition of ideal I

primdecZM(I)
 compute the primary decomposition of module I

minAssZ(I)
 compute the minimal associated primes of I

radicalZ(I)
 compute the radical of I

heightZ(I)
 compute the height of I

equidimZ(I)
 compute the equidimensional part of I

intersectZ(I,J)
 compute the intersection of I and J

```

See also: Section D.4.26 [`primdec.lib`], page 828.

### D.4.28 `primitiv_lib`

**Library:** `primitiv.lib`

**Purpose:** Computing a Primitive Element

**Author:** Martin Lamm, email: `lamm@mathematik.uni-kl.de`

**Procedures:**

```

primitive(ideal i)
 find minimal polynomial for a primitive element

primitive_extra(i)
 find primitive element for two generators

splitring(f,R[,L])
 define ring extension with name R and switch to it

```



### D.4.29 realrad\_lib

**Library:** realrad.lib

**Purpose:** Computation of real radicals

**Author :** Silke Spang

**Overview:** Algorithms about the computation of the real radical of an arbitrary ideal over the rational numbers and transcendental extensions thereof

**Procedures:**

realpoly(f)

Computes the real part of the univariate polynomial f

realzero(j)

Computes the real radical of the zerodimensional ideal j

realrad(j)

Computes the real radical of an arbitrary ideal over transcendental extension of the rational numbers

### D.4.30 reesclos\_lib

**Library:** reesclos.lib

**Purpose:** procedures to compute the int. closure of an ideal

**Author:** Tobias Hirsch, email: hirsch@math.tu-cottbus.de  
Janko Boehm, email: boehm@mathematik.uni-kl.de  
Magdalen Marais, email: magdalen@aims.ac.za

**Overview:** A library to compute the integral closure of an ideal I in a polynomial ring  $R=k[x(1),\dots,x(n)]$  using the Rees Algebra  $R[It]$  of I. It computes the integral closure of  $R[It]$ , which is a graded subalgebra of  $R[t]$ . The degree-k-component is the integral closure of the k-th power of I.

In contrast to the previous version, the library uses 'normal.lib' to compute the integral closure of  $R[It]$ . This improves the performance considerably.

**Procedures:**

ReesAlgebra(I)

computes the Rees Algebra of an ideal I

normalI(I[,p[,r]])

computes the integral closure of an ideal I using  $R[It]$

### D.4.31 rstandard\_lib

**Library:** rstandard.lib

**Purpose:** Computes Janet bases and border bases for ideals

**Authors:** Shamsa Kanwal lotus\_zone16@yahoo.com  
Gerhard Pfister pfister@mathematik.uni-kl.de

**Overview:** Computing Janet bases and border bases for any ordering using the idea of r-standard bases (defined by V. Gerdt)

**References:**

- [1] A. Kehrein, M. Kreuzer, L. Robbiano: An algebrists view on border bases, in: A. Dickenstein and I. Emiris (eds.), Solving Polynomial Equations: Foundations, Algorithms and Applications, Springer, Heidelberg 2005, 169-202.
- [2] V.P. Gerdt: Involute Algorithms for Computing Groebner Bases, In Computational Commutative and Non-Computational Algebra Geometry, S.Conjocaru, G. Pfister and V. Ufnarovski (Eds.), NATO Science Series,105 Press 2005, 199-255.

**Procedures:**

`borderBasis(I)`  
 computes a border basis of the ideal I

`modBorder(I)`  
 computes a border basis of the ideal I using modular methods

`rJanet(I)`  
 computes a Janet basis of the ideal I

`modJanet(I)`  
 computes a Janet basis of the ideal I using modular methods

### D.4.32 `sagbi.lib`

**Library:** `sagbi.lib`

**Purpose:** Compute SAGBI basis (subalgebra bases analogous to Groebner bases for ideals) of a subalgebra

**Authors:** Jan Hackfeld, Jan.Hackfeld@rwth-aachen.de  
 Gerhard Pfister, pfister@mathematik.uni-kl.de  
 Viktor Levandovskyy, levandov@math.rwth-aachen.de

**Overview:** SAGBI stands for 'subalgebra bases analogous to Groebner bases for ideals'. SAGBI bases provide important tools for working with finitely presented subalgebras of a polynomial ring. Note, that in contrast to Groebner bases, SAGBI bases may be infinite.

**References:**

- Ana Bravo: Some Facts About Canonical Subalgebra Bases, MSRI Publications 51, p. 247-254

**Procedures:**

`sagbiSPoly(A [,r,m])`  
 computes SAGBI S-polynomials of A

`sagbiReduce(I,A [,t,mt])`  
 performs subalgebra reduction of I by A

`sagbi(A [,m,t])`  
 computes SAGBI basis for A

`sagbiPart(A,k[,m])`  
 computes partial SAGBI basis for A

`algebraicDependence(I, it)`

performs iterations of SAGBI for algebraic dependencies of I

See also: Section D.4.2 [`algebra_lib`], page 809.

### D.4.33 `sing4ti2_lib`

**Library:** `sing4ti2.lib`

**Purpose:** Communication Interface to 4ti2

**Authors:** Thomas Kahle , kahle@mis.mpg.de  
Anne Fruehbis-Krueger, anne@math.uni-hannover.de

**Note:** This library uses the external program 4ti2 for calculations and the standard unix tools sed and awk for conversion of the returned result

**Procedures:**

`markov4ti2(A[, i])`  
compute Markov basis of given lattice

`hilbert4ti2(A[, i])`  
compute Hilbert basis of given lattice

`graver4ti2(A[, i])`  
compute Graver basis of given lattice

### D.4.34 `symodstd_lib`

**Library:** `symodstd.lib`

**Purpose:** Procedures for computing Groebner basis of ideals being invariant under certain variable permutations.

**Author:** Stefan Steidel, steidel@mathematik.uni-kl.de

**Overview:** A library for computing the Groebner basis of an ideal in the polynomial ring over the rational numbers, that is invariant under certain permutations of the variables, using the symmetry and modular methods. More precisely let  $I = \langle f_1, \dots, f_r \rangle$  be an ideal in  $\mathbb{Q}[x(1), \dots, x(n)]$  and  $\sigma$  a permutation of order  $k$  in  $\text{Sym}(n)$  such that  $\sigma(I) = I$ . We assume that  $\sigma(\{f_1, \dots, f_r\}) = \{f_1, \dots, f_r\}$ . This can always be obtained by adding  $\sigma(f_i)$  to  $\{f_1, \dots, f_r\}$ .

To compute a standard basis of I we apply a modification of the modular version of the standard basis algorithm (improving the calculations in positive characteristic). Therefore we only allow primes  $p$  such that  $p-1$  is divisible by  $k$ . This guarantees the existence of a  $k$ -th primitive root of unity in  $\mathbb{Z}/p\mathbb{Z}$ .

**Procedures:**

`genSymId(I, sigma)`  
compute ideal J such that  $\sigma(J) = J$  and J includes I

`isSymmetric(I, sigma)`  
check if I is invariant under permutation sigma

`primRoot(p, k)`  
int describing a  $k$ -th primitive root of unity in  $\mathbb{Z}/p\mathbb{Z}$

```
eigenvalues(I,sigma)
 list of eigenvalues of generators of I under sigma

symmStd(I,sigma)
 standard basis of I using invariance of I under sigma

syModStd(I,sigma)
 SB of I using modular methods and $\sigma(I) = I$
```

### D.4.35 toric\_lib

**Library:** toric.lib

**Purpose:** Standard Basis of Toric Ideals

**Author:** Christine Theis, email: ctheis@math.uni-sb.de

**Procedures:**

```
toric_ideal(A,..)
 computes the toric ideal of A

toric_std(ideal I)
 standard basis of I by a specialized Buchberger algorithm
```

## D.5 Algebraic geometry

### D.5.1 brillnoether\_lib

**Library:** brillnoether.lib

**Purpose:** Riemann-Roch spaces of divisors on curves

**Authors:** I. Stenger: stenger@mathematik.uni-kl.de  
Janko Boehm boehm@mathematik.uni-kl.de

**Procedures:**

```
RiemannRochBN()
 Computes a vector space basis of the Riemann-Roch space of a divisor D
 on a projective curve C
```

See also: Section D.10.1 [brnoeth.lib], page 886.

### D.5.2 chern\_lib

**Library:** chern.lib

**Purpose:** Symbolic Computations with Chern classes, Computation of Chern classes

**Author:** Oleksandr Iena, o.g.yena@gmail.com

**Overview:** A toolbox for symbolic computations with Chern classes. The Aluffi's algorithms for computation of characteristic classes of algebraic varieties (Segre, Fulton, Chern-Schwartz-MacPherson classes) are implemented as well.

**References:**

- [1] Aluffi, Paolo Computing characteristic classes of projective schemes. *Journal of Symbolic Computation*, 35 (2003), 3-19. [2] Iena, Oleksandr, On symbolic computations with Chern classes: remarks on the library `chern.lib` for Singular, <http://hdl.handle.net/10993/22395>, 2015.
- [3] Lascoux, Alain, Classes de Chern d'un produit tensoriel. *C. R. Acad. Sci., Paris, Ser. A* 286, 385-387 (1978). [4] Manivel, Laurent Chern classes of tensor products, arXiv 1012.0014, 2010.

**Procedures:**

- `symm(l [,N])`  
symmetric functions in the entries of `l`
- `symNsym(f, c)`  
symmetric and non-symmetric parts of a polynomial `f`
- `CompleteHomog(N, l)`  
complete homogeneous symmetric functions
- `segre(N, c)`  
Segre classes in terms of Chern classes
- `chern(N, s)`  
Chern classes in terms of Segre classes
- `chNum(N, c)`  
the non-zero Chern numbers in degree `N` in the entries of `c`
- `chNumbers(N, c)`  
the Chern numbers in degree `N` in the entries of `c`
- `sum_of_powers(k, l)`  
the sum of `k`-th powers of the entries of `l`
- `powSumSym(c [,N])`  
the sums of powers [up to degree `N`] in terms of the elementary symmetric polynomials (entries of `l`)
- `chAll(c [,N])`  
Chern character in terms of the Chern classes
- `chAllInv(c)`  
Chern classes in terms of the Chern character
- `chHE(c)` the highest term of the Chern character
- `ChernRootsSum(a, b)`  
the Chern roots of a direct sum
- `chSum(c, C)`  
the Chern classes of a direct sum
- `ChernRootsDual(l)`  
the Chern roots of the dual vector bundle
- `chDual(c)`  
the Chern classes of the dual vector bundle
- `ChernRootsProd(l, L)`  
the Chern roots of a tensor product of vector bundles

`chProd(r, c, R, C [,N])`  
 Chern classes of a tensor product of vector bundles

`chProdE(c, C)`  
 Chern classes of a tensor product of vector bundles

`chProdL(r, c, R, C)`  
 Chern classes of a tensor product of vector bundles

`chProdLP(r, c, R, C)`  
 total Chern class of a tensor product of vector bundles

`chProdM(r, c, R, C)`  
 Chern classes of a tensor product of vector bundles

`chProdMP(r, c, R, C)`  
 total Chern class of a tensor product of vector bundles

`ChernRootsHom(l, L)`  
 the Chern roots of a Hom vector bundle

`chHom(r, c, R, C [,N])`  
 Chern classes of the Hom-vector bundle

`ChernRootsSymm(n, l)`  
 the Chern roots of the n-th symmetric power of a vector bundle with Chern roots from l

`ChernRootsWedge(n, l)`  
 the Chern roots of the n-th exterior power of a vector bundle with Chern roots from l

`chSymm(k, r, c [,p])`  
 the rank and the Chern classes of the k-th symmetric power of a vector bundle of rank r with Chern classes c

`chSymm2L(r, c)`  
 the rank and the Chern classes of the second symmetric power of a vector bundle of rank r with Chern classes c

`chSymm2LP(r, c)`  
 the total Chern class of the second symmetric power of a vector bundle of rank r with Chern classes c

`chWedge(k, r, c [,p])`  
 the rank and the Chern classes of the k-th exterior power of a vector bundle of rank r with Chern classes c

`chWedge2L(r, c)`  
 the rank and the Chern classes of the second exterior power of a vector bundle of rank r with Chern classes c

`chWedge2LP(r, c)`  
 the total Chern class of the second exterior power of a vector bundle of rank r with Chern classes c

`todd(c [,n])`  
 the Todd class

`toddE(c)` the highest term of the Todd class

**Bern**(*n*)    the second Bernoulli numbers  
**tdCf**(*n*)    the coefficients of the Todd class of a line bundle  
**tdTerms**(*n*, *f*)  
               the terms of the Todd class of a line bundle corresponding to the Chern  
               root *t*  
**tdFactor**(*n*, *t*)  
               the Todd class of a line bundle corresponding to the Chern root *t*  
**cProj**(*n*)    the total Chern class of (the tangent bundle on) the projective space  $P_n$   
**chProj**(*n*)    the Chern character of (the tangent bundle on) the projective space  $P_n$   
**tdProj**(*n*)    the Todd class of (the tangent bundle on) the projective space  $P_n$   
**eulerChProj**(*n*, *r*, *c*)  
               Euler characteristic of a vector bundle on the projective space  $P_n$  via  
               Hirzebruch-Riemann-Roch theorem  
**chNumbersProj**(*n*)  
               the Chern numbers of the projective space  $P_n$   
**classpoly**(*l*, *t*)  
               polynomial in *t* with coefficients from *l* (without constant term)  
**chernPoly**(*l*, *t*)  
               Chern polynomial (constant term 1)  
**chernCharPoly**(*r*, *l*, *t*)  
               polynomial in *t* corresponding to the Chern character (constant term *r*)  
**toddPoly**(*td*, *t*)  
               polynomial in *t* corresponding to the Todd class (constant term 1)  
**rHRR**(*N*, *ch*, *td*)  
               the main ingredient of the right-hand side of the Hirzebruch-Riemann-Roch  
               formula  
**SchurS**(*I*, *S*)  
               the Schur polynomial corresponding to partition *I* in terms of the Segre  
               classes *S*  
**SchurCh**(*I*, *C*)  
               the Schur polynomial corresponding to partition *I* in terms of the Chern  
               classes *C*  
**part**(*m*, *n*)  
               partitions of integers not exceeding *n* into *m* non-negative summands  
**dualPart**(*I* [,*N*])  
               partition dual to *I*  
**PartC**(*I*, *m*)  
               the complement of a partition with respect to *m*  
**partOver**(*n*, *J*)  
               partitions over a given partition *J* with summands not exceeding *n*

|                           |                                                                          |
|---------------------------|--------------------------------------------------------------------------|
| <code>partUnder(J)</code> | partitions under a given partition J                                     |
| <code>SegreA(I)</code>    | Segre class of the projective subscheme defined by I                     |
| <code>FultonA(I)</code>   | Fulton class of the projective subscheme defined by I                    |
| <code>CSMA(I)</code>      | Chern-Schwartz-MacPherson class of the projective subscheme defined by I |
| <code>EulerAff(I)</code>  | Euler characteristic of the affine subvariety defined by I               |
| <code>EulerProj(I)</code> | Euler characteristic of the projective subvariety defined by I           |

### D.5.3 deRham\_lib

**Library:** deRham.lib

**Purpose:** Computation of deRham cohomology

**Authors:** Cornelia Rottner, rottner@mathematik.uni-kl.de

**Overview:** A library for computing the de Rham cohomology of complements of complex affine varieties.

**References:**

- [OT] Oaku, T.; Takayama, N.: Algorithms of D-modules - restriction, tensor product, localization, and local cohomology groups}, J. Pure Appl. Algebra 156, 267-308 (2001)
- [R] Rottner, C.: Computing de Rham Cohomology, diploma thesis (2012)
- [W1] Walther, U.: Algorithmic computation of local cohomology modules and the local cohomological dimension of algebraic varieties}, J. Pure Appl. Algebra 139, 303-321 (1999)
- [W2] Walther, U.: Algorithmic computation of de Rham Cohomology of Complements of Complex Affine Varieties}, J. Symbolic Computation 29, 796-839 (2000)
- [W3] Walther, U.: Computing the cup product structure for complements of complex affine varieties, J. Pure Appl. Algebra 164, 247-273 (2001)

**Procedures:**

- `deRhamCohomology(list[,opt])`  
computes the de Rham cohomology
- `MVComplex(list)`  
computes the Mayer-Vietoris complex

### D.5.4 divisors\_lib

**Library:** divisors.lib

**Purpose:** Divisors and P-Divisors

**Authors:** Janko Boehm boehm@mathematik.uni-kl.de  
Lars Kastner kastner@math.fu-berlin.de  
Benjamin Lorenz blorenz@math.uni-frankfurt.de  
Hans Schoenemann hannes@mathematik.uni-kl.de  
Yue Ren ren@mathematik.uni-kl.de



**Overview:** We implement a class `divisor` on an algebraic variety and methods for computing with them. Divisors are represented by tuples of ideals defining the positive and the negative part. In particular, we implement the group structure on divisors, computing global sections and testing linear equivalence.

In addition to this we provide a class `formaldivisor` which implements integer formal sums of divisors (not necessarily prime). A formal divisor can be evaluated to a divisor, and a divisor can be decomposed into a formal sum.

Finally we provide a class `pdivisor` which implements polyhedral formal sums of divisors (P-divisors) where the coefficients are assumed to be polyhedra with fixed tail cone. There is a function to evaluate a P-divisor on a vector in the dual of the tail cone. The result will be a formal divisor.

**References:**

For the class `divisor` we closely follow Macaulay2's tutorial on divisors.

**Procedures:**

```

makeDivisor(ideal,ideal)
 create a divisor

divisorplus(divisor,divisor)
 add two divisors

multdivisor(int,divisor)
 multiply a divisor by an integer

negativedivisor(divisor)
 compute the negative of the divisor

normalForm(divisor)
 normal form of a divisor

isEqualDivisor(divisor,divisor)
 test whether two divisors are equal

globalSections(divisor)
 compute the global sections of a divisor

degreeDivisor(divisor)
 degree of a divisor

linearlyEquivalent(divisor,divisor)
 test whether two divisors are linearly equivalent

effective(divisor)
 compute an effective divisor linearly equivalent to a divisor

makeFormalDivisor(list)
 make a formal integer sum of divisors

evaluateFormalDivisor(formaldivisor)
 evaluate a formal sum of divisors to a divisor

formaldivisorplus(formaldivisor,formaldivisor)
 add two formal divisors

negativeformaldivisor(formaldivisor)
 compute the negative of the formal divisor

```

```

multformaldivisor(int,formaldivisor)
 multiply a formal divisor by an integer

degreeFormalDivisor(formaldivisor)
 degree of a formal divisor

makePDivisor(List)
 make a formal polyhedral sum of divisors

evaluatePDivisor(pdivisor,intvec)
 evaluate a polyhedral divisor to an integer formal divisor

pdivisorplus(pdivisor,pdivisor)
 add two polyhedral divisors

```

### D.5.5 hess\_lib

**Library:** hess.lib

**Purpose:** Riemann-Roch space of divisors on function fields and curves

**Authors:** I. Stenger: stenger@mathematik.uni-kl.de

**Overview:** Let  $f$  be an absolutely irreducible polynomial in two variables  $x,y$ . Assume that  $f$  is monic as a polynomial in  $y$ . Let  $F = \text{Quot}(k[x,y]/f)$  be the function field defined by  $f$ . Define  $O_F = \text{IntCl}(k[x],F)$  and  $O_-(F,\text{inf}) = \text{IntCl}(k[1/x],F)$ . We represent a divisor  $D$  on  $F$  by two fractional ideals  $I$  and  $J$  of  $O_F$  and  $O_-(F,\text{inf})$ , respectively. The Riemann-Roch space  $L(D)$  is then the intersection of  $I^{-1}$  and  $J^{-1}$ .

**Procedures:**

```

RiemannRochHess()
 Computes a vector space basis of the Riemann-Roch space of a divisor

```

### D.5.6 numerAlg\_lib

Todos/Issues:

- does not follow the naming convention
- syntax errors in examples
- no test suite

**Library:** NumerAlg.lib

**Purpose:** Numerical Algebraic Algorithm

**Overview:** The library contains procedures to test the inclusion, the equality of two ideals defined by polynomial systems, compute the degree of a pure  $i$ -dimensional component of an algebraic variety defined by a polynomial system, compute the local dimension of an algebraic variety defined by a polynomial system at a point computed as an approximate value. The use of the library requires to install Bertini (<http://www.nd.edu/~sommese/bertini>).

**Author:** Shawki AlRashed, rashed@mathematik.uni-kl.de; sh.shawki@yahoo.de

**Procedures:**

```

Incl(ideal I, ideal J)
 test if I contains J

```

Equal(ideal I, ideal J)  
 test if I equals to J

DegreePure(ideal I, int i)  
 computes the degree of a pure i-dimensional

NumLocalDim(ideal I, p)  
 numerical local dimension at a point computed as an approximate value

### D.5.7 numerDecom.lib

Todos/Issues:

- does not follow the naming convention
- syntax errors in examples
- no test suite

**Library:** NumDecom.lib

**Purpose:** Numerical Decomposition of Ideals

**Overview:** The library contains procedures to compute numerical irreducible decomposition, and numerical primary decomposition of an algebraic variety defined by a polynomial system. The use of the library requires to install Bertini ([@uref{http://www.nd.edu/~sommese/bertini}](http://www.nd.edu/~sommese/bertini)).

**Author:** Shawki AlRashed, rashed@mathematik.uni-kl.de; sh.shawki@yahoo.de

**Procedures:**

re2squ(ideal I)  
 reduction to square system

UseBertini(ideal H, string sv)  
 use Bertini to compute the solutions of the homotopy function

Singular2bertini(list L)  
 adopt the list to be a read file in Bertini as a start solution set

bertini2Singular(string snp, int q)  
 adopt the file of solutions of the homotopy function to be a list in SINGULAR

ReJunkUseHomo(ideal I, ideal L, list W, list w)  
 remove junk points using the homotopy function

JuReTopDim(ideal J, list w, int tt, int d)  
 remove junk points that are on top-dimensional component

JuReZeroDim(ideal J, list w, int d)  
 remove junk points from 0-dimensional component

WitSupSet(ideal I)  
 witness point super set

WitSet(ideal I)  
 witness point set

```

NumIrrDecom(ideal I)
 numerical irreducible decomposition

defl(ideal I, int d)
 deflation of ideal I

NumPrimDecom(ideal I, int d)
 numerical primary decomposition

```

### D.5.8 orbitparam\_lib

**Library:** orbitparam.lib

**Purpose:** Parametrizing orbits of unipotent actions

**Authors:** J. Boehm, boehm at mathematik.uni-kl.de  
S. Papadakis, papadak at math.ist.utl.pt

**Overview:** This library implements the theorem of Chevalley-Rosenlicht as stated in Theorem 3.1.4 of [Corwin, Greenleaf]. Given a set of strictly upper triangular  $n \times n$  matrices  $L_1, \dots, L_c$  which generate a Lie algebra as a vector space, and a vector  $v$  of size  $n$ , the function `parametrizeOrbit` constructs a parametrization of the orbit of  $v$  under the action of  $\exp(\langle L_1, \dots, L_c \rangle)$ .

To compute  $\exp$  of the Lie algebra elements corresponding to the parameters we require that the characteristic of the base field is zero or larger than  $n$ .

By determining the parameters from bottom to top this allows you to find an element in the orbit with (at least) as many zeros as the dimension of the orbit.

Note: Theorem 3.1.4 of [Corwin, Greenleaf] uses strictly lower triangular matrices.

**References:**

Laurence Corwin, Frederick P. Greenleaf: Representations of Nilpotent Lie Groups and their Applications: Volume 1, Part 1, Basic Theory and Examples, Cambridge University Press (2004).

**Procedures:**

```

tangentGens(list,matrix)
 Returns elements in the Lie algebra, which form a basis of the tangent
 space of the parametrization.

matrixExp(matrix)
 Matrix exp for nilpotent matrices.

matrixLog(matrix)
 Matrix log for unipotent matrices.

parametrizeOrbit(list,matrix)
 Returns parametrization of the orbit.

maxZeros(list,matrix)
 Determine an element in the orbit with the maximum number of zeroes.

```

### D.5.9 `paraplanecurves.lib`

**Library:** `paraplanecurves.lib`

**Purpose:** Rational parametrization of rational plane curves

**Authors:** J. Boehm, boehm at mathematik.uni-kl.de  
 W. Decker, decker at mathematik.uni-kl.de  
 S. Laplagne, slaplagn at dm.uba.ar  
 F. Seelisch, seelisch at mathematik.uni-kl.de

**Overview:** Suppose  $C = \{f(x,y,z)=0\}$  is a rational plane curve, where  $f$  is homogeneous of degree  $n$  with coefficients in  $\mathbb{Q}$  and absolutely irreducible (these conditions are checked automatically.)

After a first step, realized by a projective automorphism in the procedure `adjointIdeal`,  $C$  satisfies:

- $C$  does not have singularities at infinity  $z=0$ .
- $C$  does not contain the point  $(0:1:0)$  (that is, the dehomogenization of  $f$  with respect to  $z$  is monic as a polynomial in  $y$ ).

Considering  $C$  in the chart  $z > 0$ , the algorithm regards  $x$  as transcendental and  $y$  as algebraic and computes an integral basis in  $\mathbb{C}(x)[y]$  of the integral closure of  $\mathbb{C}[x]$  in  $\mathbb{C}(x,y)$  using the normalization algorithm from Section D.4.23 [`normal.lib`], page 824: see Section D.4.11 [`integralbasis.lib`], page 816. In a future edition of the library, also van Hoeij's algorithm for computing the integral basis will be available.

From the integral basis, the adjoint ideal is obtained by linear algebra. Alternatively, the algorithm starts with a local analysis of the singular locus of  $C$ . Then, for each primary component of the singular locus which does not correspond to ordinary multiple points or cusps, the integral basis algorithm is applied separately. The ordinary multiple points and cusps, in turn, are addressed by a straightforward direct algorithm. The adjoint ideal is obtained by intersecting all ideals obtained locally. The local variant of the algorithm is used by default.

The linear system corresponding to the adjoint ideal maps the curve birationally to a rational normal curve in  $\mathbb{P}^{n-2}$ .

Iterating the anticanonical map, the algorithm projects the rational normal curve to  $\mathbb{P}^1$  for  $n$  odd resp. to a conic  $C_2$  in  $\mathbb{P}^2$  for  $n$  even.

In case  $n$  is even, the algorithm tests whether there is a rational point on  $C_2$  and if so gives a parametrization of  $C_2$  which is defined over  $\mathbb{Q}$ . Otherwise, the parametrization given is defined over a quadratic field extension of  $\mathbb{Q}$ .

By inverting the birational map of  $C$  to  $\mathbb{P}^1$  resp. to  $C_2$ , a parametrization of  $C$  is obtained (defined over  $\mathbb{Q}$  or the quadratic field extension).

**References:**

Janko Boehm: Parametrisierung rationaler Kurven, Diploma Thesis, <http://www.math.uni-sb.de/ag/schreyer/jb/diplom%20janko%20boehm.pdf>

Janko Boehm, Wolfram Decker, Santiago Laplagne, Gerhard Pfister: Local to global algorithms for the Gorenstein adjoint ideal of a curve, *Algorithmic and Experimental Methods in Algebra, Geometry, and Number Theory*, Springer 2018

Theo de Jong: An algorithm for computing the integral closure, *Journal of Symbolic Computation* 26 (3) (1998), p. 273-277

Gert-Martin Greuel, Santiago Laplagne, Frank Seelisch: Normalization of Rings, *Journal of Symbolic Computation* 9 (2010), p. 887-901

Mark van Hoeij: An Algorithm for Computing an Integral Basis in an Algebraic Function Field, *Journal of Symbolic Computation* 18 (1994), p. 353-363, <http://www.math.fsu.edu/~hoeij/papers/comments/jsc1994.html>

**Procedures:**

`adjointIdeal(poly, [...])`  
Adjoint ideal of a plane curve

`invertBirMap(ideal, ideal)`  
Invert a birational map of algebraic varieties

`paraPlaneCurve(poly, [...])`  
Compute a rational parametrization of a rational plane curve

`rncAntiCanonicalMap(ideal)`  
Anticanonical map of a rational normal curve

`rationalPointConic(poly)`  
Finds a point on the conic. This point has either coefficients in  $\mathbb{Q}$  or in a quadratic extension field of  $\mathbb{Q}$

`mapToRatNormCurve(poly, ideal)`  
Map a plane rational curve to a rational normal curve (RNC)

`rncItProjOdd(ideal)`  
Map a RNC via successive anticanonical maps to PP1

`rncItProjEven(ideal)`  
Map a RNC via successive anticanonical maps to a conic in PP2

`paraConic(poly)`  
Compute a rational parametrization of a conic

`testParametrization(poly, ring)`  
Checks whether a given curve is parametrized by a given rational map (defined in the given ring)

`testPointConic(poly, ring)`  
Checks whether a given point (defined in the given ring) lies on the given conic.

**D.5.10 resbinomial.lib**

Todos/Issues:

formatting is inappropriate

avoid export

bad names(or should be static): identifyvars, elimrep, convertdata, lcmofall, genoutput, salida, iniD, reslist, sumlist, dividelist, createlist

**Library:** resbinomial.lib

**Purpose:** Combinatorial algorithm of resolution of singularities of binomial ideals in arbitrary characteristic. Binomial resolution algorithm of Blanco

**Authors:** R. Blanco, [mariarocio.blanco@uclm.es](mailto:mariarocio.blanco@uclm.es),  
G. Pfister, [pfister@mathematik.uni-kl.de](mailto:pfister@mathematik.uni-kl.de)

**Procedures:**

**BINresol(J)**  
 computes a E-resolution of singularities of (J) (THE SECOND PART IS NOT IMPLEMENTED YET)

**Eresol(J)**  
 computes a E-resolution of singularities of (J) in char 0

**determinecenter(L1,L2,c,n,Y,a,mb,flag,control3)**  
 computes the next blowing-up center

**Blowupcenter(L1,id,m,L2,c,n,h)**  
 makes the blowing-up

**Nonhyp(Coef,expJ,sJ,n,flag,sums)**  
 computes the ideal generated by the non hyperbolic generators of expJ

**identifyvar()**  
 identifies status of variables

**EdataList(Coef,Exp,k,n,flag)**  
 gives the E-order of each term in Exp

**EOrdList(Coef,Exp,k,n,flag)**  
 computes the E-order of an ideal (giving in the language of lists)

**maxEord(Coef,Exp,k,n,flag)**  
 computes de maximum E-order of an ideal given by Coef and Exp

**ECoeff(Coef,expP,sP,V,auxc,n,flag)**  
 Computes a simplified version of the E-Coeff ideal. The E-orders are correct, but transformations of coefficients of the generators and powers of binomials cannot be computed easily in terms of lists.

**elimrep(L)**  
 removes repeated terms from a list

**Emaxcont(Coef,Exp,k,n,flag)**  
 computes a list of hypersurfaces of E-maximal contact

**cleanunit(mon,n,flag)**  
 clean the units in a monomial mon

**resfunction(t,auxinv,nchart,n)**  
 composes the E-resolution function

**calculateI(Coef,J,c,n,Y,a,b,D)**  
 computes the order of the non monomial part of an ideal J

**Maxord(L,n)**  
 computes the maximum exponent of an exceptional monomial ideal

**Gamma(L,c,n)**  
 computes the Gamma function for an exceptional monomial ideal given by L

**convertdata(C,L,n,flag)**  
 computes the ideal corresponding to C,L

**lcmofall(nchart,mobile)**  
 computes the lcm of the denominators of the E-orders for all the charts

`computemcm(Eolist)`  
 computes the lcm of the denominators of the E-orders for one chart

`constructH(Hhist,n,flag)`  
 construct the list of exceptional divisors accumulated at this chart

`constructblwup(blwhist,n,chy,flag)`  
 construct the ideal defining the map  $K[W] \rightarrow K[W_i]$ , which gives the composition map of all the blowing up leading to this chart

`constructlastblwup(blwhist,n,chy,flag)`  
 construct the ideal defining the last blowup leading to this chart

`genoutput(chart,mobile,nchart,nsons,n,q,p)`  
 generates the output for visualization

`salida(idchart,chart,mobile,numson,previous,a,n,q)`  
 generates the output for one chart

`iniD(n)` creates a list of lists of zeros of size  $n$

`sumlist(L1,L2)`  
 sums two lists component to component

`reslist(L1,L2)`  
 subtracts two lists component to component

`multiplylist(L,a)`  
 multiplies a list by a number, component to component

`dividelist(L1,L2)`  
 divides two lists component to component

`createlist(L1,L2)`  
 creates a list of lists of two elements

### D.5.11 resgraph.lib

**Library:** resgraph.lib

**Purpose:** Visualization of Resolution Data

**Author:** A. Fruehbis-Krueger, anne@mathematik.uni-kl.de,

**Note:** This library uses the external programs surf, graphviz and imagemagick.  
 Input data is assumed to originate from resolve.lib and reszeta.lib

**Procedures:**

`InterDiv(M[,name])`  
 dual graph of resolution of a surface (uses graphviz,imagemagick)

`ResTree(L,M[,name])`  
 tree of charts of resolution (uses graphviz,imagemagick)

`finalCharts(L,...)`  
 pictures of final charts of surface (uses surf)



### D.5.12 resjung\_lib

**Library:** jung.lib

**Purpose:** Resolution of surface singularities (Desingularization) Algorithm of Jung

**Author:** Philipp Renner, philipp\_renner@web.de

**Procedures:**

`jungresolve(J[,is_noeth])`  
 computes a resolution (!not a strong one) of the surface given by the ideal J using Jungs Method,

`jungnormal(J[,is_noeth])`  
 computes a representation of J such that all it's singularities are of Hirzebruch-Jung type,

`jungfib(J[,is_noeth])`  
 computes a representation of J such that all it's singularities are quasi-ordinary

### D.5.13 resolve.lib

**Library:** resolve.lib

**Purpose:** Resolution of singularities (Desingularization) Algorithm of Villamayor

**Authors:** A. Fruehbis-Krueger, anne@mathematik.uni-kl.de,  
 G. Pfister, pfister@mathematik.uni-kl.de

**References:**

- [1] J.Kollar: Lectures on Resolution of Singularities, Princeton University Press (2007) (contains large overview over various known methods for curves and surfaces as well as a detailed description of the approach in the general case)
- [2] A.Bravo, S.Encinas, O.Villamayor: A Simplified Proof of Desingularisation and Applications, Rev. Math. Iberoamericana 21 (2005), 349-458 (description of the algorithmic proof of desingularization in characteristic zero which underlies this implementation)
- [3] A.Fruehbis-Krueger: Computational Aspects of Singularities, in J.-P. Brasselet, J.Damon et al.: Singularities in Geometry and Topology, World Scientific Publishing, 253-327 (2007) (chapter 4 contains a detailed discussion on algorithmic desingularization and efficiency aspects thereof)

**Procedures:**

`blowUp(J,C[,W,E])`  
 computes the blowing up of the variety  $V(J)$  (considered as embedded in  $V(W)$ ) in the (smooth) center  $V(C)$ ,

`blowUp2(J,C)`  
 computes the blowing up of the variety  $V(J)$  in the (possibly singular) center  $V(C)$

`Center(J[,W,E])`  
 computes 'Villamayor'-center for blow up

`resolve(J)`  
 computes the desingularization of the variety  $V(J)$

`showBO(BO)`  
 prints the content of a BO in more human readable form

`presentTree(L)`  
 prints the final charts in more human readable form

`showDataTypes()`  
 prints help text for output data types

`blowUpBO(BO,C)`  
 computes the blowing up of the variety  $V(BO[1])$  in the center  $V(C)$ . BO is a list (basic object), C is an ideal

`createBO(J,W,E)`  
 creates basic object from input data

`CenterBO(BO)`  
 computes the center for the next blow-up of the given basic object

`Delta(BO)`  
 apply the Delta-operator of [Bravo,Encinas,Villamayor]

`DeltaList(BO)`  
 list of results of  $\Delta^0$  to  $\Delta^{bmax}$

#### D.5.14 `reszeta.lib`

**Library:** `reszeta.lib`

**Purpose:** topological Zeta-function and some other applications of desingularization

**Authors:** A. Fruehbis-Krueger, [anne@mathematik.uni-kl.de](mailto:anne@mathematik.uni-kl.de),  
 G. Pfister, [pfister@mathematik.uni-kl.de](mailto:pfister@mathematik.uni-kl.de)

**References:**

[1] Fruehbis-Krueger,A., Pfister,G.: Some Applications of Resolution of Singularities from a Practical Point of View, in Computational Commutative and Non-commutative Algebraic Geometry, NATO Science Series III, Computer and Systems Sciences 196, 104-117 (2005) [2] Fruehbis-Krueger: An Application of Resolution of Singularities: Computing the topological Zeta-function of isolated surface singularities in  $(\mathbb{C}^3,0)$ , in D.Cheniot, N.Dutertre et al.(Editors): Singularity Theory, World Scientific Publishing (2007)

**Procedures:**

`intersectionDiv(L)`  
 computes intersection form and genera of exceptional divisors (isolated singularities of surfaces)

`spectralNeg(L)`  
 computes negative spectral numbers (isolated hypersurface singularity)

`discrepancy(L)`  
 computes discrepancy of given resolution

`zetaDL(L,d)`  
 computes Denef-Loeser zeta function (hypersurface singularity of dimension 2)

`collectDiv(L[,iv])`  
 identify exceptional divisors in different charts (embedded and non-embedded case)

`prepEmbDiv(L[,b])`  
 prepare list of divisors (including components of strict transform, embedded case)

`abstractR(L)`  
 pass from embedded to non-embedded resolution

`computeV(re,DL)`  
 multiplicities of divisors in pullback of volume form

`computeN(re,DL)`  
 multiplicities of divisors in total transform of resolution

### D.5.15 schubert\_lib

**Library:** schubert.lib

**Purpose:** Proceeds for Intersection Theory

**Author:** Hiep Dang, email: hiep@mathematik.uni-kl.de

**Overview:** We implement new classes (variety, sheaf, stack, graph) and methods for computing with them. An abstract variety is represented by a nonnegative integer which is its dimension and a graded ring which is its Chow ring. An abstract sheaf is represented by a variety and a polynomial which is its Chern character. In particular, we implement the concrete varieties such as projective spaces, Grassmannians, and projective bundles. An important task of this library is related to the computation of Gromov-Witten invariants. In particular, we implement new tools for the computation in equivariant intersection theory. These tools are based on the localization of moduli spaces of stable maps and Bott's formula. They are useful for the computation of Gromov-Witten invariants. In order to do this, we have to deal with moduli spaces of stable maps, which were introduced by Kontsevich, and the graphs corresponding to the fixed point components of a torus action on the moduli spaces of stable maps.

As an insightful example, the numbers of rational curves on general complete intersection Calabi-Yau threefolds in projective spaces are computed up to degree 6. The results are all in agreement with predictions made from mirror symmetry computations.

**References:**

Hiep Dang, Intersection theory with applications to the computation of Gromov-Witten invariants, Ph.D thesis, TU Kaiserslautern, 2013.

Sheldon Katz and Stein A. Stromme, Schubert-A Maple package for intersection theory and enumerative geometry, 1992.

Daniel R. Grayson, Michael E. Stillman, Stein A. Stromme, David Eisenbud and Charley Crissman, Schubert2-A Macaulay2 package for computation in intersection theory, 2010.

Maxim Kontsevich, Enumeration of rational curves via torus actions, 1995.

**Procedures:**

`makeVariety(int, ideal)`  
create a variety

`printVariety(variety)`  
print procedure for a variety

`productVariety(variety, variety)`  
make the product of two varieties

`ChowRing(variety)`  
create the Chow ring of a variety

`Grassmannian(int, int)`  
create a Grassmannian as a variety

`projectiveSpace(int)`  
create a projective space as a variety

`projectiveBundle(sheaf)`  
create a projective bundle as a variety

`integral(variety, poly)`  
degree of a 0-cycle on a variety

`makeSheaf(variety, poly)`  
create a sheaf

`printSheaf(sheaf)`  
print procedure for sheaves

`rankSheaf(sheaf)`  
return the rank of a sheaf

`totalChernClass(sheaf)`  
compute the total Chern class of a sheaf

`ChernClass(sheaf, int)`  
compute the k-th Chern class of a sheaf

`topChernClass(sheaf)`  
compute the top Chern class of a sheaf

`totalSegreClass(sheaf)`  
compute the total Segre class of a sheaf

`dualSheaf(sheaf)`  
make the dual of a sheaf

`tensorSheaf(sheaf, sheaf)`  
make the tensor of two sheaves

`symmetricPowerSheaf(sheaf, int)`  
make the k-th symmetric power of a sheaf

`quotSheaf(sheaf, sheaf)`  
make the quotient of two sheaves

`addSheaf(sheaf, sheaf)`  
make the direct sum of two sheaves

`makeGraphVE(list,list)`  
create a graph from a list of vertices and a list of edges

`printGraphG(graph)`  
print procedure for graphs

`moduliSpace(variety,int)`  
create a moduli space of stable maps as an algebraic stack

`printStack(stack)`  
print procedure for stacks

`dimStack(stack)`  
compute the dimension of a stack

`fixedPoints(stack)`  
compute the list of graphs corresponding the fixed point components of a torus action on the stack

`contributionBundle(stack,graph)`  
compute the contribution bundle on a stack at a graph

`normalBundle(stack,graph)`  
compute the normal bundle on a stack at a graph

`multipleCover(int)`  
compute the contribution of multiple covers of a smooth rational curve as a Gromov-Witten invariant

`linesHypersurface(int)`  
compute the number of lines on a general hypersurface

`rationalCurve(int,list)`  
compute the Gromov-Witten invariant corresponding the number of rational curves on a general Calabi-Yau threefold

`sumofquotients(stack,list)`  
prepare a command for parallel computation

`homog_part(poly,int)`  
compute a homogeneous component of a polynomial.

`homog_parts(poly,int,int)`  
compute the sum of homogeneous components of a polynomial

`logg(poly,int)`  
compute Chern characters from total Chern classes.

`expp(poly,int)`  
compute total Chern classes from Chern characters

`SchubertClass(list)`  
compute the Schubert classes on a Grassmannian

`dualPartition(list)`  
compute the dual of a partition

### D.5.16 sheafcoh\_lib

**Library:** sheafcoh.lib

**Purpose:** Procedures for Computing Sheaf Cohomology

**Authors:** Wolfram Decker, [decker@mathematik.uni-kl.de](mailto:decker@mathematik.uni-kl.de)  
 Christoph Lossen, [lossen@mathematik.uni-kl.de](mailto:lossen@mathematik.uni-kl.de)  
 Gerhard Pfister, [pfister@mathematik.uni-kl.de](mailto:pfister@mathematik.uni-kl.de)  
 Oleksandr Motsak, U@D, where U={motsak}, D={mathematik.uni-kl.de}

**Procedures:**

`truncate(phi,d)`  
 truncation of `coker(phi)` at `d`

`truncateFast(phi,d)`  
 truncation of `coker(phi)` at `d` (fast+ experimental)

`CM_regularity(M)`  
 Castelnuovo-Mumford regularity of `coker(M)`

`sheafCohBGG(M,l,h)`  
 cohomology of sheaf associated to `coker(M)`

`sheafCohBGG2(M,l,h)`  
 cohomology of sheaf associated to `coker(M)`, experimental version

`sheafCoh(M,l,h)`  
 cohomology of sheaf associated to `coker(M)`

`dimH(i,M,d)`  
 compute  $h^i(F(d))$ , `F` sheaf associated to `coker(M)`

`dimGradedPart()`

`displayCohom(B,l,h,n)`  
 display intmat as Betti diagram (with zero rows)

### D.5.17 JMBTest\_lib

**Library:** JMBTest.lib

**Purpose:** A library for Singular which performs JM basis test.

**Author:** Michela Ceria, email: [michela.ceria@unito.it](mailto:michela.ceria@unito.it)

**Overview:** The library performs the J-marked basis test, as described in [CR], [BCLR]. Such a test is performed via the criterion explained in [BCLR], concerning Eliahou-Kervaire polynomials (EK from now on). We point out that all the polynomials are homogeneous and they must be arranged by degree.

The fundamental steps are the following:

- construct the  $V_m$  polynomials, via the algorithm `VConstructor` explained in [CR];
- construct the Eliahou-Kervaire polynomials defined in [BCLR];
- reduce the Eliahou-Kervaire polynomials using the  $V_m$ 's;
- if it exist an Eliahou-Kervaire polynomial such that its reduction mod  $V_m$  is different from zero, the given one is not a J-Marked basis.

The algorithm terminates only if the ordering is `rp`. Anyway, the number of reduction steps is bounded.

**References:**

- [CR] Francesca Cioffi, Margherita Roggero, Flat Families by Strongly Stable Ideals and a Generalization of Groebner Bases, *J. Symbolic Comput.* 46, 1070-1084, (2011).  
 [BCLR] Cristina Bertone, Francesca Cioffi, Paolo Lella, Margherita Roggero, Upgraded methods for the effective computation of marked schemes on a strongly stable ideal, *Journal of Symbolic Computation* (2012), <http://dx.doi.org/10.1016/j.jsc.2012.07.006>

**Procedures:**

- `Minimus(ideal)`  
 minimal variable in an ideal
- `Maximus(ideal)`  
 maximal variable in an ideal
- `StartOrderingV(list, list)`  
 ordering of polynomials as in [BCLR]
- `TestJMark(list)`  
 tests whether we have a J-marked basis

See also: Section D.5.18 [JMSTest.lib], page 853.

**D.5.18 JMSTest.lib**

**Library:** JMSTest.lib

**Purpose:** A library for Singular which constructs J-Marked Schemes.

**Author:** Michela Ceria, email: [michela.ceria@unito.it](mailto:michela.ceria@unito.it)

**Overview:** The library performs the J-marked computation, as described in [BCLR]. As in JMBTest.lib we construct the V polynomials and we reduce the EK polynomials w.r.t. them, putting the coefficients as results.

The algorithm terminates only if the ordering is rp. Anyway, the number of reduction steps is bounded.

**References:**

- [CR] Francesca Cioffi, Margherita Roggero, Flat Families by Strongly Stable Ideals and a Generalization of Groebner Bases, *J. Symbolic Comput.* 46, 1070-1084, (2011).  
 [BCLR] Cristina Bertone, Francesca Cioffi, Paolo Lella, Margherita Roggero, Upgraded methods for the effective computation of marked schemes on a strongly stable ideal, *Journal of Symbolic Computation* (2012), <http://dx.doi.org/10.1016/j.jsc.2012.07.006>

**Procedures:**

- `BorelCheck(ideal, r)`  
 checks whether the given ideal is Borel
- `JMarkedScheme(list, list, list, int)`  
 computes automatically all the J-marked scheme

See also: Section D.5.17 [JMBTest.lib], page 852; Section D.5.18 [JMSTest.lib], page 853.

## D.6 Singularities

### D.6.1 alexpoly\_lib

**Library:** alexpoly.lib

**Purpose:** Resolution Graph and Alexander Polynomial

**Author:** Fernando Hernando Carrillo, hernando@agt.uva.es  
Thomas Keilen, keilen@mathematik.uni-kl.de

**Overview:** A library for computing the resolution graph of a plane curve singularity  $f$ , the total multiplicities of the total transforms of the branches of  $f$  along the exceptional divisors of a minimal good resolution of  $f$ , the Alexander polynomial of  $f$ , and the zeta function of its monodromy operator.

**Procedures:**

`resolutiongraph(f)`  
resolution graph  $f$

`totalmultiplicities(f)`  
resolution graph, total multiplicities and strict multiplicities of  $f$

`alexanderpolynomial(f)`  
Alexander polynomial of  $f$

`semigroup(f)`  
calculates generators for the semigroup of  $f$

`proximitymatrix(f)`  
calculates the proximity matrix of  $f$

`multseq2charexp(v)`  
converts multiplicity sequence to characteristic exponents

`charexp2multseq(v)`  
converts characteristic exponents to multiplicity sequence

`charexp2generators(v)`  
converts characteristic exponents to generators of the semigroup

`charexp2inter(c,e)`  
converts contact matrix and charact. exp. to intersection matrix

`charexp2conductor(v)`  
converts characteristic exponents to conductor

`charexp2poly(v,a)`  
calculates a polynomial  $f$  with characteristic exponents  $v$

`tau_es2(f)`  
equisingular Tjurina number of  $f$

### D.6.2 arcpoint\_lib

**Library:** arcpoint.lib

**Purpose:** Truncations of arcs at a singular point



**Author:** Nadine Cremer cremer@mathematik.uni-kl.de

**Overview:** An arc is given by a power series in one variable, say  $t$ , and truncating it at a positive integer  $i$  means cutting the  $t$ -powers  $> i$ . The set of arcs truncated at order  $<bound>$  is denoted  $\text{Tr}(i)$ . An algorithm for computing these sets (which happen to be constructible) is given in [Lejeune-Jalabert, M.: Courbes trac'ees sur un germe d'hypersurface, American Journal of Mathematics, 112 (1990)]. Our procedures for computing the locally closed sets contributing to the set of truncations rely on this algorithm.

**Procedures:**

`nashmult(f, bound)`

determines locally closed sets relevant for computing truncations of arcs over a hypersurface with isolated singularity defined by  $f$ . The sets are given by two ideals specifying relations between coefficients of power series in  $t$ . One of the ideals defines an open set, the other one the complement of a closed set within the open one. We consider only coefficients up to  $t^{<bound>}$ . Moreover, the sequence of Nash Multiplicities of each set is displayed

`removepower(I)`

modifies the ideal  $I$  such that the algebraic set defined by it remains the same: removes powers of variables

`idealsimplify(I, maxiter)`

further simplification of  $I$  in the above sense: reduction with other elements of  $I$ . The positive integer  $<maxiter>$  gives a bound to the number of repetition steps

`equalJinI(I, J)`

tests if two ideals  $I$  and  $J$  are equal under the assumption that  $J$  is contained in  $I$ . Returns 1 if this is true and 0 otherwise

### D.6.3 arnoldclassify\_lib

**Library:** arnoldClassify.lib

**Purpose:** Arnol'd Classifier of Singularities

**Author:** Eva Maria Hemmerling, ehemmerl@rhrk.uni-kl.de

**Overview:** A library for classifying isolated hypersurface singularities from the list of V.I. Arnol'd w.r.t. right equivalence up to corank 2. The method relies on Baciu's list of Milnor codes and Newton's rotating ruler method to distinguish the Y- and Z- singularities.

**References:**

- [AVG85] Arnold, Varchenko, Gusein-Zade: Singularities of Differentiable Maps. Vol. 1: The classification of critical points caustics and wave fronts. Birkh"auser, Boston 1985
- [Bac01] Corina Baciu: The classification of Hypersurface Singularities using the Milnor Code, Diplomarbeit, Universit"at Kaiserslautern, 2001.
- [GP12] Greuel, Pfister: A SINGULAR Introduction to Commutative Algebra, Springer Science and Business Media, 2012
- [Hem17] Eva Maria Hemmerling: Algorithmic Arnol'd Classification in SINGULAR, Master Thesis, TU Kaiserslautern, 2017.

**Procedures:**

`arnoldListAllSeries()`  
list of all singularity series up to corank 2

`arnoldShowSeries(S)`  
data defining a singularity series S

`arnoldNormalForm(S,#)`  
normalform for a singularity series S

`arnoldClassify(f)`  
singularity class of a power series f

`arnoldCorank(f)`  
corank of singularity defined by f

`arnoldDeterminacy(f,#)`  
upper bound for the determinacy of f

`arnoldMilnorCode(f,#)`  
Milnor Code of a singularity f

`arnoldMorseSplit(f,#)`  
result of Splitting Lemma applied to f

See also: Section D.6.4 [`classify_lib`], page 856; Section D.6.19 [`realclassify_lib`], page 867.

**D.6.4 `classify_lib`**

**Library:** `classify.lib`

**Purpose:** Arnold Classifier of Singularities

**Author:** Kai Krueger, `krueger@mathematik.uni-kl.de`

**Overview:** A library for classifying isolated hypersurface singularities w.r.t. right equivalence, based on the determinant of singularities by V.I. Arnold.

**Procedures:**

`basicinvariants(f)`  
computes Milnor number, determinacy-bound and corank of

`classify(f)`  
normal form of polynomial f determined with Arnold's method

`corank(f)`  
computes the corank of f (i.e. of the Hessian of f)

`Hcode(v)` coding of intvec v according to the number repetitions

`init_debug([n])`  
print trace and debugging information depending on int n

`internalfunctions()`  
display names of internal procedures of this library

`milnorcode(f[,e])`  
Hilbert polynomial of [e-th] Milnor algebra coded with Hcode

`morsesplit(f)`  
residual part of f after applying the splitting lemma

`quickclass(f)`  
     normal form of  $f$  determined by invariants (milnorcode)  
`singularity(s, [])`  
     normal form of singularity given by its name  $s$  and index  
`A_L(s/f)` shortcut for `quickclass(f)` or `normalform(s)`  
`normalform(s)`  
     normal form of singularity given by its name  $s$   
`debug_log(lev, [])`  
     print trace and debugging information w.r.t level>@DeBug  
`swap(a,b)`  
     swaps the arguments  
`modality(f)`  
     modality of the singularity  
`complexSingType(f)`  
     complex type of the singularity as a string  
`prepRealclassify(f)`  
     the modality and the complex type of the singularity at once

See also: Section D.6.19 [realclassify\_lib], page 867.

### D.6.5 classify2\_lib

**Library:** classify2.lib

**Purpose:** Classification of isolated singularities

**Authors:** Janko Boehm, email: boehm@mathematik.uni-kl.de  
 Magdaleen Marais, email: magdaleen.marais@up.ac.za  
 Gerhard Pfister, email: pfister@mathematik.uni-kl.de

**Overview:** We classify isolated singularities of corank  $\leq 2$  and modality  $\leq 2$  with respect to right-equivalence over the complex numbers according to Arnold's list. We determine the type and, for positive modality, the parameter.

V.I. Arnold has described normal forms and has developed a classifier for, in particular, all isolated hypersurface singularities over the complex numbers up to modality 2. Building on a series of 105 theorems, this classifier determines the type of the given singularity. However, for positive modality, this does not fix the right equivalence class of the singularity, since the values of the moduli parameters are not specified.

This library implements an alternative classification algorithm for isolated hypersurface singularities of corank and modality up to two. For a singularity given by a polynomial over the rationals, the algorithm determines its right equivalence class by specifying a polynomial representative in Arnold's list of normal forms. In particular, the algorithm also determines values for the moduli parameters.

The implementation is based on the paper

Janko Boehm, Magdaleen Marais, Gerhard Pfister: A Classification Algorithm for Complex Singularities of Corank and Modality up to Two, Singularities and Computer Algebra - Festschrift for Gert-Martin Greuel on the Occasion of his 70th Birthday, Springer 2017, <http://arxiv.org/abs/1604.04774>, [https://doi.org/10.1007/978-3-319-28829-1\\_2](https://doi.org/10.1007/978-3-319-28829-1_2)

There are functions for determining a normal form equation and for determining the complex type of the singularity.

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**Procedures:**

`complexClassify(I)`  
 classifier returning a normal form equation

`complexType(I)`  
 classifier returning the type and modality

See also: Section D.6.4 [`classify_lib`], page 856; Section D.6.19 [`realclassify_lib`], page 867.

### D.6.6 `classify_aeq_lib`

**Library:** `classifyAeq.lib`

**Purpose:** Simple Space Curve singularities in characteristic 0

**Authors:** Faira Kanwal Janjua fairakanwaljanjua@gmail.com  
 Gerhard Pfister pfister@mathematik.uni-kl.de Khawar Mehmood  
 khawar1073@gmail.com

**Overview:** A library for classifying the simple singularities with respect to A equivalence in characteristic 0. Simple Surface singularities in characteristic 0 have been classified by Bruce and Gaffney [4] resp. Gibson and Hobbs [1] with respect to A equivalence. If the input is one of the simple singularities in [1] it returns a normal form otherwise a zero ideal (i.e not simple).

**References:**

- [1] Gibson,C.G; Hobbs,C.A.:Simple SIngularities of Space Curves. Math.Proc. Comb.Phil.Soc.(1993),113,297.
- [2] Hefez,A;Hernandes,M.E.:Standard bases for local rings of branches and their modules of differentials. Journal of Symbolic Computation 42(2007) 178-191. [3] Hefez,A;Hernandes,M.E.:The Analytic Classification Of Plane Branches. Bull.Lond Math Soc.43.(2011) 2,289-298. [4] Bruce, J.W.,Gaffney, T.J.: Simple singularities of mappings  $(C, 0) \rightarrow (C^2, 0)$ . J. London Math. Soc. (2) 26 (1982), 465-474.
- [5] Ishikawa,G; Janeczko,S.: The Complex Symplectic Moduli Spaces of Unimodal Parametric Plane Curve Singularities. Insitute of Mathematics of the Polish Academy of Sciences,Preprint 664(2006)

**Procedures:**

`sagbiAlg(G)`  
 Compute the Sagbi-basis of the Algebra.

`sagbiMod(I,A)`  
 Compute the Sagbi- basis of the Module.

`semiGroup(G)`  
 Compute the Semi-Group of the Algebra provided the input is Sagbi Bases of the Algebra.

`semiMod(I,A)`  
 Compute the Semi-Module provided that the input are the Sagbi Bases of the Algebra resp.Module.

`planeCur(I)`  
 Compute the type of the Simple Plane Curve singularity.

`spaceCur(I)`  
 Compute the type of the simple Space Curve singularity.

`HHnormalForm(I)`  
 computes for the parametrization defined by I normal form, semi group, semi module of differentials, Zariski number and moduli

`modSagbiAlg(G)`  
 modular variant of sagbiAlg

`classSpaceCurve(I)`  
 normal form of I if I is simple

### D.6.7 classifyceq\_lib

**Library:** classifyCeq.lib

**Purpose:** simple hypersurface singularities in characteristic  $p > 0$

**Authors:** Deeba Afzal deebafzal@gmail.com  
 Faira Kanwal Janjua fairakanwaljanjua@gmail.com

**Overview:** A library for classifying the simple singularities with respect to contact equivalence in characteristic  $p > 0$ . Simple hypersurface singularities in characteristic  $p > 0$  were classified by Greuel and Kroening [1] with respect to contact equivalence. The classifier we use has been proposed in [2].

**References:**

- [1] Greuel, G.-M.; Kroening, H.: Simple singularities in positive characteristic. Math.Z. 203, 339-354 (1990).
- [2] Afzal,D.;Binyamin,M.A.;Janjua,F.K.: On the classification of simple singularities in positive characteristic.

**Procedures:**

`classifyCeq(f)`  
 simple hypersurface singularities in characteristic  $p > 0$

### D.6.8 classifyci\_lib

**Library:** classifyci.lib

**Purpose:** Isolated complete intersection singularities in characteristic 0

**Authors:** Gerhard Pfister pfister@mathematik.uni-kl.de  
 Deeba Afzal deebafzal@gmail.com

**Overview:** A library for classifying isolated complete intersection singularities for the base field of characteristic 0 and for computing weierstrass semigroup of the space curve. Isolated complete intersection singularities were classified by M.Giusti [1] for the base field of characteristic 0. Algorithm for the semigroup of a space curve singularity is given in [2].

**References:**

- [1] Giusti,M:Classification des singularities isolees simples d'intersections completes, C.R.Acad.Sci.Paris Ser.A-B 284(1977),167-169.
- [2] Castellanos,A.,Castellanos,J.,2005:Algorithm for the semigroup of a space curve singularity. Semigroup Forum 70,44-66.

**Procedures:**

- `classifyicis(I)`  
Isolated simple complete intersection singularities for the base field of charateristic 0
- `Semigroup(I)`  
Weierstrass semigroup of the space curve given by equations

**D.6.9 classifyMapGerms.lib****Library:** classifyMapGerms.lib**Authors:** Gerhard Pfister, pfister@mathematik.uni-kl.de  
Deeba Afzal, deebafzal@gmail.com  
Shamsa Kanwal, lotus\_zone16@yahoo.com**Overview:** A library for computing the standard basis of the tangent space at the orbit of an algebraic group action. The tangent space is usually described as the sum of two modules over different rings. It computes the standard basis using modular methods and parallel modular methods. It also computes the normal form of the germ given by Riegers classification.**References:**

- [1] Idrees N.; Pfister, G.; Steidel, S.: Parallelization of modular algorithms. J. Symbolic Comput. 46(2011), no. 6, 672-684.
- [2] Gibson,C.G; Hobbs,C.A.: Simple SIngularities of Space Curves. Math.Proc. Comb.Phil.Soc.(1993),113,297.
- [3] Bruce, J.W.,Gaffney, T.J.: Simple singularities of mappings  $(C, 0) \rightarrow (C^2, 0)$ . J. London Math. Soc. (2) 26 (1982), 465-474.
- [4] Rieger, J. H.: Families of maps from the plane to the plane. J. London Math. Soc. (2)36(1987), no. 2. 351-369.

**Procedures:**

- `coDimMap(I)`  
computes a bound of the A-determinacy of the map germ defined by I
- `coDim(M,N,I,b)`  
computes the K-vectorspace dimension of  $A^r/M+N+\maxideal(b)*A^r$
- `vStd(M,N,I,b)`  
computes a standard basis of  $M+N+\maxideal(b)*A^r$
- `modVStd(M,N,I,b)`  
computes a standard basis of  $M+N+\maxideal(bound)*A^r$  (modular)
- `modVStd0(M,N,I,b)`  
computes a standard basis of  $M+N+\maxideal(bound)*A^r$  (parallel)
- `classifySimpleMaps(I)`  
computes the normal form of a germ in Riegers classification

`classifySimpleMaps1(I)`  
 computes the normal form of a germ in Riegers classification

`classifyUnimodalMaps(I)`  
 computes the normal form of a germ in Riegers classification

### D.6.10 curvepar.lib

**Library:** curvepar.lib

**Purpose:** Resolution of space curve singularities, semi-group

**Author:** Gerhard Pfister email: pfister@mathematik.uni-kl.de Nil Sahin email:  
 e150916@metu.edu.tr  
 Maryna Viazovska email: viazovsk@mathematik.uni-kl.de

**Procedures:**

`BlowingUp(f,I,l)`  
 BlowingUp of  $V(I)$  at the point 0;

`CurveRes(I)`  
 Resolution of  $V(I)$

`CurveParam(I)`  
 Parametrization of algebraic branches of  $V(I)$

`WSemigroup(X,b)`  
 Weierstrass semigroup of the curve

`primparam(x,y,c)`  
 HN matrix of parametrization  $(x(t),y(t))$

`MultiplicitySequence(I)`  
 Multiplicity sequences of the branches of plane curve  $V(I)$

`CharacteristicExponents(I)`  
 Characteristic exponents of the branches of plane curve  $V(I)$

`IntersectionMatrix(I)`  
 Intersection Matrix of the branches of plane curve  $V(I)$

`ContactMatrix(I)`  
 Contact Matrix of the branches of plane curve  $V(I)$

`plainInvariants(I)`  
 Invariants of the branches of plane curve  $V(I)$

See also: Section D.6.21 [spcurve.lib], page 869.

### D.6.11 deform.lib

**Library:** deform.lib

**Purpose:** Miniversal Deformation of Singularities and Modules

**Author:** Bernd Martin, email: martin@math.tu-cottbus.de

**Procedures:**

`versal(Fo[,d,any])`  
 miniversal deformation of isolated singularity Fo

`mod_versal(Mo,I,[,d,any])`  
 miniversal deformation of module Mo modulo ideal I

`lift_kbase(N,M)`  
 lifting N into standard kbase of M

`lift_rel_kb(N,M[,kbM,p])`  
 relative lifting N into a kbase of M

### D.6.12 equising\_lib

**Library:** equising.lib

**Purpose:** Equisingularity Stratum of a Family of Plane Curves

**Author:** Christoph Lossen, lossen@mathematik.uni-kl.de  
 Andrea Mindnich, mindnich@mathematik.uni-kl.de

**Procedures:**

`tau_es(f)`  
 codim of mu-const stratum in semi-universal def. base

`esIdeal(f)`  
 (Wahl's) equisingularity ideal of f

`esStratum(F[,m,L])`  
 equisingularity stratum of a family F

`isEquising(F[,m,L])`  
 tests if a given deformation is equisingular

`control_Matrix(M)`  
 computes list of blowing-up data

### D.6.13 gmssing\_lib

**Library:** gmssing.lib

**Purpose:** Gauss-Manin System of Isolated Singularities

**Author:** Mathias Schulze, mschulze at mathematik.uni-kl.de

**Overview:** A library for computing invariants related to the Gauss-Manin system of an isolated hypersurface singularity.

**References:**

- [Sch01] M. Schulze: Algorithms for the Gauss-Manin connection. J. Symb. Comp. 32,5 (2001), 549-564.
- [Sch02] M. Schulze: The differential structure of the Brieskorn lattice. In: A.M. Cohen et al.: Mathematical Software - ICMS 2002. World Scientific (2002).
- [Sch03] M. Schulze: Monodromy of Hypersurface Singularities. Acta Appl. Math. 75 (2003), 3-13.
- [Sch04] M. Schulze: A normal form algorithm for the Brieskorn lattice. J. Symb. Comp. 38,4 (2004), 1207-1225.

**Procedures:**

`gmsring(t,s)`  
 Gauss-Manin system of t with variable s



`gmsnf(p,K)`  
 Gauss-Manin normal form of  $p$

`gmscoeffs(p,K)`  
 Gauss-Manin basis representation of  $p$

`bernstein(t)`  
 Bernstein-Sato polynomial of  $t$

`monodromy(t)`  
 Jordan data of complex monodromy of  $t$

`spectrum(t)`  
 singularity spectrum of  $t$

`sppairs(t)`  
 spectral pairs of  $t$

`vfilt(t)`  $V$ -filtration of  $t$  on Brieskorn lattice

`wvfilt(t)`  
 weighted  $V$ -filtration of  $t$  on Brieskorn lattice

`tmatrix(t)`  
 matrix of  $t$  w.r.t. good basis of Brieskorn lattice

`endvfilt(V)`  
 endomorphism  $V$ -filtration on Jacobian algebra

`sppnf(a,w[,m])`  
 spectral pairs normal form of  $(a,w[,m])$

`spprint(spp)`  
 print spectral pairs  $spp$

`spadd(sp1,sp2)`  
 sum of spectra  $sp1$  and  $sp2$

`spsub(sp1,sp2)`  
 difference of spectra  $sp1$  and  $sp2$

`spmul(sp0,k)`  
 linear combination of spectra  $sp$

`spissemicont(sp[,opt])`  
 semicontinuity test of spectrum  $sp$

`spsemicont(sp0,sp[,opt])`  
 semicontinuous combinations of spectra  $sp0$  in  $sp$

`spmilnor(sp)`  
 Milnor number of spectrum  $sp$

`spgeomgenus(sp)`  
 geometrical genus of spectrum  $sp$

`spgamma(sp)`  
 gamma invariant of spectrum  $sp$

See also: Section 7.5.4 [`dmod_lib`], page 395; Section D.6.14 [`gmspoly_lib`], page 864; Section D.6.17 [`monodromy_lib`], page 866; Section D.6.22 [`spectrum_lib`], page 870.

### D.6.14 gmspoly\_lib

**Library:** gmspoly.lib

**Purpose:** Gauss-Manin System of Tame Polynomials

**Author:** Mathias Schulze, mschulze at mathematik.uni-kl.de

**Overview:** A library for computing the Gauss-Manin system of a cohomologically tame polynomial  $f$ . Schulze's algorithm [Sch05], based on Sabbah's theory [Sab98], is used to compute a good basis of (the Brieskorn lattice of) the Gauss-Manin system and the differential operation of  $f$  in terms of this basis. In addition, there is a test for tameness in the sense of Broughton. Tame polynomials can be considered as an affine algebraic analogue of local analytic isolated hypersurface singularities. They have only finitely many critical points, and those at infinity do not give rise to atypical values in a sense depending on the precise notion of tameness considered. Well-known notions of tameness like tameness, M-tameness, Malgrange-tameness, and cohomological tameness, and their relations, are reviewed in [Sab98,8]. For ordinary tameness, see Broughton [Bro88,3]. Sabbah [Sab98] showed that the Gauss-Manin system, the D-module direct image of the structure sheaf, of a cohomologically tame polynomial carries a similar structure as in the isolated singularity case, coming from a Mixed Hodge structure on the cohomology of the Milnor (typical) fibre (see gmssing.lib). The data computed by this library encodes the differential structure of the Gauss-Manin system, and the Mixed Hodge structure of the Milnor fibre over the complex numbers. As a consequence, it yields the Hodge numbers, spectral pairs, and monodromy at infinity.

**References:**

- [Bro88] S. Broughton: Milnor numbers and the topology of polynomial hypersurfaces. *Inv. Math.* 92 (1988) 217-241.
- [Sab98] C. Sabbah: Hypergeometric periods for a tame polynomial. [arXiv.org math.AG/9805077](https://arxiv.org/abs/math/9805077).
- [Sch05] M. Schulze: Good bases for tame polynomials. *J. Symb. Comp.* 39,1 (2005), 103-126.

**Procedures:**

`isTame(f)`  
                   test whether the polynomial  $f$  is tame

`goodBasis(f)`  
                   good basis of Brieskorn lattice of cohom. tame polynomial  $f$

See also: Section D.6.13 [gmssing\_lib], page 862.

### D.6.15 hnoether\_lib

**Library:** hnoether.lib

**Purpose:** Hamburger-Noether (Puiseux) Expansion

**Authors:** Martin Lamm, lamm@mathematik.uni-kl.de  
 Christoph Lossen, lossen@mathematik.uni-kl.de

**Overview:** A library for computing the Hamburger-Noether expansion (analogue of Puiseux expansion over fields of arbitrary characteristic) of a reduced plane curve singularity following [Campillo, A.: Algebroid curves in positive characteristic, Springer LNM 813

(1980)].

The library contains also procedures for computing the (topological) numerical invariants of plane curve singularities.

**Procedures:**

`hnexpansion(f [, "ess"])`  
Hamburger-Noether (HN) expansion of  $f$

`develop(f [, n])`  
HN expansion of irreducible plane curve germs

`extdevelop(hne, n)`  
extension of the H-N expansion  $hne$  of  $f$

`param(hne [, s])`  
parametrization of branches described by HN data

`displayHNE(hne)`  
display HN expansion as an ideal

`invariants(hne)`  
invariants of  $f$ , e.g. the characteristic exponents

`displayInvariants(hne)`  
display invariants of  $f$

`multsequence(hne)`  
sequence of multiplicities

`displayMultsequence(hne)`  
display sequence of multiplicities

`intersection(hne1, hne2)`  
intersection multiplicity of two local branches

`is_irred(f)`  
test whether  $f$  is irreducible as power series

`delta(f)` delta invariant of  $f$

`newtonpoly(f)`  
(local) Newton polygon of  $f$

`is_NND(f)`  
test whether  $f$  is Newton non-degenerate

`stripHNE(hne)`  
reduce amount of memory consumed by  $hne$

`puiseux2generators(m, n)`  
convert Puiseux pairs to generators of semigroup

`separateHNE(hne1, hne2)`  
number of quadratic transf. needed for separation

`squarefree(f)`  
a squarefree divisor of the polynomial  $f$

`allsquarefree(f, 1)`  
the maximal squarefree divisor of the polynomial  $f$

`further_hn_proc()`  
show further procedures useful for interactive use

**D.6.16 kskernel\_lib****Library:** kskernel.lib**Purpose:** procedures for computing the kernel of the kodaira-spencer map**Author:** Tetyana Povalyaeva, povalyae@mathematik.uni-kl.de**Procedures:**

- KSkerr**(p,q)  
kernel of the Kodaira-Spencer map of a versal deformation of an irreducible plane curve singularity
- KSconvert**(M)  
kernel of the Kodaira-Spencer map in quasihomogeneous variables T with corresponding negative degrees
- KSlinear**(M)  
matrix of linear terms of the kernel of the Kodaira-Spencer map
- KScoef**(i,j,P,Q,qq)  
coefficient of the given term in the matrix of kernel of the Kodaira-Spencer map
- StringF**(i,j,p,q)  
expression in variables T(i) with non-resolved brackets for the further computation of coefficient in the matrix of kernel of the Kodaira-Spencer map

**D.6.17 mondromy\_lib****Library:** mondromy.lib**Purpose:** Monodromy of an Isolated Hypersurface Singularity**Author:** Mathias Schulze, email: mschulze@mathematik.uni-kl.de

**Overview:** A library to compute the monodromy of an isolated hypersurface singularity. It uses an algorithm by Brieskorn (manuscripta math. 2 (1970), 103-161) to compute a connection matrix of the meromorphic Gauss-Manin connection up to arbitrarily high order, and an algorithm of Gerard and Levelt (Ann. Inst. Fourier, Grenoble 23,1 (1973), pp. 157-195) to transform it to a simple pole.

**Procedures:**

- detadj**(U)  
determinant and adjoint matrix of square matrix U
- invunit**(u,n)  
series inverse of polynomial u up to order n
- jacoblift**(f)  
lifts  $f^\kappa$  in jacob(f) with minimal kappa
- mondromyB**(f[,opt])  
monodromy of isolated hypersurface singularity f
- H2basis**(f)  
basis of Brieskorn lattice  $H^2$

See also: Section D.6.14 [gmspoly\_lib], page 864; Section D.6.13 [gmssing\_lib], page 862.

### D.6.18 qhmoduli\_lib

**Library:** qhmoduli.lib

**Purpose:** Moduli Spaces of Semi-Quasihomogeneous Singularities

**Author:** Thomas Bayer, email: bayert@in.tum.de

**Procedures:**

`ArnoldAction(f, [G, w])`  
Induced action of  $G$ - $f$  on  $T_{..}$

`ModEqn(f)`  
Equations of the moduli space for principal part  $f$

`QuotientEquations(G,A,I)`  
Equations of Variety( $I$ )/ $G$  w.r.t. action 'A'

`StabEqn(f)`  
Equations of the stabilizer of  $f$ .

`StabEqnId(I, w)`  
Equations of the stabilizer of the qhom. ideal  $I$ .

`StabOrder(f)`  
Order of the stabilizer of  $f$ .

`UpperMonomials(f, [w])`  
Upper basis of the Milnor algebra of  $f$ .

`Max(data)`  
maximal integer contained in 'data'

`Min(data)`  
minimal integer contained in 'data'

### D.6.19 realclassify\_lib

**Library:** realclassify.lib

**Purpose:** Classification of real singularities

**Author:** Janko Boehm, boehm@mathematik.uni-kl.de  
Magdalen Marais, magdalen@aims.ac.za  
Andreas Steenpass, steenpass@mathematik.uni-kl.de

**Overview:** A library for classifying isolated hypersurface singularities over the reals w.r.t. right equivalence, based on the determinant of singularities by V.I. Arnold. This library is based on classify2.lib by the first and second author and G. Pfister, but handles the real case, while classify2.lib does the complex classification.

**References:**

Arnold, Varchenko, Gusein-Zade: Singularities of Differentiable Maps. Vol. 1: The classification of critical points caustics and wave fronts. Birkh"auser, Boston 1985

J. Boehm, M.S. Marais, A. Steenpass: The Classification of Real Singularities Using Singular. Part III: Unimodal Singularities of Corank 2, <https://arxiv.org/abs/1512.09028>

Greuel, Lossen, Shustin: Introduction to singularities and deformations. Springer, Berlin 2007

M.S. Marais, A. Steenpass: The Classification of Real Singularities Using SINGULAR. Part I: Splitting Lemma and Simple Singularities, *J. Symb. Comput.* 68 (2015), 61-71

M.S. Marais, A. Steenpass: The Classification of Real Singularities Using SINGULAR. Part II: The Structure of the Equivalence Classes of the Unimodal Singularities, *J. Symb. Comput.* 74 (2016), 346-366

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**Procedures:**

`realclassify(f)`  
real classification of singularities of modality 0 and 1 up to stable equivalence

`realmorsesplit(f)`  
splitting lemma in the real case

`milnornumber(f)`  
Milnor number

`determinacy(f)`  
an upper bound for the determinacy

`addnondegeneratevariables(f)`  
find a right equivalent normal form by adding the non-degenerate variables

### D.6.20 `sing_lib`

**Library:** `sing.lib`

**Purpose:** Invariants of Singularities

**Authors:** Gert-Martin Greuel, email: [greuel@mathematik.uni-kl.de](mailto:greuel@mathematik.uni-kl.de)  
Bernd Martin, email: [martin@math.tu-cottbus.de](mailto:martin@math.tu-cottbus.de)

**Procedures:**

`codim(id1, id2)`  
vector space dimension of `id2/id1` if finite

`deform(i)`  
infinitesimal deformations of ideal `i`

`dim_slocus(i)`  
dimension of singular locus of ideal `i`

`is_active(f, id)`  
is polynomial `f` an active element mod `id`? (`id` ideal/module)

`is_ci(i)` is ideal `i` a complete intersection?

`is_is(i)` is ideal `i` an isolated singularity?

`is_reg(f, id)`  
is polynomial `f` a regular element mod `id`? (`id` ideal/module)

`is_regs(i[,id])`  
 are gen's of ideal  $i$  regular sequence modulo  $id$ ?  
`locstd(i)`  
 SB for local degree ordering without cancelling units  
`milnor(i)`  
 milnor number of ideal  $i$ ; (assume  $i$  is ICIS in  $nf$ )  
`nf_icis(i)`  
 generic combinations of generators; get ICIS in  $nf$   
`slocus(i)`  
 ideal of singular locus of ideal  $i$   
`qhspectrum(f,w)`  
 spectrum numbers of  $w$ -homogeneous polynomial  $f$   
`Tjurina(i)`  
 SB of Tjurina module of ideal  $i$  (assume  $i$  is ICIS)  
`tjurina(i)`  
 Tjurina number of ideal  $i$  (assume  $i$  is ICIS)  
`T_1(i)`  $T^1$ -module of ideal  $i$   
`T_2(i)`  $T^2$ -module of ideal  $i$   
`T_12(i)`  $T^1$ - and  $T^2$ -module of ideal  $i$   
`tangentcone(id)`  
 compute tangent cone of  $id$

### D.6.21 `spcurve_lib`

**Library:** `spcurve.lib`

**Purpose:** Deformations and Invariants of CM-codim 2 Singularities

**Author:** Anne Fruehbis-Krueger, [anne@mathematik.uni-kl.de](mailto:anne@mathematik.uni-kl.de)

**Procedures:**

`isCMcod2(i)`  
 presentation matrix of the ideal  $i$ , if  $i$  is CM  
`CMtype(i)`  
 Cohen-Macaulay type of the ideal  $i$   
`matrixT1(M,n)`  
 1st order deformation  $T1$  in matrix description  
`semiCMcod2(M,T1)`  
 semiuniversal deformation of maximal minors of  $M$   
`discr(sem,n)`  
 discriminant of semiuniversal deformation  
`qhmatrix(M)`  
 weights if  $M$  is quasihomogeneous  
`relweight(N,W,a)`  
 relative matrix weight of  $N$  w.r.t. weights  $(W,a)$

`posweight(M,T1,i)`  
 deformation of `coker(M)` of non-negative weight

`KSpencerKernel(M)`  
 kernel of the Kodaira-Spencer map

### D.6.22 `spectrum_lib`

**Library:** `spectrum.lib`

**Purpose:** Singularity Spectrum for Nondegenerate Singularities

**Author:** S. Endrass

**Procedures:**

`spectrumnd(poly f[,1])`  
 spectrum of nondegenerate isolated singularity `f`

### D.6.23 `surfacesignature_lib`

**Library:** `surfacesignature.lib`

**Purpose:** signature of surface singularity

**Authors:** Gerhard Pfister `pfister@mathematik.uni-kl.de`  
 Muhammad Ahsan Banyamin `ahsanbanyamin@gmail.com`  
 Stefan Steidel `steidel@mathematik.uni-kl.de`

**Overview:** A library for computing the signature of irreducible surface singularity. The signature of a surface singularity is defined in [3]. The algorithm we use has been proposed in [9].

Let  $g$  in  $C[x,y]$  define an isolated curve singularity at 0 in  $C^2$  and  $f:=z^N+g(x,y)$ . The zero-set  $V:=V(f)$  in  $C^3$  of  $f$  has an isolated singularity at 0. For a small  $\epsilon>0$  let  $V_\epsilon:=V(f-\epsilon)$  in  $C^3$  be the Milnor fibre of  $(V,0)$  and  $s: H_2(V_\epsilon,R) \times H_2(V_\epsilon,R) \rightarrow R$  be the intersection form (cf. [1],[7]).  $H_2(V_\epsilon,R)$  is an  $m$ -dimensional  $R$ -vector space,  $m$  the Milnor number of  $(V,0)$  (cf. [1],[4],[5],[6]), and  $s$  is a symmetric bilinear form. Let  $\text{sigma}(f)$  be the signature of  $s$ , called the signature of the surface singularity  $(V,0)$ . Formulaes to compute the signature are given by Nemethi (cf. [8],[9]) and van Doorn, Steenbrink (cf. [2]).

We have implemented three approaches using Puiseux expansions, the resolution of singularities resp. the spectral pairs of the singularity.

**References:**

- [1] Arnold, V.I.; Gusein-Zade, S.M.; Varchenko, A.N.: Singularities of Differentiable Mappings. Vol. 1,2, Birkh"auser (1988). [2] van Doorn, M.G.M.; Steenbrink, J.H.M.: A supplement to the monodromy theorem. Abh. Math. Sem. Univ. Hamburg 59, 225-233 (1989). [3] Durfee, A.H.: The Signature of Smoothings of Complex Surface Singularities. Mathematische Annalen 232, 85-98 (1978). [4] de Jong, T.; Pfister, G.: Local Analytic Geometry. Vieweg (2000). [5] Kerner, D.; Nemethi, A.: The Milnor fibre signature is not semi-continuous. arXiv:0907.5252 (2009). [6] Kulikov, V.S.: Mixed Hodge Structures and Singularities. Cambridge Tracts in Mathematics 132, Cambridge University Press (1998). [7] Nemethi, A.: The real Seifert form and the spectral pairs of isolated hypersurface singularities. Compositio Mathematica 98, 23-41 (1995). [8] Nemethi, A.: Dedekind sums and the signature of  $f(x,y)+z^N$ . Selecta Mathematica, New series, Vol. 4, 361-376 (1998).



[9] Nemethi, A.: The Signature of  $f(x,y)+z^N$ . Proceedings of Real and Complex Singularities (C.T.C. Wall's 60th birthday meeting, Liverpool (England), August 1996), London Math. Soc. Lecture Notes Series 263, 131–149 (1999).

**Procedures:**

`signatureBrieskorn(a1,a2,a3)`  
signature of singularity  $x^{a1}+y^{a2}+z^{a3}$

`signaturePuisseux(N,f)`  
signature of singularity  $z^N+f(x,y)=0$ ,  $f$  irred.

`signatureNemethi(N,f)`  
signature of singularity  $z^N+f(x,y)=0$

## D.7 Invariant theory

### D.7.1 `finvar.lib`

**Library:** `finvar.lib`

**Purpose:** Invariant Rings of Finite Groups

**Author:** Agnes E. Heydtmann, contact via Wolfram Decker: [decker@mathematik.uni-kl.de](mailto:decker@mathematik.uni-kl.de) Simon A. King, email: [simon.king@nuigalway.ie](mailto:simon.king@nuigalway.ie)

**Overview:** A library for computing polynomial invariants of finite matrix groups and generators of related varieties. The algorithms are based on B. Sturmfels, G. Kemper, S. King and W. Decker et al..

**Procedures:**

`invariant_ring()`  
generators of the invariant ring (i.r.)

`invariant_ring_random()`  
generators of the i.r., randomized alg.

`primary_invariants()`  
primary invariants (p.i.)

`primary_invariants_random()`  
primary invariants, randomized alg.

`invariant_algebra_reynolds()`  
minimal generating set for the invariant ring of a finite matrix group, in the non-modular case

`invariant_algebra_perm()`  
minimal generating set for the invariant ring or a permutation group, in the non-modular case

`cyclotomic()`  
cyclotomic polynomial

`group_reynolds()`  
finite group and Reynolds operator (R.o.)

`molien()` Molien series (M.s.)

`reynolds_molien()`  
Reynolds operator and Molien series

`partial_molien()`  
partial expansion of Molien series

`evaluate_reynolds()`  
image under the Reynolds operator

`invariant_basis()`  
basis of homogeneous invariants of a degree

`invariant_basis_reynolds()`  
as `invariant_basis()`, with R.o.

`primary_char0()`  
primary invariants (p.i.) in char 0

`primary_charp()`  
primary invariants in char p

`primary_char0_no_molien()`  
p.i., char 0, without Molien series

`primary_charp_no_molien()`  
p.i., char p, without Molien series

`primary_charp_without()`  
p.i., char p, without R.o. or Molien series

`primary_char0_random()`  
primary invariants in char 0, randomized

`primary_charp_random()`  
primary invariants in char p, randomized

`primary_char0_no_molien_random()`  
p.i., char 0, without M.s., randomized

`primary_charp_no_molien_random()`  
p.i., char p, without M.s., randomized

`primary_charp_without_random()`  
p.i., char p, without R.o. or M.s., random.

`power_products()`  
exponents for power products

`secondary_char0()`  
secondary invariants (s.i.) in char 0

`irred_secondary_char0()`  
irreducible s.i. in char 0

`secondary_charp()`  
s.i. in char p, with Molien series and Reynolds operator

`secondary_no_molien()`  
s.i., without Molien series but with Reynolds operator

`irred_secondary_no_molien()`  
irreducible s.i., without Molien series but with Reynolds operator

`secondary_and_irreducibles_no_molien()`  
 s.i. & irreducible s.i., without M.s.

`secondary_not_cohen_macaulay()`  
 s.i. when the invariant ring is not Cohen-Macaulay

`orbit_variety()`  
 ideal of the orbit variety

`rel_orbit_variety()`  
 ideal of a relative orbit variety (new version)

`relative_orbit_variety()`  
 ideal of a relative orbit variety (old version)

`image_of_variety()`  
 ideal of the image of a variety orbit\_sums orbit sums of a set of monomials  
 under the action of a permutation group

### D.7.2 ainvar\_lib

**Library:** ainvar.lib

**Purpose:** Invariant Rings of the Additive Group

**Authors:** Gerhard Pfister (email: pfister@mathematik.uni-kl.de), Gert-Martin Greuel (email: greuel@mathematik.uni-kl.de)

**Procedures:**

`invariantRing(m..)`  
 compute ring of invariants of  $(K,+)$ -action given by  $m$

`derivate(m,f)`  
 derivation of  $f$  with respect to the vector field  $m$

`actionIsProper(m)`  
 tests whether action defined by  $m$  is proper

`reduction(p,I)`  
 SAGBI reduction of  $p$  in the subring generated by  $I$

`completeReduction()`  
 complete SAGBI reduction

`localInvar(m,p..)`  
 invariant polynomial under  $m$  computed from  $p,\dots$

`furtherInvar(m..)`  
 compute further invariants of  $m$  from the given ones

`sortier(id)`  
 sorts generators of  $id$  by increasing leading terms

### D.7.3 rinvar\_lib

**Library:** rinvar.lib

**Purpose:** Invariant Rings of Reductive Groups

**Author:** Thomas Bayer, tbayer@in.tum.de  
<http://wwwmayr.informatik.tu-muenchen.de/personen/bayert/> Current Address: Institut fuer Informatik, TU Muenchen

**Overview:** Implementation based on Derksen's algorithm. Written in the scope of the diploma thesis (advisor: Prof. Gert-Martin Greuel) 'Computations of moduli spaces of semi-quasihomogenous singularities and an implementation in Singular'

**Procedures:**

`HilbertSeries(I, w)`  
 Hilbert series of the ideal I w.r.t. weight w

`HilbertWeights(I, w)`  
 weighted degrees of the generators of I

`ImageVariety(I, F)`  
 ideal of the image variety  $F(\text{variety}(I))$

`ImageGroup(G, F)`  
 ideal of G w.r.t. the induced representation

`InvariantRing(G, Gaction)`  
 generators of the invariant ring of G

`InvariantQ(f, G, Gaction)`  
 decide if f is invariant w.r.t. G

`LinearizeAction(G, Gaction)`  
 linearization of the action 'Gaction' of G

`LinearActionQ(action, s, t)`  
 decide if action is linear in  $\text{var}(s..nvars)$

`LinearCombinationQ(base, f)`  
 decide if f is in the linear hull of 'base'

`MinimalDecomposition(f, s, t)`  
 minimal decomposition of f (like coef)

`NullCone(G, act)`  
 ideal of the nullcone of the action 'act' of G

`ReynoldsImage(RO, f)`  
 image of f under the Reynolds operator 'RO'

`ReynoldsOperator(G, Gaction)`  
 Reynolds operator of the group G

`SimplifyIdeal(I[,m,s])`  
 simplify the ideal I (try to reduce variables)

See also: Section D.6.18 [qhmoduli.lib], page 867; Section D.8.10 [zeroset.lib], page 882.

### D.7.4 invar.lib

**Library:** invar.lib

**Purpose:** Procedures to compute invariant rings of  $SL(n)$  and torus groups

**Author:** Harm Derksen, hderksen@math.unibas.ch

**Procedures:**

- `SL(n)` sets the current group to  $SL_n$
- `torus(n)` sets the current group to an  $n$ -dimensional torus
- `torusrep(list m)`  
representation of a torus given by the weights  $m[1], m[2], \dots$
- `finiterep(<list>)`  
representation of a by a list of matrices
- `sympower(m,d)`  
computes the  $d$ -th symmetric power of a representation  $m$
- `invar(m)` computes the invariant ring of the representation  $m$ .
- `SLreynolds(f)`  
applies the Reynolds operator to  $f$
- `torusreynolds(f)`  
applies the Reynolds operator to  $f$  if the group is a torus or a finite group.

**D.7.5 stratify\_lib****Library:** stratify.lib**Purpose:** Algorithmic Stratification for Unipotent Group-Actions**Author:** Anne Fruehbis-Krueger, anne@mathematik.uni-kl.de**Procedures:**

- `prepMat(M,wr,ws,step)`  
list of submatrices corresp. to given filtration
- `stratify(M,wr,ws,step)`  
algorithmic stratification (main procedure)

**D.8 Symbolic-numerical solving****D.8.1 ffsolve\_lib****Library:** ffsolve.lib**Purpose:** multivariate equation solving over finite fields**Author:** Gergo Gyula Borus, borisz@borisz.net**Procedures:**

- `ffsolve()`  
finite field solving using heuristically chosen method
- `PEsolve()`  
solve system of multivariate equations over finite field
- `simplesolver()`  
solver using modified exhausting search
- `GBsolve()`  
multivariate solver using Groebner-basis

```

XLsolve()
 multivariate polynomial solver using linearization

ZZsolve()
 solve system of multivariate equations over finite field

```

## D.8.2 interval\_lib

**Library:** interval.lib  
**Purpose:** implements interval arithmetic on polynomials  
**Authors:** Dominik Bendle  
 Clara Petroll

**Overloads:**

```

// intervals
[intervalGet indexing
= intervalAssign assigning
== intervalEqual equality
print intervalPrint pretty print
+ intervalAdd addition
- intervalNegate negation (unary)
- intervalSubtract subtraction
* intervalMultiply multiplication
/ intervalDivide division
^ intervalPotentiate potentiation

// boxes
= boxSet assigning
[boxGet indexing
== boxEqual equality
print boxPrint printing
- boxSubtract subtraction
intersect boxIntersect intersection

// intervalmatrices
[ivmatGet indexing
print ivmatPrint printing
nrows ivmatNrows number of rows
ncols ivmatNcols number of columns
det determinant determinant
* ivmatMultiply matrix multiplication

```

**Procedures:**

```

length2()
 length/size if interval

bounds2()
 construct interval for given bounds.

intervalmatrixInit()
 initialises an interval matrix

unitMatrix2()
 unit matrix

```

`applyMatrix()`  
 apply matrix to box  
`ivmatGaussian2()`  
 Gaussian elimination on matrices  
`evalPolyAtBox2()`  
 evaluate interval extension of polynomial  
`exclusionTest()`  
 first version of our exclusion test

### D.8.3 `presolve.lib`

**Library:** `presolve.lib`

**Purpose:** Pre-Solving of Polynomial Equations

**Author:** Gert-Martin Greuel, email: [greuel@mathematik.uni-kl.de](mailto:greuel@mathematik.uni-kl.de),

**Procedures:**

`degreepart(id,d1,d2)`  
 elements of `id` of total degree  $\geq d1$  and  $\leq d2$ , and rest  
`elimlinearpart(id)`  
 linear part eliminated from `id`  
`elimpart(id[,n])`  
 partial elimination of vars [among first `n` vars]  
`elimpartanyr(i,p)`  
 factors of `p` partially eliminated from `i` in any ring  
`fastelim(i,p[.])`  
 fast elimination of factors of `p` from `i` [options]  
`findvars(id)`  
 variables occurring/not occurring in `id`  
`hilbvec(id[,c,o])`  
 intvec of Hilberseries of `id` [in char `c` and ord `o`]  
`linearpart(id)`  
 elements of `id` of total degree  $\leq 1$   
`tolessvars(id[,])`  
 maps `id` to new basering having only vars occurring in `id`  
`solvelinearpart(id)`  
 reduced std-basis of linear part of `id`  
`sortandmap(id[.])`  
 map to new basering with vars sorted w.r.t. complexity  
`sortvars(id[n1,p1..])`  
 sort vars w.r.t. complexity in `id` [different blocks]  
`valvars(id[.])`  
 valuation of vars w.r.t. to their complexity in `id`  
`idealSplit(id,tF,fS)`  
 a list of ideals such that their intersection has the same radical as `id`

### D.8.4 solve\_lib

**Library:** solve.lib

**Purpose:** Complex Solving of Polynomial Systems

**Author:** Moritz Wenk, email: wenk@mathematik.uni-kl.de  
Wilfred Pohl, email: pohl@mathematik.uni-kl.de

**Procedures:**

laguerre\_solve(p, [..])  
find all roots of univariate polynomial p

solve(i, [..])  
all roots of 0-dim. ideal i using triangular sets

ures\_solve(i, [..])  
find all roots of 0-dimensional ideal i with resultants

mp\_res\_mat(i, [..])  
multipolynomial resultant matrix of ideal i

interpolate(p,v,d)  
interpolate polynomial from evaluation points i and results j

fglm\_solve(i, [..])  
find roots of 0-dim. ideal using FGLM and lex\_solve

lex\_solve(i,p, [..])  
find roots of reduced lexicographic standard basis

simplexOut(1)  
prints solution of simplex in nice format

triangLf\_solve(l, [..])  
find roots using triangular sys. (factorizing Lazard)

triangM\_solve(l, [..])  
find roots of given triangular system (Moeller)

triangL\_solve(l, [..])  
find roots using triangular system (Lazard)

triang\_solve(l,p, [..])  
find roots of given triangular system

### D.8.5 triang\_lib

**Library:** triang.lib

**Purpose:** Decompose Zero-dimensional Ideals into Triangular Sets

**Author:** D. Hillebrand

**Procedures:**

triangL(G)  
Decomposition of (G) into triangular systems (Lazard).

triangLfak(G)  
Decomp. of (G) into tri. systems plus factorization.



`triangM(G[, .])`  
 Decomposition of (G) into triangular systems (Moeller).

`triangMH(G[, .])`  
 Decomp. of (G) into tri. syst. with disjoint varieties.

### D.8.6 ntsolve\_lib

**Library:** ntsolve.lib  
**Purpose:** Real Newton Solving of Polynomial Systems  
**Authors:** Wilfred Pohl, email: pohl@mathematik.uni-kl.de  
 Dietmar Hillebrand

**Procedures:**

`nt_solve(G, ini, [ . .])`  
 find one real root of 0-dimensional ideal G

`triMNewton(G, a, [ . .])`  
 find one real root for 0-dim triangular system G

### D.8.7 recover\_lib

**Library:** recover.lib  
**Purpose:** Hybrid numerical/symbolical algorithms for algebraic geometry  
**Author:** Adrian Koch (kocha at rhrk.uni-kl.de)

**Overview:** In this library you'll find implementations of some of the algorithms presented in the paper listed below: Bertini is used to compute a witness set of a given ideal I. Then a lattice basis reduction algorithm is used to recover exact results from the inexact numerical data. More precisely, we obtain elements of prime components of I, the radical of I, or an elimination ideal of I.

NOTE that Bertini may create quite a lot of files in the current directory (or overwrite files which have the same names as the files it wants to create). It also prints information to the screen.

The usefulness of the results of the exactness recovery algorithms heavily depends on the quality of the witness set and the quality of the lattice basis reduction algorithm. The procedures requiring a witness set as part of their input use a simple, unsophisticated version of the LLL algorithm.

**References:**

Daniel Bates, Jonathan Hauenstein, Timothy McCoy, Chris Peterson, and Andrew Sommese; Recovering exact results from inexact numerical data in algebraic geometry; Published in Experimental Mathematics 22(1) on pages 38-50 in 2013

**Procedures:**

`substAll(v, p)`  
 poly: ring variables in v substituted by elements of p

`veronese(d, p)`  
 ideal: image of p under the degree d Veronese embedding

`getRelations(p, . . .)`  
 list of ideals: homogeneous polynomial relations between components of p

`getRelationsRadical(p,...)`  
 modified version of `getRelations`

`gaussRowWithoutPerm(M)`  
 matrix: a row-reduced form of M

`gaussColWithoutPerm(M)`  
 matrix: a column-reduced form of M

`getWitnessSet()`  
 extracts the witness set from the file "main\_data" produced by Bertini

`writeBertiniInput(J)`  
 writes the input-file for bertini with the polynomials in J as functions

`num_prime_decom(I,...)`  
 is supposed to compute a prime decomposition of the radical of I

`num_prime_decom1(P,...)`  
 is supposed to compute a prime decomposition for the ideal represented by the witness point set P

`num_radical_via_decom(I,...)`  
 compute elements of the radical of I by using `num_prime_decom`

`num_radical_via_randlincom(I,...)`  
 computes elements of the radical of I by using a different method

`num_radical1(P,...)`  
 computes elements of the radical via `num_prime_decom1`

`num_radical2(P,...)`  
 computes elements of the radical using a different method

`num_elim(I,f,...)`  
 computes elements of the elimination ideal of I w.r.t. the variables specified by f

`num_elim1(P,...,v)`  
 computes elements of the elimination ideal of the ideal represented by the witness point set P (w.r.t. the variables specified in v)

`realLLL(M)`  
 simple version of the LLL-algorithm; works only over real numbers

### D.8.8 rootisolation.lib

**Library:** rootisolation.lib

**Purpose:** implements an algorithm for real root isolation using interval arithmetic

**Authors:** Dominik Bendle (bendle@rhrk.uni-kl.de)  
 Janko Boehm (boehm@mathematik.uni-lk.de), supervisor Fachpraktikum  
 Clara Petroll (petroll@rhrk.uni-kl.de)

**Overview:** In this library the interval arithmetic from `interval.so` is used. The new type `ivmat`, a matrix consisting of intervals, is implemented as `newstruct`. There are various functions for computations with interval matrices implemented, such as Gaussian elimination for interval matrices.

Interval arithmetic, the interval Newton Step and exclusion methods are used to implement the procedure `rootIsolation`, an algorithm which finds boxes containing elements of the vanishing locus of an ideal. This algorithm is specialised for zero-dimensional radical ideals. The theory about the interval Newton Step is detailed in [2].

Note that interval arithmetic and the aforementioned procedures are intended for rational or real polynomial rings.

#### References:

- [1] Cloud, Kearfott, Moore: Introduction to Interval Analysis, Society for Industrial and Applied Mathematics, 2009
- [2] Eisenbud, Grayson, Herzog, Stillman, Vasconcelos: Computational Methods in Commutative Algebra and Algebraic Geometry, Springer Verlag Berlin-Heidelberg, 3. edition 2004
- [3] Andrew J. Sommese and Charles W. Wampler: The Numerical Solution of Systems of Polynomials - Arising in Engineering and Science, World Scientific Publishing Co. Pte. Ltd., 2005

#### Overloads:

```
[ivmatGet indexing
 print ivmatPrint printing
 nrows ivmatNrows number of rows
 ncols ivmatNcols number of columns
 * ivmatMultiplyGeneral matrix multiplication
```

#### Procedures:

```
bounds(a,#)
 creates a new interval with given bounds

length(I)
 returns Euclidean length of interval

boxSet(B,i,I)
 returns box B with B[j]==I

ivmatInit(m,n)
 returns m x n [0,0]-matrix

ivmatSet(A,i,j,I)
 returns matrix A where A[i][j]=I

unitMatrix(m)
 returns m x m unit matrix where 1 = [1,1]

ivmatGaussian(M)
 computes M-1 using Gaussian elimination for intervals

evalPolyAtBox(f,B)
 returns evaluation of polynomial at a box

evalJacobianAtBox(A,B)
 jacobian matrix of A where polynomials are evaluated at B

rootIsolationNoPreprocessing(I,L,e)
 computes boxes containing unique roots of I lying in L
```

`rootIsolation(I,B,e)`  
 slims down input box B and calls `rootIsolationNoPreprocessing`

`rootIsolationPrimdec(I,B,e)`  
 runs a primary decomposition `primdecGTZ` before root isoation

### D.8.9 signcond\_lib

**Library:** signcond.lib

**Purpose:** Routines for computing realizable sign conditions

**Author:** Enrique A. Tobis, etobis@dc.uba.ar

**Overview:** Routines to determine the number of solutions of a multivariate polynomial system which satisfy a given sign configuration.

**References:**

Basu, Pollack, Roy, "Algorithms in Real Algebraic Geometry", Springer, 2003.

**Procedures:**

`signcnd(P,I)`  
 The sign conditions realized by polynomials of P on a V(I)

`psigncnd(P,l)`  
 Pretty prints the output of `signcnd(l)`

`firstoct(I)`  
 The number of elements of V(I) with every coordinate > 0

### D.8.10 zeroset\_lib

**Library:** zeroset.lib

**Purpose:** Procedures for roots and factorization

**Author:** Thomas Bayer, email: tbayer@mathematik.uni-kl.de,  
<http://wwwmayr.informatik.tu-muenchen.de/personen/bayert/>  
 Current address: Hochschule Ravensburg-Weingarten

**Overview:** Algorithms for finding the zero-set of a zero-dim. ideal in  $\mathbb{Q}(a)[x_1, \dots, x_n]$ , roots and factorization of univariate polynomials over  $\mathbb{Q}(a)[t]$  where a is an algebraic number. Written in the scope of the diploma thesis (advisor: Prof. Gert-Martin Greuel) 'Computations of moduli spaces of semiquasihomogeneous singularities and an implementation in Singular'. This library is meant as a preliminary extension of the functionality of Singular for univariate factorization of polynomials over simple algebraic extensions in characteristic 0.

**Note:** Subprocedures with postfix 'Main' require that the ring contains a variable 'a' and no parameters, and the ideal 'mpoly', where 'minpoly' from the basering is stored.

**Procedures:**

`Quotient(f,g)`  
 quotient q of f w.r.t. g (in  $f = q \cdot g + \text{remainder}$ )

`remainder(f,g)`  
 remainder of the division of f by g

`roots(f)` computes all roots of  $f$  in an extension field of  $\mathbb{Q}$   
`sqfrNorm(f)`  
norm of  $f$  ( $f$  must be squarefree)  
`zeroSet(I)`  
zero-set of the 0-dim. ideal  $I$   
`egcdMain(f, g)`  
gcd over an algebraic extension field of  $\mathbb{Q}$   
`factorMain(f)`  
factorization of  $f$  over an algebraic extension field  
`invertNumberMain(c)`  
inverts an element of an algebraic extension field  
`quotientMain(f, g)`  
quotient of  $f$  w.r.t.  $g$   
`remainderMain(f, g)`  
remainder of the division of  $f$  by  $g$   
`rootsMain(f)`  
computes all roots of  $f$ , might extend the ground field  
`sqfrNormMain(f)`  
norm of  $f$  ( $f$  must be squarefree)  
`containedQ(data, f)`  
 $f$  in data ?  
`sameQ(a, b)`  
 $a == b$  (list  $a, b$ )

## D.9 Visualization

### D.9.1 graphics\_lib

**Library:** graphics.lib

**Purpose:** Procedures to use Graphics with Mathematica

**Author:** Christian Gorzel, gorzelc@math.uni-muenster.de

**Procedures:**

`staircase(fname, I)`  
Mathematica text for displaying staircase of  $I$   
`mathinit()`  
string for loading Mathematica's ImplicitPlot  
`mplot(fname, I[# s])`  
Mathematica text for various plots

## D.9.2 latex.lib

**Library:** latex.lib

**Purpose:** Typesetting of Singular-Objects in LaTeX2e

**Author:** Christian Gorzel, gorzelc@math.uni-muenster.de

**Global variables:**

TeXwidth, TeXnofrac, TeXbrack, TeXproj, TeXaligned, TeXreplace, NoDollars are used to control the typesetting. Call `texdemo()`; to obtain a LaTeX2e file `texlibdemo.tex` explaining the features of `latex.lib` and its global variables.

TeXwidth (int) -1, 0, 1..9, >9: controls breaking of long polynomials

TeXnofrac (int) flag: write  $1/2$  instead of  $\frac{1}{2}$

TeXbrack (string) "{", "(", "<", "|", empty string:

controls brackets around ideals and matrices

TeXproj (int) flag: write ":" instead of "," in vectors

TeXaligned (int) flag: write maps (and ideals) aligned

TeXreplace (list) list entries = 2 strings: replacing symbols

NoDollars (int) flag: suppresses surrounding \$ signs

**Procedures:**

`closetex(fnm)`

writes closing line for LaTeX-document

`opentex(fnm)`

writes header for LaTeX-file fnm

`tex(fnm)` calls LaTeX2e for LaTeX-file fnm

`texdemo([n])`

produces a file explaining the features of this lib

`texfactorize(fnm,f)`

creates string in LaTeX-format for factors of polynomial f

`texmap(fnm,m,r1,r2)`

creates string in LaTeX-format for map  $m:r1 \rightarrow r2$

`texname(fnm,s)`

creates string in LaTeX-format for identifier

`texobj(l)`

creates string in LaTeX-format for any (basic) type

`texpoly(f,n[,l])`

creates string in LaTeX-format for poly

`texproc(fnm,p)`

creates string in LaTeX-format of text from proc p

`texring(fnm,r[,l])`

creates string in LaTeX-format for ring/qring

`rmx(s)` removes .aux and .log files of LaTeX-files

`xdvi(s)` calls xdvi for dvi-files (parameters in square brackets [] are optional) (Procedures with file output assume sufficient write permissions when trying to append existing or create new files.)

### D.9.3 surf\_lib

**Library:** surf.lib

**Purpose:** Procedures for Graphics with Surf

**Author:** Hans Schoenemann, Frank Seelisch

**Note:** Using this library requires the program `surf` to be installed. You can download `surf` either from <http://sourceforge.net/projects/surf> or from <ftp://jim.mathematik.uni-kl.de/pub/Math/Singular/utills/>. The procedure `surfer` requires the program `surfer` (version 1.4.1 or newer) to be installed. You can download `surfer` from <http://imaginary.org/program/surfer>

**Procedures:**

`plot(I)` plots plane curves and surfaces

`surfer(I)`  
plots surfaces interactively

See also: Section D.9.4 [`surfex_lib`], page 885.

### D.9.4 surfex\_lib

**Library:** surfex.lib

**Purpose:** Procedures for visualizing and rotating surfaces.

**Author:** Oliver Labs  
This library uses the program `surf`  
(written by Stefan Endrass and others)  
and `surfex` (written by Oliver Labs and others, mainly Stephan Holzer).

**Note:** It is still an alpha version (see <http://www.AlgebraicSurface.net>)  
This library requires the program `surfex`, `surf` and `java` to be installed. The software is used for producing raytraced images of surfaces. You can download `surfex` from <http://www.surfex.AlgebraicSurface.net>  
`surfex` is a front-end for `surf` which aims to be easier to use than the original tool.

**Procedures:**

`plotRotated(poly, coord)`  
Plot the surface given by the polynomial `p` with the coordinates `coords(list)`

`plotRot(poly)`  
Similar to `plotRotated`, but guesses automatically which coordinates should be used

`plotRotatedList(varieties, coords)`  
Plot the varieties given by the list `varieties` with the coordinates `coords`

`plotRotatedDirect(varieties)`  
Plot the varieties given by the list `varietiesList`

`plotRotatedListFromSpecifyList(varietiesList)`  
Plot the varieties given by the list `varietiesList`

See also: Section D.9.3 [`surf_lib`], page 885.

## D.10 Coding theory

### D.10.1 brnoeth.lib

**Library:** brnoeth.lib

**Purpose:** Brill-Noether Algorithm, Weierstrass-SG and AG-codes

**Authors:** Jose Ignacio Farran Martin, ignfar@eis.uva.es  
Christoph Lossen, lossen@mathematik.uni-kl.de

**Overview:** Implementation of the Brill-Noether algorithm for solving the Riemann-Roch problem and applications to Algebraic Geometry codes. The computation of Weierstrass semi-groups is also implemented. The procedures are intended only for plane (singular) curves defined over a prime field of positive characteristic. For more information about the library see the end of the file brnoeth.lib.

**Procedures:**

`Adj_div(f)`  
computes the conductor of a curve

`NSplaces(h,A)`  
computes non-singular places with given degrees

`BrillNoether(D,C)`  
computes a vector space basis of the linear system  $L(D)$

`Weierstrass(P,m,C)`  
computes the Weierstrass semigroup of  $C$  at  $P$  up to  $m$

`extcurve(d,C)`  
extends the curve  $C$  to an extension of degree  $d$

`AGcode_L(G,D,E)`  
computes the evaluation AG code with divisors  $G$  and  $D$

`AGcode_Omega(G,D,E)`  
computes the residual AG code with divisors  $G$  and  $D$

`prepSV(G,D,F,E)`  
preprocessing for the basic decoding algorithm

`decodeSV(y,K)`  
decoding of a word with the basic decoding algorithm

`closed_points(I)`  
computes the zero-set of a zero-dim. ideal in 2 vars

`dual_code(C)`  
computes the dual code

`sys_code(C)`  
computes an equivalent systematic code

`permute_L(L,P)`  
applies a permutation to a list



### D.10.2 decodegb\_lib

**Library:** decodegb.lib

**Purpose:** Decoding and min distance of linear codes with GB

**Author:** Stanislav Bulygin, bulygin@mathematik.uni-kl.de

**Overview:** In this library we generate several systems used for decoding cyclic codes and finding their minimum distance. Namely, we work with the Cooper's philosophy and generalized Newton identities. The origideal method of quadratic equations is worked out here as well. We also (for comparison) enable to work with the system of Fitzgerald-Lax. We provide some auxiliary functions for further manipulations and decoding. For an overview of the methods mentioned above Section C.8 [Decoding codes with Groebner bases], page 776. For the vanishing ideal computation the algorithm of Farr and Gao is implemented.

**Procedures:**

```

sysCRHT(..)
 generates the CRHT-ideal as in Cooper's philosophy
sysCRHTMindist(..)
 CRHT-ideal to find the minimum distance in the binary case
sysNewton(..)
 generates the ideal with the generalized Newton identities
sysBin(..)
 generates Bin system using Waring function
encode(x,g)
 encodes given message x with the given generator matrix g
syndrome(h,c)
 computes a syndrome w.r.t. the given check matrix
sysQE(..)
 generates the system of quadratic equations for decoding
errorInsert(..)
 inserts errors in a word
errorRand(y,num,e)
 inserts random errors in a word
randomCheck(m,n,e)
 generates a random check matrix
genMDSMat(n,a)
 generates an MDS (actually an RS) matrix
mindist(check)
 computes the minimum distance of a code
decode(rec)
 decoding of a word using the system of quadratic equations
decodeRandom(..)
 a procedure for manipulation with random codes

```

```

decodeCode(..)
 a procedure for manipulation with the given code

vanishId(points)
 computes the vanishing ideal for the given set of points

sysFL(..)
 generates the Fitzgerald-Lax system

decodeRandomFL(..)
 manipulation with random codes via Fitzgerald-Lax

```

## D.11 System and Control theory

### D.11.1 Control theory background

Control systems are usually described by differential (or difference) equations, but their properties of interest are most naturally expressed in terms of the system trajectories (the set of all solutions to the equations). This is formalized by the notion of the system *behavior*. On the other hand, the manipulation of linear system equations can be formalized using algebra, more precisely module theory. The relationship between modules and behaviors is very rich and leads to deep results on system structure.

The key to the module-behavior correspondence is a property of some signal spaces that are modules over the ring of differential (or difference) operators, namely, *the injective cogenerator property*. This property makes it possible to translate any statement on the solution spaces that can be expressed in terms of images and kernels, to an equivalent statement on the modules. Thus analytic properties can be identified with algebraic properties, and conversely, the results of manipulating the modules using computer algebra can be re-translated and interpreted using the language of systems theory. This duality (*algebraic analysis*) is widely used in behavioral systems and control theory today.

For instance, a system is **controllable** (a fundamental property for any control system) if and only if the associated module is torsion-free. This concept can be refined by the so-called controllability degrees. The strongest form of controllability (*flatness*) corresponds to a projective (or even free) module.

Controllability means that one can switch from one system trajectory to another without violating the system law (concatenation of trajectories). For one-dimensional systems (ODE) that evolve in time, this is usually interpreted as switching from a given past trajectory to a desired future trajectory. Thus the system can be forced to behave in an arbitrarily prescribed way.

The extreme case opposed to controllability is **autonomy**: autonomous systems evolve independently according to their law, without being influenceable from the outside. Again, the property can be refined in terms of autonomy degrees.

### D.11.2 control.lib

**Library:** control.lib

**Purpose:** Algebraic analysis tools for System and Control Theory

**Authors:** Oleksandr Iena, yena@mathematik.uni-kl.de  
 Markus Becker, mbecker@mathematik.uni-kl.de  
 Viktor Levandovskyy, levandov@mathematik.uni-kl.de

**Support:** Forschungsschwerpunkt 'Mathematik und Praxis' (Project of Dr. E. Zerz and V. Levandovskyy), University of Kaiserslautern

**Procedures:**

`control(R)`  
analysis of controllability-related properties of  $R$  (using Ext modules)

`controlDim(R)`  
analysis of controllability-related properties of  $R$  (using dimension)

`autonom(R)`  
analysis of autonomy-related properties of  $R$  (using Ext modules)

`autonomDim(R)`  
analysis of autonomy-related properties of  $R$  (using dimension)

`leftKernel(R)`  
a left kernel of  $R$

`rightKernel(R)`  
a right kernel of  $R$

`leftInverse(R)`  
a left inverse of  $R$

`rightInverse(R)`  
a right inverse of  $R$

`colrank(M)`  
a column rank of  $M$  as of matrix

`genericity(M)`  
analysis of the genericity of parameters

`canonize(L)`  
Groebnerification for modules in the output of control or autonomy procs

`iostruct(R)`  
computes an IO-structure of behavior given by a module  $R$

`findTorsion(R, I)`  
generators of the submodule of a module  $R$ , annihilated by the ideal  $I$

`controlExample(s)`  
set up an example from the mini database inside of the library

`view()` well-formatted output of lists, modules and matrices

### D.11.3 jacobson\_lib

**Library:** jacobson.lib

**Purpose:** Algorithms for Smith and Jacobson Normal Form

**Author:** Kristina Schindelar, Kristina.Schindelar@math.rwth-aachen.de,  
Viktor Levandovskyy, levandov@math.rwth-aachen.de

**Overview:** We work over a ring  $R$ , that is an Euclidean principal ideal domain. If  $R$  is commutative, we suppose  $R$  to be a polynomial ring in one variable. If  $R$  is non-commutative, we suppose  $R$  to have two variables, say  $x$  and  $d$ . We treat then the basering as the Ore

localization of  $R$  with respect to the mult. closed set  $S = K[x]$  without 0. Thus, we treat basering as principal ideal ring with  $d$  a polynomial variable and  $x$  an invertible one.

Note, that in computations no division by  $x$  will actually happen.

Given a rectangular matrix  $M$  over  $R$ , one can compute unimodular (that is invertible) square matrices  $U$  and  $V$ , such that  $U*M*V=D$  is a diagonal matrix. Depending on the ring, the diagonal entries of  $D$  have certain properties.

We call a square matrix  $D$  as before 'a weak Jacobson normal form of  $M$ '. It is known, that over the first rational Weyl algebra  $K(x)\langle d \rangle$ ,  $D$  can be further transformed into a diagonal matrix  $(1,1,\dots,1,f,0,\dots,0)$ , where  $f$  is in  $K(x)\langle d \rangle$ . We call such a form of  $D$  the strong Jacobson normal form. The existence of strong form is not guaranteed if one works with algebra, which is not rational Weyl algebra.

#### References:

- [1] N. Jacobson, 'The theory of rings', AMS, 1943.
- [2] Manuel Avelino Insua Hermo, 'Varias perspectivas sobre las bases de Groebner : Forma normal de Smith, Algorithme de Berlekamp y algebras de Leibniz'. PhD thesis, Universidad de Santiago de Compostela, 2005.
- [3] V. Levandovskyy, K. Schindelar 'Computing Jacobson normal form using Groebner bases',  
to appear in Journal of Symbolic Computation, 2010.

#### Procedures:

```
smith(M[,eng1,eng2])
 compute the Smith Normal Form of M over commutative ring

jacobson(M[,eng])
 compute a weak Jacobson Normal Form of M over non-commutative ring

divideUnits(L)
 create ones out of units in the output of smith or jacobson
```

See also: Section D.11.2 [control.lib], page 888.

### D.11.4 findifs\_lib

**Library:** findifs.lib

**Purpose:** Tools for the finite difference schemes

**Authors:** Viktor Levandovskyy, levandov@math.rwth-aachen.de

**Overview:** We provide the presentation of difference operators in a polynomial, semi-factorized and a nodal form. Running `findifs_example()`; will demonstrate, how we generate finite difference schemes of linear PDEs from given approximations.

Theory: The method we use have been developed by V. Levandovskyy and Bernd Martin. The computation of a finite difference scheme of a given single linear partial differential equation with constant coefficients with a given approximation rules boils down to the computation of a Groebner basis of a submodule of a free module with respect to the ordering, eliminating module components.

Support: SpezialForschungsBereich F1301 of the Austrian FWF

#### Procedures:

`findifs_example()`  
contains a guided explanation of our approach

`decoef(P,n)`  
decompose polynomial P into summands with respect to the number n

`difpoly2tex(S,P[,Q])`  
present the difference scheme in the nodal form

`exp2pt(P[,L])`  
convert a polynomial M into the TeX format, in nodal form

`texcoef(n)`  
converts the number n into TeX

`npar(n)` search for 'n' among the parameters and returns its number

`magnitude(P)`  
compute the square of the magnitude of a complex expression

`replace(s,what,with)`  
replace in s all the substrings with a given string

`xchange(w,a,b)`  
exchange two substrings in a given string

See also: Section D.15.4 [`finitediff_lib`], page 924; Section D.9.2 [`latex_lib`], page 884.

## D.12 Teaching

The libraries in this section are intended to be used for teaching purposes but not for serious computations.

### D.12.1 aksaka\_lib

**Library:** aksaka.lib

**Purpose:** Procedures for primality testing after Agrawal, Saxena, Kayal

**Authors:** Christoph Mang

**Overview:** Algorithms for primality testing in polynomial time based on the ideas of Agrawal, Saxena and Kayal.

**Procedures:**

`fastExpt(a,m,n)`  
 $a^m$  for numbers a,m; if  $a^k > n+1$  is returned

`log2(n)` logarithm to basis 2 of n

`PerfectPowerTest(n)`  
checks if there are a,b>1, so that  $a^b=n$

`wurzel(r)`  
square root of number r

`euler(r)` phi-function of Euler

`coeffmod(f,n)`  
polynomial f modulo number n (coefficients mod n)

powerpolyX(q,n,a,r)  
 (polynomial a)<sup>q</sup> modulo (poly r,number n)  
 ask(n) ASK-Algorithm; deterministic Primality test

### D.12.2 atkins\_lib

**Library:** atkins.lib

**Purpose:** Procedures for teaching cryptography

**Author:** Stefan Steidel, steidel@mathematik.uni-kl.de

**Note:** The library contains auxiliary procedures to compute the elliptic curve primality test of Atkin and the Atkin's Test itself. The library is intended to be used for teaching purposes but not for serious computations. Sufficiently high printlevel allows to control each step, thus illustrating the algorithms at work.

**Procedures:**

newTest(L,D)  
 checks if number D already exists in list L

bubblesort(L)  
 sorts elements of the list L

disc(N,k)  
 generates a list of negative discriminants

Cornacchia(d,p)  
 computes solution (x,y) for  $x^2+d*y^2=p$

CornacchiaModified(D,p)  
 computes solution (x,y) for  $x^2+|D|*y^2=4p$

maximum(L)  
 computes the maximal number contained in L

sqr(w,k) computes the square root of w w.r.t. accuracy k

expo(z,k)  
 computes  $\exp(z)$

jOfT(t,k)  
 computes the j-invariant of t

round(r) rounds r to the nearest number out of Z

HilbertClassPoly(D,k)  
 computes the Hilbert Class Polynomial

rootsModp(p,P)  
 computes roots of the polynomial P modulo p

wUnit(D) computes the number of units in  $\mathbb{Q}(\sqrt{D})$

Atkin(N,K,B)  
 tries to prove that N is prime

### D.12.3 crypto\_lib

**Library:** crypto.lib

**Purpose:** Procedures for teaching cryptography

**Authors:** Gerhard Pfister, pfister@mathematik.uni-kl.de  
David Brittinger, dativ@gmx.net

**Overview:** The library contains procedures to compute the discrete logarithm, primality-tests, factorization included elliptic curves. The library is intended to be used for teaching purposes but not for serious computations. Sufficiently high printlevel allows to control each step, thus illustrating the algorithms at work.

**Procedures:**

`decimal(s)`  
number corresponding to the hexadecimal number  $s$

`eexgcdN(L)`  
 $T$  with  $\sum_i L[i]*T[i]=T[n+1]=\gcd(L[1],\dots,L[n])$

`lcmN(a,b)`  
compute  $\text{lcm}(a,b)$

`powerN(m,d,n)`  
compute  $m^d \bmod n$

`chineseRem(T,L)`  
compute  $x$  such that  $x = T[i] \bmod L[i]$

`Jacobi(a,n)`  
the generalized Legendre symbol of  $a$  and  $n$

`primList(n)`  
the list of all primes  $\leq n$

`primL(q)` first primes  $p_1,\dots,p_r$  such that  $q < p_1 * \dots * p_r$

`intPart(x)`  
the integral part of a rational number

`intRoot(m)`  
the integral part of the square root of  $m$

`squareRoot(a,p)`  
the square root of  $a$  in  $\mathbb{Z}/p$ ,  $p$  prime

`solutionsMod2(M)`  
basis solutions of  $Mx=0$  over  $\mathbb{Z}/2$

`powerX(q,i,I)`  
 $q$ -th power of the  $i$ -th variable modulo  $I$

`babyGiant(b,y,p)`  
discrete logarithm  $x: b^x=y \bmod p$

`rho(b,y,p)`  
discrete logarithm  $x: b^x=y \bmod p$

`MillerRabin(n,k)`  
probabilistic primality-test of Miller-Rabin

`SolowayStrassen(n,k)`  
 probabilistic primality-test of Soloway-Strassen

`PocklingtonLehmer(N, [])`  
 primality-test of Pocklington-Lehmer

`PollardRho(n,k,a, [])`  
 Pollard's rho factorization

`pFactor(n,B,P)`  
 Pollard's p-factorization

`quadraticSieve(n,c,B,k)`  
 quadratic sieve factorization

`isOnCurve(N,a,b,P)`  
 P is on the curve  $y^2z=x^3+axz^2+bz^3$  over  $Z/N$

`ellipticAdd(N,a,b,P,Q)`  
 P+Q, addition on elliptic curves

`ellipticMult(N,a,b,P,k)`  
 k\*P on elliptic curves

`ellipticRandomCurve(N)`  
 generates  $y^2z=x^3+axz^2+bz^3$  over  $Z/N$  randomly

`ellipticRandomPoint(N,a,b)`  
 random point on  $y^2z=x^3+axz^2+bz^3$  over  $Z/N$

`countPoints(N,a,b)`  
 number of points of  $y^2=x^3+ax+b$  over  $Z/N$

`ellipticAllPoints(N,a,b)`  
 points of  $y^2=x^3+ax+b$  over  $Z/N$

`ShanksMestre(q,a,b, [])`  
 number of points of  $y^2=x^3+ax+b$  over  $Z/N$

`Schoof(N,a,b)`  
 number of points of  $y^2=x^3+ax+b$  over  $Z/N$

`generateG(a,b,m)`  
 m-th division polynomial of  $y^2=x^3+ax+b$  over  $Z/N$

`factorLenstraECM(N,S,B, [])`  
 Lenstra's factorization

`ECPP(N)` primality-test of Goldwasser-Kilian

`calculate_ordering(num1, primitive, mod1)`  
 Calculates x so that  $\text{primitive}^x \equiv \text{num1} \pmod{\text{mod1}}$

`is_primitive_root(primitive, mod1)`  
 Checks if primitive is a primitive root modulo mod1

`find_first_primitive_root(mod1)`  
 Returns the first primitive root modulo mod1, starting with 1

`binary_add(binary_list)`  
 Adds a 1 to a binary encoded list



```

inverse_modulus(num,mod1)
 Finds a t so that $t \cdot \text{num} = 1 \pmod{\text{mod1}}$
is_prime(n)
 Checks if n is prime proc find_biggest_index(a) Returns the index of the
 biggest element of a
find_index(a,e)
 Returns the list index of element e in list a. Returns 0 if e is not in a
subset_sum01(list knapsack, int solution)
 solves the subset-sum-knapsack-problem by calculating all subsets and
 choosing the right solution
subset_sum02(list knapsack, int sol)
 solves the subset-sum-knapsack-problem with a naive greedy algorithm
unbounded_knapsack(list knapsack, list profit, int capacity)
 solves the unbounded_knapsack-problem, needing a list of knapsack
 weights, a list of profits and a capacity
multidimensional_knapsack(matrix m, list capacities, list profits)
 solves the multidimensional_knapsack-problem by using the PECH algo-
 rithm, needing a weight matrix m, a list of capacities and a list of profits
naccache_stern_generation(int key, int primenum)
 generates a hard knapsack for the Naccache-Stern Kryptosystem for given
 key and prime modulus
naccache_stern_encryption(list knapsack, list message, int primenum)
 encrypts a message with the Naccache-Stern Kryptosystem, using a hard
 knapsack, a message encoded as binary list and a prime modulus
naccache_stern_decryption(list knapsack, int key, int primenum, int
message)
 decrypts a message with the Naccache-Stern Kryptosystem, using the easy
 knapsack, the key, the prime modulus and the message encoded as integer
m_merkle_hellman_transformation(list knapsack, int primitive, int mod1)
 generates a hard knapsack for the multiplicative Merkle-Hellman Kryp-
 tosystem for a given easy knapsack and a primitive root for a modulus
 mod1
m_merkle_hellman_encryption(list knapsack, list message)
 encrypts a message with the multiplicative Merkle-Hellman Kryptosystem,
 using a hard knapsack and a message encoded as binary list
m_merkle_hellman_decryption(list knapsack, bigint primitive, bigint mod1,
int message)
 decrypts a message with the multiplicative Merkle-Hellman Kryp-
 tosystem, using the easy knapsack, the key given by the primitive
 root, the modulus mod1 and the message encoded as integer
merkle_hellman_transformation(list knapsack, int key, int mod1 generates
a hard knapsack for the Merkle-Hellman Kryptosystem for a given easy
knapsack , a multiplier key and a modulus mod1
merkle_hellman_encryption(list knapsack, list message)
 encrypts a message with the Merkle-Hellman Kryptosystem, using a hard
 knapsack and a message encoded as binary list

```

`merkle_hellman_decryption(list knapsack, int key, int mod1, int message)`  
 decrypts a message with the multiplicative Merkle-Hellman Kryptosystem, using the hard knapsack, the key, the modulus mod1 and the message encoded as integer

`super_increasing_knapsack(int ksize)`  
 Creates the smallest super-increasing knapsack of given size ksize

`h_increasing_knapsack(int ksize, int h)`  
 Creates the smallest h-increasing knapsack of given size ksize and h

`injective_knapsack(int ksize, int kmaxelement)`  
 Creates all list of all injective knapsacks of given size ksize and maximal element kmaxelement

`calculate_max_sum(list a)`  
 Calculates the maximal sum of a given knapsack a

`set_is_injective(list a)`  
 Checks if knapsack a is injective

`is_h_injective(list a, int h)`  
 Checks if knapsack a is h-injective

`is_fix_injective(list a)`  
 Checks if knapsack a is fix-injective

`three_elements(list out, int iterations)`  
 Creates the smallest injective knapsack with a given injective\_knapsack by using the three-elements-algorithm with a given number of iterations

#### D.12.4 hyperel\_lib

**Library:** hyperel.lib

**Author:** Markus Hochstetter, markushochstetter@gmx.de

**Note:** The library provides procedures for computing with divisors in the jacobian of hyperelliptic curves. In addition procedures are available for computing the rational representation of divisors and vice versa. The library is intended to be used for teaching and demonstrating purposes but not for efficient computations.

**Procedures:**

`ishyper(h, f)`  
 test, if  $y^2+h(x)y=f(x)$  is hyperelliptic

`isoncurve(P, h, f)`  
 test, if point P is on C:  $y^2+h(x)y=f(x)$

`chinrestp(b, moduli)`  
 compute polynom x, s.t.  $x=b[i] \bmod \text{moduli}[i]$

`norm(a, b, h, f)`  
 norm of  $a(x)-b(x)y$  in  $\text{IF}[C]$

`multi(a, b, c, d, h, f)`  
 $(a(x)-b(x)y)*(c(x)-d(x)y)$  in  $\text{IF}[C]$  ratrep (P,h,f) returns polynomials a,b, s.t.  $\text{div}(a,b)=P$

`divisor(a,b,h,f,[])`  
 computes divisor of  $a(x)-b(x)y$   
`gcddivisor(p,q)`  
 gcd of the divisors p and q  
`semidiv(D,h,f)`  
 semireduced divisor of the pair of polys  $D[1], D[2]$   
`cantoradd(D,Q,h,f)`  
 adding divisors of the hyperell. curve  $y^2+h(x)y=f(x)$   
`cantorred(D,h,f)`  
 returns reduced divisor which is equivalent to D  
`double(D,h,f)`  
 computes  $2*D$  on  $y^2+h(x)y=f(x)$   
`cantormult(m,D,h,f)`  
 computes  $m*D$  on  $y^2+h(x)y=f(x)$

### D.12.5 teachstd\_lib

**Library:** teachstd.lib

**Purpose:** Procedures for teaching standard bases

**Author:** G.-M. Greuel, greuel@mathematik.uni-kl.de

**Note:** The library is intended to be used for teaching purposes, but not for serious computations. Sufficiently high printlevel allows to control each step, thus illustrating the algorithms at work. The procedures are implemented exactly as described in the book 'A SINGULAR Introduction to Commutative Algebra' by G.-M. Greuel and G. Pfister (Springer 2002).

#### Procedures:

`ecart(f)` ecart of f  
`tail(f)` tail of f  
`sameComponent(f,g)`  
 test for same module component of `lead(f)` and `lead(g)`  
`leadmonomial(f)`  
 leading monomial as polynomial (also for vectors)  
`monomialLcm(m,n)`  
 lcm of monomials m and n as polynomial (also for vectors)  
`spoly(f[,1])`  
 s-polynomial of f [symmetric form]  
`minEcart(T,h)`  
 element g from T of minimal ecart s.t.  $LM(g)|LM(h)$   
`NFMora(i)`  
 normal form of i w.r.t Mora algorithm  
`prodcrit(f,g[,o])`  
 test for product criterion

`chaincrit(f,g,h)`  
 test for chain criterion

`pairset(G)`  
 pairs form G neither satisfying `prodcrit` nor `chaincrit`

`updatePairs(P,S,h)`  
 pairset P enlarged by not useless pairs (h,f), f in S

`standard(id)`  
 standard basis of ideal/module

`localstd(id)`  
 local standard basis of id using Lazard's method

### D.12.6 `weierstr_lib`

**Library:** `weierstr.lib`

**Purpose:** Procedures for the Weierstrass Theorems

**Author:** G.-M. Greuel, greuel@mathematik.uni-kl.de

**Procedures:**

`weierstrDiv(g,f,d)`  
 perform Weierstrass division of g by f up to degree d

`weierstrPrep(f,d)`  
 perform Weierstrass preparation of f up to degree d

`lastvarGeneral(f)`  
 make f general of finite order w.r.t. last variable

`generalOrder(f)`  
 compute integer b s.t. f is x<sub>n</sub>-general of order b

### D.12.7 `rootsmr_lib`

**Library:** `rootsmr.lib`

**Purpose:** Counting the number of real roots of polynomial systems

**Author:** Enrique A. Tobis, etobis@dc.uba.ar

**Overview:** Routines for counting the number of real roots of a multivariate polynomial system. Two methods are implemented: deterministic computation of the number of roots, via the signature of a certain bilinear form (`nrRootsDeterm`); and a rational univariate projection, using a pseudorandom polynomial (`nrRootsProbab`). It also includes a command to verify the correctness of the pseudorandom answer.

**References:**

Basu, Pollack, Roy, "Algorithms in Real Algebraic Geometry", Springer, 2003.

**Procedures:**

`nrRootsProbab(I)`  
 Number of real roots of 0-dim ideal (probabilistic)

`nrRootsDeterm(I)`  
 Number of real roots of 0-dim ideal (deterministic)

`symsignature(m)`  
 Signature of the symmetric matrix `m`

`sturmquery(h,B,I)`  
 Sturm query of `h` on  $V(I)$

`matbil(h,B,I)`  
 Matrix of the bilinear form on  $R/I$  associated to `h`

`matmult(f,B,I)`  
 Matrix of multiplication by `f` (`m.f`) on  $R/I$  in the basis `B`

`tracemult(f,B,I)`  
 Trace of `m.f` (`B` is an ordered basis of  $R/I$ )

`coords(f,B,I)`  
 Coordinates of `f` in the ordered basis `B`

`randcharpoly(B,I,n)`  
 Pseudorandom charpoly of univ. projection, `n` optional

`verify(p,B,i)`  
 Verifies the result of `randcharpoly`

`randlinpoly(n)`  
 Pseudorandom linear polynomial, `n` optional

`powersums(f,B,I)`  
 Powersums of the roots of a char polynomial

`symmfunc(S)`  
 Symmetric functions from the powersums `S`

`univarpoly(l)`  
 Polynomial with coefficients from `l`

`qbase(i)` Like `kbase`, but the monomials are ordered

### D.12.8 rootsur\_lib

**Library:** rootsur.lib

**Purpose:** Counting number of real roots of univariate polynomial

**Author:** Enrique A. Tobis, etobis@dc.uba.ar

**Overview:** Routines for bounding and counting the number of real roots of a univariate polynomial, by means of several different methods, namely Descartes' rule of signs, the Budan-Fourier theorem, Sturm sequences and Sturm-Habicht sequences. The first two give bounds on the number of roots. The other two compute the actual number of roots of the polynomial. There are several wrapper functions, to simplify the application of the aforesaid theorems and some functions to determine whether a given polynomial is univariate.

**References:**

Basu, Pollack, Roy, "Algorithms in Real Algebraic Geometry", Springer, 2003.

**Procedures:**

`isuni(p)` Checks whether a polynomial is univariate

|                                  |                                                                         |
|----------------------------------|-------------------------------------------------------------------------|
| <code>whichvariable(p)</code>    | The only variable of a univariate monomial (or 0)                       |
| <code>varsigns(p)</code>         | Number of sign changes in a list                                        |
| <code>boundBuFou(p, a, b)</code> | Bound for number of real roots of polynomial p in interval (a,b)        |
| <code>boundposDes(p)</code>      | Bound for the number of positive real roots of polynomial p             |
| <code>boundDes(p)</code>         | Bound for the number of real roots of polynomial p                      |
| <code>allrealst(p)</code>        | Checks whether all the roots of a polynomial are real (via Sturm)       |
| <code>maxabs(p)</code>           | A bound for the maximum absolute value of a root of a poly              |
| <code>allreal(p)</code>          | Checks whether all the roots of a polynomial are real (via St-Ha)       |
| <code>sturm(p, a, b)</code>      | Number of real roots of a polynomial on an interval (via Sturm)         |
| <code>sturmseq(p)</code>         | Sturm sequence of a polynomial                                          |
| <code>sturmha(p, a, b)</code>    | Number of real roots of a polynomial in (a,b) (via Sturm-Habicht)       |
| <code>sturmhaseq(p)</code>       | A Sturm-Habicht Sequence of a polynomial                                |
| <code>reverse(l)</code>          | Reverses a list                                                         |
| <code>nrroots(p)</code>          | The number of real roots of p                                           |
| <code>isparam(p)</code>          | Returns 0 if and only if the polynomial has non-parametric coefficients |

## D.13 Tropical Geometry

### D.13.1 cimonom\_lib

**Library:** cimonom.lib

**Purpose:** Determines if the toric ideal of an affine monomial curve is a complete intersection

**Authors:** I.Bermejo, ibermejo@ull.es  
 I.Garcia-Marco, igarcia@ull.es  
 J.-J.Salazar-Gonzalez, jjsalaza@ull.es

**Overview:** A library for determining if the toric ideal of an affine monomial curve is a complete intersection with NO NEED of computing explicitly a system of generators of such ideal. It also contains procedures to obtain the minimum positive multiple of an integer which is in a semigroup of positive integers. The procedures are based on a paper by Isabel Bermejo, Ignacio Garcia and Juan Jose Salazar-Gonzalez: 'An algorithm to check whether the toric ideal of an affine monomial curve is a complete intersection', Preprint.

**Procedures:**

`BelongSemig(n, v[, sup])`  
checks whether  $n$  is in the semigroup generated by  $v$ ;

`MinMult(a, b)`  
computes  $k$ , the minimum positive integer such that  $k*a$  is in the semigroup of positive integers generated by the elements in  $b$ .

`CompInt(d)`  
checks whether  $I(d)$  is a complete intersection or not.

See also: Section C.6.4 [Integer programming], page 775.

### D.13.2 gfan\_lib

**Library:** gfan.lib

**Purpose:** Interface to gfan and gfanlib for computations in convex geometry

**Authors:** Anders N. Jensen, email: jensen@imf.au.dk  
Yue Ren, email: ren@mathematik.uni-kl.de  
Frank Seelisch

**Procedures:**

`fullSpace(n)`  
cone, the ambient space of dimension  $n$

`origin(n)`  
cone, the origin in an ambient space of dimension  $n$

`positiveOrthant(n)`  
cone, the positive orthant of dimension  $n$

`ambientDimension(c)`  
the dimension of the ambient space the input lives in

`canonicalizeCone(c)`  
a unique representation of the cone  $c$

`codimension(c)`  
the codimension of the input

`coneViaPoints()`  
define a cone

`coneViaInequalities()`  
define a cone

`coneLink(c, w)`  
the link of  $c$  around  $w$

`containsAsFace(c,d)`  
is  $d$  a face of  $c$

`containsInSupport(c,d)`  
is  $d$  contained in  $c$

`containsPositiveVector(c)`  
contains a vector with only positive entries?

`containsRelatively(c,p)`  
 $p$  in  $c$ ?

`convexHull(c1,c2)`  
convex hull

`convexIntersection(c1,c2)`  
convex hull

`dimension(c)`  
dimension of  $c$

`dualCone(c)`  
the dual of  $c$

`equations(c)`  
defining equations of  $c$

`faceContaining(c,w)`  
the face of  $c$  containing  $w$  in its relative interior

`facets(c)`  
the facets of  $c$

`generatorsOfLinealitySpace(c)`  
generators of the lineality space of  $c$

`generatorsOfSpan(c)`  
generators of the span of  $c$

`getLinearForms(c)`  
linear forms previously stored in  $c$

`getMultiplicity(c)`  
multiplicity previously stored in  $c$

`inequalities(c)`  
inequalities of  $c$

`isFullSpace(c)`  
is the entire ambient space?

`isOrigin(c)`  
is the origin?

`isSimplicial(c)`  
is simplicial?

`linealityDimension(c)`  
the dimension of the lineality space of  $c$

`linealitySpace(c)`  
the lineality space of  $c$



**negatedCone(c)**  
 the negative of  $c$

**polytopeViaInequalities()**  
**polytopeViaPoints()**

**quotientLatticeBasis(c)**  
 basis of  $Z^n$  intersected with the span of  $c$  modulo  $Z^n$  intersected with the lineality space of  $c$

**randomPoint(c)**  
 a random point in the relative interior of  $c$

**rays(c)** generators of the rays of  $c$

**relativeInteriorPoint(c)**  
 point in the relative interior of  $c$

**semigroupGenerator(c)**  
 generator of  $Z^n$  intersected with  $c$  modulo  $Z^n$  intersected with the lineality space of  $c$

**setLinearForms(c)**  
 stores linear forms in  $c$

**setMultiplicity(c)**  
 stores a multiplicity in  $c$

**span(c)** unique irredundant equations of  $c$

**uniquePoint(c)**  
 a unique point in  $c$  stable under reflections at coordinate hyperplanes

**containsInCollection(f,c)**  
 $f$  contains  $c$ ?

**emptyFan(n)**  
 empty fan in ambient dimension  $n$

**fanViaCones(L)**  
 fan generated by the cones in  $L$

**fullFan(n)**  
 full fan in ambient dimension  $n$

**fVector(f)**  
 the  $f$ -Vector of  $f$

**getCone(f,d,i[,m])**  
 the  $i$ -th cone of dimension  $d$  in  $f$

**insertCone(f,c[,b])**  
 inserts the cone  $c$  into  $f$

**isCompatible(f,c)**  
 $f$  and  $c$  live in the same ambient space

**isPure(f)**  
 all maximal cones of  $f$  are of the same dimension

**nmaxcones(f)**  
 number of maximal cones in  $f$

`ncones(f)`  
 number of cones in  $f$   
`numberOfConesOfDimension(f,d[,m])`  
 the number of cones in dimension  $d$   
`removeCone(f,c[,b])`  
 removes the cone  $c$   
`dualPolytope(p)`  
 the dual of  $p$   
`newtonPolytope(f)`  
 convex hull of all exponent vectors of  $f$   
`vertices(p)`  
 vertices of  $p$   
`onesVector(n)`  
 intvec of length  $n$  with all entries 1

### D.13.3 gitfan\_lib

**Library:** gitfan.lib

**Purpose:** Compute GIT-fans.

**Authors:** Janko Boehm, boehm at mathematik.uni-kl.de  
 Simon Keicher, keicher at mail.mathematik.uni-tuebingen.de  
 Yue Ren, ren at mathematik.uni-kl.de

**Overview:** This library allows you to calculate GIT-fans, torus orbits and GKZ-fans.

It provides features to make use of symmetries of the torus action under consideration. The main procedure is `GITfan` which can be directly applied to an ideal and a grading matrix encoding the torus action, and returns a fan, the associated GIT-fan. We also provide various procedures implementing substeps of the algorithm to deal with large computations.

The library uses the package 'gfanlib' by Anders N. Jensen.

For notation, background, and algorithms see [BKR16].

Functions produce debug output if `printlevel` is positive.

Elements of the symmetric group  $S_n$  of type permutation can be created by the function `permutationFromIntvec`.

The images of  $1, \dots, n$  can be obtained by `permutationToIntvec`. Composition of permutations can be done by the `*`-Operator, also powers can be computed in the usual way.

**References:**

[BKR16] J. Boehm, S. Keicher, Y. Ren: Computing GIT-Fans with Symmetry and the Mori Chamber Decomposition of  $M_{0,6}$ , <https://arxiv.org/abs/1603.09241>

**Types:** permutation; Permutation in map representation.

**Procedures:**

`isAface(ideal,intvec)`  
 Checks whether the given face is an a-face.

`afaces(ideal)`  
Returns a list of intvecs that correspond to the set of all a-faces, optionally for given list of simplex faces.

`fullDimImages(list,intmat)`  
Finds the afaces which have a full-dimensional projection.

`minimalAfaces(list)`  
compute the minimal a-faces among the a-faces with full dimensional projection.

`orbitCones(list,intmat)`  
Returns the list of all orbit cones.

`GITcone(list,bigintmat)`  
Returns the GIT-cone containing the given weight vector.

`GITfan(ideal,intmat)`  
Compute GIT-fan.

`GITfanFromOrbitCones(list,intmat,cone)`  
Compute GIT-fan from orbit cones.

`GITfanParallel(list,intmat,cone)`  
Compute GIT-fan in parallel from orbit cones.

`GKZfan(intmat)`  
Returns the GKZ-fan of the matrix  $Q$ .

`movingCone(intmat)`  
Compute the moving cone.

`computeAfaceOrbits(list,list)`  
Compute orbits of a-faces under a permutation group action.

`minimalAfaceOrbits(list)`  
Find the minimal a-face orbits.

`orbitConeOrbits(list,intmat)`  
Project the a-face orbits to orbit cone orbits.

`minimalOrbitConeOrbits(list)`  
Find the minimal orbit cone orbits.

`intersectOrbitsWithMovingCone(list,cone)`  
Intersect orbit cone orbits with moving cone.

`groupActionOnQImage(list,intmat)`  
Determine the induced group action in the target of the grading matrix.

`groupActionOnHashes(list,list)`  
Determine the induced group action on the set of orbit cones.

`storeActionOnOrbitConeIndices(list,string)`  
Write the group action on the set of orbit cones to a file.

`permutationFromIntvec(intvec)`  
Create a permutation from an intvec of images.

`permutationToIntvec(permutation)`  
Return the intvec of images.

`evaluateProduct(list,list)`  
 Evaluate a list of products of group elements in terms of a given representation of the elements as permutations.

`GITfanSymmetric(list,intmat,cone,list)`  
 Compute GIT-fan from orbit cones by determining a minimal representing set for the orbits of maximal dimensional GIT-cones.

`GITfanParallelSymmetric(list,intmat,cone,list)`  
 Compute GIT-fan in parallel from orbit cones by determining a minimal representing set for the orbits of maximal dimensional GIT-cones.

`bigintToBinary(bigint,int)`  
 Convert bigint into a sparse binary representation specifying the indices of the one-entries

`binaryToBigint(intvec)`  
 Convert sparse binary representation specifying the indices of the one-entries to bigint

`applyPermutationToIntvec(intvec,permutation)`  
 Apply permutation to a set of integers represented as an intvec

`hashToCone(bigint,list)`  
 Convert a bigint hash to a GIT-cone

`hashesToFan(list hashes,list OC)`

`gitCone(ideal,bigintmat,bigintmat)`  
 Returns the GIT-cone around the given weight vector w

### D.13.4 polymake\_lib

**Library:** polymake.lib

**Purpose:** Computations with polytopes and fans, interface to polymake and TOPCOM

**Author:** Thomas Markwig, email: keilen@mathematik.uni-kl.de  
 Yue Ren, email: ren@mathematik.uni-kl.de

**Warning:** Most procedures will not work unless polymake or topcom is installed and if so, they will only work with the operating system LINUX! For more detailed information see IMPORTANT NOTE respectively consult the help string of the procedures.

The conventions used in this library for polytopes and fans, e.g. the length and labeling of their vertices resp. rays, differs from the conventions used in polymake and thus from the conventions used in the polymake extension polymake.so of Singular. We recommend to use the newer polymake.so whenever possible.

**Important note:**

Even though this is a Singular library for computing polytopes and fans such as the Newton polytope or the Groebner fan of a polynomial, most of the hard computations are NOT done by Singular but by the program

- polymake by Ewgenij Gawrilow, TU Berlin and Michael Joswig, TU Darmstadt (see <http://www.polymake.org/>),
- respectively (only in the procedure triangulations) by the program
- topcom by Joerg Rambau, Universitaet Bayreuth (see <http://www.rambau.wm.uni-bayreuth.de/TOPCOM/>);

this library should rather be seen as an interface which allows to use a (very limited) number of options which polymake respectively topcom offers to compute with polytopes and fans and to make the results available in Singular for further computations; moreover, the user familiar with Singular does not have to learn the syntax of polymake or topcom, if the options offered here are sufficient for his purposes.

Note, though, that the procedures concerned with planar polygons are independent of both, polymake and topcom.

**Procedures using polymake:**

`polymakePolytope()`  
 computes the vertices of a polytope using polymake

`newtonPolytopeP()`  
 computes the Newton polytope of a polynomial

`newtonPolytopeLP()`  
 computes the lattice points of the Newton polytope

**Procedures using topcom:**

`triangulations()`  
 computes all triangulations of a marked polytope

`secondaryPolytope()`  
 computes the secondary polytope of a marked polytope

**Procedures concerned with planar polygons:**

`cycleLength()`  
 computes the cycleLength of cycle

`splitPolygon()`  
 splits a marked polygon into vertices, facets, interior points

`eta()`      computes the eta-vector of a triangulation

`findOrientedBoundary()`  
 computes the boundary of a convex hull

`cyclePoints()`  
 computes lattice points connected to some lattice point

`latticeArea()`  
 computes the lattice area of a polygon

`picksFormula()`  
 computes the ingredients of Pick's formula for a polygon

`ellipticNF()`  
 computes the normal form of an elliptic polygon

`ellipticNFDB()`  
 displays the 16 normal forms of elliptic polygons

**Procedures using libpolymake:**

`boundaryLatticePoints()`

`ehrhartPolynomialCoeff()`

`fVectorP()`

`facetVertexLatticeDistances()`

```
facetWidth()
facetWidths()
gorensteinIndex()
gorensteinVector()
hStarVector()
hVector()
hilbertBasis()
interiorLatticePoints()
isBounded()
isCanonical()
isCompressed()
isGorenstein()
isLatticeEmpty()
isNormal()
isReflexive()
isSmooth()
isTerminal()
isVeryAmple()
latticeCodegree()
latticeDegree()
latticePoints()
latticeVolume()
maximalFace()
maximalValue()
minimalFace()
minimalValue()
minkowskiSum()
nBoundaryLatticePoints()
nHilbertBasis()
nInteriorLatticePoints()
nLatticePoints()
normalFan()
vertexAdjacencyGraph()
vertexEdgeGraph()
visual()
```

### D.13.5 realizationMatroids\_lib

**Library:** realizationMatroids.lib

**Purpose:** Deciding Relative Realizability for Tropical Fan Curves in 2-Dimensional Matroidal Fans

**Authors:** Anna Lena Winstel, winstel@mathematik.uni-kl.de

**Overview:** In tropical geometry, one question to ask is the following: given a one-dimensional balanced polyhedral fan  $C$  which is set theoretically contained in the tropicalization  $\text{trop}(Y)$  of an algebraic variety  $Y$ , does there exist a curve  $X$  in  $Y$  such that  $\text{trop}(X) = C$ ? This equality of  $C$  and  $\text{trop}(X)$  denotes an equality of both, the fans  $\text{trop}(X)$  and  $C$  and their weights on the maximal cones. The relative realization space of  $C$  with respect to  $Y$  is the space of all algebraic curves in  $Y$  which tropicalize to  $C$ .

This library provides procedures deciding relative realizability for tropical fan curves, i.e. one-dimensional weighted balanced polyhedral fans, contained in two-dimensional matroidal fans  $\text{trop}(Y)$  where  $Y$  is a projective plane.

**Notation:** If  $Y$  is a projective plane in  $(n-1)$ -dimensional projective space, we consider  $\text{trop}(Y)$  in  $\mathbb{R}^n/\langle 1 \rangle$ . Moreover, for the relative realization space of  $C$  with respect to  $Y$  we only consider algebraic curves of degree  $\deg(C)$  in  $Y$  which tropicalize to  $C$ .

**Procedures:**

`realizationDim(I,C)`

For a given tropical fan curve  $C$  in  $\text{trop}(Y)$ , where  $Y = V(I)$  is a projective plane, this routine returns the dimension of the relative realization space of  $C$  with respect to  $Y$ , that is the space of all algebraic curves of degree  $\deg(C)$  in  $Y$  which tropicalize to  $C$ . If the realization space is empty, the output is set to -1.

`irrRealizationDim(I,C)`

This routine returns the dimension of the irreducible relative realization space of the tropical fan curve  $C$  with respect to  $Y = V(I)$ , that is the space of all irreducible algebraic curves of degree  $\deg(C)$  in  $Y$  which tropicalize to  $C$ . If the irreducible relative realization space is empty, the output is set to -1.

`realizationDimPoly(I,C)`

If  $C$  is a tropical fan curve contained in the tropicalization  $\text{trop}(Y)$  of the projective plane  $Y = V(I)$  such that the relative realization space  $M$  of  $C$  is non-empty, this routine returns the tuple  $(\dim(M),f)$  where  $f$  is an example of a homogeneous polynomial of degree  $\deg(C)$  cutting out a curve  $X$  in  $Y$  which tropicalizes to  $C$ . If  $M$  is empty, the output is set to -1.

### D.13.6 tropical\_lib

**Library:** tropical.lib

**Purpose:** Computations in Tropical Geometry

**Authors:** Anders Jensen Needergard, email: jensen@math.tu-berlin.de  
 Hannah Markwig, email: hannah@uni-math.gwdg.de  
 Thomas Markwig, email: keilen@mathematik.uni-kl.de  
 Yue Ren, email: ren@mathematik.uni-kl.de

**Warning:** - `tropicalLifting` will only work with LINUX and if in addition `gfan` is installed.  
 - `drawTropicalCurve` and `drawTropicalNewtonSubdivision` will only display the tropical curve with LINUX and if in addition `latex` and `xdg-open` are installed.  
 - For `tropicalLifting` in the definition of the basering the parameter  $t$  from the Puiseux series field  $C\{\{t\}\}$  must be defined as a variable, while for all other procedures it must be defined as a parameter.

**Theory:** Fix some base field  $K$  and a bunch of lattice points  $v_0, \dots, v_m$  in the integer lattice  $Z^n$ , then this defines a toric variety as the closure of  $(K^*)^n$  in the projective space  $P^m$ , where the torus is embedded via the map sending a point  $x$  in  $(K^*)^n$  to the point  $(x^{v_0}, \dots, x^{v_m})$ .

The generic hyperplane sections are just the images of the hypersurfaces in  $(K^*)^n$  defined by the polynomials  $f = a_0 x^{v_0} + \dots + a_m x^{v_m} = 0$ . Some properties of these hypersurfaces can be studied via tropicalisation.

For this we suppose that  $K = C\{\{t\}\}$  is the field of Puiseux series over the field of complex numbers (or any other field with a valuation into the real numbers). One associates to the hypersurface given by  $f = a_0 x^{v_0} + \dots + a_m x^{v_m}$  the tropical hypersurface defined by the tropicalisation  $\text{trop}(f) = \min\{\text{val}(a_0) + \langle v_0, x \rangle, \dots, \text{val}(a_m) + \langle v_m, x \rangle\}$ .

Here,  $\langle v, x \rangle$  denotes the standard scalar product of the integer vector  $v$  in  $Z^n$  with the vector  $x = (x_1, \dots, x_n)$  of variables, so that  $\text{trop}(f)$  is a piecewise linear function on  $R^n$ . The corner locus of this function (i.e. the points at which the minimum is attained a least twice) is the tropical hypersurface defined by  $\text{trop}(f)$ .

The theorem of Newton-Kapranov states that this tropical hypersurface is the same as if one computes pointwise the valuation of the hypersurface given by  $f$ . The analogue holds true if one replaces one equation  $f$  by an ideal  $I$ . A constructive proof of the theorem is given by an adapted version of the Newton-Puiseux algorithm. The hard part is to find a point in the variety over  $C\{\{t\}\}$  which corresponds to a given point in the tropical variety.

It is the purpose of this library to provide basic means to deal with tropical varieties. Of course we cannot represent the field of Puiseux series over  $C$  in its full strength, however, in order to compute interesting examples it will be sufficient to replace the complex numbers  $C$  by the rational numbers  $Q$  and to replace Puiseux series in  $t$  by rational functions in  $t$ , i.e. we replace  $C\{\{t\}\}$  by  $Q(t)$ , or sometimes even by  $Q[t]$ . Note, that this in particular forbids rational exponents for the  $t$ 's.

Moreover, in Singular no negative exponents of monomials are allowed, so that the integer vectors  $v_i$  will have to have non-negative entries. Shifting all exponents by a fixed integer vector does not change the tropicalisation nor does it change the toric variety. Thus this does not cause any restriction.

If, however, for some reason you prefer to work with general  $v_i$ , then you have to pass right away to the tropicalisation of the equations, wherever this is allowed – these are linear polynomials where the constant coefficient corresponds to the valuation of the original coefficient and where the non-constant coefficient correspond to the exponents of the monomials, thus they may be rational numbers respectively negative numbers: e.g. if  $f = t^{1/2} x^{-2} y^3 + 2t x y + 4$  then  $\text{trop}(f) = \min\{1/2 - 2x + 3y, 1 + x + y, 0\}$ .

The main tools provided in this library are as follows:

- `tropicalLifting` implements the constructive proof of the Theorem of Newton-Kapranov and constructs a point in the variety over  $C\{\{t\}\}$  corresponding to a given point in the

corresponding tropical variety associated to an ideal  $I$ ; the generators of  $I$  have to be in the polynomial ring  $Q[t, x_1, \dots, x_n]$  considered as a subring of  $C\{\{t\}\}[x_1, \dots, x_n]$ ; a solution will be constructed up to given order; note that several field extensions of  $Q$  might be necessary throughout the intermediate computations; the procedures use the external program `gfan`



- `puiseuxExpansion` computes a Newton-Puiseux expansion of a plane curve singularity
- `drawTropicalCurve` visualises a tropical plane curve either given by a polynomial in  $\mathbb{Q}(t)[x,y]$  or by a list of linear polynomials of the form  $ax+by+c$  with  $a,b$  in  $\mathbb{Z}$  and  $c$  in  $\mathbb{Q}$ ; latex must be installed on your computer
- `tropicalJInvariant` computes the tropical j-invariant of a tropical elliptic curve
- `jInvariant` computes the j-invariant of an elliptic curve
- `weierstrassForm` computes the Weierstrass form of an elliptic curve

**Procedures for tropical lifting:**

- `tropicalLifting()`  
computes a point in the tropical variety
- `displayTropicalLifting()`  
displays the output of `tropicalLifting`
- `puiseuxExpansion()`  
computes a Newton-Puiseux expansion in the plane
- `displayPuisseuxExpansion()`  
displays the output of `puiseuxExpansion`

**Procedures for drawing tropical curves:**

- `tropicalCurve()`  
computes a tropical curve and its Newton subdivision
  - `drawTropicalCurve()`  
produces a post script image of a tropical curve
  - `drawNewtonSubdivision()`  
produces a post script image of a Newton subdivision
- PROCEDURES FOR J-INVARIANTS:
- `tropicalJInvariant()`  
computes the tropical j-invariant of a tropical curve
  - `weierstrassForm()`  
computes the Weierstrass form of a cubic polynomial
  - `jInvariant()`  
computes the j-invariant of a cubic polynomial

**General procedures:**

- `conicWithTangents()`  
computes a conic through five points with tangents
- `tropicalise()`  
computes the tropicalisation of a polynomial
- `tropicaliseSet()`  
computes the tropicalisation several polynomials
- `tInitialForm()`  
computes the tInitial form of a polynomial in  $\mathbb{Q}[t,x_1,\dots,x_n]$
- `tInitialIdeal()`  
computes the tInitial ideal of an ideal in  $\mathbb{Q}[t,x_1,\dots,x_n]$

`initialForm()`  
 computes the initial form of poly in  $\mathbb{Q}[x_1, \dots, x_n]$

`initialIdeal()`  
 computes the initial ideal of an ideal in  $\mathbb{Q}[x_1, \dots, x_n]$

**Procedures for latex conversion:**

`texNumber()`  
 outputs the texcommand for the leading coefficient of poly

`texPolynomial()`  
 outputs the texcommand for the polynomial poly

`texMatrix()`  
 outputs the texcommand for the matrix

`texDrawBasic()`  
 embeds output of `texDrawTropical` in a `texdraw` environment

`texDrawTropical()`  
 computes the `texdraw` commands for a tropical curve

`texDrawNewtonSubdivision()`  
 computes `texdraw` commands for a Newton subdivision

`texDrawTriangulation()`  
 computes `texdraw` commands for a triangulation

**Auxiliary procedures:**

`radicalMemberShip()`  
 checks radical membership

`tInitialFormPar()`  
 computes the t-initial form of poly in  $\mathbb{Q}(t)[x_1, \dots, x_n]$

`tInitialFormParMax()`  
 same as `tInitialFormPar`, but uses maximum

`solveTInitialFormPar()`  
 displays approximated solution of a 0-dim ideal

`detropicalise()`  
 computes the detropicalisation of a linear form

`tDetropicalise()`  
 computes the detropicalisation of a linear form

`dualConic()`  
 computes the dual of an affine plane conic

`parameterSubstitute()`  
 substitutes in the polynomial the parameter `t` by `t^N`

`tropicalSubst()`  
 makes certain substitutions in a tropical polynomial

`randomPolyInT()`  
 computes a polynomial with random coefficients

`cleanTmp()`  
 clears `/tmp` from files created by other procedures

**Procedures from binary library:**

`groebnerCone()`  
 constructs the Groebner cone with respect to a weight vector

`maximalGroebnerCone()`  
 constructs the Groebner cone with respect to the current ordering

`homogeneitySpace()`  
 constructs the homogeneity space

`initial()`  
 constructs the initial form resp. ideal

`tropicalVariety()`  
 computes the tropical variety of a poly or ideal

`groebnerFan()`  
 computes the Groebner fan of a poly or ideal

`groebnerComplex()`  
 computes the Groebner complex of a poly or ideal

**D.13.7 tropicalNewton\_lib**

**Library:** tropicalNewton.lib

**Purpose:** Computations in Tropical Geometry using Newton Polygon methods

**Authors:** Tommy Hofman, email: thofmann@mathematik.uni.kl.de Yue Ren, email: reny@cs.bgu.ac.il

**Overview:** This libraries contains algorithms for computing

- non-trivial points on tropical varieties,
- zero-dimensional tropical varieties,
- one-codimensional links of tropical varieties

based on Newton polygon methods.

**References:**

Hofmann, Ren: Computing tropical points and tropical links, arXiv:1611.02878 (WARNING: this library follows the max convention instead and triangular sets follow the definition of the Singular book)

**Procedures:**

`setUniformizingParameter()`  
 sets the uniformizingParameter

`val()` returns valuation of element in ground field

`newtonPolygonNegSlopes()`  
 returns negative of the Newton Polyong slopes

`cccMatrixToPositiveIntvec()`  
 helper function to turn a computed valuation vector into a usable weight vector in Singular

`tropicalPointNewton()`  
 computes point on tropical variety

```

switchRingsAndComputeInitialIdeal()
 switches rings and computes initial ideal

tropicalVarietyNewton()
 computes tropical variety of zero-dimensional ideal

tropicalLinkNewton()
 computes tropical variety that is polyhedral fan and has codimension one
 lineality space

```

See also: [tropicalVariety], page 913; Section D.13.6 [tropical.lib], page 909.

## D.14 Miscellaneous libraries

### D.14.1 arr.lib

**Library:** arr.lib

**Purpose:** a library of algorithms for arrangements of hyperplanes

**Authors:** Randolph Scholz (rscholz@rhrk.uni-kl.de),  
Patrick Serwene (serwene@mathematik.uni-kl.de),  
Lukas Kuehne (lf.kuehne@gmail.com)

**Overloads:**

```

// OPERATORS
= arrAdd assignment
+ arrAdd union of two arrs
[arrGet access to a single/multiple hyperplane(s) - arrMinus deletes given hyperplanes
from the arr <= arrLEQ comparison
>= arrGEQ comparison
== arrEQ comparison
!= arrNEQ comparison
< arrLNEQ comparison
> arrGNEQ comparison
// TYPECASTING
matrix arr2mat coeff matrix
poly arr2poly defining polynomial
// OTHER
variables arrVariables ideal generated by the variables the arr depends on nvars arrN-
vars number of variables the arr depends on delete arrDelete deletes hyperplanes by
indices print arrPrint prints the arr on the screen
// IDEAL INHERITED FUNCTIONS
homog arrHomog checks if arrangement is homogeneous simplify arrSimplify simplifies
arrangement size arrSize number of planes
subst arrSubst substitute variables
// MULTI-ARRANGEMENTS
= multarrAdd assignment of multarr
+ multarrAdd union of multarr
poly multarr2poly defining polynomial
size multarrSize number of hyperplanes with mult. print multarrPrint displays multiarr
delete multarrDelete deletes hyperplane

```

**Procedures:**

`arrSet(arr A, int k, poly p)`  
 replaces the k-th Hyperplane with poly p

`type2arr(#)`  
 converts general input to 'arr' using `arrAdd`.

`mat2arr(matrix M)`  
 affine arrangement from coeff matrix

`mat2carr(matrix M)`  
 central arrangement from coeff matrix

`arrPrintMatrix(arr A)`  
 readable output as a coeff matrix

`varMat(intvec v)`  
 matrix of the corresponding ring\_variables

`varNum(def u)`  
 number of given variable (enh. version of `varNum` in `dmod.lib`)

`arrSwapVar(arr A, i, j)`  
 swaps two variables in the arrangement

`arrLastVar(arr A)`  
 ring\_variable of largest index used in arrangement

`arrCenter(arr A)`  
 computes center of an arrangement

`arrCentral(arr A)`  
 checks if arrangement is central

`arrCentered(arr A)`  
 checks if arrangement is centered

`arrCentralize(arr A)`  
 makes centered arrangement central

`arrCoordChange(A, T, #)`  
 performs coordinate change

`arrCoordNormalize(A, v)`  
 performs projection onto coordinate hyperplane

`arrCone(arr A, var)`  
 coned arrangement

`arrDecone(arr A, int k)`  
 deconed arrangement

`arrLocalize(arr A, intvec v)`  
 localization of an arrangement onto a flat

`arrRestrict(arr A, intvec v)`  
 restricted arrangement onto a flat

`arrIsEssential(arr A)`  
 checks if arrangement is essential

`arrEssentialize(arr A)`  
essentialized arrangement

`arrBoolean(int v)`  
boolean arrangement

`arrBraid(int v)`  
braid arrangement

`arrTypeB(int v)`  
type B arrangement

`arrTypeD(int v)`  
type D arrangement

`arrRandom(d,m,n)`  
random (affine) arrangement

`arrRandomCentral(d,m,n)`  
random central arrangement

`arrEdelmanReiner()`  
Edelman-Reiner arrangement

`arrOrlikSolomon(arr A)`  
Orlik-Solomon algebra of the arrangement

`arrDer(A)`  
module of derivation

`arrIsFree(A)`  
checks if arrangement is free

`arrExponents(A)`  
exponents of a (free) arrangement

`arr2multarr(arr A, intvec v)`  
converts normal arrangement to multiarrangement

`multarr2arr(multarr A)`  
converts multiarrangement to normal arrangement

`multarrRestrict(arr A, v)`  
restriction of A (as arr) to a flat with multiplicities

`multarrMultRestrict(A, int k)`  
restriction of A (as multarr) to a hyperplane with multiplicities

`arrFlats(arr A)`  
intersection lattice

`arrLattice(arr A)`  
computes the intersection lattice / poset

`moebius(arrposet P)`  
computes moebius values

`arrCharPoly(arr A)`  
characteristic polynomial

`arrPoincare(arr A)`  
poincare polynomial of the arrangement

`arrChambers(arr A)`  
 number of chambers of the arrangement

`arrBoundedChambers(arr A)`  
 number of bounded chambers of the arrangement

`printMoebius(arr A)`  
 displays the moebius values of all the flats in the poset

### D.14.2 customstd\_lib

**Library:** customstd.lib

**Purpose:** Load customstd.so

**Authors:** Hans Schoenemann, hannes at mathematik.uni-kl.de  
 Yue Ren, ren at mathematik.uni-kl.de

**Overview:** This library offers customly modified standard bases algorithms in order to increase the performance of other algorithms. If you require a customly modified standard bases algorithm, please contact the authors.

**Procedures:**

`monomialabortstd(ideal)`  
`satstd(ideal, [...])`

### D.14.3 phindex\_lib

**Library :** phindex.lib

**Purpose:** Procedures to compute the index of real analytic vector fields

**Author:** Victor Castellanos

**Note:** To compute the Poincare-Hopf index of a real analytic vector field with an algebraically isolated singularity at 0 (w. an a. i. s), we use the algebraic formula for the degree of the real analytic map germ found by Eisenbud-Levine in 1997. This result was also proved by Khimshiashvili. If the isolated singularity is non algebraically isolated and the vector field has similar reduced complex zeroes of codimension 1, we use a formula as the Eisenbud-Levine found by Victor Castellanos, in both cases is necessary to use a local order (ds,...). To compute the signature of a quadratic form (or symmetric matrix) we use the method of Lagrange.

**Procedures:**

`signatureL(M[,n])`  
 signature of symmetric matrix M, method of Lagrange.

`signatureLqf(h[,n])`  
 signature of quadratic form h, method of Lagrange.

`PH_ais(I)`  
 P-H index of real analytic vector field I w. an a. i. s.

`PH_nais(I)`  
 P-H index of real analytic vector field I w. a non a. i. s.

### D.14.4 polybori\_lib

**Library:** polybori.lib

**Purpose:** A Singular Library Interface for PolyBoRi

**Authors:** Maximilian Kammermeier: Max0791@gmx.de  
Susanne Scherer: sscherer90@yahoo.de

**Overview:** A library for using PolyBoRi in the SINGULAR interface, with procedures that convert structures (polynomials, rings, ideals) in both directions. Therefore, it is possible to compute boolean groebner basis via [boolean\_std], page 919. Polynomials can be converted to zero-suppressed decision diagrams (zdd) and vice versa.

For usability it defines the PolyBoRi types `bideal`, `bpoly`, and `bring` which are equivalent to Singular's `ideal`, `poly`, and `ring`, as well as `bset` which corresponds to the type `zdd` introduced here. In addition `bvar(i)` constructs the Boolean variable corresponding to `var(i)` from current `ring`;

For convenience, the corresponding types can be converted explicitly or implicitly while assigning. Also several SINGULAR operators were overloaded: `bring` comes with `nvars`, `bpoly` implements `lead`, `leadmonom` and `leadcoef`. Objects of this type may be added and multiplied, too. Finally, `bideal` yields `std` and `size` as well as addition and element access.

Hence, by using these types PolyBoRi functionality can be carried out seamlessly in SINGULAR:

```
> LIB "polybori.lib";
> ring r0=2,x(1..4),lp;
> def x=bvar; // enforce Boolean variables

> bpoly f1=x(1)+x(4);
> bpoly f2=x(1)+x(3)*x(1);
> bideal bI=list(f1,f2);

> std(bI);
_[1] = x(1) + x(4)
_[2] = x(3)*x(4) + x(4)
```

**Note:** For using this library SINGULAR's `python` interface must be available on your system. Please `./configure --with-python` when building SINGULAR for this purpose.

There are prebuilt binary packages for PolyBoRi available from <http://polybori.sf.net/>.

For building your own PolyBoRi please ensure that you have `scons` and a development version of the boost libraries installed on you system. Then you may execute the following commands in a `bash`-style shell to build PolyBoRi available to `python`:

```
PBDIR=/path/to/custom/polybori
wget http://downloads.sf.net/project/polybori/polybori/\
0.8.2/polybori-0.8.2.tar.gz
tar -xvzf polybori-0.8.2.tar.gz
cd polybori-0.8.2
scons install PREFIX=$PBDIR PYINSTALLPREFIX=$PBDIR/python
```



```
export PYTHONPATH=$PBDIR/python:$PYTHONPATH
```

**References:**

See <http://polybori.sf.net> for details about PolyBoRi.

**Procedures:**

```
boolean_std(bideal)
 Singular ideal of boolean groebner basis of I

boolean_poly_ring(ring)
 convert ring

boolean_constant(int[, bring])
 convert constant

boolean_poly(poly[, int, bring])
 convert polynomial

direct_boolean_poly(poly[, bring])
 convert polynomial direct

recursive_boolean_poly(poly[, bring])
 convert polynomial recursively

boolean_ideal(ideal[, bring])
 convert ideal

boolean_set(zdd[, bring])
 convert zdd

from_boolean_constant(bpoly)
 convert boolean constant

from_boolean_poly(bpoly[, int])
 convert boolean polynomial

direct_from_boolean_poly(bpoly)
 convert boolean polynomial direct

recursive_from_boolean_poly(bpoly)
 convert boolean polynomial recursively

from_boolean_ideal(bpoly)
 convert to ideal

from_boolean_set(bset)
 convert to zdd

bvar(i) return i-th Boolean variable

poly2zdd(poly)
 return zdd of a given polynomial

zdd2poly(zdd)
 return polynomial representation of a given zdd

disp_zdd(zdd)
 return string with a visualization of a given
```

See also: Section 3.8 [Libraries], page 54; Section 4.23 [User defined types], page 133; Section 4.27 [pyobject], page 138.

## D.15 Experimental libraries

This sections collect libraries in the beta test phase. Everything in these libraries may change.

For the minimal requirements and guidelines see Section 3.8 [Libraries], page 54.

Comments should be send to the author of the library directly.

### D.15.1 autgradalg\_lib

**Library:** autgradalg.lib

**Purpose:** Compute automorphism groups of pointedly graded algebras and of Mori dream spaces.

**Authors:** Simon Keicher

**Overview:** This library provides a framework for computing automorphisms of integral, finitely generated algebras that are graded by a finitely generated abelian group. This library also contains functions to compute automorphism groups of Mori dream spaces. The results are ideals  $I$  such that the respective automorphism group is isomorphic to the subgroup  $V(I)$  in some  $GL(n)$ .

**Assumptions:**

\* the algebra  $R$  is given as factor algebra  $S/I$  with a graded polynomial ring  $S = KK[T_1, \dots, T_r]$ . We will always assume that the basering is  $S$  and it is given over the rationals  $QQ$  or a number field  $QQ(a)$ . \*  $R$  must be minimally presented, i.e.,  $I$  is contained in  $\langle T_1, \dots, T_r \rangle^2$ . \*  $S$  (and hence  $R$ ) are graded via 'setBaseMultigrading(Q)' from 'multigrading.lib'. The last rows of the matrix  $Q$  are interpreted over  $\mathbb{Z}/a_i\mathbb{Z}$  if the respective entry of the list  $TOR$  is  $a_i$  and has been provided as parameter to the respective function. (See the functions for more details.) \* For all  $1 \leq i \leq r$ :  $L_{w_i} = 0$  where  $w_i := \deg(T_i)$ . \* the grading is pointed, i.e., no generator has degree 0 and the cone over all generator degrees is pointed. \* For Mori dream spaces  $X$ , we assume them to be given as  $X = X(R, w)$  with the Cox ring  $R$  of  $X$  (given as the algebra  $R$  before) and an ample class  $w$  in the grading group  $K$  with the torsion entries removed.

**Note:** we require the user to execute 'LIB'gfanlib.so" before using this library.

**Procedures:**

`autKS()` : compute the automorphism group of the basering (must be a polynomial ring) as an algebraic subgroup  $V(I)$  of some  $GL(n)$

`autGradAlg(I0, TOR)`  
: compute the automorphism group of  $R$  as an algebraic subgroup  $V(I)$  of some  $GL(n)$ .

`autGenWeights()`  
: compute the automorphisms of the grading group that fix the generator degrees.

`stabilizer(I0, TOR)`  
: compute the stabilizer of the given ideal

`autXhat(I0, w, TOR)`  
: compute the automorphism group of widehat  $X$  as an algebraic subgroup  $V(I)$  of some  $GL(n)$ .

`autX(I0, w, TOR)`  
: compute the automorphism group of  $X=X(R, w)$  as an algebraic subgroup  $V(I)$  of some  $GL(n)$ .

**Note:** the following functions were taken from 'quotingsingcox.lib' by M.Donten-Bury and S.Keicher: 'hilbertBas'.

**Note:** This library comes without any warranty whatsoever. Use it at your own risk.

### D.15.2 `combinat.lib`

**Library:** `combinat.lib`

**Purpose:** Some useful functions

**Authors:** J. Boehm, boehm @ mathematik.uni-kl.de

**Overview:** Some useful basic functions from combinatorics.

**Procedures:**

```
intersectLists(list,list)
 returns a list of elements which are in both lists (no duplicates)

sublists(list)
 all sublists

member(def,list)
 test whether an element is a member of a list
```

### D.15.3 `diform.lib`

**Library:** `diform.lib`

**Purpose:** Procedures for differential forms

**Author:** Peter Chini, chini@rhrk.uni-kl.de

**Overview:** A library for computing with elements of the differential algebra over a (quotient) ring. To compute in this algebra, a non-commutative ring with additional variables  $dx_1, \dots, dx_n$  and 'exterior' relations between this variables is used. In the case of a quotient ring, the defining ideal and its image under the universal derivation are added as relations. The differential forms themselves are defined via an additional type 'diform'. Objects of this type carry as an attribute a polynomial in the differential algebra and make it available over the basering.

Additionally, the universal derivation is available as a procedure and the differentials between the graded parts of the differential algebra can be applied to differential forms. The library also supports derivations: maps from the first graded part of the differential algebra to the basering. These are defined via the type 'derivation' and there are procedures for basic arithmetic operations, evaluation and Lie-derivative.

**Procedures:**

```
diffAlgebra()
 provides the differential algebra structure and the differential forms
 dx_1, ..., dx_n

diffAlgebraStructure()
 generates the structure of the differential algebra from the basering

diffAlgebraGens()
 defines the differential forms dx_1, ..., dx_n
```

`diffAlgebraUnivDerIdeal(ideal)`  
computes the image of an ideal under the universal derivation

`diffAlgebraChangeOrd(list)`  
returns a ring with the structure of the differential algebra but changed monomial ordering

`diffAlgebraListGen(int)`  
returns a list of the generators of the differential algebra or of a graded part of it

`diformFromPoly(poly)`  
constructs differential forms of degree 0 from polynomials

`diformCoef(diform)`  
computes the representation as a linear combination of the generators

`diformGenToString(diform)`  
casts a generator of the differential algebra to a string

`diformHomogDecomp(df)`  
list of differential forms: homogeneous decomposition

`diformToString(diform)`  
casts a differential form to a string

`diformPrint(diform)`  
prints differential forms

`diformIsGen(diform)`  
decides, whether a given differential form is a generator of the differential algebra

`diformAdd(diform,diform)`  
adds two differential forms

`diformSub(diform,diform)`  
subtracts one differential form from the other

`diformNeg(diform)`  
returns the negative of a differential form

`diformMul(diform,diform)`  
multiplies two differential forms

`diformDiv(diform,diform)`  
computes the quotient of two differential forms

`diformEqu(diform,diform)`  
compares two differential forms

`diformNeq(diform,diform)`  
returns the negation of comparing two differential forms

`diformIsBigger(diform,diform)`  
tests if a given differential form is greater than another one

`diformIsSmaller(diform,diform)`  
tests if a given differential form is smaller than another one

`diformDeg(diform)`  
returns the degree of a given differential form

`diformIsHomog(diform)`  
checks if the given differential form is homogeneous

`diformIsHomogDeg(diform,int)`  
checks if the given differential form is homogeneous of given degree

`diformListCont(list,diform)`  
checks if a given differential form is in a given list

`diformListSort(list)`  
sorts lists of differential forms and special lists of lists

`diformUnivDer(diform)`  
computes the image of an polynomial under the universal derivation

`diformDiff(diform)`  
computes the image of an differential form under the differential

`derivationFromList(list)`  
constructs a derivation from a given list

`derivationCheckList(list)`  
checks the form of a given structure list for a derivation

`derivationFromPoly(poly)`  
creates a derivation from a polynomial

`derivationConstructor(def)`  
constructs a derivation from arbitrary input

`derivationToString(derivation)`  
casts a derivation to a string

`derivationPrint(derivation)`  
prints a derivation

`derivationAdd(derivation,derivation)`  
computes the sum of two derivations

`derivationSub(derivation,derivation)`  
subtracts two derivations

`derivationNeg(derivation)`  
negates a given derivation

`derivationMul(derivation,derivation)`  
multiplies two derivations componentwise

`derivationEqu(derivation,derivation)`  
compares two derivations

`derivationNeq(derivation,derivation)`  
returns the negation of comparing two derivations

`derivationEval(derivation,diform)`  
evaluates a derivation at a given differential form of degree 1

`derivationContractionGen(derivation,diform)`  
computes the contraction and applies it to a generator

`derivationContraction(derivation,diform)`  
computes the contraction and applies it to a differential form

`derivationLie(derivation,diform)`  
 returns the Lie-derivative applied to a differential form

### D.15.4 finitediff\_lib

#### Issues:

- installation of `qepcadfilter.pl` needs to be solved
- tests for (nearly) all procedures are missing
- global variables needs to be cleaned
- temporary files needs to be cleaned
- temporary file names need to be unique (think about multiple instances)
- pollution of global Top namespace must be solved
- `u` is not a good name for a procedure

**Library:** `finitediff.lib`

**Purpose:** procedures to compute finite difference schemes for linear differential equations

**Author:** Christian Dingler

**Overview:** Using `qepcad/qepcadsystem` from this library requires the program `qepcad` to be installed. You can download `qepcad` from <http://www.usna.edu/CS/qepcadweb/B/QEPCAD.html>

#### Procedures:

`visualize(f)`  
 shows a scheme in index-notation

`u(D[,#])` gives some vector; depends on `@derivatives`

`scheme([v1,...,vn])`  
 computes the finite difference scheme defined by `v1,...,vn`

`laxfrT(Ut,U,space)`  
 Lax-Friedrich-approximation for the time-direction

`laxfrX(Ux,U,space)`  
 Lax-Friedrich-approximation for the space-direction

`forward(U1,U2,VAR)`  
 forward-approximation

`backward(U1,U2,VAR)`  
 backward-approximation

`central1st(U1,U2,VAR)`  
 central-approximation of first order

`central2nd(U1,U2,VAR)`  
 central-approximation of second order

`trapezoid(U1,U2,VAR)`  
 trapezoid-approximation

`midpoint(U1,U2,VAR)`  
 midpoint-approximation

`pyramid(U1,U2,VAR)`  
 pyramid-approximation  
`setinitials(variable,der[,#])`  
 constructs and sets the basering for further computations  
`errormap(f)`  
 performs the Fouriertransformation of a poly  
`matrixsystem(M,A)`  
 gives the scheme of a pde-system as one matrix  
`timestep(M)`  
 gives the several timelevels of a scheme derived from a pde-system  
`fouriersystem(M,A)`  
 performs the Fouriertransformation of a matrix scheme  
`PartitionVar(f,n)`  
 partitions a poly into the var(n)-part and the rest  
`ComplexValue(f)`  
 computes the complex value of f, var(1) being the imaginary unit  
`VarToPar(f)`  
 substitute var(i) by par(i)  
`ParToVar(f)`  
 substitute par(i) by var(i)  
`qepcad(f)`  
 ask QEPCAD for equivalent constraints to  $f < 1$   
`qepcadsystem(l)`  
 ask QEPCAD for equivalent constraints to all eigenvals of some matrices  
 being  $< 1$

### D.15.5 goettsche\_lib

**Library:** goettsche.lib

**Purpose:** Drezet's formula for the Betti numbers of the moduli space of Kronecker modules; Goettsche's formula for the Betti numbers of the Hilbert scheme of points on a surface; Nakajima's and Yoshioka's formula for the Betti numbers of the punctual Quot-schemes on a plane or, equivalently, of the moduli spaces of the framed torsion-free planar sheaves; Macdonald's formula for the symmetric product

**Author:** Oleksandr Iena, o.g.yena@gmail.com

**References:**

- [1] Drezet, Jean-Marc Cohomologie des varié'te's de modules de hauteur nulle. *Mathematische Annalen*: 281, 43-85, (1988).
- [2] Goettsche, Lothar, The Betti numbers of the Hilbert scheme of points on a smooth projective surface. *Mathematische Annalen*: 286, 193-208, (1990).
- [3] Macdonald, I. G., The Poincare polynomial of a symmetric product, *Mathematical proceedings of the Cambridge Philosophical Society*: 58, 563-568, (1962).
- [4] Nakajima, Hiraku; Lectures on instanton counting, CRM Proceedings and Lecture Notes, Yoshioka, Kota Volume 88, 31-101, (2004).

**Procedures:**

- `GoettscheF(z, t, n, b)`  
The Goettsche's formula up to n-th degree
- `PPolyH(z, n, b)`  
Poincare Polynomial of the Hilbert scheme of n points on a surface
- `BettiNumsH(n, b)`  
Betti numbers of the Hilbert scheme of n points on a surface
- `NakYoshF(z, t, r, n)`  
The Nakajima-Yoshioka formula up to n-th degree
- `PPolyQp(z, n, b)`  
Poincare Polynomial of the punctual Quot-scheme of rank r on n planar points
- `BettiNumsQp(n, b)`  
Betti numbers of the punctual Quot-scheme of rank r on n planar points
- `MacdonaldF(z, t, n, b)`  
The Macdonald's formula up to n-th degree
- `PPolyS(z, n, b)`  
Poincare Polynomial of the n-th symmetric power of a variety
- `BettiNumsS(n, b)`  
Betti numbers of the n-th symmetric power of a variety
- `PPolyN(t, q, m, n)`  
Poincare Polynomial of the moduli space of Kronecker modules N (q; m, n)
- `BettiNumsN(q, m, n)`  
Betti numbers of the moduli space of Kronecker modules N (q; m, n)

**D.15.6 graal\_lib****Library:** graal.lib**Purpose:** localization at prime ideals and their associated graded rings**Author:** Magdaleen Marais, magdaleen@aims.ac.za  
Yue Ren, ren@mathematik.uni-kl.de**Overview:** This library is on a computational treatment of localizations at prime ideals and their associated graded rings based on a work of Mora. Not only does it construct a ring isomorphic to the localization of an affine coordinate ring at a prime ideal, the algorithms in this library aim to exploit the topology in the localization by computing first and foremost in the associated graded ring and lifting the result to the localization afterwards. Features include a check for regularity and the resolution of ideals.**References:**

- Mora, Teo: La queste del Saint Gr<sub>a</sub>(A.L): A computational approach to local algebra
- Marais, Magdaleen and Ren, Yue: Mora's holy graal: Algorithms for computing in localizations at prime ideals



**Procedures:**

`graalMixed(ideal L[,int t])`  
 construct `graalBearer`

`dimensionOfLocalization(def L)`  
 dimension of the localization  $A_L$  of  $A$  at  $L$

`systemOfParametersOfLocalization(def L)`  
 system of parameter of the localization  $A_L$  of  $A$  at  $L$

`isLocalizationRegular(def L)`  
 test if localization  $A_L$  of  $A$  at  $L$  is regular

`warkedPreimageStd(warkedModule wM)`  
 std for `warkedModule`

`resolutionInLocalization(ideal I, def L)`  
 the resolution of  $I \cdot A_L$

**D.15.7 GND\_lib****Library :** GND.lib**Author :** Adrian Popescu, popescu@mathematik.uni-kl.de**Overview :**

A method to compute the General Neron Desingularization in the frame of one dimensional local domains

**References:**

[1] A. Popescu, D. Popescu, "A method to compute the General Neron Desingularization in the frame of one dimensional local domains", [arxiv.org/abs/1508.05511](https://arxiv.org/abs/1508.05511)

**Procedures:**

`desingularization()`

**D.15.8 gradedModules\_lib****Library:** gradedModules.lib**Purpose:** Operations with graded modules/matrices/resolutions**Authors:** Oleksandr Motsak <U@D>, where U=motsak, D=mathematik.uni-kl.de  
 Hanieh Keneshlou <hkeneshlou@yahoo.com>

**Overview:** The library contains several procedures for constructing and manipulating graded modules/matrices/resolutions. Basics about graded objects can be found in [DL]. Throughout this library graded objects are graded maps, that is, matrices with polynomials, together with grading weights for source and destination. Graded modules are implicitly given as coker of a graded map. Note that in special cases we may also consider submodules in  $S^r$  generated by columns of a graded polynomial matrix (or a graded map).

**Note:** set `assumeLevel` to positive integer value in order to auto-check all assumptions. We denote the current basering by  $S$ .

**References:**

[DL] Decker, W., Lossen, Ch.: Computing in Algebraic Geometry, Springer, 2006

**Procedures:**

**grobj**(M,w[,d])  
 construct a graded object (map) given by matrix M

**grtest**(A)  
 check whether A is a valid graded object

**grdeg**(M) compute graded degrees of columns of the map M

**grview**(M)  
 view the graded structure of map M

**grshift**(M,d)  
 shift graded module coker(M) by +d

**grzero**() presentation of  $S(0)^1$

**grtwist**(r,d)  
 presentation of  $S(d)^r$

**grtwists**(v)  
 presentation of  $S(v[1])+\dots+S(v[\text{size}(v)])$

**grsum**(M,N)  
 direct sum of two graded modules coker(M) + coker(N)

**grpower**(M,p)  
 direct p-th power of graded module coker(M)

**grtranspose**(M)  
 un-ordered graded transpose of map M

**grgens**(M)  
 try to compute submodule generators of coker(M)

**grpres**(F)  
 presentation of submodule generated by columns of F

**grorder**(M)  
 reorder cols/rows of M for correct graded-block-structure

**grtranspose1**(M)  
 reordered graded transpose of map M

**TestGRRes**(n,I)  
 compute/order/transpose a graded resolution of ideal I

**KeneshlouMatrixPresentation**(v)  
 build some presentation with intvec v

**grsyz**(M) syzygy of  $\text{Im}(M)$

**grres**(M,l[,m])  
 resolution of  $\text{Im}(M)$  of length l... minimal?

**grlift**(A,B)  
 graded lift, gens!

**grprod**(A,B)  
 composition of graded maps (product of matrices?)

**grgroebner**(M)  
 Groebner Basis of  $\text{Im}(M)$  as a graded object

`grconcat(M,N)`  
 sum of maps into the same target module  
`grrndmat(s,d[,p,b])`  
 generate random matrix compatible with src and dst gradings  
`grrndmap(S,D[,p,b])`  
 generate random 0-deg homomorphism  $\text{src}(S) \rightarrow \text{src}(D)$   
`grrndmap2(S,D[,p,b])`  
 generate random 0-deg homomorphism  $\text{dst}(S) \rightarrow \text{dst}(D)$   
`grlifting(A,B)`  
 RND! chain lifting  
`grlifting2(A,B)`  
 RND! chain lifting  
`mappingcone(M,N)`  
 mapping cone?  
`grlifting3(A,B)`  
 RND! chain lifting? probably wrong one  
`mappingcone3(A,B)`  
 mapping cone3?  
`grrange(M)`  
 get the row-weightings  
`grneg(A)` graded object given by  $-A$   
`matrixpres(a)`  
 matrix presentation of direct sum of  $\Omega^{\{a[i]\}}(i)$

### D.15.9 maxlike\_lib

**Library:** maxlike.lib

**Purpose:** Procedures to compute maximum likelihood estimates

**Author:** Adrian Koch (kocha at rhrk.uni-kl.de)

**References:**

Lior Pachter, Bernd Sturmfels; Algebraic Statistics for Computational Biology; published by Cambridge University Press

**Procedures:**

`likeIdeal(I,u)`  
 the likelihood ideal with respect to I and u  
`logHessian(I,u)`  
 modified Hessian of the loglikelihood function  
`getMaxPoints(Iu,H,prec,[..])`  
 maximum likelihood estimates  
`maxPoints(I,u,prec,[..])`  
 maximum likelihood estimates, combines the procedures above  
`maxPointsProb(I,u,prec,[..])`  
 maximum likelihood estimates and probability distributions

### D.15.10 methods\_lib

**Library:** methods.lib

**Purpose:** installing methods in Singular

**Authors:** J. Boehm, boehm @ mathematik.uni-kl.de

**Overview:** Methods select the function to execute by the types of the input tuple. The central function is installMethod, which takes a hashtable associating a tuple of input types to function names and creates a corresponding procedure.

HashTables are lists with arbitrary index sets. They can be created by the command hashTable. Their size can be determined by the command size. Values can be extracted by selectKey or the \* operator. HashTables can also be added using addHashTables or the + operator.

Methods can be added with the + operator.

**Types:** Method the class of all methods  
HashTable the class of all hash tables

**Procedures:**

### D.15.11 modules\_lib

**Library:** modules.lib

**Purpose:** Modules

**Authors:** J. Boehm, boehm@mathematik.uni-kl.de  
D. Wienholz wienholz@mathematik.uni-kl.de  
C. Koenen koenen@rhrk.uni-kl.de  
M. Mayer mayer@mathematik.uni-kl.de

**Overview:** This library is used for the computation of graded free resolutions with an own graduation of the monomials. For these Resolution is a new class of modules needed. These modules, can be computed via the image, kernel, cokernel of a matrix or the subquotient of two matrices. The used matrices also have a free module as source and target, with graded generators if the matrix is homogeneous. A matrix of this new form is created by a normal matrix, source, target and the graduation, if the matrix is homogeneous, are done automatically. With this matrices it is then possible to compute the new class of modules.

This library also offers the opportunity to create R-module-homomorphisms between two modules. For these homomorphisms the kernel can be computed and will be returned as a module of the new class.

This is experimental work in progress!!!

**Types:** Matrix the class of matrices with source and target in form of free modules FreeModule  
free modules represented with the ring and degree Resolution class of graded resolutions

Module modules represented by either the image, coker, kernel of a matrix or the subquotient of two matrices Vector element of a Module

Ideal same as ideal, but with its own basering saved, used to compute resolutions  
Homomorphism class of R-module-homomorphisms

**Procedures:**

id(int n) return a nxn identity Matrix

```
zero(int n,int m)
 return a nxm zero Matrix

freeModule(ring,int,list)
 creating a graded free module

makeMatrix(matrix,#int)
 creating a Matrix with graded target and source if the matrix is homo-
 geneous. If # is set to 1, makeMatrix ignores the grading of source &
 target.

makeIdeal(ideal)
 creates an Ideal from an given ideal, is used to compute a resolution of the
 ideal

Target(Matrix)
 return target of the Matrix

Source(Matrix)
 return source of the Matrix

printMatrix(Matrix)
 print a Matrix

printFreeModule(FreeModule)
 print a FreeModule

printResolution(Resolution)
 print a Resolution

printModule(Module)
 print a Module

printHom(Homomorphism)
 print a Homomorphism

mRes(Module/Ideal,#int)
 return a minimized graded Resolution

sRes(Module/Ideal,#int)
 return a graded Resolution computed with Schreyer's method

Res(Module/Ideal,#int)
 return a graded Resolution

Betti(Resolution)
 return the Betti-Matrix of the Resolution

printBetti(Resolution)
 prints the Betti-matrix of the Resolution

SetDeg(list/intvec)
 sets an own graduation for the monomials

Deg(poly)
 same as deg, but can be used with an own graduation

Degree(FreeModule)
 return list with degrees of the module

Degrees(Module)
 return list with degrees of the module
```

`subquotient(Matrix,Matrix)`  
return a Module, the subquotient of the two Matrices

`coker(Matrix)`  
return a Module, the cokernel of the Matrix

`image(Matrix)`  
return a Module, the image of the Matrix

`Ker(Matrix)`  
return a Module, the kernel of the Matrix

`compareModules(Module,Module)`  
return 0 or 1, compares the two Modules up to isomorphism

`addModules(Module,Module)`  
return a Module, sum of the two Modules

`homomorphism(matrix,Module,Module)`  
creates a R-Modul-Homomorphism

`target(Homomorphism)`  
return a Module, target of the Homomorphism

`source(Homomorphism)`  
return a Module, source of the Homomorphism

`compareMatrix(Matrix,Matrix)`  
return 0 or 1, compares two Matrices

`freeModule2Module(FreeModule)`  
converts a FreeModule into a Module

`makeVector(vector,Module)`  
creates Vector in the given Module

`netVector(Vector)`  
prints Vector

`netMatrix(Matrix)`  
prints Matrix

`presentation(Module)`  
converts M as a Subquotient to the Coker of a matrix C

`tensorMatrix(Matrix,Matrix)`  
computes tensorproduct of two Matrices

`tensorModule(Module,Module)`  
computes tensorproduct of two Modules

`tensorModFreemod(Module,FreeModule)`  
computes tensorproduct of Module and FreeModule

`tensorFreemodMod(FreeModule,Module)`  
computes tensorproduct of FreeModule and Module

`tensorFreeModule(FreeModule,FreeModule)`  
computes tensorproduct of two FreeModules

`tensorProduct(def,def)`  
computes tensorproduct

`pruneModule(Module)`  
simplifies the presentation of a Module  
`hom(Module,Module)`  
computes  $\text{Hom}(M,N)$   
`kerHom(Homomorphism)`  
computes the kernel of a Homomorphism  
`interpret(Vector)`  
interprets the Vector in some Module or abstract space  
`interpretInv(def,Module)`  
interprets a Vector or Homomorphism into the given Module  
`reduceIntChain(Module,#int)`  
reduces a chain of interpretations to minimal size or # steps  
`interpretElem(Vector,#int)`  
interpret a Vector with # steps or until can't interpret further  
`interpretList(list,#int)`  
interpret a list of Vectors as far as possible  
`compareVectors(Vector,Vector)`  
compares two Vectors with regard to the relations of their Module  
`simplePrune(Module)`  
simplify module

### D.15.12 modwalk\_lib

**Library:** modwalk.lib

**Purpose:** Groebner basis conversion

**Authors:** S. Oberfranz oberfran@mathematik.uni-kl.de

**Overview:** A library for converting Groebner bases of an ideal in the polynomial ring over the rational numbers using modular methods. The procedures are inspired by the following paper:  
Elizabeth A. Arnold: Modular algorithms for computing Groebner bases. Journal of Symbolic Computation 35, 403-419 (2003).

**Procedures:**

`modWalk(I,#)`  
standard basis conversion of I by Groebner Walk using modular methods  
`modrWalk(I,radius,#)`  
standard basis conversion of I by Random Walk using modular methods  
`modfWalk(I,#)`  
standard basis conversion of I by Fractal Walk using modular methods  
`modfrWalk(I,radius,#)`  
standard basis conversion of I by Random Fractal Walk using modular methods

See also: Section D.4.9 [grwalk\_lib], page 815; Section D.15.18 [rwalk\_lib], page 940; Section D.15.21 [swalk\_lib], page 942.

**D.15.13 multigrading\_lib**

Todos/Issues:

See <http://code.google.com/p/convex-singular/wiki/Multigrading>

**Library:** multigrading.lib

**Purpose:** Multigraded Rings

**Authors:** Benjamin Bechtold, benjamin.bechtold@gmail.com  
 Rene Birkner, rbirkner@math.fu-berlin.de  
 Lars Kastner, lkastner@math.fu-berlin.de  
 Simon Keicher, keicher@mail.mathematik.uni-tuebingen.de  
 Oleksandr Motsak, U@D, where U={motsak}, D={mathematik.uni-kl.de}  
 Anna-Lena Winz, anna-lena.winz@math.fu-berlin.de

**Overview:** This library allows one to virtually add multigradings to Singular: grade multivariate polynomial rings with arbitrary (fin. gen. Abelian) groups. For more see <http://code.google.com/p/convex-singular/wiki/Multigrading> For theoretical references see:

E. Miller, B. Sturmfels: 'Combinatorial Commutative Algebra' and  
 M. Kreuzer, L. Robbiano: 'Computational Commutative Algebra'.

**Note:** 'multiDegBasis' relies on 4ti2 for computing Hilbert Bases. All groups are finitely generated Abelian

**Procedures:**

```

setBaseMultigrading(M,L)
 attach multiweights/grading group matrices to the basering

getVariableWeights([R])
 get matrix of multidegrees of vars attached to a ring

getGradingGroup([R])
 get grading group attached to a ring

getLattice([R[,choice]])
 get grading group' lattice attached to a ring (or its NF)

createGroup(S,L)
 create a group generated by S, with relations L

createQuotientGroup(L)
 create a group generated by the unit matrix with relations L

createTorsionFreeGroup(S)
 create a group generated by S which is torsionfree

printGroup(G)
 print a group

isGroup(G)
 test whether G is a valid group

isGroupHomomorphism(L1,L2,A)
 test whether A defines a group homomorphism from L1 to L2

isGradedRingHomomorphism(R,f,A)
 test graded ring homomorph

```



`createGradedRingHomomorphism(R,f,A)`  
create a graded ring homomorph

`setModuleGrading(M,v)`  
attach multiweights of units to a module and return it

`getModuleGrading(M)`  
get multiweights of module units (attached to M)

`isSublattice(A,B)`  
test whether A is a sublattice of B

`imageLattice(P,L)`  
computes an integral basis for  $P(L)$

`intRank(A)`  
computes the rank of the intmat A

`kernelLattice(P)`  
computes an integral basis for the kernel of the linear map P.

`latticeBasis(B)`  
computes an integral basis of the lattice B

`preimageLattice(P,L)`  
computes an integral basis for the preimage of the lattice L under the linear map P.

`projectLattice(B)`  
computes a linear map of lattices having the primitive span of B as its kernel.

`intersectLattices(A,B)`  
computes an integral basis for the intersection of the lattices A and B.

`isIntegralSurjective(P)`  
test whether the map P of lattices is surjective.

`isPrimitiveSublattice(A)`  
test whether A generates a primitive sublattice.

`intInverse(A)`  
computes the integral inverse matrix of the intmat A

`integralSection(P)`  
for a given linear surjective map P of lattices this procedure returns an integral section of P.

`primitiveSpan(A)`  
computes a basis for the minimal primitive sublattice that contains the given vectors (by A).

`factorgroup(G,H)`  
create the group  $G \bmod H$

`productgroup(G,H)`  
create the group  $G \times H$

`multiDeg(A)`  
compute the multidegree of A

**multiDegBasis(d)**  
 compute all monomials of multidegree d

**multiDegPartition(p)**  
 compute the multigraded-homogeneous components of p

**isTorsionFree()**  
 test whether the current multigrading is free

**isPositive()**  
 test whether the current multigrading is positive

**isZeroElement(p)**  
 test whether p has zero multidegree

**areZeroElements(M)**  
 test whether an integer matrix M considered as a collection of columns has zero multidegree

**isHomogeneous(a)**  
 test whether 'a' is multigraded-homogeneous

**equalMultiDeg(e1,e2[,V])**  
 test whether  $e1=e2$  in the current multigrading

**multiDegGroebner(M)**  
 compute the multigraded GB/SB of M

**multiDegSyzygy(M)**  
 compute the multigraded syzygies of M

**multiDegModulo(I,J)**  
 compute the multigraded 'modulo' module of I and J

**multiDegResolution(M,l[,m])**  
 compute the multigraded resolution of M

**multiDegTensor(m,n)**  
 compute the tensor product of multigraded modules m,n

**multiDegTor(i,m,n)**  
 compute the  $Tor_i(m,n)$  for multigraded modules m,n

**defineHomogeneous(p)**  
 get a grading group wrt which p becomes homogeneous

**pushForward(f)**  
 find the finest grading on the image ring, homogenizing f

**gradiator(h)**  
 coarsens grading of the ring until h becomes homogeneous

**hermiteNormalForm(A)**  
 compute the Hermite Normal Form of a matrix

**smithNormalForm(A,#)**  
 compute matrices D,P,Q with  $D=P*A*Q$  and D is the smith normal form of A

**hilbertSeries(M)**  
 compute the multigraded Hilbert Series of M

**lll(A)** applies LLL(.) of lll.lib which only works for lists on a matrix A

**D.15.14 nets\_lib****Library:** net.lib**Purpose:** Net structures for pretty printing**Authors:** J. Boehm, boehm@mathematik.uni-kl.de  
M. Mueller, mkmuelle@mathematik.uni-kl.de  
H. Rombach, rombach@mathematik.uni-kl.de  
M. Stein, maxstein77@web.de**Overview:** Nets are arrays of characters, which are printed in a matrix format. They can be concatenated horizontally and vertically. When concatenating horizontally, empty rows are filled with spaces. All Singular types can be converted to a Net by applying the command net.**Types:** Net The class of all nets**Procedures:**

catNets(Net,Net)  
horizontal concatenation

net(def) general procedure to generate a net from a Singular object

netBigIntMat(bigintmat)  
procedure to generate a net from a bigintmat

netBigIntMatShort(bigintmat)  
procedure to generate a net from a bigintmat

netCoefficientRing(ring)  
procedure to generate a net from a Singular coefficient ring

netIdeal(ideal)  
procedure to generate a net from an ideal

netInt(int)  
procedure to generate a net from a integer

netBigInt(bigint)  
procedure to generate a net from a bigint

netIntMat(intmat)  
procedure to generate a net from an intmat

netIntMatShort(intmat)  
procedure to generate a net from an intmat

netIntVector(vector)  
procedure to generate a net from an intvec

netIntVectorShort(vector)  
procedure to generate a net from an intvec

netNumber(number)  
procedure to generate a net from a number

netList(list)  
procedure to generate a net from a list

`netMap(map)`  
     procedure to generate a net from a map  
`netMap2(map)`  
     procedure to generate a net from a map  
`netmatrix(matrix)`  
     procedure to generate a net from a matrix  
`netmatrixShort(matrix)`  
     procedure to generate a net from a matrix  
`netPoly(poly)`  
     procedure to generate a net from a poly  
`netPrimePower(int,int)`  
     procedure to generate a net from a prime power  
`netRing(ring)`  
     procedure to generate a net from a polynomial ring  
`netString(string)`  
     procedure to generate a net from a string  
`netvector(vector)`  
     procedure to generate a net from a vector  
`netvectorShort(vector)`  
     procedure to generate a net from a vector  
`stackNets(Net,Net)`  
     vertical concatenation

### D.15.15 pfd\_lib

**Library:** pfd.lib

**Purpose:** Multivariate Partial Fraction Decomposition

**Author:** Marcel Wittmann, e-mail: mwittman@mathematik.uni-kl.de

**Overview:** This Library implements an algorithm based on the work of E. K. Leinartas to write rational functions in mutiple variables as a sum of functions with "smaller" numerators and denominators.

This can be used to shorten the IBP reduction coefficients of multi-loop Feynman integrals. For this application, we also provide a procedure that applies the algorithm to all entries of a matrix of rational functions given as one (possibly very big) txt-file. If you use the library pfd.lib, please cite the corresponding paper [J. Boehm, M. Wittmann, Z. Wu, Y. Xu, Y. Zhang: 'IBP reduction coefficients made simple' (preprint 2020).

**Procedures:**

`pfd()`      calculate a partial fraction decomposition of a rational function  
`checkpfd()`  
     test if a decomposition is equal to a rational function given by numerator/denominator polynomials  
`evaluatepfd()`  
     substitute values in a partial fraction decomposition gotten from `pfd`

`displaypfd()`  
 print a decomposition gotten as output of `pfd`  
`displaypfd_long()`  
 like `display`, but denominators are written out  
`getStringpfd()`  
 turn a decomposition gotten from `pfd` into one string  
`getStringpfd_indexed()`  
 like `getStringpfd`, but writes the denominator factors just as `q1, q2, ...`  
`readInputTXT()`  
 read a matrix of rational functions from a txt-file  
`pfdMat()` apply `pfd` to a matrix of rational functions in parallel (using Section D.2.7  
 [`parallel.lib`], page 797) and save result as easy-to-read txt-files.  
`checkpfdMat()`  
 test output files of `pfdMat` for correctness

### D.15.16 `polyclass.lib`

**Library:** `polyclass.lib`

**Purpose:** Data types for normal form equations

**Authors:** Janko Boehm, email: [boehm@mathematik.uni-kl.de](mailto:boehm@mathematik.uni-kl.de)  
 Magdaleen Marais, email: [magdaleen.marais@up.ac.za](mailto:magdaleen.marais@up.ac.za)  
 Gerhard Pfister, email: [pfister@mathematik.uni-kl.de](mailto:pfister@mathematik.uni-kl.de)

**Overview:** This library implements a ring independent polynomial type used for the return value in `classify2.lib` and `realclassify.lib`. You can use `+`, `*` and `==` for addition, multiplication and comparison. The key over contains the base ring of the polynomial, the key value its value as a polynomial of type `poly`. The constructor can be called by assigning a polynomial of type `poly` to a polynomial of type `Poly` via `=`.

Moreover the library implements a class `NormalFormEquation` consisting out of a string type, an integer `milnorNumber`, a `Poly normalFormEquation`, and integer modality, a list of numbers parameters, a list variables, an integer `corank`, in the real case, an integer `inertiaIndex`, a list of open intervals represented as lists consisting out of two rationals used to select a real root of the minimal polynomial (which is stored in the variable `minpoly` of the polynomial ring containing `normalFormEquation`, that is, in `normalFormEquation.in`), or if no minimal polynomial is defined then an interval containing the rational parameter value.

**Acknowledgements:** This research was supported by the Staff Exchange Bursary Programme of the University of Pretoria, DFG SPP 1489, DFG TRR 195. The financial assistance of the National Research Foundation (NRF), South Africa, towards this research is hereby acknowledged. Opinions expressed and conclusions arrived at are those of the author and are not necessarily to be attributed to the National Research Foundation, South Africa.

**Procedures:**

`makePoly(f)`  
 constructor for ring independent polynomial type `Poly`  
`printPoly(f)`  
 print routine for polynomial type `Poly`

```
printNormalFormEquation(F)
 print routine for normal form equations
```

See also: Section D.6.5 [classify2\_lib], page 857; Section D.6.19 [realclassify\_lib], page 867.

### D.15.17 ringgb\_lib

**Library:** ringgb.lib

**Purpose:** Functions for coefficient rings

**Author:** Oliver Wienand, email: wienand@mathematik.uni-kl.de

**Procedures:**

```
findZeroPoly(f)
 finds a vanishing polynomial for reducing f

zeroReduce(f)
 normal form of f concerning the ideal of vanishing polynomials

testZero(poly f)
 tests f defines the constant zero function

noElements(def r)
 the number of elements of the coefficient ring, if of type (integer, ...)
```

### D.15.18 rwalk\_lib

**Library:** rwalk.lib

**Purpose:** Groebner Walk Conversion Algorithms

**Author:** Stephan Oberfranz

**Procedures:**

```
prwalk(ideal, int, int [, intvec, intvec])
 standard basis of ideal via Random Perturbation Walk algorithm

rwalk(ideal, int [, intvec, intvec])
 standard basis of ideal via Random Walk algorithm

frandwalk(ideal, int [, intvec, intvec])
 standard basis of ideal via Random Fractal Walk algorithm
```

See also: Section D.4.9 [grwalk.lib], page 815; Section D.15.21 [swalk\_lib], page 942.

### D.15.19 sets\_lib

**Library:** sets.lib

**Purpose:** Sets in Singular

**Authors:** J. Boehm, boehm @ mathematik.uni-kl.de  
 D. Wienholz, wienholz @ mathematik.uni-kl.de  
 S. Zillien, zillien @ rhrk.uni-kl.de

**Overview:** We implement the new class `set` and all basic methods needed to work with sets. A set is generated from a list. After the generating of a set, the adding of an element or the union of two sets, automatically every double element is removed to secure that no element occurs in a set more than once.

There is a comparison operator, we access the operator via the function `isEqual`. This function `isEqual` can be used to compare two elements of the same type (`Set`, `list`, `int`, `bigint`, `string`, `intmat`, `bigintmat`, `intvec`, `ring`, `map`, `poly`, `matrix`, `ideal`, `module`, `vector`, `resolution`) and also works for comparing of `int`, `bigint` and number with each other, similarly for `matrix`, `bigintmat` and `intmat`.

The function `size` can be used to determine the number of elements.

The `+` operator is used for the union, the `*` operator for the intersection.

The operators `<` and `>` can be used for inclusion tests.

The `print` function can be used for printing sets.

Note that the implementation of the underlying data structure and algorithms is very trivial and will at some point be replaced with something more efficient.

**Types:** `Set` The class of all sets

**Procedures:**

```

set(list)
 general procedure to generate a set from a list

union(Set,Set)
 union of sets

intersectionSet(Set,Set)
 intersection of sets

complement(Set,Set)
 complement of sets

isElement(def,Set)
 test whether an object is in a set

isSubset(Set,Set)
 test whether a set is a subset of another set

isSuperset(Set,Set)
 test whether a set is a superset of another set

addElement(Set,def)
 adds an element to the set

```

### D.15.20 stanleyreisner\_lib

**Library:** `stanleyreisner.lib`

**Purpose:** Deformations of Stanley-Reiser ideals

**Authors:**

**Overview:** Firstly, we implement the graded pieces has certain degree of cotangent modules `T1` and `T2` for a general Stanley-Reiser ring. And the graded pieces of homomorphisms are represented by lists of integers.

**Types:** Homomorphism class of homomorphisms

**Procedures:**

`T1(ideal)`  
compute first order deformations

`T2(ideal)`  
compute second order deformations

`makeQPoly(poly)`  
create a QPoly

`fPiece(ideal,poly,poly)`  
create a FirstOrderDeformation

`sPiece(ideal,poly,poly)`  
create a SecondOrderDeformation

`makeLinks(ideal,poly,poly)`  
create a FirstOrderDeformation of the links

**D.15.21 swalk\_lib****Library:** swalk.lib**Purpose:** Sagbi Walk Conversion Algorithm**Author:** Junaid Alam Khan junaidalamkhan@gmail.com**Overview:** A library for computing the Sagbi basis of subalgebra through Sagbi walk algorithm.

**Theory:** The concept of SAGBI ( Subalgebra Analog to Groebner Basis for Ideals) is defined in [L. Robbiano, M. Sweedler: Subalgebra Bases, volume 42, volume 1430 of Lectures Note in Mathematics series, Springer-Verlag (1988),61-87]. The Sagbi Walk algorithm is the subalgebra analogue to the Groebner Walk algorithm which has been proposed in [S. Collart, M. Kalkbrener and D.Mall: Converting bases with the Grobner Walk. J. Symbolic Computation 24 (1997), 465-469].

**Procedures:**

`swalk(ideal[,intvec])`  
Sagbi basis of subalgebra via Sagbi walk algorithm

`rswalk(ideal,int,int[,intvec])`  
Sagbi basis of subalgebra via Random Sagbi Walk Algorithm

See also: Section D.4.9 [grwalk\_lib], page 815; Section D.15.18 [rwalk\_lib], page 940.

**D.15.22 systhreads\_lib****Library:** systhreads.lib**Purpose:** Primitives for Singular's multi-threaded objects**Author:** Reimer Behrends

**Overview:** This library implements basic functionality for shared objects in a multi-threaded system, such as channels, shared tables & lists, and synchronization variables.



**D.15.23 VecField\_lib****Library:** VecField.lib**Purpose:** vector fields, with algorithms for jordan and diagonal forms**Authors:** Adrian Rettich, rettich@mathematik.uni-kl.de  
Raul Epure, epure@mathematik.uni-kl.de**References:**

[1] Kyoji Saito, Quasihomogene isolierte Singularitaeten von Hyperflaechen, 1971

**Overview:** Implements a class VecField, represented by a vector. For example, 'VecField V = [x3,xy]' declares the vector field  $v = x^3 dx + xy dy$ . Instead of a vector, an nx1 matrix is also accepted. The vector can be recovered as V.vec.

Supports coordinate transformations (via maps), which are represented by tracking a map 'V.coordinates' which maps the standard coordinates to those in which V is currently represented. V.dimension stores the vector field's dimension, which is just nvars(basing), and V.lin yields the linear part of V. You may set an additional parameter V.precision, which dictates the degree to which operations on the vector field should be exact.

The default precision is 1. Precision is preserved across transformations, additions, and all other manipulations of vector fields.

**Procedures:**

```

applyVecField(VecField V, ..., [int n])
 apply V to a poly p / an ideal I as an operator; you can also use 'V*p'/'V*I'.
 If an integer n is passed, consider V up to degree n.

changeCoordinates(VecField V, map psi)
 transform V by psi; you can also use 'V*phi'

jordanVecField(VecField V)
 transform V s.t. the linear part is in Jordan normal form

diagonalizeVecFieldLin(list l)
 l a list of VecFields. Change coordinates s.t. all linear parts are diagonal
 simultaneously.

SaitoBase(VecField V)
 algorithm to find a basis where the semisimple and nilpotent parts are
 easily read off

diagonalizeVecField(list l)
 diagonalize all VecFields in l simultaneously

vecFieldToMatrix(VecField V, ideal W)
 matrix representation of V in the basis W

decomposeVecField(...)
 split a vectorfield V / all entries of a list of vectorfields l into semisimple
 and nilpotent components

diagonalizeMatrixSimul(list l)
 find transformation which simultaneously diagonalizes all matrices in l

invertAlgebraMorphism(map p, int n)
 return inverse of p exact up to degree n

```

## 8 Release Notes

### 8.1 News and changes

#### NEWS in SINGULAR 4-2-0

##### News for version 4-2-0

Syntax changes:

renamed `poly.lib` to `polylib.lib` (Section D.2.8 [`polylib.lib`], page 798)

New libraries:

`interval.lib`: interval arithmetic (Section D.8.2 [`interval.lib`], page 876)

`maxlike.lib`: algebraic statistics (Section D.15.9 [`maxlike.lib`], page 929)

`nchilbert.lib`: Hilbert series for LetterPlace algebras (Section 7.5.13 [`nchilbert.lib`], page 514)

`polyclass.lib`: class of polynomials (Section D.15.16 [`polyclass.lib`], page 939)

`recover.lib`: Hybrid numerical/symbolical algorithms (Section D.8.7 [`recover.lib`], page 879)

`redcgs.lib`: Reduced Comprehensive Groebner Systems (Section D.2.9 [`redcgs.lib`], page 799)

`ringgb.lib`: coefficient rings (Section D.15.17 [`ringgb.lib`], page 940)

`sets.lib`: Sets (Section D.15.19 [`sets.lib`], page 940)

`stanleyreisner.lib`: T1 and T2 for a general Stanley-Reiser ring (Section D.15.20 [`stanleyreisner.lib`], page 941)

`systhreads.lib`: multi-threaded objects (Section D.15.22 [`systhreads.lib`], page 942)

Changed libraries:

`classify_aeq.lib`: new procedure `classSpaceCurve` (Section D.6.6 [`classify_aeq.lib`], page 858)

`grobcov.lib`: new version (Section D.2.4 [`grobcov.lib`], page 791)

`modular.lib`: parallel version for verification via `system("verifyGB",I)`

New commands:

`system("verifyGB",I)`: test, if I is a Groebner basis (using parallel processes)

Letterplace: `modulo,syz,lift,liftstd, rightStd` (Section 7.7 [LETTERPLACE], page 613)

Changes in the kernel/build system:

update for using FLINT 2.6.x and for FLINT 2.7.0

Singular can be build with NTL or FLINT or both (if non is available, `factroize` and `gcd` will not work.)

## News for version 4-1-3

### New libraries:

- invar.lib: Invariant theory Section D.7.4 [invar.lib], page 874
- moddiq.lib: ideal quotient and saturation Section D.4.14 [moddiq.lib], page 817
- ncModslimgb.lib: modular Groebner bases for G-algebras Section 7.5.18 [ncModslimgb.lib], page 550

### Changed libraries:

- chern.lib: new version (Section D.5.2 [chern.lib], page 834)
- grobcov.lib: new version (Section D.2.4 [grobcov.lib], page 791), new functions [ConsLevels], page 795, [Levels], page 796, [GroblLevels], page 796, [DifConsLCSets], page 796

### Changes in the kernel/build system:

- improved gcd and multiplication via FLINT
- improved lift (and related)
- port to polymake 3.5.x
- rational functions via flint (Section 5.1.44 [flintQ], page 183)
- free algebra over Z (Section 7.7 [LETTERPLACE], page 613)
- adaptions/functions for Singular.jl(<https://github.com/oscar-system/Singular.jl>)

## News for version 4-1-2

### New libraries:

- arnoldclassify.lib: Arnol'd Classifier of Singularities (Section D.6.3 [arnoldclassify.lib], page 855)
- diffform.lib: Procedures for differential forms (Section D.15.3 [diffform.lib], page 921)
- dmodideal.lib: Algorithms for Bernstein-Sato ideals of morphisms (Section 7.5.6 [dmodideal.lib], page 440)
- fpalgebras.lib: Generation of various algebras in the letterplace case (Section 7.10.2 [fpalgebras.lib], page 634)
- ncrat.lib: non-commutative rational functions (Section 7.10.6 [ncrat.lib], page 668)

### Changed libraries:

- freegb.lib: lpDivision, lpPrint (Section 7.10.4 [freegb.lib], page 652)
- fpadim.lib (Section 7.10.1 [fpadim.lib], page 629)
- schreyer.lib: deprecated
- goettsche.lib: new, extended version (The Nakajima-Yoshioka formula up to n-th degree, Poincare Polynomial of the punctual Quot-scheme of rank r on n planar points Betti numbers of the punctual Quot-scheme of rank r on n planar points)(Section D.15.5 [goettsche.lib], page 925)
- grobcov.lib: small bug fix (Section D.2.4 [grobcov.lib], page 791)

### Changes in the kernel/build system:

- integrated xalloc into omalloc: (`./configure --disable-omalloc`)
- improved heuristic for `det` (Section 5.1.23 [det], page 170)
- improved reading of long polynomials

improved groebner bases over  $\mathbb{Z}$  coefficients  
code for free algebras (letterplace rings) rewritten (using now the standard `+, -, *, ^, std, ...`) (Section 7.7 [LETTERPLACE], page 613)  
new commands `rightstd` (Section 7.8.6 [rightstd (letterplace)], page 623)  
extended `twostd` to LETTERPLACE (Section 7.8.9 [twostd (letterplace)], page 624, Section 7.3.29 [twostd (plural)], page 358)  
pseudo type `polyBucket`  
new type `smatrix`: sparse matrix (experimental) (Section 4.20 [smatrix], page 127).  
extended `coef` to ideals (Section 5.1.11 [coef], page 162).  
error and signal handling in `libSingular` (Section 8.3 [libSingular], page 953).  
updated `gfanlib` to version 0.6.2  
port to NTL 11 (needs C++11: `gcc6` or `-std=c++11`), which does not conflict with `polymake` (needs C++14)

## News for version 4-1-1

New syntax:

`alias`: may be used as a prefix to a variable declaration. Can only be used in procedure headings. (Section 3.5.1 [General command syntax], page 41).

New command:

`fres`: improved version of `sres`: computes a (not necessarily minimal) free resolution of the input ideal/module, using Schreyer's algorithm. (Section 5.1.48 [fres], page 186, Section 5.1.147 [sres], page 264).

Extended commands:

pseudo ordering `L` allows setting of limits for exponents in polynomials (Section B.2.9 [Pseudo ordering L], page 765, Section 5.1.2 [attrib], page 154 for `maxExp`)

`%,mod`: also for poly operands (Section 4.16.3 [poly operations], page 119).

`delete`: extended to `intvec`, `ideal`, `module` (Section 5.1.21 [delete], page 169).

`syz` (Section 5.1.154 [syz], page 275), `lift` (Section 5.1.80 [lift], page 209), `liftstd` (Section 5.1.81 [liftstd], page 209), `intersect` (Section 5.1.65 [intersect], page 199): with a specified GB algorithm

New libraries:

`classify2.lib`: Classification of isolated singularities of corank  $\leq 2$  and modality  $\leq$  wrt. right equivalence over the complex numbers according to Arnold's list. (Section D.6.5 [classify2.lib], page 857)

`goettsche.lib`: Goettsche's formula for the Betti numbers of the Hilbert scheme of points on a surface, Macdonald's formula for the symmetric product (Section D.15.5 [goettsche.lib], page 925)

`combinat.lib`, `modules.lib`, `methods.lib`, `nets.lib`: a more mathematical view of modules (Section D.15.2 [combinat.lib], page 921: combinatorics), (Section D.15.10 [methods.lib], page 930: construct procedures), (Section D.15.11 [modules.lib], page 930: free resolutions), (Section D.15.14 [nets.lib], page 937: pretty printing)

`ncHilb.lib`: Hilbert series of non-commutative monomial algebras (Section 7.10.5 [ncHilb.lib], page 665)

`realclassify.lib`: Classification of real singularities (Section D.6.19 [realclassify.lib], page 867)

rootisolation.lib: real root isolation using interval arithmetic (Section D.8.8 [rootisolation.lib], page 880)

rstandard.lib: Janet bases and border bases for ideals (Section D.4.31 [rstandard.lib], page 831)

Changed libraries:

chern.lib: new version (Section D.5.2 [chern.lib], page 834)

gitfan.lib: new (incompatible) version (Section D.13.3 [gitfan.lib], page 904)

grobcov.lib: new version (Section D.2.4 [grobcov.lib], page 791)

Changes in the kernel/build system:

port to polymake 3.x.x

port to NTL 10 with threads (needs also C++11: gcc6 or -std=c++11)

p\_Invers is only a helper for p\_Series: now static

p\_Divide is now p\_MDivide, pDivide/p\_Divide is a new routine

## News for version 4-1-0

Syntax changes:

new (additional) form of ring definitions: (for example `ring R=QQ[x,y,z];`) (Section 3.3.2 [General syntax of a ring declaration], page 32)

new (additional) form of multi-indices: (for example `i(1,2,3,4,5)`) (Section 3.5.3 [Names], page 44)

changed behaviour of `charstr` (Section 5.1.7 [charstr], page 160)

new data type `cring` to describe the coefficient rings, to be used for the new definitions for (polynomial) rings (Section 3.3.2 [General syntax of a ring declaration], page 32)

new command `ring_list` to access the parts used to construct polynomial rings (Section 5.1.136 [ring\_list], page 252, Section 5.1.135 [ringlist], page 250)

extended polynomial ring construction: also from lists produced by `ring_list`

new attribute `ring_cf` for `ring` (Section 5.1.2 [attrib], page 154)

printing of rings changed to match `cring` names (Section 5.1.7 [charstr], page 160)

New libraries:

new library: `classifyMapGerms.lib`: standard basis of the tangent space at the orbit of an algebraic group action (Section D.6.9 [classifyMapGerms.lib], page 860)

new library: `ffmodstd.lib`: Groebner bases of ideals in polynomial rings over algebraic function fields (Section D.4.8 [ffmodstd.lib], page 813)

new library: `nfmodsyzy.lib`: syzygy modules of submodules of free modules over algebraic number fields (Section D.4.21 [nfmodsyzy.lib], page 823)

new library: `curveInv.lib`: invariants of curves (Section D.4.5 [curveInv.lib], page 811)

new library: `gfan.lib`: interface to `gfanlib` (Section D.13.2 [gfan.lib], page 901)

extended library: interface to `polymake` merged into Section D.13.4 [polymake.lib], page 906

new library: `tropicalNewton.lib`: Newton polygon methods in tropical geometry (Section D.13.7 [tropicalNewton.lib], page 913)

new library: `schubert.lib`: some procedures for intersection theory (Section D.5.15 [schubert.lib], page 849)

Changed libraries:

classify\_aeq.lib: new procedures (Section D.6.6 [classify\_aeq\_lib], page 858)  
 grobcov.lib: new version (Section D.2.4 [grobcov\_lib], page 791)  
 ncfactor.lib: factorization in some noncommutative algebras (Section 7.5.12 [ncfactor\_lib], page 480) with new routine ncfactor (Section 7.5.12.1 [ncfactor], page 480)  
 primdec.lib: new option "subsystem" (Section D.4.26 [primdec\_lib], page 828)

#### Changes in the kernel:

improved mapping of polynomials/ideals/...  
 port to gcc 6  
 port to gfanlib 0.6 (requires C++11, i.e. gcc >=4.3)  
 port to NTL 10  
 port to polymake 3.0  
 port to readline 7  
 Section 5.1.138 [sba], page 254 works for global orderings, also for coefficient types  $Z$  and  $Z/m$   
 Section 5.1.149 [std], page 266 works for all orderings, also for coefficient types  $Z$  and  $Z/m$  with local/mixed orderings  
 Section 5.1.36 [factorize], page 178 works for polynomial rings over  $\mathbb{Z}$

#### Experimental stuff:

module Section D.14.2 [customstd\_lib], page 917: modify `std` ([satstd], page 917)

## News for version 4-0-3

#### New libraries:

new library: brillnoether.lib: Riemann-Roch spaces of divisors on curves (Section D.5.1 [brillnoether\_lib], page 834)  
 new library: chern.lib: Chern classes (Section D.5.2 [chern\_lib], page 834)  
 new library: ffgroebner.lib: Groebner bases of ideals in polynomial rings over algebraic function fields (Section D.4.8 [ffgroebner\_lib], page 813)  
 new library: GND.lib: General Neron Desingularization (Section D.15.7 [GND\_lib], page 927)  
 new library: graal.lib: localization at prime ideals (Section D.15.6 [graal\_lib], page 926)  
 new library: hess.lib: Riemann-Roch space of divisors (Section D.5.5 [hess\_lib], page 840)

#### Changed libraries:

renamed algemodstd\_lib to Section D.4.20 [nfgroebner\_lib], page 822, extended to `module`  
 renamed derham\_lib to Section D.5.3 [deRham\_lib], page 838  
 grobcov.lib (grobcovK): Groebner Cover for parametric ideals (Section D.2.4 [grobcov\_lib], page 791) with new routine ConsLevels ([ConsLevels], page 795), removed AddCons AddConsP.

## News for version 4-0-2

#### New commands:

align (Section 5.1.1 [align], page 154)  
 branchTo (Section 4.17.3 [procs with different argument types], page 122)  
 -> (Section 4.17.2 [proc expression], page 122)

#### Change in ring handling:

`typeof( qring )` returns "ring"

New libraries:

`algemodstd.lib`: Groebner bases of ideals in polynomial rings over algebraic number fields (renamed to Section D.4.20 [`nfmodstd.lib`], page 822)

`arr.lib`: arrangements of hyperplanes (Section D.14.1 [`arr.lib`], page 914)

`brillnoether.lib`: Riemann-Roch spaces of divisors on curve (Section D.5.1 [`brillnoether.lib`], page 834)

`hess.lib`: Riemann-Roch space of divisors on function fields and curves (Section D.5.5 [`hess.lib`], page 840)

`gradedModules.lib`: graded modules/matrices/resolutions (Section D.15.8 [`gradedModules.lib`], page 927)

Changed libraries:

revised `polymake` interface (`polymake.so`)

revised `gfanlib` interface (`gfanlib.so`)

`Presolve::findvars` ([`findvars`], page 877, Section 5.1.164 [`variables`], page 280)

`Ring::addvarsTo` ([`addvarsTo`], page 804)

`Ring::addNvarsTo` ([`addNvarsTo`], page 804)

`Ring::hasAlgExtensionCoefficient` ([`hasAlgExtensionCoefficient`], page 804)

`Schreyer::s_res` (`s_res`)

`grobcov.lib` (`grobcovK`) (Section D.2.4 [`grobcov.lib`], page 791) with new routines `AddCons` `AddConsP`.

`normaliz.lib` (for `normaliz >=2.8`) (Section D.4.24 [`normaliz.lib`], page 825)

renamed `groebnerFan` to `groebnerFanP` in `polymake.lib` (Section D.13.4 [`polymake.lib`], page 906)

renamed `fVector` to `fVectorP` in `polymake.lib` (Section D.13.4 [`polymake.lib`], page 906, `polymakeInterface.lib`)

## News for version 4-0-1

Version 4-0-1 is a bug fix release.

New feature: attribute `ring_cf` for `ring` (Section 5.1.2 [`attrib`], page 154)

## News for version 4-0-0

Version 4-0-0 is a milestone release of Singular. The new release series 4 aims for an entirely modularized architecture simplifying connectivity with other systems and paving the way for parallel computations. As a first step in modularization, the new release features an internal structural separation of coefficient rings and polynomial rings. This allows for a flexible integration of new coefficient rings.

SINGULAR 4-0-0's list of new functionality and significant improvements further extends that of the 3-1-6/7 prerelease series.

New functionality

de Rham cohomology of complements of algebraic varieties (Section D.5.3 [`deRham.lib`], page 838)

Gromov-Witten numbers of elliptic curves (Section D.4.7 [`ellipticcovers.lib`], page 812)

classification of isolated complete intersection singularities in characteristic 0 (Section D.6.8 [classifyci\_lib], page 859)

parametrization of orbits of unipotent actions (Section D.5.8 [orbitparam\_lib], page 842)

F5-like Groebner basis algorithm (Section 5.1.138 [sba], page 254)

element-wise application of functions to data structures (Section 5.2.1 [apply], page 285)

support for debugging libraries (Section 3.9.1 [ASSUME], page 67)

#### Improved functionality

Groebner cover for parametric ideals (Section D.2.4 [grobpcov\_lib], page 791)

normalization of affine rings (Section D.4.23 [normal\_lib], page 824)

classification of real singularities (Section D.6.19 [realclassify\_lib], page 867)

GIT-fans (Section D.13.3 [gitfan\_lib], page 904)

algebraic/transcendental field extensions

Chapter 7 [Non-commutative subsystem], page 312

an abstraction layer for parallel computations (Section D.2.7 [parallel\_lib], page 797)

run-time loading of supplementary kernel code (Section A.1.9 [Dynamic modules], page 701)

interpreter language support for name spaces (Section 4.15 [package], page 117)

#### Availability

SINGULAR is available as source code and for Linux, Mac OS X, Windows, FreeBSD and SunOS-5.

## 8.2 Singular 3 and Singular 4

The purpose of this section is to describe new features and changes between Singular 3-1-7 and Singular 4.\* (formerly known as Spielwiese) both for developers and Singular users. In what follows we will refer to the systems as Singular 3 and Singular 4.

### 8.2.1 Version schema for Singular

SINGULAR version is of the form `a.b.c.d` which may also be written as `a-b-c-d` where `a`, `b`, `c` and `d` are numbers:

`a` is changed with major, incompatible changes

`b` is changed with incompatible changes (of some commands/libraries)

`c` is changed with compatible changes (i.e. new commands, extended options, new algorithms, etc.)

`d` is changed with each release (i.e. with bug fixes, etc.)

SINGULAR does also have "unofficial" build originating from a code version between "official" version: such builds display "Development version `a.b.c`" in the header while "official" versions show "version `a.b.c`". Also the manual describes version `a-b-c`. To get the complete version number, use `system("version");` or use `SINGULAR_VERSION` in C.

### 8.2.2 Notes for Singular users



## Coefficient rings

To allow for easy integration of new coefficient rings into Singular, the way coefficient rings are being handled has been redesigned.

In general, the user syntax has not changed, however there are some changes in the behaviour of Singular:

- setting `minpoly` results in changing the current coefficient domain and clears all previously defined variables of that ring
- Minor changes in the output of coefficient ring description. Moreover the output of elements of certain rings has been improved (for example, reals).
- Algebraic and transcendental extensions of rationals and finite fields have been reimplemented. In particular, the heuristics for clearing denominators and factoring out content have been changed. In some cases this leads to a different, mathematically equivalent results of Groebner bases and related computations. For example a Groebner basis element may differ by a unit.
- Most notably, due to the redesign of the coefficient rings, if the user sets the minimal polynomial all variables dependent on the current ring are deleted.

## Ring-dependent options

Formally global Singular Section 5.1.110 [option], page 230 now belong to individual polynomial rings. This includes:

- `intStrategy`
- `redTail`
- `redThrough`

Also the following settings now belong to individual (currently active) polynomial rings:

- `short`
- `minpoly`
- `noether`

Hence setting these options only affects the current ring. Be aware of this when switching between different rings, since the options affect the result of various computations (in particular Groebner bases).

## Path names

- The tree structure of the binary Singular distribution has been changed. The typical tree now looks as show at <https://github.com/Singular/Singular/wiki/Sw-tree>
- Accordingly Singular search paths (where Singular searches for libraries, dynamic modules, etc.) have been changed. You can display them by calling Singular by `Singular -v`.
- currently, multi-arch installations of Singular 4 aere not possible.

## Library versioning

Due to switching from Subversion to GIT revision control system for the Singular source code, library version variables (displayed when loading a library) have changed.

## New orderings for modules

The now can assign weights to module components, when defining a monomial ordering. For example

```
ring R = 0, (x,y,z), (am(1,2,3, 10,20,30,40), dp, C);
deg(x*gen(1));
↳ 11
```

will assign weights 1,2,3 to x,y,z respectively, and weights 10,20,30,40,0,0,... to components of any free module defined over R. This ordering will first sort by this weighted degree, then by dp on the ring monomials and then will give priority to the large component index.

## Future benefits of Singular 4

The redesign of Singular will allow us to provide new features in the future, for example:

- Interpreter type for coefficient rings.
- User defined coefficient rings.
- Improved syntax for defining polynomial rings.

### 8.2.3 Notes for developers

There has been an extensive process of refactoring, redesign and modularization of Singular to facilitate easier maintenance and future development:

- Build System : automake, libfac has been integrated into Factory
- Removed MP (Multi protocol) in favor of SSI links.
- Separation/modularization into libraries and packages
- For easy integration of new coefficient rings, we defined a generic interface for coefficient rings and a supporting framework for making them accessible to the user.

In particular we have separated everything related to coefficient rings into a separate library `libcoeffs`. Dependency tree between restructured packages is show at <https://www.singular.uni-kl.de/dox/singular.png>

In order to use `libSingular` as a C++ library, see Section 8.3 [`libSingular`], page 953.

### 8.2.4 Building Singular

The user can build and install Singular with the following standard UNIX-like procedure:

- Download and extract the latest official source package (.tar.gz).
- Run the configure script, for instance, `./configure`.
- Build Singular by running `make`.
- Install Singular by running `make install`.

In contrast to Singular 3, there are now many more configuration options.

All possible options for configure can be seen by running the configure script with option `--help`. On a multicore compute consider running `make` with the option `-j [cores]`.

### 8.2.5 Side-by-side installation

Due to choosing paths according to FS standards it is no longer possible to have a side-by-side installation of different Singular versions or versions for different architectures.

## 8.3 libSingular

libSingular is the C++-library version of SINGULAR.

Singular/libsingular.h is the main include file, -lSingular the link parameter,

lib/pkgconfig/Singular.pc provides all parameters in the pkgconfig format.

It contains all parts of SINGULAR with the following exceptions:

1. memory allocation functions for GMP (see mmInit in Singular/testths.cc)
2. signal handlers (see init\_signals in Singular/cntrlc.cc).  
At least a handler for SIGCHLD must be installed for the commands Section 5.1.168 [waitfirst], page 282, Section 5.1.167 [waitall], page 281 and the routines from Section D.2.7 [parallel\_lib], page 797, Section D.4.16 [modstd\_lib], page 818, Section D.4.15 [modnormal\_lib], page 818, Section D.2.13 [tasks\_lib], page 804.  
If the child was started by libSingular the handler has to call sig\_chld\_hdl from Singular/links/ssiLink.cc or implement something similar (call s1Close(1) for ssi links).
3. error handlers for factory, NTL (see init\_signals in Singular/cntrlc.cc).

## 8.4 Download instructions

SINGULAR is available as source and binary program for most common hard- and software platforms. Instructions to download and install SINGULAR can be found at

<https://www.singular.uni-kl.de/index.php/singular-download.html>.

Release versions of SINGULAR are also available from our FTP site

<ftp://jim.mathematik.uni-kl.de/pub/Math/Singular/src/4-1-2/>.

## 8.5 Used environment variables

SINGULAR needs to find some files (dynamic modules, libraries, help files). Usually they are found relative to the location of the (main) executable (after following symlinks). This can be changed by setting the following environment variables.

SINGULAR\_EXECUTABLE (should usually not be set)

the complete filename of the main executable, usually derived from the command line (inspecting also PATH, following symlinks).

If SINGULAR\_EXECUTABLE cannot be found, \$prefix/bin/Singular is assumed.

For libSingular: SINGULAR\_EXECUTABLE is set to the argument of siInit (it must exist).

SINGULAR\_BIN\_DIR

the directory of the main executable, usually derived from \$SINGULAR\_EXECUTABLE

SINGULAR\_ROOT\_DIR

the root of the singular tree, default: \$SINGULAR\_BIN\_DIR/..

SINGULAR\_DATA\_DIR

the root of the singular data files, default: \$SINGULAR\_BIN\_DIR/./share

SINGULARPATH

the directories for libraries and optional dynamic modules (separated by ;), default:

\$SINGULAR\_DATA\_DIR/singular/LIB

\$SINGULAR\_ROOT\_DIR/share/singular/LIB

\$SINGULAR\_BIN\_DIR/./share/singular/LIB

```

$SINGULAR_DATA_DIR/factory
$SINGULAR_ROOT_DIR/share/factory
$SINGULAR_BIN_DIR/LIB
$SINGULAR_BIN_DIR/./factory
$SINGULAR_BIN_DIR/MOD
$SINGULAR_ROOT_DIR/lib/singular/MOD
$SINGULAR_ROOT_DIR/libexec/singular/MOD
$prefix/lib/singular/MOD
$prefix/libexec/singular/MOD
$SINGULAR_BIN_DIR
SINGULAR_PROCS_DIR
the directories for dynamic modules (separated by ;), default:
$SINGULAR_BIN_DIR/MOD
$SINGULAR_ROOT_DIR/lib/singular/MOD
$SINGULAR_ROOT_DIR/libexec/singular/MOD
$prefix/lib/singular/MOD
$prefix/libexec/singular/MOD
SINGULAR_INFO_FILE
singular.hlp, default: $SINGULAR_DATA_DIR/info/singular.hlp
SINGULAR_IDX_FILE
the help index, default: $SINGULAR_DATA_DIR/singular/singular.idx
SINGULAR_HTML_DIR
the directory of the manual as html files, default: $SINGULAR_DATA_DIR/singular/html
SINGULAR_URL
the URL of the manual, default: https://www.singular.uni-kl.de/Manual/

```

The effective list of directories/files can be printed by `Singular -v`, see Section 3.1.6 [Command line options], page 19.

Depending on the used functions, these environment variables apply also to `libSingular`.

## 8.6 Unix installation instructions

Install binaries: <https://www.singular.uni-kl.de/index.php/singular-download/install-linuxunix>  
or build it yourself:

Install the necessary packages:

```

libtool
gnu make
gcc, g++
libreadline
gmp, mpfr
ntl
libcdd

```

Install flint 2.5 (or newer): `./configure --with-gmp=/usr --prefix=$HOME/tmp`  
`make && make install`

```
Install Singular ./configure --with-flint=$HOME/tmp --enable-gfanlib --
prefix=$HOME/Singular4
make && make install
($prefix/bin/Singular is the main executatble)
(optional) install 4ti2
(optional) install surf/surfer
(optional) install normaliz 2.8 (or newer)
```

See also <https://github.com/Singular/Singular/wiki/Step-by-Step-Installation-Instructions-for-Singular> which includes instructions adapted for debian and fedora based systems.

## 8.7 Windows installation instructions

Singular relies on Cygwin as its environment under Windows. There is a 32bit and a 64bit version of Cygwin.

<https://www.singular.uni-kl.de/index.php/singular-download/install-windows.html>

## 8.8 Macintosh installation instructions

**Installation of the provided binaries** <https://www.singular.uni-kl.de/index.php/singular-download/install-os-x.html>

If your Mac refuses to open Singular because of an "unidentified developer": Open System Preferences. Go to the Security & Privacy tab. Click on the lock and enter your password so you can make changes. Change the setting for 'Allow apps downloaded from' to 'App Store and identified developers'.

You may also check <https://support.apple.com/en-en/guide/mac-help/mh40616/mac>

## 9 Index

- !**  
 ! ..... 43  
 != ..... 43, 86
- #**  
 # ..... 43
- \$**  
 \$ ..... 43
- %**  
 % ..... 43, 74, 84, 91, 115, 119
- &**  
 && ..... 43, 87
- (**  
 ( ..... 42  
 () ..... 105
- )**  
 ) ..... 42
- \***  
 \* ..... 43, 74, 76, 79, 84, 90, 91, 108, 111, 115, 119  
 \*\* ..... 43
- - ..... 42, 74, 76, 84, 90, 91, 108, 115, 119, 132  
 - ..... 42  
 -allow-net ..... 19  
 -batch ..... 21  
 -browser ..... 19  
 -echo ..... 19  
 -emacs ..... 21  
 -emacs-dir ..... 21  
 -emacs-load ..... 21  
 -execute ..... 20  
 -help ..... 19  
 -min-time ..... 20  
 -MPhost ..... 21  
 -MPport ..... 21  
 -no-out ..... 20  
 -no-rc ..... 20  
 -no-shell ..... 20  
 -no-stdlib ..... 20  
 -no-tty ..... 20  
 -no-warn ..... 20  
 -quiet ..... 20  
 -random ..... 20  
 -sdb ..... 19  
 -singular ..... 21  
 -ticks-per-sec ..... 21  
 -user-option ..... 20  
 -> ..... 122  
 -b ..... 21  
 -c ..... 20  
 -d ..... 19  
 -e ..... 19  
 -h ..... 19  
 -q ..... 20  
 -r ..... 20  
 -u ..... 20  
 -v ..... 20
- .**  
 ..... 43  
 .singularrc file ..... 22  
 .singularrc file, no loading ..... 20
- /**  
 / ..... 43, 72, 91, 108, 115, 119, 132  
 // ..... 43
- :**  
 : ..... 43, 91  
 :: ..... 43, 117
- ;**  
 ; ..... 43
- =**  
 = ..... 42  
 == ..... 43, 86, 108, 115, 119, 125, 130, 132
- ?**  
 ? ..... 43, 191
- [**  
 [ ..... 42  
 [] ..... 111, 119, 130, 132
- ]**  
 ] ..... 42

|    |                                                                    |
|----|--------------------------------------------------------------------|
| '  |                                                                    |
| '  | 43                                                                 |
| -  |                                                                    |
| -  | 43                                                                 |
| {  |                                                                    |
| {  | 42                                                                 |
|    |                                                                    |
|    | 43, 87                                                             |
| }  |                                                                    |
| }  | 42                                                                 |
| ~  |                                                                    |
| ~  | 43, 297                                                            |
| "  |                                                                    |
| "  | 43                                                                 |
| +  |                                                                    |
| +  | 42, 74, 76, 79, 84, 90, 91, 102, 108, 111, 115, 119, 125, 130, 132 |
| ++ | 42                                                                 |
| >  |                                                                    |
| >  | 43, 119, 132                                                       |
| >= | 43, 86, 115, 119, 130, 132                                         |
| ^  |                                                                    |
| ^  | 43, 74, 79, 115, 119                                               |
| \  |                                                                    |
| \  | 43                                                                 |
| <  |                                                                    |
| <  | 43, 119, 132, 182                                                  |
| <= | 43, 86, 115, 119, 130, 132                                         |
| <> | 43, 86, 108, 115, 119, 125, 130, 132                               |

**A**

|                                           |     |
|-------------------------------------------|-----|
| a, ordering                               | 764 |
| A.L                                       | 857 |
| A.Z                                       | 790 |
| A.Z.L                                     | 790 |
| absfact.lib                               | 809 |
| absfact.lib                               | 809 |
| absFactorize                              | 809 |
| absFactorizeBCG                           | 809 |
| absolute factorization                    | 809 |
| absPrimdecGTZ                             | 829 |
| absPrimdecGTZE                            | 829 |
| abstractR                                 | 849 |
| absValue                                  | 790 |
| Access to elements of a user defined type | 135 |
| actionIsProper                            | 873 |
| addcol                                    | 806 |
| addcores                                  | 803 |
| addElement                                | 941 |
| addLeftFractions                          | 589 |
| addModules                                | 932 |
| addnondegeneratevariables                 | 868 |
| addNvarsTo                                | 804 |
| addRat                                    | 533 |
| addrow                                    | 806 |
| addSheaf                                  | 850 |
| addvarsTo                                 | 804 |
| ademRelations                             | 636 |
| ADGT                                      | 795 |
| Adj_div                                   | 886 |
| adjoint                                   | 808 |
| Adjoint ideal                             | 844 |
| adjointIdeal                              | 844 |
| admissibleSub                             | 554 |
| afaces                                    | 905 |
| affine code                               | 780 |
| AG codes                                  | 756 |
| AGcode.L                                  | 886 |
| AGcode.Omega                              | 886 |
| ainvar.lib                                | 873 |
| ainvar.lib                                | 873 |
| aksaka.lib                                | 891 |
| aksaka.lib                                | 891 |
| Alexander polynomial                      | 854 |
| alexanderpolynomial                       | 854 |
| alexpoly.lib                              | 854 |
| alexpoly.lib                              | 854 |
| alg_kernel                                | 810 |
| algDependent                              | 810 |
| algebra.lib                               | 809 |
| algebra.containment                       | 809 |
| algebra.lib                               | 809 |
| algebraic dependence                      | 596 |
| Algebraic dependence                      | 725 |
| Algebraic geometry                        | 834 |
| Algebraic Geometry codes                  | 886 |

- algebraic statistics . . . . . 929
- algebraicDependence . . . . . 833
- algorithm of Bigatti, La Scala and Robbiano . . . . . 775
- algorithm of Conti and Traverso . . . . . 773
- algorithm of Di Biase and Urbanke . . . . . 774
- algorithm of Hosten and Sturmfels . . . . . 774
- algorithm of Pottier . . . . . 774
- alias . . . . . 41
- align . . . . . 154
- all.lib . . . . . 787
- all.lib . . . . . 787
- allDoubleExt . . . . . 603
- allExtOfLeft . . . . . 601
- allExtOfRight . . . . . 602
- allowing net access . . . . . 19
- allPositive . . . . . 380
- allprint . . . . . 796
- allreal . . . . . 900
- allrealst . . . . . 900
- allsquarefree . . . . . 865
- AltVarEnd . . . . . 574
- AltVarStart . . . . . 573
- ambientDimension . . . . . 901
- and . . . . . 87, 305
- Ann . . . . . 828
- annfalphal . . . . . 445
- annfs . . . . . 397
- annfs0 . . . . . 407
- annfs2 . . . . . 408
- annfsBMI . . . . . 404
- annfsLogIdeal . . . . . 441
- annfsParamBM . . . . . 403
- annfspecial . . . . . 397
- annfsRB . . . . . 409
- annihilator of polynomial . . . . . 415
- annihilator of rational function . . . . . 415, 519
- annihilatorMultiFs . . . . . 442
- annil . . . . . 821
- annPoly . . . . . 415
- annRat . . . . . 416
- annRatSyz . . . . . 526
- Appel function . . . . . 415
- Appel hypergeometric function . . . . . 415
- appelF1 . . . . . 432
- appelF2 . . . . . 433
- appelF4 . . . . . 433
- appendWeight2Ord . . . . . 556
- Applications . . . . . 752
- apply . . . . . 285
- applyAdF . . . . . 390
- applyMatrix . . . . . 877
- applyPermutationToIntvec . . . . . 906
- applyVecField . . . . . 943
- arcpoint.lib . . . . . 854
- arcpoint.lib . . . . . 854
- areEqualLeftFractions . . . . . 591
- areZeroElements . . . . . 936
- argument, default . . . . . 52
- argument, optional . . . . . 52
- Arnol'd . . . . . 855
- ArnoldAction . . . . . 867
- arnoldClassify . . . . . 856
- arnoldclassify.lib . . . . . 855
- arnoldclassify.lib . . . . . 855
- arnoldCorank . . . . . 856
- arnoldDeterminacy . . . . . 856
- arnoldListAllSeries . . . . . 856
- arnoldMilnorCode . . . . . 856
- arnoldMorseSplit . . . . . 856
- arnoldNormalForm . . . . . 856
- arnoldShowSeries . . . . . 856
- arr.lib . . . . . 914
- arr.lib . . . . . 914
- arr2multarr . . . . . 916
- arrange . . . . . 411
- arrBoolean . . . . . 916
- arrBoundedChambers . . . . . 917
- arrBraid . . . . . 916
- arrCenter . . . . . 915
- arrCentered . . . . . 915
- arrCentral . . . . . 915
- arrCentralize . . . . . 915
- arrChambers . . . . . 917
- arrCharPoly . . . . . 916
- arrCone . . . . . 915
- arrCoordChange . . . . . 915
- arrCoordNormalize . . . . . 915
- arrDecone . . . . . 915
- arrDer . . . . . 916
- arrEdelmanReiner . . . . . 916
- arrEssentialize . . . . . 916
- arrExponents . . . . . 916
- arrFlats . . . . . 916
- arrIsEssential . . . . . 915
- arrIsFree . . . . . 916
- arrLastVar . . . . . 915
- arrLattice . . . . . 916
- arrLocalize . . . . . 915
- arrOrlikSolomon . . . . . 916
- arrPoincare . . . . . 916
- arrPrintMatrix . . . . . 915
- arrRandom . . . . . 916
- arrRandomCentral . . . . . 916
- arrRestrict . . . . . 915
- arrSet . . . . . 915
- arrSwapVar . . . . . 915
- arrTypeB . . . . . 916
- arrTypeD . . . . . 916
- ASCII . . . . . 790
- ASCII links . . . . . 95
- ask . . . . . 892
- assignment, custom . . . . . 136
- Assignments for user defined types . . . . . 136
- AssocTanToEnv . . . . . 795



- assPrimes . . . . . 810
  - assprimeszerodim.lib . . . . . 810
  - assprimeszerodim.lib . . . . . 810
  - ASSUME . . . . . 67
  - Atkin . . . . . 892
  - atkins.lib . . . . . 892
  - atkins.lib . . . . . 892
  - attrib . . . . . 154
  - autGenWeights . . . . . 920
  - autGradAlg . . . . . 920
  - autgradalg.lib . . . . . 920
  - autgradalg.lib . . . . . 920
  - Authors . . . . . 3
  - autKS . . . . . 920
  - automorphism group . . . . . 921
  - automorphisms . . . . . 921
  - autonom . . . . . 889
  - autonomDim . . . . . 889
  - autX . . . . . 920
  - autXhat . . . . . 920
  - awalk1 . . . . . 815
  - awalk2 . . . . . 815
- B**
- b-function . . . . . 371
  - babyGiant . . . . . 893
  - Background . . . . . 4
  - backward . . . . . 924
  - bareiss . . . . . 156
  - base2str . . . . . 790
  - basing . . . . . 30, 77
  - Basic programming . . . . . 691
  - basicinvariants . . . . . 856
  - baumslagGroup . . . . . 637
  - baumslagSolitar . . . . . 636
  - BelongSemig . . . . . 901
  - belongSemigroup . . . . . 811
  - BerlekampMassey . . . . . 814
  - Bern . . . . . 837
  - bernstein . . . . . 863
  - Bernstein operator . . . . . 396
  - Bernstein-Sato ideal . . . . . 441
  - Bernstein-Sato polynomial . . . . . 371, 396, 863
  - Bernstein-Sato polynomial for variety . . . . . 447
  - bernsteinBM . . . . . 400
  - bernsteinLift . . . . . 400
  - BernsteinSatoIdeal . . . . . 443
  - bertini . . . . . 841, 842, 879
  - bertini2Singular . . . . . 841
  - beti . . . . . 157
  - Betti . . . . . 931
  - beti (plural) . . . . . 329
  - Betti number . . . . . 768
  - BettiNumsH . . . . . 926
  - BettiNumsN . . . . . 926
  - BettiNumsQp . . . . . 926
  - BettiNumsS . . . . . 926
  - Betty number . . . . . 926
  - bFactor . . . . . 435
  - BFBoundsBudur . . . . . 444
  - bfct . . . . . 371
  - bfctAnn . . . . . 373
  - bfctBound . . . . . 526
  - bfctIdeal . . . . . 375
  - bfctOneGB . . . . . 374
  - bfctSyz . . . . . 372
  - bfctVarAnn . . . . . 448
  - bfctVarIn . . . . . 447
  - bfun.lib . . . . . 370
  - bfun.lib . . . . . 370
  - Bigatti-La Scala-Robbiano algorithm . . . . . 775
  - bigint . . . . . 73
  - bigint declarations . . . . . 73
  - bigint expressions . . . . . 73
  - bigint operations . . . . . 74
  - bigint related functions . . . . . 74
  - bigintmat . . . . . 74
  - bigintmat declarations . . . . . 74
  - bigintmat expressions . . . . . 75
  - bigintmat operations . . . . . 76
  - bigintmat type cast . . . . . 75
  - bigintToBinary . . . . . 906
  - bimodules . . . . . 366
  - Bimodules . . . . . 628
  - bimodules.lib . . . . . 365
  - bimodules.lib . . . . . 365
  - binary\_add . . . . . 894
  - binaryToBigint . . . . . 906
  - binomial . . . . . 790
  - binomials2intmat . . . . . 827
  - BINresol . . . . . 845
  - bistd . . . . . 366
  - biszygies . . . . . 366
  - Biszygy . . . . . 628
  - bitrinity . . . . . 367
  - blackbox . . . . . 138, 144
  - block . . . . . 47, 285
  - BlowingUp . . . . . 861
  - blowUp . . . . . 847
  - blowup0 . . . . . 812
  - blowUp2 . . . . . 847
  - blowUpBO . . . . . 848
  - Blowupcenter . . . . . 845
  - boolean expressions . . . . . 86
  - boolean operations . . . . . 87
  - boolean\_constant . . . . . 919
  - boolean\_ideal . . . . . 919
  - boolean\_poly . . . . . 919
  - boolean\_poly\_ring . . . . . 919
  - boolean\_set . . . . . 919
  - boolean\_std . . . . . 919
  - border basis . . . . . 832
  - borderBasis . . . . . 832

- |                                       |               |                           |          |
|---------------------------------------|---------------|---------------------------|----------|
| BorelCheck                            | 853           | cccMatrixToPositiveIntvec | 913      |
| Bott's formula                        | 851           | cdd                       | 1        |
| boundaryLatticePoints                 | 907           | cddlib                    | 1        |
| boundBuFou                            | 900           | CenCharDec                | 478      |
| boundDes                              | 900           | center                    | 381, 385 |
| boundposDes                           | 900           | Center                    | 847      |
| bounds                                | 881           | CenterBO                  | 848      |
| bounds2                               | 876           | centerRed                 | 384      |
| boxSet                                | 881           | centerVS                  | 383      |
| bracket                               | 270, 306, 330 | central.lib               | 381      |
| Branches of space curve singularities | 738           | central_lib               | 381      |
| branchTo                              | 122           | central1st                | 924      |
| break                                 | 286           | central2nd                | 924      |
| break point                           | 297           | centralize                | 381      |
| breakpoint                            | 286           | centralizer               | 381, 386 |
| Briancon-Maisonobe algorithm          | 396           | centralizerRed            | 383      |
| Brieskorn lattice                     | 863, 864, 866 | centralizerVS             | 382      |
| Brill-Noether algorithm               | 886           | centralizeSet             | 382      |
| BrillNoether                          | 886           | CentralQuot               | 477      |
| brillnoether.lib                      | 834           | CentralSaturation         | 478      |
| brillnoether_lib                      | 834           | cf_class                  | 155      |
| brnoeth.lib                           | 886           | cgs                       | 790      |
| brnoeth_lib                           | 886           | cgsdr                     | 794      |
| browser, command line option          | 19            | chaincrit                 | 898      |
| browser, setting the                  | 270           | chAll                     | 835      |
| browsers                              | 16            | chAllInv                  | 835      |
| browsers, setting the                 | 270           | Change of rings           | 11       |
| BSidealFromAnn                        | 442           | changechar                | 803      |
| bubblesort                            | 892           | changeCoordinates         | 943      |
| Buchberger algorithm for toric ideals | 775           | changeDenominator         | 825      |
| Budur-Mustata-Saito approach          | 447           | changeord                 | 803      |
| Building Singular                     | 952           | changeordTo               | 804      |
| buildtree                             | 801           | changes                   | 944      |
| buildtreetoMaple                      | 801           | changevar                 | 803      |
| busadj                                | 808           | char                      | 159      |
| bvar                                  | 919           | char_series               | 159      |
| <b>C</b>                              |               |                           |          |
| C programming language                | 304           | Characteristic sets       | 768      |
| c, module ordering                    | 762           | characteristic variety    | 415, 519 |
| C, module ordering                    | 762           | CharacteristicExponents   | 861      |
| calculate_max_sum                     | 896           | charexp2conductor         | 854      |
| calculate_ordering                    | 894           | charexp2generators        | 854      |
| calculateI                            | 845           | charexp2inter             | 854      |
| cancelunit, option                    | 232           | charexp2multseq           | 854      |
| canonicalizeCone                      | 901           | charexp2poly              | 854      |
| canonize                              | 889           | charInfo                  | 430      |
| canonMap                              | 815           | charpoly                  | 808      |
| cantodiffcgs                          | 801           | charstr                   | 160      |
| cantoradd                             | 897           | charVariety               | 429      |
| cantormult                            | 897           | chDual                    | 835      |
| cantorred                             | 897           | checkFactor               | 410      |
| cardGroup                             | 811           | checkpfd                  | 938      |
| case                                  | 305           | checkpfdMat               | 939      |
| Category string                       | 55            | checkRoot                 | 405      |
| catNets                               | 937           | chern                     | 835      |
|                                       |               | chern.lib                 | 834      |
|                                       |               | chern_lib                 | 834      |
|                                       |               | chernCharPoly             | 837      |

|                                              |          |                                                                    |     |
|----------------------------------------------|----------|--------------------------------------------------------------------|-----|
| ChernClass                                   | 850      | classpoly                                                          | 837 |
| chernPoly                                    | 837      | classSpaceCurve                                                    | 859 |
| ChernRootsDual                               | 835      | cleanTmp                                                           | 912 |
| ChernRootsHom                                | 836      | cleanunit                                                          | 845 |
| ChernRootsProd                               | 835      | cleardenom                                                         | 161 |
| ChernRootsSum                                | 835      | close                                                              | 161 |
| ChernRootsSymm                               | 836      | closed_points                                                      | 886 |
| ChernRootsWedge                              | 836      | closetex                                                           | 884 |
| Chevalley-Rosenlicht theorem                 | 842      | closureFrac                                                        | 825 |
| chHE                                         | 835      | CM_regularity                                                      | 852 |
| chHom                                        | 836      | CMtype                                                             | 869 |
| chineseRem                                   | 893      | Code                                                               | 777 |
| chinrem                                      | 160      | Codes and the decoding problem                                     | 777 |
| chinrempoly                                  | 822      | codim                                                              | 868 |
| chinrestp                                    | 896      | coDim                                                              | 860 |
| chNum                                        | 835      | codimension                                                        | 901 |
| chNumbers                                    | 835      | coDimMap                                                           | 860 |
| chNumbersProj                                | 837      | coding theory                                                      | 756 |
| ChowRing                                     | 850      | Coding theory                                                      | 886 |
| chProd                                       | 836      | coef                                                               | 162 |
| chProdE                                      | 836      | coefficient field                                                  | 113 |
| chProdL                                      | 836      | Coefficient rings                                                  | 951 |
| chProdLP                                     | 836      | coefficient rings, ring of integers, zero divisors, p-adic numbers | 36  |
| chProdM                                      | 836      | coefficients, long                                                 | 696 |
| chProdMP                                     | 836      | coeffmod                                                           | 891 |
| chProj                                       | 837      | coeffs                                                             | 163 |
| chSum                                        | 835      | coHom                                                              | 545 |
| chSymm                                       | 836      | coker                                                              | 932 |
| chSymm2L                                     | 836      | collectDiv                                                         | 849 |
| chSymm2LP                                    | 836      | colrank                                                            | 889 |
| chWedge                                      | 836      | colred                                                             | 807 |
| chWedge2L                                    | 836      | combinat.lib                                                       | 921 |
| chWedge2LP                                   | 836      | combinat_lib                                                       | 921 |
| cimonom.lib                                  | 900      | comma                                                              | 306 |
| cimonom_lib                                  | 900      | Command line options                                               | 19  |
| cisimplicial.lib                             | 810      | command,custom                                                     | 135 |
| cisimplicial_lib                             | 810      | command-line option, setting value of                              | 274 |
| classification                               | 855, 858 | command-line option, value of                                      | 273 |
| Classification of hypersurface singularities | 741      | command-line options, print all values of                          | 273 |
| classify                                     | 856      | command-line options, short help                                   | 19  |
| classify.lib                                 | 856      | Commands                                                           | 154 |
| classify_aeq.lib                             | 858      | commands (letterplace)                                             | 619 |
| classify_aeq_lib                             | 858      | commands (plural)                                                  | 329 |
| classify_lib                                 | 856      | Commands for user defined types                                    | 135 |
| classify2.lib                                | 857      | comment                                                            | 43  |
| classify2_lib                                | 857      | commRing                                                           | 534 |
| classifyCeq                                  | 859      | Commutative algebra                                                | 809 |
| classifyceq.lib                              | 859      | Commutative Algebra                                                | 706 |
| classifyceq_lib                              | 859      | compareMatrix                                                      | 932 |
| classifyci.lib                               | 859      | compareModules                                                     | 932 |
| classifyci_lib                               | 859      | compareTasks                                                       | 805 |
| classifyicis                                 | 860      | compareVectors                                                     | 933 |
| classifyMapGerms.lib                         | 860      | CompDecomp                                                         | 369 |
| classifyMapGerms_lib                         | 860      | CompInt                                                            | 901 |
| classifySimpleMaps                           | 860      | complement                                                         | 941 |
| classifySimpleMaps1                          | 861      | CompleteHomog                                                      | 835 |
| classifyUnimodalMaps                         | 861      |                                                                    |     |

- completeReduction . . . . . 873
- complex . . . . . 30
- complexClassify . . . . . 858
- complexSingType . . . . . 857
- complexType . . . . . 858
- ComplexValue . . . . . 925
- compregb.lib . . . . . 790
- compregb\_lib . . . . . 790
- comprehensive Groebner system . . . . . 790
- Comprehensive Groebner Systems . . . . . 791, 799
- compress . . . . . 806
- computeAfaceOrbits . . . . . 905
- computeConstant . . . . . 813
- computeGromovWitten . . . . . 813
- computemcm . . . . . 846
- computeN . . . . . 849
- computeV . . . . . 849
- Computing Groebner and Standard Bases . . . . . 702
- concat . . . . . 806
- cone . . . . . 137
- coneLink . . . . . 901
- coneViaInequalities . . . . . 901
- coneViaPoints . . . . . 901
- conicWithTangents . . . . . 911
- ConsLevels . . . . . 795
- constructblwup . . . . . 846
- constructH . . . . . 846
- constructlastblwup . . . . . 846
- ContactMatrix . . . . . 861
- containedQ . . . . . 883
- containsAsFace . . . . . 902
- containsInCollection . . . . . 903
- containsInSupport . . . . . 902
- containsPositiveVector . . . . . 902
- containsRelatively . . . . . 902
- content . . . . . 798
- contentSB, option . . . . . 232
- Conti-Traverso algorithm . . . . . 773
- continue . . . . . 286, 306
- contract . . . . . 165
- contraHom . . . . . 546
- contributionBundle . . . . . 851
- contributors . . . . . 270
- Contributors . . . . . 3
- control . . . . . 889
- Control structures . . . . . 285
- Control theory . . . . . 888
- control.lib . . . . . 888
- control\_lib . . . . . 888
- control\_Matrix . . . . . 862
- controlDim . . . . . 889
- controlExample . . . . . 889
- convertdata . . . . . 845
- convertLeftToRightFraction . . . . . 587
- convertRightToLeftFraction . . . . . 586
- convexHull . . . . . 902
- convexIntersection . . . . . 902
- convloc . . . . . 412
- Cooper philosophy . . . . . 778
- coords . . . . . 899
- copyright . . . . . 1
- copyTask . . . . . 805
- corank . . . . . 856
- Cornacchia . . . . . 892
- CornacchiaModified . . . . . 892
- cornerMonomials . . . . . 828
- countPoints . . . . . 894
- cProj . . . . . 837
- cpu . . . . . 270, 273
- crcgs . . . . . 801
- create\_ring . . . . . 165, 786
- createBO . . . . . 848
- createGradedRingHomomorphism . . . . . 935
- createGroup . . . . . 934
- createlist . . . . . 846
- createQuotientGroup . . . . . 934
- createTask . . . . . 805
- createTorsionFreeGroup . . . . . 934
- Crep . . . . . 794
- CRHT-ideal . . . . . 778
- cring . . . . . 72
- cring declarations . . . . . 72
- cring expressions . . . . . 72
- cring operations . . . . . 72
- cring related functions . . . . . 73
- Critical points . . . . . 728
- crossprod . . . . . 166
- crypto.lib . . . . . 893
- crypto\_lib . . . . . 893
- crystallographicGroupC2MM . . . . . 641
- crystallographicGroupCM . . . . . 640
- crystallographicGroupP1 . . . . . 637
- crystallographicGroupP2 . . . . . 639
- crystallographicGroupP2GG . . . . . 640
- crystallographicGroupP2MM . . . . . 639
- crystallographicGroupP3 . . . . . 643
- crystallographicGroupP31M . . . . . 643
- crystallographicGroupP3M1 . . . . . 644
- crystallographicGroupP4 . . . . . 641
- crystallographicGroupP4GM . . . . . 642
- crystallographicGroupP4MM . . . . . 642
- crystallographicGroupP6 . . . . . 645
- crystallographicGroupP6MM . . . . . 645
- crystallographicGroupPG . . . . . 638
- crystallographicGroupPM . . . . . 638
- CSMA . . . . . 838
- cup . . . . . 815
- cupproduct . . . . . 815
- curve singularities . . . . . 854, 865
- curve singularity . . . . . 811
- curveColengthDerivations . . . . . 811
- curveConductorMult . . . . . 811
- curveDeligneNumber . . . . . 811
- curveDeltaInv . . . . . 811

- curveInv.lib ..... 811
  - curveInv\_lib ..... 811
  - curvepar.lib ..... 861
  - curvepar\_lib ..... 861
  - CurveParam ..... 861
  - CurveRes ..... 861
  - Curves ..... 844
  - custom assignment ..... 136
  - custom command ..... 135
  - custom type ..... 133
  - Customization of the Emacs interface ..... 27
  - customstd.lib ..... 917
  - customstd\_lib ..... 917
  - cycleLength ..... 907
  - cyclePoints ..... 907
  - cyclic ..... 798
  - cyclic code ..... 777
  - Cyclic code ..... 888
  - Cyclic roots ..... 699
  - cyclotomic ..... 871
  - CYGWIN and ESingular ..... 26
- D**
- D-integration ..... 415
  - D-localization ..... 415, 519
  - D-module ..... 371, 396, 415, 441, 447, 519, 597
  - D-module structure ..... 396, 441, 447
  - D-restriction ..... 415
  - Data types ..... 72
  - Data types (plural) ..... 313
  - datetime ..... 167, 785
  - DBM links ..... 99
  - dbprint ..... 167
  - debug\_log ..... 857
  - debugger ..... 68
  - debugging library code ..... 68
  - Debugging tools ..... 67
  - debugLib, option ..... 232
  - decIvar ..... 821
  - decimal ..... 893
  - Decker, Wolfram ..... 3
  - Declaration of objects of a user defined type ..... 134
  - decode ..... 887
  - decodeCode ..... 888
  - decodegb.lib ..... 887
  - decodegb\_lib ..... 887
  - decodeRandom ..... 887
  - decodeRandomFL ..... 888
  - decodeSV ..... 886
  - Decoding ..... 888
  - Decoding codes with Groebner bases ..... 776
  - Decoding method based on quadratic equations ... 781
  - decoding, decoding problem ..... 777
  - decoef ..... 891
  - decomposeVecField ..... 943
  - decomposition ..... 938
  - decomposition of modules ..... 821
  - decomposition, numerical ..... 842
  - def ..... 77
  - def declarations ..... 77
  - default argument ..... 52
  - defined ..... 167
  - defineHomogeneous ..... 936
  - Definition of a user defined type ..... 133
  - defl ..... 842
  - deform ..... 868
  - deform.lib ..... 861
  - deform\_lib ..... 861
  - Deformations ..... 733
  - Deformations, T1 and T2 ..... 731
  - defRes, option ..... 232
  - defring ..... 803
  - defringp ..... 803
  - defrings ..... 803
  - deg ..... 168
  - Deg ..... 931
  - degBound ..... 297
  - degree ..... 169, 309
  - Degree ..... 931
  - degree lexicographical ordering ..... 761
  - degree of a polynomial ..... 309
  - degree reverse lexicographical ordering ..... 761
  - degreeDivisor ..... 839
  - degreeFormalDivisor ..... 840
  - degreepart ..... 877
  - DegreePure ..... 841
  - Degrees ..... 931
  - delete ..... 102, 169, 204
  - deleteGenerator ..... 438
  - deleteSublist ..... 790
  - deligne number ..... 811
  - delta ..... 865
  - Delta ..... 848
  - DeltaList ..... 848
  - deltaLoc ..... 825
  - Demo mode ..... 27
  - denom\_list ..... 271
  - denominator ..... 170
  - depth ..... 815
  - Depth ..... 719
  - depthIdeal ..... 822
  - deRham.lib ..... 838
  - deRham\_lib ..... 838
  - deRhamCohom ..... 427
  - deRhamCohomIdeal ..... 428
  - deRhamCohomology ..... 838
  - derivate ..... 873
  - derivationAdd ..... 923
  - derivationCheckList ..... 923
  - derivationConstructor ..... 923
  - derivationContraction ..... 923
  - derivationContractionGen ..... 923

|                            |     |                              |                  |
|----------------------------|-----|------------------------------|------------------|
| derivationEqu              | 923 | diformNeq                    | 922              |
| derivationEval             | 923 | diformPrint                  | 922              |
| derivationFromList         | 923 | diformSub                    | 922              |
| derivationFromPoly         | 923 | diformToString               | 922              |
| derivationLie              | 924 | diformUnivDer                | 923              |
| derivationMul              | 923 | diffRat                      | 534              |
| derivationNeg              | 923 | difpoly2tex                  | 891              |
| derivationNeq              | 923 | dim                          | 171              |
| derivationPrint            | 923 | dim (plural)                 | 331              |
| derivationSub              | 923 | dim_slocus                   | 868              |
| derivationToString         | 923 | dimension                    | 902              |
| desingularization          | 927 | dimensionOfLocalization      | 927              |
| det                        | 170 | dimGradedPart                | 852              |
| det_B                      | 808 | dimH                         | 852              |
| detadj                     | 866 | dimMon                       | 820              |
| determinacy                | 868 | dimStack                     | 851              |
| determinecenter            | 845 | direct_boolean_poly          | 919              |
| detropicalise              | 912 | direct_from_boolean_poly     | 919              |
| develop                    | 865 | disc                         | 892              |
| Di Biase-Urbanke algorithm | 774 | discr                        | 869              |
| diag                       | 806 | discrepancy                  | 848              |
| diag_test                  | 808 | discrim                      | 795              |
| diagInvariants             | 826 | disp_zdd                     | 919              |
| diagonalizeMatrixSimul     | 943 | DISPLAY environment variable | 17               |
| diagonalizeVecField        | 943 | displayCohom                 | 852              |
| diagonalizeVecFieldLin     | 943 | displayHNE                   | 865              |
| DifConsLCSets              | 796 | displayInvariants            | 865              |
| diff                       | 171 | displayMultsequence          | 865              |
| diffAlgebra                | 921 | displaypfd                   | 939              |
| diffAlgebraChangeOrd       | 922 | displaypfd_long              | 939              |
| diffAlgebraGens            | 921 | displayPuiseuxExpansion      | 911              |
| diffAlgebraListGen         | 922 | displayTropicalLifting       | 911              |
| diffAlgebraStructure       | 921 | distributed computing        | 797, 802, 805    |
| diffAlgebraUnivDerIdeal    | 922 | Distributed computing        | 797              |
| differential algebra       | 924 | div                          | 74, 84, 119, 308 |
| differential forms         | 924 | dividelist                   | 846              |
| differentials              | 924 | divideUnits                  | 890              |
| diform.lib                 | 921 | division                     | 172              |
| diform_lib                 | 921 | division (plural)            | 332              |
| diformAdd                  | 922 | divisor                      | 897              |
| diformCoef                 | 922 | divisorplus                  | 839              |
| diformDeg                  | 922 | divisors.lib                 | 838              |
| diformDiff                 | 923 | divisors_lib                 | 838              |
| diformDiv                  | 922 | DLoc                         | 417              |
| diformEqu                  | 922 | DLoc0                        | 418              |
| diformFromPoly             | 922 | Dlocalization                | 520              |
| diformGenToString          | 922 | dmod.lib                     | 395              |
| diformHomogDecomp          | 922 | dmod_lib                     | 395              |
| diformIsBigger             | 922 | dmodAction                   | 531              |
| diformIsGen                | 922 | dmodActionRat                | 532              |
| diformIsHomog              | 923 | dmodapp.lib                  | 414              |
| diformIsHomogDeg           | 923 | dmodapp_lib                  | 414              |
| diformIsSmaller            | 922 | dmodGeneralAssumptionCheck   | 528              |
| diformListCont             | 923 | dmodideal.lib                | 440              |
| diformListSort             | 923 | dmodideal_lib                | 440              |
| diformMul                  | 922 | dmodloc.lib                  | 519              |
| diformNeg                  | 922 | dmodloc_lib                  | 519              |

- |                                         |     |                                       |          |
|-----------------------------------------|-----|---------------------------------------|----------|
| dmoddoublext                            | 546 | Elimination                           | 709      |
| dmodvar.lib                             | 446 | elimlinearpart                        | 877      |
| dmodvar.lib                             | 446 | elimpart                              | 877      |
| Documentation Tool                      | 62  | elimpartanyr                          | 877      |
| double                                  | 897 | elimrep                               | 845      |
| doubleExt                               | 603 | elimRing                              | 812      |
| downloading                             | 953 | elimWeight                            | 557      |
| dp, global ordering                     | 761 | elliptic curves                       | 813      |
| Dp, global ordering                     | 761 | ellipticAdd                           | 894      |
| drawNewtonSubdivision                   | 911 | ellipticAllPoints                     | 894      |
| drawTropicalCurve                       | 911 | ellipticcovers.lib                    | 812      |
| ds, local ordering                      | 761 | ellipticcovers.lib                    | 812      |
| Ds, local ordering                      | 761 | ellipticMult                          | 894      |
| DsingularLocus                          | 523 | ellipticNF                            | 907      |
| dsum                                    | 806 | ellipticNFDB                          | 907      |
| dual_code                               | 886 | ellipticRandomCurve                   | 894      |
| dualCone                                | 902 | ellipticRandomPoint                   | 894      |
| dualConic                               | 912 | else                                  | 291      |
| dualPart                                | 837 | Emacs                                 | 22       |
| dualPartition                           | 851 | Emacs, a quick guide                  | 23       |
| dualPolytope                            | 904 | Emacs, customization of Singular mode | 27       |
| dualSheaf                               | 850 | Emacs, editing Singular input files   | 28       |
| dump                                    | 173 | Emacs, important commands             | 29       |
| dyckGroup1                              | 646 | Emacs, overview                       | 23       |
| dyckGroup2                              | 646 | Emacs, running Singular under         | 25       |
| dyckGroup3                              | 647 | Emacs, Singular demo mode             | 27       |
| Dynamic loading                         | 71  | Emacs, user interface                 | 22       |
| Dynamic modules                         | 701 | Emaxcont                              | 845      |
| <b>E</b>                                |     |                                       |          |
| ecart                                   | 897 | embedMat                              | 577      |
| echo                                    | 298 | emptyFan                              | 903      |
| ECoef                                   | 845 | encode                                | 887      |
| ECPP                                    | 894 | endvfil                               | 863      |
| Edatalist                               | 845 | ensureLeftNcfrac                      | 539      |
| Editing input                           | 18  | ensureRightNcfrac                     | 541      |
| Editing SINGULAR input files with Emacs | 28  | Enumerative geometry                  | 851      |
| eexgcdN                                 | 893 | envelop                               | 794      |
| effective                               | 839 | envelope                              | 334      |
| egcdMain                                | 883 | enveloping algebra                    | 271, 366 |
| ehrhartPolynomialCoeff                  | 907 | environment variable, DISPLAY         | 17       |
| ehrhartRing                             | 826 | environment variables                 | 953      |
| eigenvals                               | 808 | EOrdlist                              | 845      |
| eigenvalue                              | 242 | Equal                                 | 841      |
| eigenvalues                             | 834 | equalJinI                             | 855      |
| elemSymmId                              | 798 | equalMultiDeg                         | 936      |
| elim                                    | 812 | equations                             | 902      |
| elim.lib                                | 812 | equidim                               | 829      |
| elim.lib                                | 812 | equidimMax                            | 829      |
| elim1                                   | 812 | equidimMaxEHV                         | 829      |
| elim2                                   | 812 | equidimZ                              | 830      |
| eliminate                               | 174 | equiRadical                           | 829      |
| eliminate (plural)                      | 333 | equising.lib                          | 862      |
| eliminateNC                             | 552 | equising.lib                          | 862      |
| elimination                             | 552 | equisingular Tjurina number           | 854      |
|                                         |     | Eresol                                | 845      |
|                                         |     | ERROR                                 | 175      |
|                                         |     | error recovery                        | 15       |
|                                         |     | errorInsert                           | 887      |

- |                                         |          |
|-----------------------------------------|----------|
| errormap                                | 925      |
| errorRand                               | 887      |
| esIdeal                                 | 862      |
| ESingular, CYGWIN                       | 26       |
| esStratum                               | 862      |
| eta                                     | 907      |
| euler                                   | 891      |
| EulerAff                                | 838      |
| eulerChProj                             | 837      |
| EulerProj                               | 838      |
| eval                                    | 174      |
| evalJacobianAtBox                       | 881      |
| evalPolyAtBox                           | 881      |
| evalPolyAtBox2                          | 877      |
| evaluate_reynolds                       | 872      |
| evaluateFormalDivisor                   | 839      |
| evaluatePDivisor                        | 840      |
| evaluatepfd                             | 938      |
| evaluateProduct                         | 906      |
| Evaluation of logical expressions       | 305      |
| evalutateIntegral                       | 813      |
| exactness recovery                      | 879      |
| example                                 | 175      |
| Examples                                | 691      |
| Examples of ring declarations           | 31       |
| Examples of use of Letterplace          | 613      |
| Examples of use of Letterplace over $Z$ | 614      |
| exclusionTest                           | 877      |
| execute                                 | 176      |
| exit                                    | 295      |
| exp2pt                                  | 891      |
| Experimental libraries                  | 920      |
| expo                                    | 892      |
| export                                  | 287      |
| exportNuminvs                           | 826      |
| exportto                                | 288      |
| expp                                    | 851      |
| expression list                         | 72, 313  |
| Ext                                     | 816      |
| Ext, computation of                     | 716      |
| ext-module                              | 597      |
| Ext_R                                   | 815      |
| extcurve                                | 886      |
| extdevelop                              | 865      |
| extendedTensor                          | 557      |
| extendGC                                | 794      |
| extendpoly                              | 794      |
| extending                               | 803      |
| extendWeyl                              | 528      |
| Exterior                                | 564      |
| exteriorBasis                           | 807      |
| exteriorPower                           | 807      |
| extgcd                                  | 176      |
| Extra weight vector                     | 764      |
| extractS                                | 446      |
| <b>F</b>                                |          |
| faceContaining                          | 902      |
| facets                                  | 902      |
| facetVertexLatticeDistances             | 907      |
| facetWidth                              | 908      |
| facetWidths                             | 908      |
| facFirstShift                           | 489      |
| facFirstWeyl                            | 486      |
| facGBIdeal                              | 828      |
| facShift                                | 488      |
| facstd                                  | 177      |
| facSubWeyl                              | 487      |
| factmodd                                | 177      |
| factor                                  | 237      |
| factorgroup                             | 935      |
| factorH                                 | 791      |
| factorial                               | 791      |
| factorization                           | 237, 809 |
| Factorization                           | 720      |
| factorize                               | 178, 237 |
| factorLenstraECM                        | 894      |
| factorMain                              | 883      |
| factory                                 | 1        |
| facvar                                  | 801      |
| facWeyl                                 | 485      |
| FamElementsAtEnvCompPoints              | 795      |
| fan                                     | 137, 908 |
| fanViaCones                             | 903      |
| farey                                   | 179      |
| fareypoly                               | 814      |
| fastelim                                | 877      |
| fastExpt                                | 891      |
| fastHC, option                          | 230      |
| fetch                                   | 180      |
| fetch (plural)                          | 335      |
| fetchall                                | 803      |
| Feynman graph                           | 813      |
| ffmodStd                                | 815      |
| ffmodstd.lib                            | 813      |
| ffmodstd.lib                            | 813      |
| ffsolve                                 | 875      |
| ffsolve.lib                             | 875      |
| ffsolve.lib                             | 875      |
| fglm                                    | 181, 267 |
| fglm_solve                              | 878      |
| fglmquot                                | 182      |
| fibonacci                               | 791      |
| fibonacciGroup                          | 647      |
| field                                   | 113      |
| file, singularrc                        | 22       |
| filecmd                                 | 182      |
| filtration                              | 597      |
| finalcases                              | 801      |
| finalCharts                             | 846      |
| find                                    | 182      |
| find_first_primitive_root               | 894      |



- find\_index . . . . . 895
  - findAuto . . . . . 453
  - findifs.lib . . . . . 890
  - findifs\_example . . . . . 891
  - findifs\_lib . . . . . 890
  - findimAlgebra . . . . . 564
  - findInvo . . . . . 451
  - findInvoDiag . . . . . 452
  - findOrientedBoundary . . . . . 907
  - findTorsion . . . . . 889
  - finduni . . . . . 183
  - findvars . . . . . 877
  - findZeroPoly . . . . . 940
  - finite field . . . . . 113, 875
  - Finite fields . . . . . 707
  - finiteDiagInvariants . . . . . 826
  - finitediff.lib . . . . . 924
  - finitediff\_lib . . . . . 924
  - finitely presented algebra . . . . . 629, 634, 649, 653, 665
  - finitely presented algebra, standard finitely presented algebra . . . . . 625
  - finitely presented group . . . . . 634
  - finitenessTest . . . . . 810
  - finiterep . . . . . 875
  - finvar.lib . . . . . 871
  - finvar\_lib . . . . . 871
  - first index is 1 . . . . . 307
  - First steps . . . . . 6
  - firstoct . . . . . 882
  - fitting . . . . . 816
  - Fitzgerald-Lax method . . . . . 780
  - fixedPoints . . . . . 851
  - fl2poly . . . . . 437
  - flatten . . . . . 806
  - flatteningStrat . . . . . 816
  - flintQ . . . . . 183
  - Float . . . . . 184
  - Flow control . . . . . 47
  - for . . . . . 290
  - Formal Checker . . . . . 62
  - formal linear representations . . . . . 669
  - formaldivisorplus . . . . . 839
  - Formatting output . . . . . 699
  - forward . . . . . 924
  - fourier . . . . . 434
  - fouriersystem . . . . . 925
  - fpadim.lib . . . . . 629
  - fpadim\_lib . . . . . 629
  - fpalgebras.lib . . . . . 634
  - fpalgebras\_lib . . . . . 634
  - fpaprops.lib . . . . . 649
  - fpaprops\_lib . . . . . 649
  - fPiece . . . . . 942
  - fprintf . . . . . 184, 785
  - fracStatus . . . . . 582
  - framed sheaves . . . . . 926
  - frandwalk . . . . . 940
  - free associative algebra . . . . . 634, 653
  - free associative algebra, tensor algebra . . . . . 625
  - Free associative algebras . . . . . 625
  - free noncommutative Groebner basis . . . . . 653
  - Free resolution . . . . . 712
  - Free resolution, graded . . . . . 715
  - freeAlgebra (letterplace) . . . . . 619
  - freegb.lib . . . . . 652
  - freegb\_lib . . . . . 652
  - freemodule . . . . . 186
  - freeModule . . . . . 931
  - freeModule2Module . . . . . 932
  - freerank . . . . . 798
  - fres . . . . . 186
  - from\_boolean\_constant . . . . . 919
  - from\_boolean\_ideal . . . . . 919
  - from\_boolean\_poly . . . . . 919
  - from\_boolean\_set . . . . . 919
  - frwalk . . . . . 187
  - fullDimImages . . . . . 905
  - fullFan . . . . . 903
  - fullSerreRelations . . . . . 635
  - fullSpace . . . . . 901
  - FultonA . . . . . 838
  - Functionality and release notes of Letterplace . . . . . 617
  - Functions . . . . . 154
  - Functions (letterplace) . . . . . 619
  - further\_hn\_proc . . . . . 865
  - furtherInvar . . . . . 873
  - Future benefits of Singular 4 . . . . . 952
  - fVector . . . . . 903
  - fVectorP . . . . . 907
  - fwalk . . . . . 815
- ## G
- G-algebra . . . . . 360, 514
  - G-algebra, setup . . . . . 361
  - G\_a -Invariants . . . . . 744
  - galois field . . . . . 113
  - Gamma . . . . . 845
  - Gauss-Manin connection . . . . . 769, 866
  - Gauss-Manin system . . . . . 863, 864
  - gauss\_col . . . . . 806
  - gauss\_nf . . . . . 808
  - gauss\_row . . . . . 806
  - gaussColWithoutPerm . . . . . 880
  - gaussred . . . . . 808
  - gaussred\_pivot . . . . . 808
  - gaussRowWithoutPerm . . . . . 880
  - GBsolve . . . . . 875
  - GBWeight . . . . . 419
  - gcd . . . . . 188
  - gcddivisor . . . . . 897
  - gcdMon . . . . . 819
  - Gelfand-Kirillov dimension . . . . . 649
  - Gelfand-Kirillov dimension, G-Algebra . . . . . 518

- gen . . . . . 188
- General command syntax . . . . . 41
- General concepts . . . . . 15
- general error-locator polynomial . . . . . 778
- general neron . . . . . 927
- General purpose . . . . . 787
- General syntax of a ring declaration . . . . . 32
- general weighted lexicographical ordering . . . . . 761
- general weighted reverse lexicographical ordering . . 761
- general.lib . . . . . 790
- general.lib . . . . . 790
- Generalized Hilbert Syzygy Theorem . . . . . 363
- Generalized Newton identities . . . . . 779
- generalOrder . . . . . 898
- generateG . . . . . 894
- generatorsOfLinealitySpace . . . . . 902
- generatorsOfSpan . . . . . 902
- genericid . . . . . 802
- genericity . . . . . 889
- genericmat . . . . . 806
- genMDSMat . . . . . 887
- genoutput . . . . . 846
- genSymId . . . . . 833
- genus . . . . . 824
- Geometric genus . . . . . 844
- Geometric Invariant Theory . . . . . 746
- German Umlaute . . . . . 304
- getArguments . . . . . 805
- getCommand . . . . . 805
- getCone . . . . . 903
- getcores . . . . . 803
- getdump . . . . . 189
- getenv . . . . . 271
- getGradingGroup . . . . . 934
- getLattice . . . . . 934
- getLinearForms . . . . . 902
- getMaxPoints . . . . . 929
- getModuleGrading . . . . . 935
- getMultiplicity . . . . . 902
- getOneVar . . . . . 825
- getRelations . . . . . 879
- getRelationsRadical . . . . . 880
- getResult . . . . . 805
- getSmallest . . . . . 825
- getState . . . . . 805
- getStringpfd . . . . . 939
- getStringpfd\_indexed . . . . . 939
- Getting started . . . . . 6
- getVariableWeights . . . . . 934
- getWitnessSet . . . . . 880
- gfan.lib . . . . . 901
- gfan.lib . . . . . 901
- gfanlib . . . . . 1
- GIT-Fans . . . . . 746
- gitCone . . . . . 906
- GITcone . . . . . 905
- GITfan . . . . . 905
- gitfan.lib . . . . . 904
- gitfan.lib . . . . . 904
- GITfanFromOrbitCones . . . . . 905
- GITfanParallel . . . . . 905
- GITfanParallelSymmetric . . . . . 906
- GITfanSymmetric . . . . . 906
- GKdim . . . . . 457
- gkdim.lib . . . . . 457
- gkdim.lib . . . . . 457
- GKExp . . . . . 518
- GKZfan . . . . . 905
- GKZsystem . . . . . 475
- global . . . . . 155
- global Bernstein-Sato ideal . . . . . 441
- global Bernstein-Sato polynomial . . . . . 371, 396
- global Bernstein-Sato polynomial for variety . . . . . 447
- global homological dimension . . . . . 649
- Global orderings . . . . . 761
- globalSections . . . . . 839
- GMP . . . . . 1, 953
- gmscoeffs . . . . . 863
- gmsnf . . . . . 863
- gmspoly.lib . . . . . 864
- gmspoly.lib . . . . . 864
- gmsring . . . . . 862
- gmssing.lib . . . . . 862
- gmssing.lib . . . . . 862
- GND.lib . . . . . 927
- GND.lib . . . . . 927
- Goettsche's formula . . . . . 926
- goettsche.lib . . . . . 925
- goettsche.lib . . . . . 925
- GoettscheF . . . . . 926
- good basis . . . . . 863, 864
- goodBasis . . . . . 864
- gorensteinIndex . . . . . 908
- gorensteinVector . . . . . 908
- graal.lib . . . . . 926
- graal.lib . . . . . 926
- graalMixed . . . . . 927
- graded algebra . . . . . 921
- Graded commutative algebras . . . . . 611
- graded Hilbert series . . . . . 665
- graded modules, graded homomorphisms, syzygies . . . . . 927
- graded modules, handling of . . . . . 715
- graded Weyl algebra . . . . . 371
- graded-module, graded-resolution, homogenous-matrix, R-module-homomorphism . . . . . 930
- gradedModules.lib . . . . . 927
- gradedModules.lib . . . . . 927
- gradeNumber . . . . . 600
- gradiator . . . . . 936
- graphics.lib . . . . . 883
- graphics.lib . . . . . 883
- graphviz . . . . . 1

- Grassmannian . . . . . 850  
 graver4ti2 . . . . . 833  
 grconcat . . . . . 929  
 grdeg . . . . . 928  
 Greuel, Gert-Martin . . . . . 3  
 grgens . . . . . 928  
 groebner . . . . . 928  
 grlift . . . . . 928  
 grlifting . . . . . 929  
 grlifting2 . . . . . 929  
 grlifting3 . . . . . 929  
 grneg . . . . . 929  
 Grob1Levels . . . . . 796  
 grobcov . . . . . 793  
 grobcov.lib . . . . . 791  
 grobcov\_lib . . . . . 791  
 grobj . . . . . 928  
 groebner . . . . . 189, 702, 785  
 Groebner Bases . . . . . 702  
 Groebner bases for two-sided ideals in free associative  
   algebras . . . . . 627  
 Groebner bases in free associative algebras . . . . . 627  
 Groebner bases in G-algebras . . . . . 361  
 Groebner bases, decodeGB . . . . . 888  
 Groebner bases, slim . . . . . 706  
 groebner basis computations . . . . . 190  
 Groebner basis conversion . . . . . 704  
 Groebner fan . . . . . 908  
 Groebner-Shirshov bases . . . . . 627  
 Groebner-Shirshov bases in free associative algebras  
   . . . . . 627  
 groebnerComplex . . . . . 913  
 groebnerCone . . . . . 913  
 groebnerFan . . . . . 913  
 Groebnerwalk . . . . . 815, 933, 940, 942  
 Gromov-Witten invariants . . . . . 813  
 Gromov-Witten invariants. . . . . 851  
 gromovWitten . . . . . 813  
 grorder . . . . . 928  
 ground field . . . . . 113  
 group\_reynolds . . . . . 871  
 groupActionOnHashes . . . . . 905  
 groupActionOnQImage . . . . . 905  
 growth of algebra . . . . . 649  
 grpower . . . . . 928  
 grpres . . . . . 928  
 grprod . . . . . 928  
 grrange . . . . . 929  
 gres . . . . . 928  
 grndmap . . . . . 929  
 grndmap2 . . . . . 929  
 grndmat . . . . . 929  
 grshift . . . . . 928  
 grsum . . . . . 928  
 grsyz . . . . . 928  
 grtest . . . . . 928  
 grtranspose . . . . . 928  
 grtranspose1 . . . . . 928  
 grtwist . . . . . 928  
 grtwists . . . . . 928  
 grview . . . . . 928  
 grwalk.lib . . . . . 815  
 grwalk\_lib . . . . . 815  
 grzero . . . . . 928  
 GTZmod . . . . . 821  
 GTZopt . . . . . 821  
 gwalk . . . . . 815  
 Gweights . . . . . 559
- ## H
- h\_increasing\_knapsack . . . . . 896  
 H2basis . . . . . 866  
 Hamburger-Noether expansion . . . . . 854, 865  
 Handling graded modules . . . . . 715  
 hardware platform . . . . . 273  
 hasAlgExtensionCoefficient . . . . . 804  
 hasCommutativeVars . . . . . 804  
 hasGlobalOrdering . . . . . 804  
 hashesToFan . . . . . 906  
 hashToCone . . . . . 906  
 hasLeftDenom . . . . . 536  
 hasMixedOrdering . . . . . 804  
 hasNumericCoeffs . . . . . 804  
 hasRightDenom . . . . . 537  
 hasTransExtensionCoefficient . . . . . 804  
 Hcode . . . . . 856  
 headStand . . . . . 807  
 heightZ . . . . . 830  
 help . . . . . 191  
 help browsers . . . . . 16  
 help browsers, dummy . . . . . 16  
 help browsers, emacs . . . . . 16  
 help browsers, html . . . . . 16  
 help browsers, setting command to use . . . . . 17  
 help browsers, setting the . . . . . 270  
 Help string . . . . . 53  
 help, accessing over the net . . . . . 19  
 help, online help system . . . . . 15  
 Hensel . . . . . 177  
 hermiteNormalForm . . . . . 936  
 hess.lib . . . . . 840  
 hess.lib . . . . . 840  
 hessenberg . . . . . 271, 808  
 HHnormalForm . . . . . 859  
 highcorner . . . . . 192  
 hilb . . . . . 193  
 Hilbert function . . . . . 269, 514, 766  
 Hilbert polynomial . . . . . 514  
 Hilbert scheme . . . . . 926  
 Hilbert series . . . . . 514, 629, 766  
 Hilbert-driven GB algorithm . . . . . 704  
 hilbert4ti2 . . . . . 833

- hilbertBasis . . . . . 908
  - HilbertClassPoly . . . . . 892
  - hilbertSeries . . . . . 936
  - HilbertSeries . . . . . 874
  - HilbertWeights . . . . . 874
  - hilbPoly . . . . . 799
  - hilbvec . . . . . 877
  - hnexpansion . . . . . 865
  - hnoether.lib . . . . . 864
  - hnoether.lib . . . . . 864
  - holonomic rank . . . . . 519
  - holonomicRank . . . . . 522
  - hom . . . . . 933
  - Hom . . . . . 816
  - hom\_kernel . . . . . 816
  - HomJJ . . . . . 824
  - homog . . . . . 194
  - homog\_part . . . . . 851
  - homog\_parts . . . . . 851
  - homogeneitySpace . . . . . 913
  - homogfacFirstQWeyl . . . . . 491
  - homogfacFirstQWeyl.all . . . . . 510
  - homogfacNthQWeyl . . . . . 490
  - homogfacNthQWeyl.all . . . . . 492
  - homogfacNthWeyl . . . . . 490
  - homolog.lib . . . . . 815
  - homolog.lib . . . . . 815
  - homology . . . . . 816
  - homomorphism . . . . . 932
  - Hosten-Sturmfels algorithm . . . . . 774
  - How to enter and exit . . . . . 15
  - How to use this manual . . . . . 4
  - howto, download . . . . . 953
  - howto, install on Macintosh . . . . . 955
  - howto, install on Unix . . . . . 954
  - howto, install on Windows . . . . . 955
  - hres . . . . . 195
  - hStarVector . . . . . 908
  - html, default help . . . . . 16
  - hVector . . . . . 908
  - hybrid algorithms . . . . . 879
  - hyperel.lib . . . . . 896
  - hyperel.lib . . . . . 896
  - hyperplane arrangement . . . . . 396
  - Hypersurface singularities, classification of . . . . . 741
  - hypersurface singularity . . . . . 866
- I**
- id . . . . . 930
  - id2mod . . . . . 798
  - ideal . . . . . 78
  - ideal (plural) . . . . . 313
  - ideal declarations . . . . . 78
  - ideal declarations (plural) . . . . . 313
  - ideal expressions . . . . . 78
  - ideal expressions (plural) . . . . . 314
  - Ideal membership . . . . . 766
  - ideal operations . . . . . 79
  - ideal operations (plural) . . . . . 315
  - ideal related functions . . . . . 80
  - ideal related functions (plural) . . . . . 316
  - ideal, toric . . . . . 773
  - ideals . . . . . 309
  - idealsimplify . . . . . 855
  - idealSplit . . . . . 877
  - identifier . . . . . 310
  - Identifiers, syntax of . . . . . 44
  - identifyvar . . . . . 845
  - if . . . . . 291
  - image . . . . . 932
  - image\_of\_variety . . . . . 873
  - ImageGroup . . . . . 874
  - imageLattice . . . . . 935
  - ImageVariety . . . . . 874
  - imap . . . . . 195
  - imap (plural) . . . . . 336
  - Imap, option . . . . . 232
  - imapall . . . . . 803
  - impart . . . . . 196
  - Implemented algorithms . . . . . 36
  - importfrom . . . . . 291
  - iMult . . . . . 825
  - IN . . . . . 206
  - inCenter . . . . . 388
  - inCentralizer . . . . . 388
  - Incl . . . . . 840
  - indepSet . . . . . 196
  - Index . . . . . 956
  - indexed names . . . . . 42, 44
  - indices, multi . . . . . 44
  - indSet . . . . . 821
  - inequalities . . . . . 902
  - infinitely presented algebra . . . . . 665
  - info . . . . . 17
  - Info string . . . . . 55
  - inForm . . . . . 422
  - infRedTail, option . . . . . 231
  - iniD . . . . . 846
  - init\_debug . . . . . 856
  - initial . . . . . 913
  - initial form . . . . . 371
  - initial ideal . . . . . 371
  - initial ideal approach . . . . . 371, 447
  - initialForm . . . . . 912
  - initialIdeal . . . . . 912
  - initialIdealW . . . . . 421
  - initialMalgrange . . . . . 420
  - injective\_knapsack . . . . . 896
  - inout.lib . . . . . 796
  - inout.lib . . . . . 796
  - input . . . . . 48
  - insert . . . . . 102, 197

- insertCone . . . . . 903
- insertGenerator . . . . . 438
- instructions, downloading . . . . . 953
- instructions, Macintosh installation . . . . . 955
- instructions, Unix installation . . . . . 954
- instructions, Windows installation . . . . . 955
- inSubring . . . . . 809
- int . . . . . 82
- int declarations . . . . . 82
- int expressions . . . . . 83
- int operations . . . . . 84
- int related functions . . . . . 85
- intclMonIdeal . . . . . 826
- intclToricRing . . . . . 826
- integer division . . . . . 308
- integer programming . . . . . 775
- integral . . . . . 850
- integral closure . . . . . 826
- integralBasis . . . . . 817
- integralbasis.lib . . . . . 816
- integralbasis.lib . . . . . 816
- integralIdeal . . . . . 425
- integralModule . . . . . 426
- integralSection . . . . . 935
- integration of D-module . . . . . 415
- Interactive use . . . . . 15
- InterDiv . . . . . 846
- interface, Emacs . . . . . 22
- interiorLatticePoints . . . . . 908
- internalfunctors . . . . . 856
- interpolate . . . . . 878
- interpolation . . . . . 198
- interpret . . . . . 933
- interpretElem . . . . . 933
- interpretInv . . . . . 933
- interpretList . . . . . 933
- interred . . . . . 199
- Interrupting SINGULAR . . . . . 18
- intersect . . . . . 199
- intersect (plural) . . . . . 337
- intersectElim, option . . . . . 232
- intersection . . . . . 865
- Intersection theory . . . . . 851
- intersectionDiv . . . . . 848
- IntersectionMatrix . . . . . 861
- intersectionSet . . . . . 941
- intersectionValRingIdeals . . . . . 826
- intersectionValRings . . . . . 826
- intersectLattices . . . . . 935
- intersectLists . . . . . 921
- intersectMon . . . . . 819
- intersectOrbitsWithMovingCone . . . . . 905
- intersectpar . . . . . 795
- intersectSyz, option . . . . . 232
- IntersectWithSub . . . . . 479
- intersectZ . . . . . 830
- interval arithmetic . . . . . 882
- interval.lib . . . . . 876
- interval.lib . . . . . 876
- intervalmatrixInit . . . . . 876
- intInverse . . . . . 935
- intmat . . . . . 88
- intmat declarations . . . . . 88
- intmat expressions . . . . . 88
- intmat operations . . . . . 90
- intmat related functions . . . . . 91
- intmat type cast . . . . . 89
- intmat2mons . . . . . 827
- intPart . . . . . 893
- intprog.lib . . . . . 817
- intprog.lib . . . . . 817
- intRank . . . . . 935
- Introduction . . . . . 4
- intRoot . . . . . 893
- intRoots . . . . . 436
- intStrategy, option . . . . . 231
- intvec . . . . . 91
- intvec declarations . . . . . 91
- intvec expressions . . . . . 91
- intvec operations . . . . . 92
- intvec related functions . . . . . 93
- invar . . . . . 875
- invar.lib . . . . . 874
- invar.lib . . . . . 874
- Invariant theory . . . . . 871
- Invariant Theory . . . . . 744
- invariant\_algebra\_perm . . . . . 871
- invariant\_algebra\_reynolds . . . . . 871
- invariant\_basis . . . . . 872
- invariant\_basis\_reynolds . . . . . 872
- invariant\_ring . . . . . 871
- invariant\_ring\_random . . . . . 871
- InvariantQ . . . . . 874
- invariantRing . . . . . 873
- InvariantRing . . . . . 874
- invariants . . . . . 811, 865
- Invariants of a finite group . . . . . 745
- Invariants of plane curve singularities . . . . . 736
- inverse . . . . . 807
- inverse of a matrix via its LU-decomposition . . . . . 213
- inverse\_B . . . . . 807
- inverse\_L . . . . . 807
- inverse\_modulus . . . . . 895
- inverseFourier . . . . . 434
- invertAlgebraMorphism . . . . . 943
- invertBirMap . . . . . 844
- invertLeftFraction . . . . . 592
- invertNfrac . . . . . 543
- invertNumberMain . . . . . 883
- involut.lib . . . . . 450
- involut.lib . . . . . 450
- involution . . . . . 455
- invunit . . . . . 866
- iostruct . . . . . 889

|                           |          |                          |     |
|---------------------------|----------|--------------------------|-----|
| irred_secondary_char0     | 872      | isIntegralSurjective     | 935 |
| irred_secondary_no_molien | 872      | isInvertibleLeftFraction | 592 |
| irreddecMon               | 820      | isInvertibleNcfrac       | 542 |
| irrRealizationDim         | 909      | isInvolution             | 456 |
| is_active                 | 868      | isirreducibleMon         | 820 |
| is_bijjective             | 810      | isLatticeEmpty           | 908 |
| is_cenBimodule            | 547      | isLocalizationRegular    | 927 |
| is_cenSubbimodule         | 547      | isLocallyFree            | 816 |
| is_ci                     | 868      | isMonomial               | 819 |
| is_complex                | 806      | isNC                     | 571 |
| is_fix_injective          | 896      | isNormal                 | 908 |
| is_h_injective            | 896      | isoncurve                | 896 |
| is_injective              | 810      | isOnCurve                | 894 |
| is_irred                  | 865      | isOneFraction            | 594 |
| is_is                     | 868      | isOneNcfrac              | 538 |
| is_nested                 | 822      | isOrderingShiftInvariant | 655 |
| is_NND                    | 865      | isOrigin                 | 902 |
| is_NP                     | 822      | isparam                  | 900 |
| is_prime                  | 895      | isPositive               | 936 |
| is_primitive_root         | 894      | isprimaryMon             | 820 |
| is_pure                   | 605      | isprimeMon               | 820 |
| is_reg                    | 868      | isPrimitiveSublattice    | 935 |
| is_regs                   | 869      | isPure                   | 903 |
| is_surjective             | 810      | isPureTensor             | 369 |
| is_zero                   | 798      | isQuotientRing           | 804 |
| isAface                   | 904      | isRational               | 413 |
| isAntiEndo                | 457      | isReflexive              | 908 |
| isartinianMon             | 820      | isReg                    | 816 |
| isBounded                 | 908      | isSB                     | 155 |
| isCanonical               | 908      | IsSCA                    | 575 |
| isCartan                  | 389      | isSimplicial             | 902 |
| isCentral                 | 570      | isSmooth                 | 908 |
| isCI                      | 155, 811 | isSublattice             | 935 |
| isCM                      | 155, 816 | isSubModule              | 804 |
| isCMcod2                  | 869      | isSubset                 | 941 |
| isCommutative             | 572      | isSuperset               | 941 |
| isCompatible              | 903      | isSymmetric              | 833 |
| isCompressed              | 908      | isTame                   | 864 |
| isDenom                   | 548      | isTerminal               | 908 |
| isElement                 | 941      | isTorsionFree            | 936 |
| isEqualDivisor            | 839      | isTwoSidedGB             | 370 |
| isEquising                | 862      | isuni                    | 899 |
| isFlat                    | 816      | isUpperTriangular        | 555 |
| isFreeAlgebra             | 653      | isVeryAmple              | 908 |
| isFsat                    | 432      | isWeyl                   | 572 |
| isFullSpace               | 902      | isZeroElement            | 936 |
| isgenericMon              | 820      | isZeroFraction           | 593 |
| isGorenstein              | 908      | isZeroNcfrac             | 537 |
| isGradedRingHomomorphism  | 934      | ivmatGaussian            | 881 |
| isGroup                   | 934      | ivmatGaussian2           | 877 |
| isGroupHomomorphism       | 934      | ivmatInit                | 881 |
| isHolonomic               | 411      | ivmatSet                 | 881 |
| isHomog                   | 155      |                          |     |
| isHomogeneous             | 936      |                          |     |
| ishyper                   | 896      |                          |     |
| isInS                     | 581      |                          |     |
| isInt                     | 439      |                          |     |

**J**

|                                |     |
|--------------------------------|-----|
| J-marked schemes               | 852 |
| J-marked schemes, Borel ideals | 853 |
| jacob                          | 200 |
| Jacobi                         | 893 |
| jacoblift                      | 866 |
| Jacobson                       | 890 |
| Jacobson form                  | 890 |
| Jacobson normal form           | 890 |
| Jacobson.lib                   | 889 |
| Jacobson.lib                   | 889 |
| janet                          | 201 |
| janet basis                    | 832 |
| jet                            | 201 |
| jInvariant                     | 911 |
| JMarkedScheme                  | 853 |
| JMBTest.lib                    | 852 |
| JMBTest.lib                    | 852 |
| JMSConst.lib                   | 853 |
| JMSConst.lib                   | 853 |
| jOft                           | 892 |
| jordan                         | 808 |
| jordanbasis                    | 808 |
| jordanmatrix                   | 808 |
| jordannf                       | 808 |
| jordanVecField                 | 943 |
| jungfib                        | 847 |
| jungnormal                     | 847 |
| jungresolve                    | 847 |
| JuReTopDim                     | 841 |
| JuReZeroDim                    | 841 |

**K**

|                                |     |
|--------------------------------|-----|
| K-basis                        | 629 |
| K-dimension                    | 629 |
| katsura                        | 798 |
| kbase                          | 203 |
| kbase (plural)                 | 337 |
| keepring                       | 293 |
| KeneshlouMatrixPresentation    | 928 |
| Ker                            | 932 |
| kerHom                         | 933 |
| kernel                         | 203 |
| Kernel of module homomorphisms | 724 |
| kernelLattice                  | 935 |
| kill                           | 204 |
| killall                        | 791 |
| killattrib                     | 204 |
| killTask                       | 805 |
| kmemory                        | 791 |
| kohom                          | 816 |
| kontrahom                      | 816 |
| koszul                         | 205 |
| KoszulHomology                 | 816 |
| Kronecker module               | 926 |

|                   |     |
|-------------------|-----|
| Kronecker product | 275 |
| KScoef            | 866 |
| KSconvert         | 866 |
| KSkcr             | 866 |
| kskernel.lib      | 866 |
| kskernel.lib      | 866 |
| KSlinear          | 866 |
| KSpencerKernel    | 870 |

**L**

|                                   |               |
|-----------------------------------|---------------|
| L, ordering                       | 765           |
| l-adic numbers                    | 36            |
| laguerre                          | 205           |
| laguerre_solve                    | 878           |
| lastvarGeneral                    | 898           |
| latex.lib                         | 884           |
| latex.lib                         | 884           |
| latticeArea                       | 907           |
| latticeBasis                      | 935           |
| latticeCodegree                   | 908           |
| latticeDegree                     | 908           |
| latticePoints                     | 908           |
| latticeVolume                     | 908           |
| laxfrT                            | 924           |
| laxfrX                            | 924           |
| lazy, option                      | 231           |
| lcm                               | 798           |
| lcmMon                            | 819           |
| lcmN                              | 893           |
| lcmofall                          | 845           |
| lead                              | 206           |
| leadcoef                          | 207           |
| leadexp                           | 207           |
| leadmonom                         | 208           |
| leadmonomial                      | 897           |
| Left and two-sided Groebner bases | 748           |
| left annihilator ideal            | 396, 441      |
| Left ideal membership (plural)    | 362           |
| Left normal form                  | 362           |
| leftInverse                       | 889           |
| leftKernel                        | 889           |
| leftOre                           | 584           |
| Leinartas                         | 938           |
| length                            | 881           |
| length, option                    | 231           |
| length2                           | 876           |
| LengthSym                         | 608           |
| LengthSymElement                  | 608           |
| letplaceGBasis                    | 657           |
| letterplace                       | 216, 246      |
| Letterplace                       | 613           |
| LETTERPLACE                       | 613           |
| Letterplace correspondence        | 628           |
| Letterplace Groebner basis        | 629, 649, 653 |
| Letterplace libraries             | 629           |

- LETTERPLACE libraries . . . . . 629
- Letterplace ring . . . . . 266
- Levels . . . . . 796
- lex\_solve . . . . . 878
- Lexicographic Groebner bases, computation of . . . . . 704
- lexicographical ordering . . . . . 761
- LIB . . . . . 208
- LIB commands . . . . . 57
- lib2doc . . . . . 62
- libcdd . . . . . 1
- libfac . . . . . 1
- libparse . . . . . 69
- Libraries . . . . . 54, 692
- Libraries in the SINGULAR Documentation . . . . . 55
- library . . . . . 919
- Library versioning . . . . . 951
- library, gitfan, GIT, geometric invariant theory, quotients . . . . . 906
- library, info string . . . . . 61
- library, polybori.lib . . . . . 919
- library, ringgb.lib . . . . . 940
- library, template . . . . . 58
- library, template.lib . . . . . 61
- LIBs . . . . . 785
- libSingular . . . . . 953
- Lie bracket . . . . . 330
- lieBracket . . . . . 658
- lift . . . . . 209, 366
- lift (letterplace) . . . . . 620
- lift (plural) . . . . . 338
- lift\_kbase . . . . . 862
- lift\_rel\_kb . . . . . 862
- liftenvelope . . . . . 368
- liftstd . . . . . 209
- liftstd (letterplace) . . . . . 621
- liftstd (plural) . . . . . 339
- likeIdeal . . . . . 929
- likelihood ideal . . . . . 929
- Limitations . . . . . 304
- linalg.lib . . . . . 807
- linalg.lib . . . . . 807
- linealityDimension . . . . . 902
- linealitySpace . . . . . 902
- linear algebra . . . . . 156, 213, 214
- Linear algebra . . . . . 805
- Linear code . . . . . 888
- linear interreduction . . . . . 371
- linear\_relations . . . . . 807
- LinearActionQ . . . . . 874
- LinearCombinationQ . . . . . 874
- linearCombinations . . . . . 392
- LinearizeAction . . . . . 874
- linearlyEquivalent . . . . . 839
- linearMapKernel . . . . . 391
- linearpart . . . . . 877
- linesHypersurface . . . . . 851
- link . . . . . 94, 700
- link declarations . . . . . 94
- link expressions . . . . . 94
- link related functions . . . . . 95
- linReduce . . . . . 378
- linReduceIdeal . . . . . 379
- linSyzSolve . . . . . 380
- list . . . . . 101
- list declarations . . . . . 101
- list expressions . . . . . 101
- list operations . . . . . 102
- list related functions . . . . . 103
- listvar . . . . . 210
- lll . . . . . 936
- LLL . . . . . 271
- load . . . . . 294
- Loading a library . . . . . 66
- loadLib, option . . . . . 232
- loadProc, option . . . . . 232
- local methods . . . . . 817
- local names . . . . . 54
- Local orderings . . . . . 761
- local rings, computing in . . . . . 695
- local weighted lexicographical ordering . . . . . 761
- local weighted reverse lexicographical ordering . . . . . 761
- localInvar . . . . . 873
- localization . . . . . 695
- localization of D-module . . . . . 415, 519
- localstd . . . . . 898
- locAtZero . . . . . 825
- locNormal . . . . . 817
- locnormal.lib . . . . . 817
- locnormal.lib . . . . . 817
- locStatus . . . . . 578
- locstd . . . . . 869
- locus . . . . . 794
- locusdg . . . . . 794
- locusto . . . . . 795
- log2 . . . . . 891
- logarithmic annihilator ideal . . . . . 396, 441
- logg . . . . . 851
- logHessian . . . . . 929
- Long coefficients . . . . . 696
- LOT algorithm . . . . . 396
- lp, global ordering . . . . . 761
- lpCalcSubstDegBound . . . . . 652
- lpDegBound . . . . . 653
- lpDivision . . . . . 654
- lpGBPres2Poly . . . . . 655
- lpGLDimBound . . . . . 651
- lpHilbert . . . . . 632
- lpIsPrime . . . . . 650
- lpIsSemiPrime . . . . . 650
- lpKDim . . . . . 631
- lpKDimCheck . . . . . 630
- lpMonomialBasis . . . . . 631
- lpNcgenCount . . . . . 654
- lpNoetherian . . . . . 649



- lprint . . . . . 796  
 lpSickleDim . . . . . 633  
 lpSubstitute . . . . . 651  
 lpVarBlockSize . . . . . 653  
 lres . . . . . 212  
 ls, local ordering . . . . . 761  
 lsum . . . . . 813  
 LU-decomposition of a matrix of numbers . . . . . 213  
 ludcomp . . . . . 213  
 luinverse . . . . . 213  
 lusolve . . . . . 214
- M**
- M, ordering . . . . . 762  
 m\_merkle\_hellman\_decryption . . . . . 895  
 m\_merkle\_hellman\_encryption . . . . . 895  
 m\_merkle\_hellman\_transformation . . . . . 895  
 Macdonald's formula . . . . . 926  
 MacdonaldF . . . . . 926  
 Macintosh installation . . . . . 955  
 magnitude . . . . . 891  
 makeDivisor . . . . . 839  
 makeFormalDivisor . . . . . 839  
 makeGraph . . . . . 813  
 makeGraphVE . . . . . 851  
 makeHeisenberg . . . . . 563  
 makeIdeal . . . . . 931  
 makeLetterplaceRing . . . . . 656  
 makeLinks . . . . . 942  
 makeMalgrange . . . . . 450  
 makeMatrix . . . . . 931  
 makeModElimRing . . . . . 577  
 makePDivisor . . . . . 840  
 makePoly . . . . . 939  
 makeQPoly . . . . . 942  
 makeQsl2 . . . . . 473  
 makeQsl3 . . . . . 473  
 makeQso3 . . . . . 472  
 makeSheaf . . . . . 850  
 makeUe6 . . . . . 470  
 makeUe7 . . . . . 470  
 makeUe8 . . . . . 471  
 makeUf4 . . . . . 469  
 makeUg2 . . . . . 469  
 makeUgl . . . . . 460  
 makeUsl . . . . . 459  
 makeUsl2 . . . . . 459  
 makeUso10 . . . . . 464  
 makeUso11 . . . . . 464  
 makeUso12 . . . . . 465  
 makeUso5 . . . . . 461  
 makeUso6 . . . . . 461  
 makeUso7 . . . . . 462  
 makeUso8 . . . . . 463  
 makeUso9 . . . . . 463  
 makeUsp1 . . . . . 465  
 makeUsp2 . . . . . 466  
 makeUsp3 . . . . . 467  
 makeUsp4 . . . . . 468  
 makeUsp5 . . . . . 468  
 makeVariety . . . . . 850  
 makeVector . . . . . 932  
 makeWeyl . . . . . 562  
 map . . . . . 103  
 map (plural) . . . . . 317  
 map declarations . . . . . 104  
 map declarations (plural) . . . . . 318  
 map expressions . . . . . 105  
 map expressions (plural) . . . . . 319  
 map operations . . . . . 105  
 map operations (plural) . . . . . 319  
 map related functions . . . . . 105  
 map related functions (plural) . . . . . 319  
 mapall . . . . . 803  
 mapIsFinite . . . . . 810  
 mappingcone . . . . . 929  
 mappingcone3 . . . . . 929  
 mapToRatNormCurve . . . . . 844  
 markov4ti2 . . . . . 833  
 mat\_rk . . . . . 808  
 mat2arr . . . . . 915  
 mat2carr . . . . . 915  
 matbil . . . . . 899  
 Mathematical background . . . . . 766  
 Mathematical background (letterplace) . . . . . 625  
 Mathematical background (plural) . . . . . 360  
 mathematical objects . . . . . 759  
 mathinit . . . . . 883  
 matmult . . . . . 899  
 matrix . . . . . 106  
 matrix declarations . . . . . 106  
 matrix diagonalization . . . . . 890  
 matrix expressions . . . . . 107  
 matrix operations . . . . . 108  
 Matrix orderings . . . . . 762  
 matrix related functions . . . . . 109  
 matrix type cast . . . . . 107  
 matrix.lib . . . . . 806  
 matrix.lib . . . . . 806  
 matrixExp . . . . . 842  
 matrixLog . . . . . 842  
 matrixpres . . . . . 929  
 matrixsystem . . . . . 925  
 matrixT1 . . . . . 869  
 max . . . . . 215, 785  
 Max . . . . . 867  
 maxabs . . . . . 900  
 maxcoef . . . . . 798  
 maxdeg . . . . . 798  
 maxdeg1 . . . . . 798  
 maxEord . . . . . 845  
 maxExp . . . . . 155  
 maxideal . . . . . 216

- maximalFace . . . . . 908
- maximalGroebnerCone . . . . . 913
- maximalValue . . . . . 908
- maximum . . . . . 892
- maximum likelihood estimate . . . . . 929
- Maximus . . . . . 853
- maxIntRoot . . . . . 530
- maxlike.lib . . . . . 929
- maxlike\_lib . . . . . 929
- Maxord . . . . . 845
- maxPoints . . . . . 929
- maxPointsProb . . . . . 929
- maxZeros . . . . . 842
- mdouble . . . . . 61
- mem, option . . . . . 232
- member . . . . . 921
- memberpos . . . . . 801
- membershipMon . . . . . 819
- memory . . . . . 216
- memory management . . . . . 216
- merkle\_hellman\_decryption . . . . . 896
- merkle\_hellman\_encryption . . . . . 895
- methods, hashtables . . . . . 930
- methods.lib . . . . . 930
- methods\_lib . . . . . 930
- midpoint . . . . . 924
- MillerRabin . . . . . 893
- milnor . . . . . 869
- Milnor code . . . . . 855
- Milnor number . . . . . 726
- milnorcode . . . . . 856
- milnornumber . . . . . 868
- min . . . . . 217, 786
- Min . . . . . 867
- minAssChar . . . . . 829
- minAssCharE . . . . . 829
- minAssGTZ . . . . . 829
- minAssGTZE . . . . . 829
- minAssZ . . . . . 830
- minbase . . . . . 218
- minbaseMon . . . . . 819
- mindeg . . . . . 798
- mindeg1 . . . . . 798
- mindist . . . . . 887
- minEcart . . . . . 897
- minimal display time, setting the . . . . . 274
- minimal representations . . . . . 669
- minimalAfaceOrbits . . . . . 905
- minimalAfaces . . . . . 905
- MinimalDecomposition . . . . . 874
- minimalFace . . . . . 908
- minimalOrbitConeOrbits . . . . . 905
- minimalValue . . . . . 908
- Minimum distance . . . . . 888
- Minimus . . . . . 853
- minIntRoot . . . . . 412
- minIntRoot2 . . . . . 530
- minipoly . . . . . 808
- minkowskiSum . . . . . 908
- minMult . . . . . 811
- MinMult . . . . . 901
- minor . . . . . 218
- minpoly . . . . . 298
- minres . . . . . 220
- minres (plural) . . . . . 340
- mirror symmetry . . . . . 813
- Miscellaneous libraries . . . . . 914
- mixed Hodge structure . . . . . 863, 864
- mod . . . . . 74, 84, 115, 119
- mod\_versal . . . . . 862
- mod2id . . . . . 798
- modality . . . . . 857
- modberlekampMassey . . . . . 814
- modBorder . . . . . 832
- modDec . . . . . 821
- moddiq.lib . . . . . 817
- moddiq\_lib . . . . . 817
- ModEqn . . . . . 867
- modfrWalk . . . . . 933
- modfWalk . . . . . 933
- modIntersect . . . . . 819
- modJanet . . . . . 832
- modNormal . . . . . 818
- modnormal.lib . . . . . 818
- modnormal\_lib . . . . . 818
- modNPos . . . . . 824
- modNpos\_test . . . . . 824
- modQuotient . . . . . 818
- modrationalInterpolation . . . . . 814
- modregCM . . . . . 824
- modrWalk . . . . . 933
- modSagbiAlg . . . . . 859
- modSat . . . . . 818
- modsatiety . . . . . 824
- modStd . . . . . 819
- modstd.lib . . . . . 818
- modstd\_lib . . . . . 818
- modSyz . . . . . 819
- modular . . . . . 797
- modular methods . . . . . 817, 818
- Modular techniques . . . . . 797
- modular.lib . . . . . 796
- modular\_lib . . . . . 796, 797
- module . . . . . 110
- module (plural) . . . . . 320
- module declarations . . . . . 110
- module declarations (plural) . . . . . 320
- module expressions . . . . . 110
- module expressions (plural) . . . . . 320
- module operations . . . . . 111
- module operations (plural) . . . . . 321
- module ordering c . . . . . 762
- module ordering C . . . . . 762
- Module orderings . . . . . 761

- module related functions . . . . . 111
  - module related functions (plural) . . . . . 321
  - module\_containment . . . . . 809
  - Modules and and their annihilator . . . . . 12
  - modules.lib . . . . . 930
  - modules.lib . . . . . 930
  - moduliSpace . . . . . 851
  - modulo . . . . . 220
  - modulo (plural) . . . . . 341
  - moduloSlim . . . . . 569
  - modVStd . . . . . 860
  - modVStd0 . . . . . 860
  - modWalk . . . . . 933
  - modwalk.lib . . . . . 933
  - modwalk.lib . . . . . 933
  - moebius . . . . . 916
  - molien . . . . . 871
  - mondim . . . . . 518
  - mondromy.lib . . . . . 866
  - mondromy.lib . . . . . 866
  - monitor . . . . . 221
  - monodromy . . . . . 863, 864
  - Monodromy . . . . . 866
  - monodromyB . . . . . 866
  - monoideal, dimension . . . . . 519
  - monomial . . . . . 221
  - monomial orderings . . . . . 695, 760
  - Monomial orderings . . . . . 760
  - monomial orderings introduction . . . . . 760
  - Monomial orderings on free algebras . . . . . 625
  - Monomial orderings, Term orderings . . . . . 34
  - monomial output . . . . . 300
  - monomial, write . . . . . 300
  - monomialabortstd . . . . . 917
  - monomialideal.lib . . . . . 819
  - monomialideal.lib . . . . . 819
  - monomialInIdeal . . . . . 529
  - monomialLcm . . . . . 897
  - monomials and precedence . . . . . 308
  - mons2intmat . . . . . 827
  - Mori dream spaces . . . . . 921
  - morsesplit . . . . . 856
  - movingCone . . . . . 905
  - mp\_res\_mat . . . . . 878
  - mplot . . . . . 883
  - mpresmat . . . . . 222
  - mprimdec.lib . . . . . 820
  - mprimdec.lib . . . . . 820
  - mrcgs . . . . . 801
  - mregular.lib . . . . . 821
  - mregular.lib . . . . . 821
  - mres . . . . . 222
  - mRes . . . . . 931
  - mres (plural) . . . . . 342
  - mstd . . . . . 223
  - msum . . . . . 61
  - mtriple . . . . . 61
  - mult . . . . . 224, 309
  - multarr2arr . . . . . 916
  - multarrMultRestrict . . . . . 916
  - multarrRestrict . . . . . 916
  - multBound . . . . . 298
  - multcol . . . . . 806
  - multdivisor . . . . . 839
  - multformaldivisor . . . . . 840
  - multi . . . . . 896
  - multi indices . . . . . 44
  - multi-graded Hilbert series . . . . . 665
  - multiDeg . . . . . 935
  - multiDegBasis . . . . . 936
  - multiDegGroebner . . . . . 936
  - multiDegModulo . . . . . 936
  - multiDegPartition . . . . . 936
  - multiDegResolution . . . . . 936
  - multiDegSyzygy . . . . . 936
  - multiDegTensor . . . . . 936
  - multiDegTor . . . . . 936
  - multidimensional\_knapsack . . . . . 895
  - multigrading, multidegree, multiweights,  
multigraded-homogeneous, integral linear algebra  
. . . . . 936
  - multigrading.lib . . . . . 934
  - multigrading.lib . . . . . 934
  - multipleCover . . . . . 851
  - multiplicity . . . . . 514
  - MultiplicitySequence . . . . . 861
  - multiplyLeftFractions . . . . . 590
  - multiplylist . . . . . 846
  - multivariate equations . . . . . 875
  - multRat . . . . . 533
  - multrow . . . . . 807
  - multseq2charexp . . . . . 854
  - multsequence . . . . . 865
  - MVComplex . . . . . 838
- ## N
- naccache\_stern\_decryption . . . . . 895
  - naccache\_stern\_encryption . . . . . 895
  - naccache\_stern\_generation . . . . . 895
  - NakYoshF . . . . . 926
  - nameof . . . . . 224
  - names . . . . . 225
  - Names . . . . . 44
  - Names in procedures . . . . . 54
  - Names, indexed . . . . . 44
  - nashmult . . . . . 855
  - nblocks . . . . . 271
  - nBoundaryLatticePoints . . . . . 908
  - nc\_algebra . . . . . 343
  - nc\_hilb . . . . . 271
  - ncalg.lib . . . . . 458
  - ncalg.lib . . . . . 458

|                            |     |                                                  |          |
|----------------------------|-----|--------------------------------------------------|----------|
| ncalgebra                  | 345 | ncrepIsRegular                                   | 684      |
| ncdecomp.lib               | 477 | ncrepMultiply                                    | 677      |
| ncdecomp.lib               | 477 | ncrepPencilCombine                               | 689      |
| ncdetection                | 455 | ncrepPencilGet                                   | 688      |
| ncExt_R                    | 544 | ncrepPrint                                       | 679      |
| ncfactor                   | 480 | ncrepRegularMinimize                             | 686      |
| ncfactor.lib               | 480 | ncrepRegularZeroMinimize                         | 685      |
| ncfactor.lib               | 480 | ncrepSubstitute                                  | 680      |
| ncfrac.lib                 | 535 | ncrepSubtract                                    | 676      |
| ncfrac.lib                 | 535 | nctools.lib                                      | 559      |
| ncgen                      | 621 | nctools.lib                                      | 559      |
| nchilb                     | 665 | ncVarsAdd                                        | 669      |
| ncHilb                     | 514 | ncVarsGet                                        | 669      |
| ncHilb.lib                 | 665 | ndcond                                           | 561      |
| ncHilb.lib                 | 665 | negatedCone                                      | 903      |
| nchilbert.lib              | 514 | negateNcfrac                                     | 542      |
| nchilbert.lib              | 514 | negative degree lexicographical ordering         | 761      |
| ncHilbertMultiplicity      | 517 | negative degree reverse lexicographical ordering | 761      |
| ncHilbertPolynomial        | 516 | negative lexicographical ordering                | 761      |
| ncHilbertSeries            | 515 | negativedivisor                                  | 839      |
| ncHom                      | 545 | negativeformaldivisor                            | 839      |
| nchomolog.lib              | 544 | net                                              | 937      |
| nchomolog.lib              | 544 | net access                                       | 19       |
| ncInit                     | 669 | netBigInt                                        | 937      |
| ncloc.lib                  | 548 | netBigIntMat                                     | 937      |
| ncloc.lib                  | 548 | netBigIntMatShort                                | 937      |
| ncmodslimb                 | 550 | netCoefficientRing                               | 937      |
| ncModslimb.lib             | 550 | netIdeal                                         | 937      |
| ncModslimb.lib             | 550 | netInt                                           | 937      |
| ncols                      | 227 | netIntMat                                        | 937      |
| ncones                     | 904 | netIntMatShort                                   | 937      |
| ncpreim.lib                | 551 | netIntVector                                     | 937      |
| ncpreim.lib                | 551 | netIntVectorShort                                | 937      |
| ncrat.lib                  | 668 | netList                                          | 937      |
| ncrat.lib                  | 668 | netMap                                           | 938      |
| ncratAdd                   | 670 | netMap2                                          | 938      |
| ncratDefine                | 670 | netmatrix                                        | 938      |
| ncratEvaluateAt            | 674 | netMatrix                                        | 932      |
| ncratFromPoly              | 673 | netmatrixShort                                   | 938      |
| ncratFromString            | 673 | netNumber                                        | 937      |
| ncratInvert                | 672 | netPoly                                          | 938      |
| ncratMultiply              | 671 | netPrimePower                                    | 938      |
| ncratPower                 | 674 | netRing                                          | 938      |
| ncratPrint                 | 673 | nets.lib                                         | 937      |
| ncratSPrint                | 672 | nets.lib                                         | 937      |
| ncratSubtract              | 671 | netString                                        | 938      |
| ncRelations                | 570 | netvector                                        | 938      |
| ncrepAdd                   | 675 | netVector                                        | 932      |
| ncrepDim                   | 680 | netvectorShort                                   | 938      |
| ncrepEvaluate              | 681 | New orderings for modules                        | 952      |
| ncrepEvaluateAt            | 682 | newline                                          | 127      |
| ncrepGet                   | 674 | news                                             | 944      |
| ncrepGetRegularMinimal     | 687 | newstruct                                        | 133, 919 |
| ncrepGetRegularZeroMinimal | 686 | newTest                                          | 892      |
| ncrepInvert                | 678 | Newton polygons                                  | 913      |
| ncrepIsDefined             | 683 | Newton polytope                                  | 908      |
| ncrepIsDefinedDim          | 682 | Newton step                                      | 882      |

- newtonDiag . . . . . 799
- newtonpoly . . . . . 865
- newtonPolygonNegSlopes . . . . . 913
- newtonPolytope . . . . . 904
- newtonPolytopeLP . . . . . 907
- newtonPolytopeP . . . . . 907
- NF . . . . . 246
- NF (letterplace) . . . . . 622
- NF (plural) . . . . . 351
- nf\_icis . . . . . 869
- nfmodStd . . . . . 823
- nfmodstd.lib . . . . . 822
- nfmodstd\_lib . . . . . 822
- nfmodSyz . . . . . 823
- nfmodsyz.lib . . . . . 823
- nfmodsyz\_lib . . . . . 823
- NFMora . . . . . 897
- nHilbertBasis . . . . . 908
- nilpotent Lie algebras . . . . . 842
- nInteriorLatticePoints . . . . . 908
- nLatticePoints . . . . . 908
- nmaxcones . . . . . 903
- noElements . . . . . 940
- noether . . . . . 299
- noether.lib . . . . . 823
- noether\_lib . . . . . 823
- noetherNormal . . . . . 810
- NoetherPosition . . . . . 822
- Non-commutative algebra . . . . . 748, 776
- Non-commutative subsystem . . . . . 312
- non-english special characters . . . . . 304
- noncommutative, rational expressions . . . . . 669
- none, option . . . . . 230
- Nonhyp . . . . . 845
- nonMonomials . . . . . 828
- nonZeroEntry . . . . . 810
- norm . . . . . 896
- normaform (up to a bound) . . . . . 272
- normal . . . . . 824
- normal form . . . . . 766
- normal.lib . . . . . 824
- normal.lib . . . . . 824
- normalBundle . . . . . 851
- normalC . . . . . 824
- normalConductor . . . . . 825
- normalFan . . . . . 908
- normalform . . . . . 857
- normalForm . . . . . 839
- normall . . . . . 831
- normaliz . . . . . 1, 826
- normaliz.lib . . . . . 825
- normaliz.lib . . . . . 825
- normalization . . . . . 817, 818, 826
- Normalization . . . . . 723
- normalize . . . . . 798
- normalizeMonoidal . . . . . 594
- normalizeRational . . . . . 595
- normalP . . . . . 824
- normalToricRing . . . . . 826
- normalToricRingFromBinomials . . . . . 826
- norTest . . . . . 825
- not . . . . . 87
- notBuckets, option . . . . . 231
- Notes for developers . . . . . 952
- Notes for Singular users . . . . . 950
- notRegularity, option . . . . . 231
- notSugar, option . . . . . 231
- notWarnSB, option . . . . . 232
- npar . . . . . 891
- npars . . . . . 227
- NPos . . . . . 824
- NPos\_test . . . . . 824
- nres . . . . . 228
- nres (plural) . . . . . 345
- nrows . . . . . 228
- nrroots . . . . . 900
- nrRootsDeterm . . . . . 898
- nrRootsProbab . . . . . 898
- nsatiety . . . . . 824
- nselect . . . . . 812
- NSplaces . . . . . 886
- nt\_solve . . . . . 879
- NTL . . . . . 1
- ntsolve.lib . . . . . 879
- ntsolve\_lib . . . . . 879
- NullCone . . . . . 874
- num\_elim . . . . . 880
- num\_elim1 . . . . . 880
- num\_prime\_decom . . . . . 880
- num\_prime\_decom1 . . . . . 880
- num\_radical\_via\_decom . . . . . 880
- num\_radical\_via\_randlincom . . . . . 880
- num\_radical1 . . . . . 880
- num\_radical2 . . . . . 880
- number . . . . . 113
- number declarations . . . . . 113
- number expressions . . . . . 114
- number operations . . . . . 115
- number related functions . . . . . 116
- number\_e . . . . . 791
- number\_pi . . . . . 791
- numberOfConesOfDimension . . . . . 904
- numerAlg.lib . . . . . 840
- numerAlg.lib . . . . . 840, 842
- numerator . . . . . 229
- numerDecom.lib . . . . . 841
- numerDecom.lib . . . . . 841
- numerical algebraic geometry . . . . . 879
- numerical irreducible decomposition . . . . . 842
- NumIrrDecom . . . . . 842
- NumLocalDim . . . . . 841
- NumPrimDecom . . . . . 842
- nvars . . . . . 229

**O**

|                         |         |
|-------------------------|---------|
| Oaku-Takayama algorithm | 396     |
| Objects                 | 45      |
| olga.lib                | 578     |
| olga.lib                | 578     |
| oneDimBelongSemigroup   | 811     |
| oneNcfrac               | 539     |
| onesVector              | 904     |
| online help             | 15      |
| open                    | 230     |
| opentex                 | 884     |
| operatorAlgebra         | 634     |
| operatorBM              | 401     |
| operatorModulo          | 402     |
| oppose                  | 347     |
| opposite                | 348     |
| opposite polynomial     | 272     |
| opposite ring           | 272     |
| option                  | 230     |
| option(warn)            | 71      |
| optionIsSet             | 804     |
| or                      | 87, 305 |
| orbit                   | 842     |
| orbit_variety           | 873     |
| orbitConeOrbits         | 905     |
| orbitCones              | 905     |
| orbitparam.lib          | 842     |
| orbitparam.lib          | 842     |
| ord                     | 234     |
| ord_test                | 803     |
| orderings               | 760     |
| orderings introduction  | 760     |
| orderings, a            | 764     |
| orderings, global       | 761     |
| orderings, L            | 765     |
| orderings, local        | 761     |
| orderings, M            | 762     |
| orderings, product      | 764     |
| ordstr                  | 235     |
| origin                  | 901     |
| orthogonalize           | 808     |
| outer                   | 806     |
| output                  | 48      |
| Output, formatting of   | 699     |
| outputting monomials    | 300     |

**P**

|                           |          |
|---------------------------|----------|
| p-adic numbers            | 36       |
| package                   | 117      |
| package declarations      | 117      |
| package related functions | 117      |
| pairset                   | 898      |
| par                       | 235      |
| par2varRing               | 785, 786 |
| paraConic                 | 844      |

|                                    |               |
|------------------------------------|---------------|
| parallel computing                 | 942           |
| parallel skeletons                 | 797           |
| parallel.lib                       | 797           |
| parallel.lib                       | 797           |
| parallelization                    | 797, 802, 805 |
| Parallelization                    | 700, 797      |
| parallelTestAND                    | 797           |
| parallelTestOR                     | 797           |
| parallelWaitAll                    | 797           |
| parallelWaitFirst                  | 797           |
| parallelWaitN                      | 797           |
| param                              | 865           |
| Parameter list                     | 52            |
| parameter, as numbers              | 113           |
| Parameters                         | 698           |
| parameterSubstitute                | 912           |
| parametric annihilator             | 396           |
| parametric annihilator for variety | 447           |
| parametrization                    | 842           |
| Parametrization                    | 844           |
| parametrizeOrbit                   | 842           |
| paraPlaneCurve                     | 844           |
| paraplanecurves.lib                | 843           |
| paraplanecurves.lib                | 843           |
| pardeg                             | 235           |
| parstr                             | 236           |
| part                               | 837           |
| PartC                              | 837           |
| partial fraction                   | 938           |
| partial_molien                     | 872           |
| partitions                         | 813           |
| PartitionVar                       | 925           |
| ParToVar                           | 925           |
| partOver                           | 837           |
| partUnder                          | 838           |
| path                               | 953           |
| path integral                      | 813           |
| Path names                         | 951           |
| paths                              | 953           |
| pause                              | 796           |
| PBW                                | 381           |
| PBW basis                          | 360           |
| PBW_eqDeg                          | 394           |
| PBW_maxDeg                         | 394           |
| PBW_maxMonom                       | 395           |
| pdivi                              | 794           |
| pdivi2                             | 801           |
| pdivisorplus                       | 840           |
| PerfectPowerTest                   | 891           |
| permcop                            | 807           |
| permrow                            | 807           |
| permutationFromIntvec              | 905           |
| permutations, sum, max, min        | 921           |
| permutationToIntvec                | 905           |
| permute                            | 813           |
| permute_L                          | 886           |
| perron                             | 596           |

- perron.lib . . . . . 596
- perron\_lib . . . . . 596
- PEsolve . . . . . 875
- pFactor . . . . . 894
- pfd . . . . . 938
- pfd.lib . . . . . 938
- pfd\_lib . . . . . 938
- pfdMat . . . . . 939
- Pfister, Gerhard . . . . . 3
- PH\_ais . . . . . 917
- PH\_nais . . . . . 917
- phindex.lib . . . . . 917
- phindex\_lib . . . . . 917
- picksFormula . . . . . 907
- pid . . . . . 272
- pIntersect . . . . . 376
- pIntersectSyz . . . . . 377
- Pipe links . . . . . 99
- plainInvariants . . . . . 861
- planeCur . . . . . 859
- plot . . . . . 885
- plotRot . . . . . 885
- plotRotated . . . . . 885
- plotRotatedDirect . . . . . 885
- plotRotatedList . . . . . 885
- plotRotatedListFromSpecifyList . . . . . 885
- PLURAL . . . . . 312
- PLURAL libraries . . . . . 365
- PLURAL LIBs . . . . . 365
- pmat . . . . . 796
- pnormalf . . . . . 794
- pnormalform . . . . . 801
- PocklingtonLehmer . . . . . 894
- pointid.lib . . . . . 827
- pointid\_lib . . . . . 827
- Polar curves . . . . . 729
- PollardRho . . . . . 894
- pollTask . . . . . 805
- polSol . . . . . 523
- polSolFiniteRank . . . . . 524
- poly . . . . . 117
- poly (plural) . . . . . 322
- poly declarations . . . . . 118
- poly declarations (plural) . . . . . 322
- poly expressions . . . . . 118
- poly expressions (plural) . . . . . 323
- poly operations . . . . . 119
- poly operations (plural) . . . . . 324
- poly related functions . . . . . 120
- poly related functions (plural) . . . . . 324
- poly2list . . . . . 436
- poly2zdd . . . . . 919
- polybori . . . . . 919
- polybori.lib . . . . . 918, 919
- polybori\_lib . . . . . 918
- polyclass.lib . . . . . 939
- polyclass\_lib . . . . . 939
- polyInterpolation . . . . . 814
- polylib.lib . . . . . 798
- polylib\_lib . . . . . 798
- polymake . . . . . 908
- polymake.lib . . . . . 906
- polymake\_lib . . . . . 906
- polymakePolytope . . . . . 907
- Polynomial data . . . . . 759
- polynomial functions . . . . . 940
- polynomial solutions . . . . . 519
- polynomial, opposite . . . . . 272
- polynomials . . . . . 939
- polytope . . . . . 138, 908
- polytopeViaInequalities . . . . . 903
- polytopeViaPoints . . . . . 903
- polyVars . . . . . 529
- pos\_def . . . . . 808
- positiveOrthant . . . . . 901
- posweight . . . . . 870
- Pottier algorithm . . . . . 774
- power . . . . . 806
- power\_products . . . . . 872
- powerN . . . . . 893
- powerpolyX . . . . . 892
- powersums . . . . . 899
- powerX . . . . . 893
- powSumSym . . . . . 835
- PPolyH . . . . . 926
- PPolyN . . . . . 926
- PPolyQp . . . . . 926
- PPolyS . . . . . 926
- preComp . . . . . 821
- Preface . . . . . 1
- preimage . . . . . 236, 552
- preimage (plural) . . . . . 349
- preimageLattice . . . . . 935
- preimageLoc . . . . . 804
- preimageNC . . . . . 553
- Prep . . . . . 794
- prepareAss . . . . . 829
- prepEmbDiv . . . . . 849
- prepMat . . . . . 875
- prepRealclassify . . . . . 857
- prepSV . . . . . 886
- presentation . . . . . 932
- presentTree . . . . . 848
- presolve.lib . . . . . 877
- presolve\_lib . . . . . 877
- primary decomposition . . . . . 829, 830
- Primary decomposition . . . . . 721
- primary decomposition of modules . . . . . 821
- primary decomposition, numerical . . . . . 842
- primary\_char0 . . . . . 872
- primary\_char0\_no\_molien . . . . . 872
- primary\_char0\_no\_molien\_random . . . . . 872
- primary\_char0\_random . . . . . 872
- primary\_charp . . . . . 872

- primary\_charp\_no\_molien . . . . . 872
- primary\_charp\_no\_molien\_random . . . . . 872
- primary\_charp\_random . . . . . 872
- primary\_charp\_without . . . . . 872
- primary\_charp\_without\_random . . . . . 872
- primary\_invariants . . . . . 871
- primary\_invariants\_random . . . . . 871
- primdec.lib . . . . . 828
- primdec.lib . . . . . 828
- PrimdecA . . . . . 820
- PrimdecB . . . . . 820
- primdecGTZ . . . . . 828
- primdecGTZE . . . . . 828
- primdecint.lib . . . . . 830
- primdecint.lib . . . . . 830
- primdecMon . . . . . 820
- primdecSY . . . . . 829
- primdecSYE . . . . . 829
- primdecZ . . . . . 830
- primdecZM . . . . . 830
- prime . . . . . 237
- primeClosure . . . . . 824
- primecoeffs . . . . . 791
- primefactors . . . . . 237
- primes . . . . . 791
- primitiv.lib . . . . . 830
- primitiv.lib . . . . . 830
- primitive . . . . . 830
- primitive\_extra . . . . . 830
- primitiveSpan . . . . . 935
- primL . . . . . 893
- primList . . . . . 893
- primparam . . . . . 861
- primRoot . . . . . 833
- primTest . . . . . 821
- principal intersection . . . . . 371
- print . . . . . 238
- printBetti . . . . . 931
- printf . . . . . 240, 785
- printFreeModule . . . . . 931
- printGraph . . . . . 813
- printGraphG . . . . . 851
- printGroup . . . . . 934
- printHom . . . . . 931
- printlevel . . . . . 299
- printMatrix . . . . . 931
- printModule . . . . . 931
- printMoebius . . . . . 917
- printNormalFormEquation . . . . . 940
- printPoly . . . . . 939
- printResolution . . . . . 931
- printSheaf . . . . . 850
- printStack . . . . . 851
- printTask . . . . . 805
- printVariety . . . . . 850
- proc . . . . . 121
- proc declaration . . . . . 121
- proc expression . . . . . 122
- Procedure definition . . . . . 50
- procedure, ASCII help . . . . . 61
- procedure, ASCII/TeXinfo help . . . . . 62
- procedure, texinfo help . . . . . 61
- Procedure-specific commands . . . . . 54
- Procedures . . . . . 50
- Procedures and libraries . . . . . 10, 692
- Procedures in a library . . . . . 57
- procedures, help string . . . . . 53
- procedures, static . . . . . 50
- procs with different argument types . . . . . 122
- prodcrit . . . . . 897
- product . . . . . 791
- Product orderings . . . . . 764
- productgroup . . . . . 935
- productVariety . . . . . 850
- Programming . . . . . 691
- progress watch . . . . . 231
- projective dimension . . . . . 597
- projective limes . . . . . 36
- projectiveBundle . . . . . 850
- projectiveDimension . . . . . 597
- projectiveSpace . . . . . 850
- projectLattice . . . . . 935
- prompt . . . . . 15
- prompt, option . . . . . 232
- propagator . . . . . 813
- prot, option . . . . . 231
- protocol of computations . . . . . 231
- proximitymatrix . . . . . 854
- prune . . . . . 241
- pruneModule . . . . . 933
- prwalk . . . . . 940
- Pseudo ordering L . . . . . 765
- psigncmd . . . . . 882
- PtoCrep . . . . . 794
- Puiseux expansion . . . . . 854, 865
- Puiseux pairs . . . . . 736
- puiseux2generators . . . . . 865
- puiseuxExpansion . . . . . 911
- pure tensor . . . . . 366
- purelist . . . . . 605
- purity . . . . . 597
- purityFiltration . . . . . 598
- purityfiltration.lib . . . . . 597
- purityfiltration.lib . . . . . 597
- purityTriang . . . . . 599
- pushForward . . . . . 936
- pwalk . . . . . 815
- pyobject . . . . . 138, 919
- pyobject declarations . . . . . 139
- pyobject expressions . . . . . 139
- pyobject operations . . . . . 140
- pyobject related functions . . . . . 142
- pyramid . . . . . 925
- python\_eval . . . . . 143



python\_import . . . . . 143  
python\_run . . . . . 144

## Q

qbase . . . . . 899  
qepcad . . . . . 925  
qepcadsystem . . . . . 925  
qhmatrix . . . . . 869  
qhmoduli.lib . . . . . 867  
qhmoduli\_lib . . . . . 867  
qhspectrum . . . . . 869  
qhweight . . . . . 241  
qmatrix.lib . . . . . 606  
qmatrix\_lib . . . . . 606  
qminor . . . . . 607  
QQ . . . . . 30, 72  
qrds . . . . . 242  
qring . . . . . 124, 309  
qring (plural) . . . . . 324  
qring declaration . . . . . 126  
qring declaration (plural) . . . . . 325  
qring related functions (plural) . . . . . 325  
qringNF, option . . . . . 231  
qslimgb . . . . . 785, 786  
Qso3Casimir . . . . . 475  
quadraticSieve . . . . . 894  
quantMat . . . . . 606  
quickclass . . . . . 857  
quit . . . . . 295  
Quot-scheme . . . . . 926  
quote . . . . . 242  
quotient . . . . . 243  
Quotient . . . . . 882  
quotient (plural) . . . . . 350  
QuotientEquations . . . . . 867  
quotientLatticeBasis . . . . . 903  
quotientMain . . . . . 883  
quotientMon . . . . . 820  
quotSheaf . . . . . 850

## R

rad\_con . . . . . 798  
radical . . . . . 829  
radicalEHV . . . . . 829  
radicalMemberShip . . . . . 912  
radicalMon . . . . . 820  
radicalZ . . . . . 830  
randcharpoly . . . . . 899  
randlinpoly . . . . . 899  
random . . . . . 244, 272  
random number generator, seed . . . . . 272  
random.lib . . . . . 802  
random\_lib . . . . . 802  
randomBinomial . . . . . 802

randomCheck . . . . . 887  
randomid . . . . . 802  
randomLast . . . . . 802  
randommat . . . . . 802  
randomPoint . . . . . 903  
randomPolyInT . . . . . 912  
rank . . . . . 155, 244  
rankSheaf . . . . . 850  
ratgb.lib . . . . . 609  
ratgb\_lib . . . . . 609  
Rational curves . . . . . 844  
rational functions . . . . . 669  
rational solutions . . . . . 519  
rationalCurve . . . . . 851  
rationalPointConic . . . . . 844  
ratSol . . . . . 525  
ratstd . . . . . 609  
rays . . . . . 903  
rcgs . . . . . 801  
rcolon . . . . . 668  
re2squ . . . . . 841  
read . . . . . 245  
reading, option . . . . . 232  
readInputTXT . . . . . 939  
readline . . . . . 1  
readNmzData . . . . . 827  
real . . . . . 30  
real root isolation . . . . . 882  
real roots, univariate polynomial . . . . . 900  
real roots, univariate projection . . . . . 899  
real roots, sign conditions . . . . . 882  
realclassify . . . . . 868  
realclassify.lib . . . . . 867  
realclassify\_lib . . . . . 867  
realizationDim . . . . . 909  
realizationDimPoly . . . . . 909  
realizationMatroids.lib . . . . . 908  
realizationMatroids\_lib . . . . . 908  
realLLL . . . . . 880  
realmorsesplit . . . . . 868  
realpoly . . . . . 831  
realrad . . . . . 831  
realrad.lib . . . . . 831  
realrad\_lib . . . . . 831  
realzero . . . . . 831  
recover.lib . . . . . 879  
recover\_lib . . . . . 879  
recursive\_boolean\_poly . . . . . 919  
recursive\_from\_boolean\_poly . . . . . 919  
redcgs.lib . . . . . 799  
redcgs\_lib . . . . . 799  
redefine, option . . . . . 232  
redSB, option . . . . . 231  
redspec . . . . . 801  
redTail, option . . . . . 231  
redThrough, option . . . . . 231  
reduce . . . . . 246, 381

- reduce (letterplace) . . . . . 622
- reduce (plural) . . . . . 351
- reduce (up to a bound) . . . . . 272
- Reduced Comprehensive Groebner Systems . . . . . 799
- reduced standard basis . . . . . 231
- reduceIntChain . . . . . 933
- reduction . . . . . 873
- ReesAlgebra . . . . . 831
- reesclos.lib . . . . . 831
- reesclos.lib . . . . . 831
- reference . . . . . 144
- reference declarations . . . . . 146
- reference expressions . . . . . 147
- reference operations . . . . . 149
- reference related functions . . . . . 151
- References . . . . . 783
- References (plural) . . . . . 364
- References and history of Letterplace . . . . . 618
- regCM . . . . . 824
- regIdeal . . . . . 821
- regMonCurve . . . . . 822
- regularity . . . . . 247, 768
- reiffen . . . . . 411
- ReJunkUseHomo . . . . . 841
- rel\_orbit\_variety . . . . . 873
- relations . . . . . 596
- relative\_orbit\_variety . . . . . 873
- relativeInteriorPoint . . . . . 903
- Release Notes . . . . . 944
- Release Notes (letterplace) . . . . . 689
- relweight . . . . . 869
- remainder . . . . . 882
- remainderMain . . . . . 883
- removeCone . . . . . 904
- removepower . . . . . 855
- repart . . . . . 248
- replace . . . . . 891
- representation, math objects . . . . . 759
- res . . . . . 248, 785
- Res . . . . . 931
- resbinomial.lib . . . . . 844
- resbinomial.lib . . . . . 844
- reservedName . . . . . 249
- resfunction . . . . . 845
- resgraph.lib . . . . . 846
- resgraph.lib . . . . . 846
- resjung.lib . . . . . 847
- resjung.lib . . . . . 847
- reslist . . . . . 846
- resolution . . . . . 123, 597
- Resolution . . . . . 13
- resolution (plural) . . . . . 326
- resolution declarations . . . . . 123
- resolution declarations (plural) . . . . . 326
- resolution expressions . . . . . 123
- resolution expressions (plural) . . . . . 327
- resolution graph . . . . . 854
- Resolution of singularities . . . . . 743
- resolution related functions . . . . . 124
- resolution related functions (plural) . . . . . 327
- resolution, computation of . . . . . 248
- Resolution, free . . . . . 712
- resolution, hilbert-driven . . . . . 195
- resolution, La Scala's method . . . . . 212
- resolutiongraph . . . . . 854
- resolutionInLocalization . . . . . 927
- resolve . . . . . 848
- resolve.lib . . . . . 847
- resolve.lib . . . . . 847
- resources.lib . . . . . 802
- resources.lib . . . . . 802
- ResTree . . . . . 846
- restriction of . . . . . 415
- restrictionIdeal . . . . . 423
- restrictionModule . . . . . 424
- resultant . . . . . 222, 250
- reszeta.lib . . . . . 848
- reszeta.lib . . . . . 848
- return . . . . . 295
- return type of procedures . . . . . 307
- returnSB, option . . . . . 230
- reverse . . . . . 900
- reverse lexicographical ordering . . . . . 761
- reynolds\_molien . . . . . 872
- ReynoldsImage . . . . . 874
- ReynoldsOperator . . . . . 874
- rho . . . . . 893
- rHRR . . . . . 837
- RiemannRochBN . . . . . 834
- RiemannRochHess . . . . . 840
- Right Groebner bases and syzygies . . . . . 750
- Right ideal membership (plural) . . . . . 362
- rightInverse . . . . . 889
- rightKernel . . . . . 889
- rightModulo . . . . . 568
- rightNF . . . . . 568
- rightNFWeyl . . . . . 535
- rightOre . . . . . 585
- rightStd . . . . . 567
- rightstd (letterplace) . . . . . 623
- ring . . . . . 124
- ring (plural) . . . . . 327
- ring declarations . . . . . 124
- ring declarations (plural) . . . . . 327
- ring operations . . . . . 125
- ring operations (plural) . . . . . 328
- ring related functions . . . . . 125
- ring related functions (plural) . . . . . 328
- ring theory . . . . . 649
- ring, opposite . . . . . 272
- Ring-dependent options . . . . . 951
- ring.lib . . . . . 803
- ring\_cf . . . . . 155
- ring.lib . . . . . 803

- ring\_list . . . . . 252
  - ringgb.lib . . . . . 940
  - ringgb.lib, functions for coefficient rings . . . . . 940
  - ringgb\_lib . . . . . 940
  - ringlist . . . . . 250
  - ringlist (plural) . . . . . 352
  - Rings and orderings . . . . . 30
  - Rings and standard bases . . . . . 7
  - ringtensor . . . . . 803
  - ringweights . . . . . 803
  - rinvar.lib . . . . . 873
  - rinvar\_lib . . . . . 873
  - rJanet . . . . . 832
  - rm\_unitcol . . . . . 807
  - rm\_unitrow . . . . . 807
  - rMacaulay . . . . . 796
  - rmNmzFiles . . . . . 827
  - rmx . . . . . 884
  - rncAntiCanonicalMap . . . . . 844
  - rncItProjEven . . . . . 844
  - rncItProjOdd . . . . . 844
  - root of Bernstein-Sato polynomial . . . . . 396
  - rootIsolation . . . . . 882
  - rootisolation.lib . . . . . 880
  - rootisolation\_lib . . . . . 880
  - rootIsolationNoPreprocessing . . . . . 881
  - rootIsolationPrimdec . . . . . 882
  - rootofUnity . . . . . 804
  - roots . . . . . 883
  - rootsMain . . . . . 883
  - rootsModp . . . . . 892
  - rootsmr.lib . . . . . 898
  - rootsmr\_lib . . . . . 898
  - rootsur.lib . . . . . 899
  - rootsur\_lib . . . . . 899
  - round . . . . . 892
  - rowred . . . . . 807
  - rowShift . . . . . 155, 157
  - rp, global ordering . . . . . 761
  - rstandard.lib . . . . . 831
  - rstandard\_lib . . . . . 831
  - rswalk . . . . . 942
  - rtimer . . . . . 303
  - Running SINGULAR under Emacs . . . . . 25
  - rvalue . . . . . 305
  - rvar . . . . . 253
  - rwalk . . . . . 940
  - rwalk.lib . . . . . 940
  - rwalk\_lib . . . . . 940
- S**
- sa\_poly\_reduce . . . . . 387
  - sa\_reduce . . . . . 387
  - sagbi . . . . . 832
  - sagbi.lib . . . . . 832
  - sagbi\_lib . . . . . 832
  - sagbiAlg . . . . . 858
  - sagbiMod . . . . . 858
  - sagbiPart . . . . . 832
  - sagbiReduce . . . . . 832
  - sagbiSPoly . . . . . 832
  - SaitoBase . . . . . 943
  - salida . . . . . 846
  - sameComponent . . . . . 897
  - sameQ . . . . . 883
  - Sannfs . . . . . 398
  - SannfsBFCT . . . . . 406
  - Sannfslog . . . . . 399
  - SannfsVar . . . . . 449
  - sat . . . . . 812
  - satiety . . . . . 822
  - satstd . . . . . 917
  - Saturation . . . . . 707
  - sba . . . . . 254
  - SCA . . . . . 611
  - scalarProd . . . . . 381
  - scheme . . . . . 924
  - Schönemann, Hans . . . . . 3
  - Schoof . . . . . 894
  - Schubert calculus . . . . . 851
  - schubert.lib . . . . . 849
  - schubert\_lib . . . . . 849
  - SchubertClass . . . . . 851
  - SchurCh . . . . . 837
  - SchurS . . . . . 837
  - SDB breakpoint . . . . . 69
  - SDB debugger . . . . . 69
  - sdb, source code debugger . . . . . 68
  - SDLoc . . . . . 417
  - secondary fan . . . . . 908
  - secondary polytope . . . . . 908
  - secondary\_and\_irreducibles\_no\_molien . . . . . 873
  - secondary\_char0 . . . . . 872
  - secondary\_charp . . . . . 872
  - secondary\_no\_molien . . . . . 872
  - secondary\_not\_cohen\_macaulay . . . . . 873
  - secondaryPolytope . . . . . 907
  - segre . . . . . 835
  - SegreA . . . . . 838
  - select . . . . . 812
  - select1 . . . . . 812
  - semaphore . . . . . 272, 803
  - semaphores . . . . . 802
  - semi-prime ideal . . . . . 649
  - semiCMcod2 . . . . . 869
  - semidiv . . . . . 897
  - semigroup . . . . . 854
  - semiGroup . . . . . 858
  - Semigroup . . . . . 860
  - semigroupGenerator . . . . . 903
  - semiMod . . . . . 859
  - sep . . . . . 829
  - separateHNE . . . . . 865

- separator . . . . . 820
- Serre relations . . . . . 634
- serreRelations . . . . . 635
- set . . . . . 941
- set, intersectionSet, union, complement, equality,  
  isEqual . . . . . 941
- set\_is\_injective . . . . . 896
- setBaseMultigrading . . . . . 934
- setcores . . . . . 803
- SetDeg . . . . . 931
- setenv . . . . . 272
- setglobalrings . . . . . 801
- setinitials . . . . . 925
- setLetterplaceAttributes . . . . . 658
- setLinearForms . . . . . 903
- setModuleGrading . . . . . 935
- setMultiplicity . . . . . 903
- setNmzDataPath . . . . . 827
- setNmzExecPath . . . . . 826
- setNmzFilename . . . . . 827
- setNmzOption . . . . . 826
- setring . . . . . 255
- sets.lib . . . . . 940
- sets\_lib . . . . . 940
- Setting up a G-algebra . . . . . 361
- setUniformizingParameter . . . . . 913
- sh . . . . . 273
- ShanksMestre . . . . . 894
- shared . . . . . 144
- shared declarations . . . . . 148
- shared expressions . . . . . 148
- shared operations . . . . . 149
- shared related functions . . . . . 151
- sheaf cohomology . . . . . 852
- sheafCoh . . . . . 852
- sheafcoh.lib . . . . . 852
- sheafcoh\_lib . . . . . 852
- sheafCohBGG . . . . . 852
- sheafCohBGG2 . . . . . 852
- short . . . . . 300
- show . . . . . 796
- showBO . . . . . 848
- showDataTypes . . . . . 848
- showgrades . . . . . 601
- showNmzOptions . . . . . 826
- showNuminvs . . . . . 826
- showrecursive . . . . . 796
- signal handler . . . . . 953
- signatureBrieskorn . . . . . 871
- signatureL . . . . . 917
- signatureLqf . . . . . 917
- signatureNemethi . . . . . 871
- signaturePuisseux . . . . . 871
- signcnd . . . . . 882
- signcond.lib . . . . . 882
- signcond\_lib . . . . . 882
- simplePrune . . . . . 933
- simplesolver . . . . . 875
- simplex . . . . . 256
- simplexOut . . . . . 878
- simplify . . . . . 258
- SimplifyIdeal . . . . . 874
- simplifyRat . . . . . 532
- sing.lib . . . . . 868
- sing\_lib . . . . . 868
- sing4ti2.lib . . . . . 833
- sing4ti2\_lib . . . . . 833
- Singular . . . . . 273
- Singular 3 and Singular 4 . . . . . 950
- SINGULAR libraries . . . . . 785
- singular locus of D-module . . . . . 519
- Singular, customization of Emacs user interface . . . . . 27
- Singular, demo mode . . . . . 27
- Singular, editing input files with Emacs . . . . . 28
- Singular, important commands of Emacs interface . . . . . 29
- Singular, running within Emacs . . . . . 25
- Singular2bertini . . . . . 841
- SINGULARHIST . . . . . 18
- singularities . . . . . 855, 858, 863
- Singularities . . . . . 854
- singularities, resolution of . . . . . 743
- singularity . . . . . 857
- Singularity Theory . . . . . 726
- SingularityDBM.lib . . . . . 855
- SingularityDBM\_lib . . . . . 855
- SingularLib . . . . . 273
- singularrc . . . . . 22
- size . . . . . 259, 309
- Skeletons for parallelization . . . . . 797
- skewmat . . . . . 806
- SL . . . . . 875
- sleep . . . . . 265
- slimgb . . . . . 260, 706
- slimgb (plural) . . . . . 354
- slocus . . . . . 869
- SLreynolds . . . . . 875
- smatrix . . . . . 127
- smith . . . . . 890
- Smith form . . . . . 890
- Smith normal form . . . . . 890
- smithNormalForm . . . . . 936
- SolowayStrassen . . . . . 894
- solutionsMod2 . . . . . 893
- solve . . . . . 205, 878
- solve a linear equation system  $A*x = b$  via the  
  LU-decomposition of A . . . . . 214
- solve.lib . . . . . 878
- solve\_IP . . . . . 817
- solve\_lib . . . . . 878
- solvelinearpart . . . . . 877
- solveTInitialFormPar . . . . . 912
- solving . . . . . 879
- Solving systems of polynomial equations . . . . . 752
- sort . . . . . 791

- sortandmap . . . . . 877
- sortier . . . . . 873
- sortIntvec . . . . . 439
- sortvars . . . . . 877
- sortvec . . . . . 261
- source . . . . . 932
- Source . . . . . 931
- Source code debugger, invocation . . . . . 19
- source code debugger, sdb . . . . . 68
- Space curve singularities, branches of . . . . . 738
- spaceCur . . . . . 859
- spadd . . . . . 863
- span . . . . . 903
- sparseHomogIdeal . . . . . 802
- sparseid . . . . . 802
- sparseInterpolation . . . . . 815
- sparsemat . . . . . 802
- sparsematrix . . . . . 802
- sparsepoly . . . . . 802
- spasetriag . . . . . 802
- spcurve.lib . . . . . 869
- spcurve.lib . . . . . 869
- Special characters . . . . . 42
- special characters, non-english . . . . . 304
- spectral pairs . . . . . 863, 864
- spectralNeg . . . . . 848
- spectrum . . . . . 863, 864
- spectrum.lib . . . . . 870
- spectrum.lib . . . . . 870
- spectrumnd . . . . . 870
- spgamma . . . . . 863
- spgeomgenus . . . . . 863
- sPiece . . . . . 942
- spissemicont . . . . . 863
- split . . . . . 796
- splitPolygon . . . . . 907
- splitring . . . . . 830
- splitting . . . . . 821
- spmilnor . . . . . 863
- spmul . . . . . 863
- spnf . . . . . 808
- spoly . . . . . 897
- sppairs . . . . . 863
- sppnf . . . . . 863
- sppprint . . . . . 863
- spprint . . . . . 808
- sprintf . . . . . 262, 785
- spsemicont . . . . . 863
- spsub . . . . . 863
- sqfrNorm . . . . . 883
- sqfrNormMain . . . . . 883
- sqr . . . . . 892
- sqrfree . . . . . 261
- squarefree . . . . . 865
- squareRoot . . . . . 893
- sres . . . . . 264
- sRes . . . . . 931
- ssi . . . . . 700
- Ssi file links . . . . . 97
- Ssi links . . . . . 96
- Ssi tcp links . . . . . 98
- StabEqn . . . . . 867
- StabEqnId . . . . . 867
- stabilizer . . . . . 920
- StabOrder . . . . . 867
- stackNets . . . . . 938
- staircase . . . . . 883
- standard . . . . . 898
- Standard bases . . . . . 766
- Standard Bases . . . . . 702
- standard.lib . . . . . 785
- standard.lib . . . . . 785
- Stanely-Reisner ideals, Stanley-Reisner rings,  
    deformations . . . . . 941
- stanleyreisner.lib . . . . . 941
- stanleyreisner.lib . . . . . 941
- startNmz . . . . . 827
- StartOrderingV . . . . . 853
- startTasks . . . . . 805
- Startup sequence . . . . . 22
- static procedures . . . . . 50
- status . . . . . 265
- std . . . . . 266, 702
- std (letterplace) . . . . . 623
- std (plural) . . . . . 355
- stdfglm . . . . . 267, 785
- stdhilb . . . . . 268, 785
- stdlocus . . . . . 795
- stopTask . . . . . 805
- storeActionOnOrbitConeIndices . . . . . 905
- stratify . . . . . 875
- stratify.lib . . . . . 875
- stratify.lib . . . . . 875
- string . . . . . 127, 309
- string declarations . . . . . 127
- string expressions . . . . . 128
- string operations . . . . . 130
- string related functions . . . . . 130
- string type cast . . . . . 128
- StringF . . . . . 866
- stripHNE . . . . . 865
- sturm . . . . . 900
- sturmha . . . . . 900
- sturmhaseq . . . . . 900
- sturmquery . . . . . 899
- sturmseq . . . . . 900
- sublists . . . . . 921
- submat . . . . . 806
- subquotient . . . . . 932
- subrInterred . . . . . 799
- subset . . . . . 801
- subset\_sum01 . . . . . 895
- subset\_sum02 . . . . . 895
- subst . . . . . 269

- subst (plural) . . . . . 357
- substAll . . . . . 879
- substitute . . . . . 799
- sugarCrit, option . . . . . 231
- sum . . . . . 791
- sum\_of\_powers . . . . . 835
- sumlist . . . . . 846
- sumofquotients . . . . . 851
- super-commutative algebras . . . . . 611
- super\_increasing\_knapsack . . . . . 896
- superCommutative . . . . . 565
- surf.lib . . . . . 885
- surf.lib . . . . . 885
- surfacesignature.lib . . . . . 870
- surfacesignature.lib . . . . . 870
- surfer . . . . . 1, 885
- surfex . . . . . 1
- surfex.lib . . . . . 885
- surfex.lib . . . . . 885
- suspend . . . . . 265
- swalk . . . . . 942
- swalk.lib . . . . . 942
- swalk.lib . . . . . 942
- swap . . . . . 857
- switch . . . . . 305
- switchRingsAndComputeInitialIdeal . . . . . 914
- sym\_gauss . . . . . 807
- Symbolic-numerical solving . . . . . 752, 875
- SymGroup . . . . . 608
- symm . . . . . 835
- symmat . . . . . 806
- symmetric product . . . . . 926
- symmetricBasis . . . . . 807
- symmetricPower . . . . . 807
- symmetricPowerSheaf . . . . . 850
- symmetries . . . . . 921
- symmfunc . . . . . 899
- symmStd . . . . . 834
- symNsym . . . . . 835
- syModStd . . . . . 834
- symodstd.lib . . . . . 833
- symodstd.lib . . . . . 833
- sympower . . . . . 875
- symsignature . . . . . 899
- syndrome . . . . . 887
- sys\_code . . . . . 886
- sysBin . . . . . 887
- sysCRHT . . . . . 887
- sysCRHTMindist . . . . . 887
- sysFL . . . . . 888
- sysNewton . . . . . 887
- sysQE . . . . . 887
- system . . . . . 270
- System and Control theory . . . . . 888
- System dependent limitations . . . . . 304
- System variables . . . . . 297
- system, - . . . . . 273
- system, -long\_option\_name . . . . . 273
- system, -long\_option\_name=value . . . . . 274
- system, absFact . . . . . 270
- system, alarm . . . . . 270
- system, blackbox . . . . . 270
- system, bracket . . . . . 270
- system, browsers . . . . . 270
- system, complexNearZero . . . . . 270
- system, contributors . . . . . 270
- system, cpu . . . . . 270, 273
- system, denom\_list . . . . . 271
- system, eigenvals . . . . . 271
- system, env . . . . . 271
- system, executable . . . . . 271
- system, getenv . . . . . 271
- system, getPrecDigits . . . . . 271
- system, gmsnf . . . . . 271
- system, HC . . . . . 271
- system, hessenberg . . . . . 271
- system, install . . . . . 271
- system, LLL . . . . . 271
- system, nblocks . . . . . 271
- system, nc\_hilb . . . . . 271
- system, neworder . . . . . 272
- system, newstruct . . . . . 272
- system, opp . . . . . 272
- system, oppose . . . . . 272
- system, pcvBasis . . . . . 272
- system, pcvCV2P . . . . . 272
- system, pcvDim . . . . . 272
- system, pcvLAddL . . . . . 272
- system, pcvMinDeg . . . . . 272
- system, pcvP2CV . . . . . 272
- system, pcvPMulL . . . . . 272
- system, pid . . . . . 272
- system, random . . . . . 272
- system, reserve . . . . . 272
- system, reservedLink . . . . . 272
- system, semaphore . . . . . 272
- system, semic . . . . . 272
- system, setenv . . . . . 272
- system, sh . . . . . 273
- system, shrinktest . . . . . 273
- system, Singular . . . . . 273
- system, SingularLib . . . . . 273
- system, spadd . . . . . 273
- system, spectrum . . . . . 273
- system, spmul . . . . . 273
- system, std\_syz . . . . . 273
- system, tensorModuleMult . . . . . 273
- system, twostd . . . . . 273
- system, uname . . . . . 273
- system, verifyGB . . . . . 273
- system, version . . . . . 273
- system, with . . . . . 273
- systemOfParametersOfLocalization . . . . . 927
- systhreads.lib . . . . . 942

- systhreads.lib ..... 942  
 syz ..... 275  
 syz (letterplace) ..... 623  
 syz (plural) ..... 357  
 Syzygies and resolutions ..... 767  
 Syzygies and resolutions (plural) ..... 362  
 Syzygy bimodule ..... 628
- T**
- T\_1 ..... 869  
 T\_12 ..... 869  
 T\_2 ..... 869  
 T1 ..... 731, 942  
 T2 ..... 731, 942  
 tab ..... 796  
 tail ..... 897  
 tame polynomial ..... 864  
 tangentcone ..... 869  
 tangentGens ..... 842  
 target ..... 932  
 Target ..... 931  
 task ..... 805  
 tasks.lib ..... 804  
 tasks.lib ..... 804  
 tau\_es ..... 862  
 tau\_es2 ..... 854  
 tdCf ..... 837  
 tDetropicalise ..... 912  
 tdFactor ..... 837  
 tdProj ..... 837  
 tdTerms ..... 837  
 Teaching ..... 891  
 teachstd.lib ..... 897  
 teachstd.lib ..... 897  
 Template for writing a library ..... 58  
 template.lib ..... 60, 61  
 template.lib ..... 58, 60  
 tensor ..... 275, 366  
 tensor algebra ..... 653  
 tensorFreemodMod ..... 932  
 tensorFreeModule ..... 932  
 tensorMatrix ..... 932  
 tensorMod ..... 816  
 tensorModFreemod ..... 932  
 tensorModule ..... 932  
 tensorProduct ..... 932  
 tensorSheaf ..... 850  
 term orderings ..... 760  
 term orderings introduction ..... 760  
 testFraction ..... 583  
 TestGRRes ..... 928  
 TestJMark ..... 853  
 testLift ..... 659  
 testLocData ..... 581  
 testNCfac ..... 486  
 testNcfrac ..... 544  
 testNcfracExamples ..... 544  
 testNcloc ..... 549  
 testNclocExamples ..... 550  
 testOlga ..... 595  
 testOlgaExamples ..... 595  
 testParametrization ..... 844  
 testPointConic ..... 844  
 testPrimary ..... 829  
 testPrimaryE ..... 829  
 testSyz ..... 662  
 testZero ..... 940  
 tetrahedronGroup ..... 648  
 tex ..... 884  
 texcoef ..... 891  
 texdemo ..... 884  
 texDrawBasic ..... 912  
 texDrawNewtonSubdivision ..... 912  
 texDrawTriangulation ..... 912  
 texDrawTropical ..... 912  
 texfactorize ..... 884  
 texmap ..... 884  
 texMatrix ..... 912  
 texname ..... 884  
 texNumber ..... 912  
 texobj ..... 884  
 texpoly ..... 884  
 texPolynomial ..... 912  
 texproc ..... 884  
 texring ..... 884  
 The online help system ..... 15  
 The SINGULAR language ..... 40  
 three\_elements ..... 896  
 timeFactorize ..... 791  
 timer ..... 300  
 timer resolution, setting the ..... 274  
 timeStd ..... 791  
 timestep ..... 925  
 tInitialForm ..... 911  
 tInitialFormPar ..... 912  
 tInitialFormParMax ..... 912  
 tInitialIdeal ..... 911  
 tjurina ..... 869  
 Tjurina ..... 869  
 Tjurina number ..... 726  
 tmatrix ..... 863  
 todd ..... 836  
 toddE ..... 836  
 toddPoly ..... 837  
 tolessvars ..... 877  
 Top 20 Emacs commands ..... 29  
 topChernClass ..... 850  
 topological invariants ..... 854  
 Tor ..... 816  
 toric ideals ..... 773  
 Toric ideals and integer programming ..... 773  
 toric ring ..... 826





|                             |     |
|-----------------------------|-----|
| variablesStandard           | 393 |
| varMat                      | 915 |
| varNum                      | 915 |
| vars2pars                   | 530 |
| varsigns                    | 900 |
| varstr                      | 280 |
| VarToPar                    | 925 |
| vdim                        | 281 |
| vdim (plural)               | 359 |
| vec2poly                    | 381 |
| VecField.lib                | 943 |
| VecField.lib                | 943 |
| vecFieldToMatrix            | 943 |
| vector                      | 131 |
| vector declarations         | 131 |
| vector expressions          | 131 |
| vector operations           | 132 |
| vector related functions    | 132 |
| verify                      | 899 |
| verify Groebner base        | 273 |
| verifyGB                    | 273 |
| veronese                    | 879 |
| versal                      | 861 |
| version                     | 273 |
| Version number              | 950 |
| Version schema for Singular | 950 |
| Version string              | 55  |
| vertexAdjacencyGraph        | 908 |
| vertexEdgeGraph             | 908 |
| vertices                    | 904 |
| vfilt                       | 863 |
| view                        | 889 |
| visual                      | 908 |
| Visualization               | 883 |
| visualize                   | 924 |
| voice                       | 303 |
| vStd                        | 860 |
| vwfilt                      | 863 |

## W

|                       |                    |
|-----------------------|--------------------|
| waitall               | 281                |
| waitAllTasks          | 805                |
| waitfirst             | 282                |
| waitTasks             | 805                |
| walk, groebner        | 815, 933, 940, 942 |
| warkedPreimageStd     | 927                |
| warn, option          | 71, 230            |
| watchdog              | 791                |
| wedge                 | 282                |
| weierstr.lib          | 898                |
| weierstr.lib          | 898                |
| Weierstrass           | 886                |
| Weierstrass semigroup | 886                |
| weierstrassForm       | 911                |
| weierstrDiv           | 898                |
| weierstrPrep          | 898                |

|                                           |               |
|-------------------------------------------|---------------|
| weight                                    | 283           |
| weight filtration                         | 863, 864      |
| weighted lexicographical ordering         | 761           |
| weighted reverse lexicographical ordering | 761           |
| weightedRing                              | 560           |
| weightKB                                  | 283, 785      |
| weightM, option                           | 232           |
| Weyl                                      | 562           |
| Weyl algebra                              | 396, 441, 447 |
| Weyl closure                              | 519           |
| WeylClosure                               | 521           |
| WeylClosure1                              | 522           |
| whichvariable                             | 900           |
| while                                     | 296           |
| Windows installation                      | 955           |
| withDim                                   | 155           |
| withHilb                                  | 155           |
| withMult                                  | 155           |
| withRes                                   | 155           |
| withSB                                    | 155           |
| WitSet                                    | 841           |
| WitSupSet                                 | 841           |
| WLCGS                                     | 795           |
| WLemma                                    | 795           |
| wp, global ordering                       | 761           |
| WP, global ordering                       | 761           |
| write                                     | 284           |
| writeBertiniInput                         | 880           |
| writeNmzData                              | 827           |
| writeNmzPaths                             | 827           |
| writing monomials                         | 300           |
| ws, local ordering                        | 761           |
| Ws, local ordering                        | 761           |
| WSemigroup                                | 861           |
| wUnit                                     | 892           |
| wurzel                                    | 891           |

## X

|         |     |
|---------|-----|
| xchange | 891 |
| xdvi    | 884 |
| XLsolve | 876 |

## Z

|             |     |
|-------------|-----|
| zdd         | 919 |
| zdd2poly    | 919 |
| zero        | 931 |
| zerodec     | 829 |
| zeroMod     | 821 |
| zeroNcfrac  | 538 |
| zeroOpt     | 821 |
| zeroRadical | 810 |
| zeroreduce  | 940 |
| zeroReduce  | 940 |
| zeroSet     | 883 |
| zeroset.lib | 882 |

|                  |        |              |     |
|------------------|--------|--------------|-----|
| zeroset.lib..... | 882    | ZZ/p.....    | 30  |
| zetaDL.....      | 849    |              |     |
| ZZ.....          | 30, 72 |              |     |
| ZZ/m.....        | 30     | ZZsolve..... | 876 |

# Table of Contents

|          |                                                           |           |
|----------|-----------------------------------------------------------|-----------|
| <b>1</b> | <b>Preface</b> .....                                      | <b>1</b>  |
| <b>2</b> | <b>Introduction, General concepts, Preface, Top</b> ..... | <b>4</b>  |
| 2.1      | Background .....                                          | 4         |
| 2.2      | How to use this manual .....                              | 4         |
| 2.3      | Getting started .....                                     | 6         |
| 2.3.1    | First steps .....                                         | 6         |
| 2.3.2    | Rings and standard bases .....                            | 7         |
| 2.3.3    | Procedures and libraries .....                            | 10        |
| 2.3.4    | Change of rings .....                                     | 11        |
| 2.3.5    | Modules and their annihilator .....                       | 12        |
| 2.3.6    | Resolution .....                                          | 13        |
| <b>3</b> | <b>General concepts</b> .....                             | <b>15</b> |
| 3.1      | Interactive use .....                                     | 15        |
| 3.1.1    | How to enter and exit .....                               | 15        |
| 3.1.2    | The SINGULAR prompt .....                                 | 15        |
| 3.1.3    | The online help system .....                              | 15        |
| 3.1.4    | Interrupting SINGULAR .....                               | 18        |
| 3.1.5    | Editing input .....                                       | 18        |
| 3.1.6    | Command line options .....                                | 19        |
| 3.1.7    | Startup sequence .....                                    | 22        |
| 3.2      | Emacs user interface .....                                | 22        |
| 3.2.1    | A quick guide to Emacs .....                              | 23        |
| 3.2.2    | Running SINGULAR under Emacs .....                        | 25        |
| 3.2.3    | Demo mode .....                                           | 27        |
| 3.2.4    | Customization of the Emacs interface .....                | 27        |
| 3.2.5    | Editing SINGULAR input files with Emacs .....             | 28        |
| 3.2.6    | Top 20 Emacs commands .....                               | 29        |
| 3.3      | Rings and orderings .....                                 | 30        |
| 3.3.1    | Examples of ring declarations .....                       | 31        |
| 3.3.2    | General syntax of a ring declaration .....                | 32        |
| 3.3.3    | Term orderings .....                                      | 34        |
| 3.3.4    | Coefficient rings .....                                   | 36        |
| 3.4      | Implemented algorithms .....                              | 36        |
| 3.5      | The SINGULAR language .....                               | 40        |
| 3.5.1    | General command syntax .....                              | 41        |
| 3.5.2    | Special characters .....                                  | 42        |
| 3.5.3    | Names .....                                               | 44        |
| 3.5.4    | Objects .....                                             | 45        |
| 3.5.5    | Type conversion and casting .....                         | 46        |
| 3.5.6    | Flow control .....                                        | 47        |
| 3.6      | Input and output .....                                    | 48        |
| 3.7      | Procedures .....                                          | 50        |
| 3.7.1    | Procedure definition .....                                | 50        |
| 3.7.2    | Parameter list .....                                      | 52        |

|          |                                         |           |
|----------|-----------------------------------------|-----------|
| 3.7.3    | Help string                             | 53        |
| 3.7.4    | Names in procedures                     | 54        |
| 3.7.5    | Procedure-specific commands             | 54        |
| 3.8      | Libraries                               | 54        |
| 3.8.1    | Libraries in the SINGULAR Documentation | 55        |
| 3.8.2    | Version string                          | 55        |
| 3.8.3    | Category string                         | 55        |
| 3.8.4    | Info string                             | 55        |
| 3.8.5    | LIB commands                            | 57        |
| 3.8.6    | Procedures in a library                 | 57        |
| 3.8.7    | template.lib                            | 58        |
| 3.8.7.1  | mdouble                                 | 61        |
| 3.8.7.2  | mtriple                                 | 61        |
| 3.8.7.3  | msum                                    | 61        |
| 3.8.8    | Formal Checker                          | 62        |
| 3.8.9    | Documentation Tool                      | 62        |
| 3.8.10   | Typesetting of help and info strings    | 63        |
| 3.8.11   | Loading a library                       | 66        |
| 3.9      | Debugging tools                         | 67        |
| 3.9.1    | ASSUME                                  | 67        |
| 3.9.2    | Tracing of procedures                   | 68        |
| 3.9.3    | Source code debugger                    | 68        |
| 3.9.4    | Break points                            | 69        |
| 3.9.5    | Printing of data                        | 69        |
| 3.9.6    | libparse                                | 69        |
| 3.9.7    | option(warn)                            | 71        |
| 3.10     | Dynamic loading                         | 71        |
| <b>4</b> | <b>Data types</b>                       | <b>72</b> |
| 4.1      | cring                                   | 72        |
| 4.1.1    | cring declarations                      | 72        |
| 4.1.2    | cring expressions                       | 72        |
| 4.1.3    | cring operations                        | 72        |
| 4.1.4    | cring related functions                 | 73        |
| 4.2      | bigint                                  | 73        |
| 4.2.1    | bigint declarations                     | 73        |
| 4.2.2    | bigint expressions                      | 73        |
| 4.2.3    | bigint operations                       | 74        |
| 4.2.4    | bigint related functions                | 74        |
| 4.3      | bigintmat                               | 74        |
| 4.3.1    | bigintmat declarations                  | 74        |
| 4.3.2    | bigintmat expressions                   | 75        |
| 4.3.3    | bigintmat type cast                     | 75        |
| 4.3.4    | bigintmat operations                    | 76        |
| 4.4      | def                                     | 77        |
| 4.4.1    | def declarations                        | 77        |
| 4.5      | ideal                                   | 78        |
| 4.5.1    | ideal declarations                      | 78        |
| 4.5.2    | ideal expressions                       | 78        |
| 4.5.3    | ideal operations                        | 79        |
| 4.5.4    | ideal related functions                 | 80        |
| 4.6      | int                                     | 82        |

|         |                          |     |
|---------|--------------------------|-----|
| 4.6.1   | int declarations         | 82  |
| 4.6.2   | int expressions          | 83  |
| 4.6.3   | int operations           | 84  |
| 4.6.4   | int related functions    | 85  |
| 4.6.5   | boolean expressions      | 86  |
| 4.6.6   | boolean operations       | 87  |
| 4.7     | intmat                   | 88  |
| 4.7.1   | intmat declarations      | 88  |
| 4.7.2   | intmat expressions       | 88  |
| 4.7.3   | intmat type cast         | 89  |
| 4.7.4   | intmat operations        | 90  |
| 4.7.5   | intmat related functions | 91  |
| 4.8     | intvec                   | 91  |
| 4.8.1   | intvec declarations      | 91  |
| 4.8.2   | intvec expressions       | 91  |
| 4.8.3   | intvec operations        | 92  |
| 4.8.4   | intvec related functions | 93  |
| 4.9     | link                     | 94  |
| 4.9.1   | link declarations        | 94  |
| 4.9.2   | link expressions         | 94  |
| 4.9.3   | link related functions   | 95  |
| 4.9.4   | ASCII links              | 95  |
| 4.9.5   | Ssi links                | 96  |
| 4.9.5.1 | Ssi file links           | 97  |
| 4.9.5.2 | Ssi tcp links            | 98  |
| 4.9.6   | Pipe links               | 99  |
| 4.9.7   | DBM links                | 99  |
| 4.10    | list                     | 101 |
| 4.10.1  | list declarations        | 101 |
| 4.10.2  | list expressions         | 101 |
| 4.10.3  | list operations          | 102 |
| 4.10.4  | list related functions   | 103 |
| 4.11    | map                      | 103 |
| 4.11.1  | map declarations         | 104 |
| 4.11.2  | map expressions          | 105 |
| 4.11.3  | map operations           | 105 |
| 4.11.4  | map related functions    | 105 |
| 4.12    | matrix                   | 106 |
| 4.12.1  | matrix declarations      | 106 |
| 4.12.2  | matrix expressions       | 107 |
| 4.12.3  | matrix type cast         | 107 |
| 4.12.4  | matrix operations        | 108 |
| 4.12.5  | matrix related functions | 109 |
| 4.13    | module                   | 110 |
| 4.13.1  | module declarations      | 110 |
| 4.13.2  | module expressions       | 110 |
| 4.13.3  | module operations        | 111 |
| 4.13.4  | module related functions | 111 |
| 4.14    | number                   | 113 |
| 4.14.1  | number declarations      | 113 |
| 4.14.2  | number expressions       | 114 |
| 4.14.3  | number operations        | 115 |
| 4.14.4  | number related functions | 116 |

|        |                                               |     |
|--------|-----------------------------------------------|-----|
| 4.15   | package                                       | 117 |
| 4.15.1 | package declarations                          | 117 |
| 4.15.2 | package related functions                     | 117 |
| 4.16   | poly                                          | 117 |
| 4.16.1 | poly declarations                             | 118 |
| 4.16.2 | poly expressions                              | 118 |
| 4.16.3 | poly operations                               | 119 |
| 4.16.4 | poly related functions                        | 120 |
| 4.17   | proc                                          | 121 |
| 4.17.1 | proc declaration                              | 121 |
| 4.17.2 | proc expression                               | 122 |
| 4.17.3 | procs with different argument types           | 122 |
| 4.18   | resolution                                    | 123 |
| 4.18.1 | resolution declarations                       | 123 |
| 4.18.2 | resolution expressions                        | 123 |
| 4.18.3 | resolution related functions                  | 124 |
| 4.19   | ring                                          | 124 |
| 4.19.1 | qring                                         | 124 |
| 4.19.2 | ring declarations                             | 124 |
| 4.19.3 | ring related functions                        | 125 |
| 4.19.4 | ring operations                               | 125 |
| 4.19.5 | qring declaration                             | 126 |
| 4.20   | smatrix                                       | 127 |
| 4.21   | string                                        | 127 |
| 4.21.1 | string declarations                           | 127 |
| 4.21.2 | string expressions                            | 128 |
| 4.21.3 | string type cast                              | 128 |
| 4.21.4 | string operations                             | 130 |
| 4.21.5 | string related functions                      | 130 |
| 4.22   | vector                                        | 131 |
| 4.22.1 | vector declarations                           | 131 |
| 4.22.2 | vector expressions                            | 131 |
| 4.22.3 | vector operations                             | 132 |
| 4.22.4 | vector related functions                      | 132 |
| 4.23   | User defined types                            | 133 |
| 4.23.1 | Definition of a user defined type             | 133 |
| 4.23.2 | Declaration of objects of a user defined type | 134 |
| 4.23.3 | Access to elements of a user defined type     | 135 |
| 4.23.4 | Commands for user defined types               | 135 |
| 4.23.5 | Assignments for user defined types            | 136 |
| 4.24   | cone                                          | 137 |
| 4.25   | fan                                           | 137 |
| 4.26   | polytope                                      | 138 |
| 4.27   | pyobject                                      | 138 |
| 4.27.1 | pyobject declarations                         | 139 |
| 4.27.2 | pyobject expressions                          | 139 |
| 4.27.3 | pyobject operations                           | 140 |
| 4.27.4 | pyobject related functions                    | 142 |
| 4.27.5 | python_eval                                   | 143 |
| 4.27.6 | python_import                                 | 143 |
| 4.27.7 | python_run                                    | 144 |
| 4.28   | reference and shared (experimental)           | 144 |
| 4.28.1 | reference declarations                        | 146 |

|          |                                        |            |
|----------|----------------------------------------|------------|
| 4.28.2   | reference expressions                  | 147        |
| 4.28.3   | shared declarations                    | 148        |
| 4.28.4   | shared expressions                     | 148        |
| 4.28.5   | reference and shared operations        | 149        |
| 4.28.6   | reference and shared related functions | 151        |
| <b>5</b> | <b>Functions and system variables</b>  | <b>154</b> |
| 5.1      | Functions                              | 154        |
| 5.1.1    | align                                  | 154        |
| 5.1.2    | attrib                                 | 154        |
| 5.1.3    | bareiss                                | 156        |
| 5.1.4    | betti                                  | 157        |
| 5.1.5    | char                                   | 159        |
| 5.1.6    | char_series                            | 159        |
| 5.1.7    | charstr                                | 160        |
| 5.1.8    | chinrem                                | 160        |
| 5.1.9    | cleardenom                             | 161        |
| 5.1.10   | close                                  | 161        |
| 5.1.11   | coef                                   | 162        |
| 5.1.12   | coeffs                                 | 163        |
| 5.1.13   | contract                               | 165        |
| 5.1.14   | create_ring                            | 165        |
| 5.1.15   | crossprod                              | 166        |
| 5.1.16   | datetime                               | 167        |
| 5.1.17   | dbprint                                | 167        |
| 5.1.18   | defined                                | 167        |
| 5.1.19   | deg                                    | 168        |
| 5.1.20   | degree                                 | 169        |
| 5.1.21   | delete                                 | 169        |
| 5.1.22   | denominator                            | 170        |
| 5.1.23   | det                                    | 170        |
| 5.1.24   | diff                                   | 171        |
| 5.1.25   | dim                                    | 171        |
| 5.1.26   | division                               | 172        |
| 5.1.27   | dump                                   | 173        |
| 5.1.28   | eliminate                              | 174        |
| 5.1.29   | eval                                   | 174        |
| 5.1.30   | ERROR                                  | 175        |
| 5.1.31   | example                                | 175        |
| 5.1.32   | execute                                | 176        |
| 5.1.33   | extgcd                                 | 176        |
| 5.1.34   | facstd                                 | 177        |
| 5.1.35   | factmodd                               | 177        |
| 5.1.36   | factorize                              | 178        |
| 5.1.37   | farey                                  | 179        |
| 5.1.38   | fetch                                  | 180        |
| 5.1.39   | fglm                                   | 181        |
| 5.1.40   | fglmquot                               | 182        |
| 5.1.41   | files, input from                      | 182        |
| 5.1.42   | find                                   | 182        |
| 5.1.43   | finduni                                | 183        |
| 5.1.44   | flintQ                                 | 183        |

|        |               |     |
|--------|---------------|-----|
| 5.1.45 | Float         | 184 |
| 5.1.46 | fprintf       | 184 |
| 5.1.47 | freemodule    | 186 |
| 5.1.48 | fres          | 186 |
| 5.1.49 | frwalk        | 187 |
| 5.1.50 | gcd           | 188 |
| 5.1.51 | gen           | 188 |
| 5.1.52 | getdump       | 189 |
| 5.1.53 | groebner      | 189 |
| 5.1.54 | help          | 191 |
| 5.1.55 | highcorner    | 192 |
| 5.1.56 | hilb          | 193 |
| 5.1.57 | homog         | 194 |
| 5.1.58 | hres          | 195 |
| 5.1.59 | imap          | 195 |
| 5.1.60 | impart        | 196 |
| 5.1.61 | indepSet      | 196 |
| 5.1.62 | insert        | 197 |
| 5.1.63 | interpolation | 198 |
| 5.1.64 | interred      | 199 |
| 5.1.65 | intersect     | 199 |
| 5.1.66 | jacob         | 200 |
| 5.1.67 | janet         | 201 |
| 5.1.68 | jet           | 201 |
| 5.1.69 | kbase         | 203 |
| 5.1.70 | kernel        | 203 |
| 5.1.71 | kill          | 204 |
| 5.1.72 | killattrib    | 204 |
| 5.1.73 | koszul        | 205 |
| 5.1.74 | laguerre      | 205 |
| 5.1.75 | lead          | 206 |
| 5.1.76 | leadcoef      | 207 |
| 5.1.77 | leadexp       | 207 |
| 5.1.78 | leadmonom     | 208 |
| 5.1.79 | LIB           | 208 |
| 5.1.80 | lift          | 209 |
| 5.1.81 | liftstd       | 209 |
| 5.1.82 | listvar       | 210 |
| 5.1.83 | lres          | 212 |
| 5.1.84 | ludecomp      | 213 |
| 5.1.85 | luinverse     | 213 |
| 5.1.86 | lusolve       | 214 |
| 5.1.87 | max           | 215 |
| 5.1.88 | maxideal      | 216 |
| 5.1.89 | memory        | 216 |
| 5.1.90 | min           | 217 |
| 5.1.91 | minbase       | 218 |
| 5.1.92 | minor         | 218 |
| 5.1.93 | minres        | 220 |
| 5.1.94 | modulo        | 220 |
| 5.1.95 | monitor       | 221 |
| 5.1.96 | monomial      | 221 |
| 5.1.97 | mpresmat      | 222 |



|         |              |     |
|---------|--------------|-----|
| 5.1.98  | mres         | 222 |
| 5.1.99  | mstd         | 223 |
| 5.1.100 | mult         | 224 |
| 5.1.101 | nameof       | 224 |
| 5.1.102 | names        | 225 |
| 5.1.103 | ncols        | 227 |
| 5.1.104 | npars        | 227 |
| 5.1.105 | nres         | 228 |
| 5.1.106 | nrows        | 228 |
| 5.1.107 | numerator    | 229 |
| 5.1.108 | nvars        | 229 |
| 5.1.109 | open         | 230 |
| 5.1.110 | option       | 230 |
| 5.1.111 | ord          | 234 |
| 5.1.112 | ordstr       | 235 |
| 5.1.113 | par          | 235 |
| 5.1.114 | pardeg       | 235 |
| 5.1.115 | parstr       | 236 |
| 5.1.116 | preimage     | 236 |
| 5.1.117 | prime        | 237 |
| 5.1.118 | primefactors | 237 |
| 5.1.119 | print        | 238 |
| 5.1.120 | printf       | 240 |
| 5.1.121 | prune        | 241 |
| 5.1.122 | qhweight     | 241 |
| 5.1.123 | qrds         | 242 |
| 5.1.124 | quote        | 242 |
| 5.1.125 | quotient     | 243 |
| 5.1.126 | random       | 244 |
| 5.1.127 | rank         | 244 |
| 5.1.128 | read         | 245 |
| 5.1.129 | reduce       | 246 |
| 5.1.130 | regularity   | 247 |
| 5.1.131 | repart       | 248 |
| 5.1.132 | res          | 248 |
| 5.1.133 | reservedName | 249 |
| 5.1.134 | resultant    | 250 |
| 5.1.135 | ringlist     | 250 |
| 5.1.136 | ring_list    | 252 |
| 5.1.137 | rvar         | 253 |
| 5.1.138 | sba          | 254 |
| 5.1.139 | setring      | 255 |
| 5.1.140 | simplex      | 256 |
| 5.1.141 | simplify     | 258 |
| 5.1.142 | size         | 259 |
| 5.1.143 | slimgb       | 260 |
| 5.1.144 | sortvec      | 261 |
| 5.1.145 | sqrfree      | 261 |
| 5.1.146 | sprintf      | 262 |
| 5.1.147 | sres         | 264 |
| 5.1.148 | status       | 265 |
| 5.1.149 | std          | 266 |
| 5.1.150 | stdfglm      | 267 |

|         |                    |     |
|---------|--------------------|-----|
| 5.1.151 | stdhilb            | 268 |
| 5.1.152 | subst              | 269 |
| 5.1.153 | system             | 270 |
| 5.1.154 | syz                | 275 |
| 5.1.155 | tensor             | 275 |
| 5.1.156 | trace              | 276 |
| 5.1.157 | transpose          | 276 |
| 5.1.158 | type               | 277 |
| 5.1.159 | typeof             | 277 |
| 5.1.160 | univariate         | 278 |
| 5.1.161 | uressolve          | 278 |
| 5.1.162 | vandermonde        | 279 |
| 5.1.163 | var                | 279 |
| 5.1.164 | variables          | 280 |
| 5.1.165 | varstr             | 280 |
| 5.1.166 | vdim               | 281 |
| 5.1.167 | waitall            | 281 |
| 5.1.168 | waitfirst          | 282 |
| 5.1.169 | wedge              | 282 |
| 5.1.170 | weight             | 283 |
| 5.1.171 | weightKB           | 283 |
| 5.1.172 | write              | 284 |
| 5.2     | Control structures | 285 |
| 5.2.1   | apply              | 285 |
| 5.2.2   | break              | 286 |
| 5.2.3   | breakpoint         | 286 |
| 5.2.4   | continue           | 286 |
| 5.2.5   | else               | 287 |
| 5.2.6   | export             | 287 |
| 5.2.7   | exportto           | 288 |
| 5.2.8   | for                | 290 |
| 5.2.9   | if                 | 291 |
| 5.2.10  | importfrom         | 291 |
| 5.2.11  | keepring           | 293 |
| 5.2.12  | load               | 294 |
| 5.2.13  | quit               | 295 |
| 5.2.14  | return             | 295 |
| 5.2.15  | while              | 296 |
| 5.2.16  | ~ (break point)    | 297 |
| 5.3     | System variables   | 297 |
| 5.3.1   | degBound           | 297 |
| 5.3.2   | echo               | 298 |
| 5.3.3   | minpoly            | 298 |
| 5.3.4   | multBound          | 298 |
| 5.3.5   | noether            | 299 |
| 5.3.6   | printlevel         | 299 |
| 5.3.7   | short              | 300 |
| 5.3.8   | timer              | 300 |
| 5.3.9   | TRACE              | 301 |
| 5.3.10  | rtimer             | 303 |
| 5.3.11  | voice              | 303 |

|          |                                                 |            |
|----------|-------------------------------------------------|------------|
| <b>6</b> | <b>Tricks and pitfalls</b>                      | <b>304</b> |
| 6.1      | Limitations                                     | 304        |
| 6.2      | System dependent limitations                    | 304        |
| 6.3      | Major differences to the C programming language | 304        |
| 6.3.1    | No rvalue of increments and assignments         | 305        |
| 6.3.2    | Evaluation of logical expressions               | 305        |
| 6.3.3    | No case or switch statement                     | 305        |
| 6.3.4    | Usage of commas                                 | 306        |
| 6.3.5    | Usage of brackets                               | 306        |
| 6.3.6    | Behavior of continue                            | 306        |
| 6.3.7    | Return type of procedures                       | 307        |
| 6.3.8    | First index is 1                                | 307        |
| 6.4      | Miscellaneous oddities                          | 308        |
| 6.5      | Identifier resolution                           | 310        |
| <b>7</b> | <b>Non-commutative subsystem</b>                | <b>312</b> |
| 7.1      | PLURAL                                          | 312        |
| 7.2      | Data types (plural)                             | 313        |
| 7.2.1    | ideal (plural)                                  | 313        |
| 7.2.1.1  | ideal declarations (plural)                     | 313        |
| 7.2.1.2  | ideal expressions (plural)                      | 314        |
| 7.2.1.3  | ideal operations (plural)                       | 315        |
| 7.2.1.4  | ideal related functions (plural)                | 316        |
| 7.2.2    | map (plural)                                    | 317        |
| 7.2.2.1  | map declarations (plural)                       | 318        |
| 7.2.2.2  | map expressions (plural)                        | 319        |
| 7.2.2.3  | map (plural) operations                         | 319        |
| 7.2.2.4  | map related functions (plural)                  | 319        |
| 7.2.3    | module (plural)                                 | 320        |
| 7.2.3.1  | module declarations (plural)                    | 320        |
| 7.2.3.2  | module expressions (plural)                     | 320        |
| 7.2.3.3  | module operations (plural)                      | 321        |
| 7.2.3.4  | module related functions (plural)               | 321        |
| 7.2.4    | poly (plural)                                   | 322        |
| 7.2.4.1  | poly declarations (plural)                      | 322        |
| 7.2.4.2  | poly expressions (plural)                       | 323        |
| 7.2.4.3  | poly operations (plural)                        | 324        |
| 7.2.4.4  | poly related functions (plural)                 | 324        |
| 7.2.5    | qring (plural)                                  | 324        |
| 7.2.5.1  | qring declaration (plural)                      | 325        |
| 7.2.5.2  | qring related functions (plural)                | 325        |
| 7.2.6    | resolution (plural)                             | 326        |
| 7.2.6.1  | resolution declarations (plural)                | 326        |
| 7.2.6.2  | resolution expressions (plural)                 | 327        |
| 7.2.6.3  | resolution related functions (plural)           | 327        |
| 7.2.7    | ring (plural)                                   | 327        |
| 7.2.7.1  | ring declarations (plural)                      | 327        |
| 7.2.7.2  | ring operations (plural)                        | 328        |
| 7.2.7.3  | ring related functions (plural)                 | 328        |
| 7.3      | Functions (plural)                              | 329        |
| 7.3.1    | beti (plural)                                   | 329        |
| 7.3.2    | bracket                                         | 330        |

|          |                                   |     |
|----------|-----------------------------------|-----|
| 7.3.3    | dim (plural)                      | 331 |
| 7.3.4    | division (plural)                 | 332 |
| 7.3.5    | eliminate (plural)                | 333 |
| 7.3.6    | envelope                          | 334 |
| 7.3.7    | fetch (plural)                    | 335 |
| 7.3.8    | imap (plural)                     | 336 |
| 7.3.9    | intersect (plural)                | 337 |
| 7.3.10   | kbase (plural)                    | 337 |
| 7.3.11   | lift (plural)                     | 338 |
| 7.3.12   | liftstd (plural)                  | 339 |
| 7.3.13   | minres (plural)                   | 340 |
| 7.3.14   | modulo (plural)                   | 341 |
| 7.3.15   | mres (plural)                     | 342 |
| 7.3.16   | nc_algebra                        | 343 |
| 7.3.17   | ncalgebra                         | 345 |
| 7.3.18   | nres (plural)                     | 345 |
| 7.3.19   | oppose                            | 347 |
| 7.3.20   | opposite                          | 348 |
| 7.3.21   | preimage (plural)                 | 349 |
| 7.3.22   | quotient (plural)                 | 350 |
| 7.3.23   | reduce (plural)                   | 351 |
| 7.3.24   | ringlist (plural)                 | 352 |
| 7.3.25   | slimgb (plural)                   | 354 |
| 7.3.26   | std (plural)                      | 355 |
| 7.3.27   | subst (plural)                    | 357 |
| 7.3.28   | syz (plural)                      | 357 |
| 7.3.29   | twostd (plural)                   | 358 |
| 7.3.30   | vdim (plural)                     | 359 |
| 7.4      | Mathematical background (plural)  | 360 |
| 7.4.1    | G-algebras                        | 360 |
| 7.4.2    | Groebner bases in G-algebras      | 361 |
| 7.4.3    | Syzygies and resolutions (plural) | 362 |
| 7.4.4    | References (plural)               | 364 |
| 7.5      | PLURAL libraries                  | 365 |
| 7.5.1    | bimodules.lib                     | 365 |
| 7.5.1.1  | bistd                             | 366 |
| 7.5.1.2  | bitrinity                         | 367 |
| 7.5.1.3  | liftenvelope                      | 368 |
| 7.5.1.4  | CompDecomp                        | 369 |
| 7.5.1.5  | isPureTensor                      | 369 |
| 7.5.1.6  | isTwoSidedGB                      | 370 |
| 7.5.2    | bfun.lib                          | 370 |
| 7.5.2.1  | bfct                              | 371 |
| 7.5.2.2  | bfctSyz                           | 372 |
| 7.5.2.3  | bfctAnn                           | 373 |
| 7.5.2.4  | bfctOneGB                         | 374 |
| 7.5.2.5  | bfctIdeal                         | 375 |
| 7.5.2.6  | pIntersect                        | 376 |
| 7.5.2.7  | pIntersectSyz                     | 377 |
| 7.5.2.8  | linReduce                         | 378 |
| 7.5.2.9  | linReduceIdeal                    | 379 |
| 7.5.2.10 | linSyzSolve                       | 380 |
| 7.5.2.11 | allPositive                       | 380 |

|       |          |                    |     |
|-------|----------|--------------------|-----|
|       | 7.5.2.12 | scalarProd         | 381 |
|       | 7.5.2.13 | vec2poly           | 381 |
| 7.5.3 |          | central_lib        | 381 |
|       | 7.5.3.1  | centralizeSet      | 382 |
|       | 7.5.3.2  | centralizerVS      | 382 |
|       | 7.5.3.3  | centralizerRed     | 383 |
|       | 7.5.3.4  | centerVS           | 383 |
|       | 7.5.3.5  | centerRed          | 384 |
|       | 7.5.3.6  | center             | 385 |
|       | 7.5.3.7  | centralizer        | 386 |
|       | 7.5.3.8  | sa_reduce          | 387 |
|       | 7.5.3.9  | sa_poly_reduce     | 387 |
|       | 7.5.3.10 | inCenter           | 388 |
|       | 7.5.3.11 | inCentralizer      | 388 |
|       | 7.5.3.12 | isCartan           | 389 |
|       | 7.5.3.13 | applyAdF           | 390 |
|       | 7.5.3.14 | linearMapKernel    | 391 |
|       | 7.5.3.15 | linearCombinations | 392 |
|       | 7.5.3.16 | variablesStandard  | 393 |
|       | 7.5.3.17 | variablesSorted    | 393 |
|       | 7.5.3.18 | PBW_eqDeg          | 394 |
|       | 7.5.3.19 | PBW_maxDeg         | 394 |
|       | 7.5.3.20 | PBW_maxMonom       | 395 |
| 7.5.4 |          | dmod_lib           | 395 |
|       | 7.5.4.1  | annfs              | 397 |
|       | 7.5.4.2  | annfspecial        | 397 |
|       | 7.5.4.3  | Sannfs             | 398 |
|       | 7.5.4.4  | Sannfslog          | 399 |
|       | 7.5.4.5  | bernsteinBM        | 400 |
|       | 7.5.4.6  | bernsteinLift      | 400 |
|       | 7.5.4.7  | operatorBM         | 401 |
|       | 7.5.4.8  | operatorModulo     | 402 |
|       | 7.5.4.9  | annfsParamBM       | 403 |
|       | 7.5.4.10 | annfsBMI           | 404 |
|       | 7.5.4.11 | checkRoot          | 405 |
|       | 7.5.4.12 | SannfsBFCT         | 406 |
|       | 7.5.4.13 | annfs0             | 407 |
|       | 7.5.4.14 | annfs2             | 408 |
|       | 7.5.4.15 | annfsRB            | 409 |
|       | 7.5.4.16 | checkFactor        | 410 |
|       | 7.5.4.17 | arrange            | 411 |
|       | 7.5.4.18 | reiffen            | 411 |
|       | 7.5.4.19 | isHolonomic        | 411 |
|       | 7.5.4.20 | convloc            | 412 |
|       | 7.5.4.21 | minIntRoot         | 412 |
|       | 7.5.4.22 | isRational         | 413 |
| 7.5.5 |          | dmodapp_lib        | 414 |
|       | 7.5.5.1  | annPoly            | 415 |
|       | 7.5.5.2  | annRat             | 416 |
|       | 7.5.5.3  | DLoc               | 417 |
|       | 7.5.5.4  | SDLoc              | 417 |
|       | 7.5.5.5  | DLoc0              | 418 |
|       | 7.5.5.6  | GBWeight           | 419 |

|          |                    |     |
|----------|--------------------|-----|
| 7.5.5.7  | initialMalgrange   | 420 |
| 7.5.5.8  | initialIdealW      | 421 |
| 7.5.5.9  | inForm             | 422 |
| 7.5.5.10 | restrictionIdeal   | 423 |
| 7.5.5.11 | restrictionModule  | 424 |
| 7.5.5.12 | integralIdeal      | 425 |
| 7.5.5.13 | integralModule     | 426 |
| 7.5.5.14 | deRhamCohom        | 427 |
| 7.5.5.15 | deRhamCohomIdeal   | 428 |
| 7.5.5.16 | charVariety        | 429 |
| 7.5.5.17 | charInfo           | 430 |
| 7.5.5.18 | isFsat             | 432 |
| 7.5.5.19 | appelF1            | 432 |
| 7.5.5.20 | appelF2            | 433 |
| 7.5.5.21 | appelF4            | 433 |
| 7.5.5.22 | fourier            | 434 |
| 7.5.5.23 | inverseFourier     | 434 |
| 7.5.5.24 | bFactor            | 435 |
| 7.5.5.25 | intRoots           | 436 |
| 7.5.5.26 | poly2list          | 436 |
| 7.5.5.27 | fl2poly            | 437 |
| 7.5.5.28 | insertGenerator    | 438 |
| 7.5.5.29 | deleteGenerator    | 438 |
| 7.5.5.30 | isInt              | 439 |
| 7.5.5.31 | sortIntvec         | 439 |
| 7.5.6    | dmodideal_lib      | 440 |
| 7.5.6.1  | annfsLogIdeal      | 441 |
| 7.5.6.2  | annihilatorMultiFs | 442 |
| 7.5.6.3  | BSidealFromAnn     | 442 |
| 7.5.6.4  | BernsteinSatoIdeal | 443 |
| 7.5.6.5  | BFBoundsBudur      | 444 |
| 7.5.6.6  | annfalphiI         | 445 |
| 7.5.6.7  | extractS           | 446 |
| 7.5.7    | dmodvar_lib        | 446 |
| 7.5.7.1  | bfctVarIn          | 447 |
| 7.5.7.2  | bfctVarAnn         | 448 |
| 7.5.7.3  | SannfsVar          | 449 |
| 7.5.7.4  | makeMalgrange      | 450 |
| 7.5.8    | involut_lib        | 450 |
| 7.5.8.1  | findInvo           | 451 |
| 7.5.8.2  | findInvoDiag       | 452 |
| 7.5.8.3  | findAuto           | 453 |
| 7.5.8.4  | ncdetection        | 455 |
| 7.5.8.5  | involution         | 455 |
| 7.5.8.6  | isInvolution       | 456 |
| 7.5.8.7  | isAntiEndo         | 457 |
| 7.5.9    | gkdim_lib          | 457 |
| 7.5.9.1  | GKdim              | 457 |
| 7.5.10   | ncalg_lib          | 458 |
| 7.5.10.1 | makeUsl2           | 459 |
| 7.5.10.2 | makeUsl            | 459 |
| 7.5.10.3 | makeUgl            | 460 |
| 7.5.10.4 | makeUso5           | 461 |

|           |                        |     |
|-----------|------------------------|-----|
| 7.5.10.5  | makeUso6               | 461 |
| 7.5.10.6  | makeUso7               | 462 |
| 7.5.10.7  | makeUso8               | 463 |
| 7.5.10.8  | makeUso9               | 463 |
| 7.5.10.9  | makeUso10              | 464 |
| 7.5.10.10 | makeUso11              | 464 |
| 7.5.10.11 | makeUso12              | 465 |
| 7.5.10.12 | makeUsp1               | 465 |
| 7.5.10.13 | makeUsp2               | 466 |
| 7.5.10.14 | makeUsp3               | 467 |
| 7.5.10.15 | makeUsp4               | 468 |
| 7.5.10.16 | makeUsp5               | 468 |
| 7.5.10.17 | makeUg2                | 469 |
| 7.5.10.18 | makeUf4                | 469 |
| 7.5.10.19 | makeUe6                | 470 |
| 7.5.10.20 | makeUe7                | 470 |
| 7.5.10.21 | makeUe8                | 471 |
| 7.5.10.22 | makeQso3               | 472 |
| 7.5.10.23 | makeQsl2               | 473 |
| 7.5.10.24 | makeQsl3               | 473 |
| 7.5.10.25 | Qso3Casimir            | 475 |
| 7.5.10.26 | GKZsystem              | 475 |
| 7.5.11    | ncdecomp_lib           | 477 |
| 7.5.11.1  | CentralQuot            | 477 |
| 7.5.11.2  | CentralSaturation      | 478 |
| 7.5.11.3  | CenCharDec             | 478 |
| 7.5.11.4  | IntersectWithSub       | 479 |
| 7.5.12    | ncfactor_lib           | 480 |
| 7.5.12.1  | ncfactor               | 480 |
| 7.5.12.2  | facWeyl                | 485 |
| 7.5.12.3  | facFirstWeyl           | 486 |
| 7.5.12.4  | testNCfac              | 486 |
| 7.5.12.5  | facSubWeyl             | 487 |
| 7.5.12.6  | facShift               | 488 |
| 7.5.12.7  | facFirstShift          | 489 |
| 7.5.12.8  | homogfacNthWeyl        | 490 |
| 7.5.12.9  | homogfacNthQWeyl       | 490 |
| 7.5.12.10 | homogfacFirstQWeyl     | 491 |
| 7.5.12.11 | homogfacNthQWeyl_all   | 492 |
| 7.5.12.12 | homogfacFirstQWeyl_all | 510 |
| 7.5.12.13 | tst_ncfactor           | 511 |
| 7.5.13    | nchilbert_lib          | 514 |
| 7.5.13.1  | ncHilb                 | 514 |
| 7.5.13.2  | ncHilbertSeries        | 515 |
| 7.5.13.3  | ncHilbertPolynomial    | 516 |
| 7.5.13.4  | ncHilbertMultiplicity  | 517 |
| 7.5.13.5  | GKExp                  | 518 |
| 7.5.13.6  | mondim                 | 518 |
| 7.5.14    | dmodloc_lib            | 519 |
| 7.5.14.1  | Dlocalization          | 520 |
| 7.5.14.2  | WeylClosure            | 521 |
| 7.5.14.3  | WeylClosure1           | 522 |
| 7.5.14.4  | holonomicRank          | 522 |

|           |                            |     |
|-----------|----------------------------|-----|
| 7.5.14.5  | DsingularLocus             | 523 |
| 7.5.14.6  | polSol                     | 523 |
| 7.5.14.7  | polSolFiniteRank           | 524 |
| 7.5.14.8  | ratSol                     | 525 |
| 7.5.14.9  | bfctBound                  | 526 |
| 7.5.14.10 | annRatSyz                  | 526 |
| 7.5.14.11 | dmodGeneralAssumptionCheck | 528 |
| 7.5.14.12 | extendWeyl                 | 528 |
| 7.5.14.13 | polyVars                   | 529 |
| 7.5.14.14 | monomialInIdeal            | 529 |
| 7.5.14.15 | vars2pars                  | 530 |
| 7.5.14.16 | minIntRoot2                | 530 |
| 7.5.14.17 | maxIntRoot                 | 530 |
| 7.5.14.18 | dmodAction                 | 531 |
| 7.5.14.19 | dmodActionRat              | 532 |
| 7.5.14.20 | simplifyRat                | 532 |
| 7.5.14.21 | addRat                     | 533 |
| 7.5.14.22 | multRat                    | 533 |
| 7.5.14.23 | diffRat                    | 534 |
| 7.5.14.24 | commRing                   | 534 |
| 7.5.14.25 | rightNFWeyl                | 535 |
| 7.5.15    | ncfrac_lib                 | 535 |
| 7.5.15.1  | hasLeftDenom               | 536 |
| 7.5.15.2  | hasRightDenom              | 537 |
| 7.5.15.3  | isZeroNcfrac               | 537 |
| 7.5.15.4  | isOneNcfrac                | 538 |
| 7.5.15.5  | zeroNcfrac                 | 538 |
| 7.5.15.6  | oneNcfrac                  | 539 |
| 7.5.15.7  | ensureLeftNcfrac           | 539 |
| 7.5.15.8  | ensureRightNcfrac          | 541 |
| 7.5.15.9  | negateNcfrac               | 542 |
| 7.5.15.10 | isInvertibleNcfrac         | 542 |
| 7.5.15.11 | invertNcfrac               | 543 |
| 7.5.15.12 | testNcfrac                 | 544 |
| 7.5.15.13 | testNcfracExamples         | 544 |
| 7.5.16    | nchomolog_lib              | 544 |
| 7.5.16.1  | ncExt_R                    | 544 |
| 7.5.16.2  | ncHom                      | 545 |
| 7.5.16.3  | coHom                      | 545 |
| 7.5.16.4  | contraHom                  | 546 |
| 7.5.16.5  | dmodoublext                | 546 |
| 7.5.16.6  | is_cenBimodule             | 547 |
| 7.5.16.7  | is_cenSubbimodule          | 547 |
| 7.5.17    | ncloc_lib                  | 548 |
| 7.5.17.1  | isDenom                    | 548 |
| 7.5.17.2  | testNcloc                  | 549 |
| 7.5.17.3  | testNclocExamples          | 550 |
| 7.5.18    | ncModslimgb_lib            | 550 |
| 7.5.18.1  | ncmodslimgb                | 550 |
| 7.5.19    | ncpreim_lib                | 551 |
| 7.5.19.1  | eliminateNC                | 552 |
| 7.5.19.2  | preimageNC                 | 553 |
| 7.5.19.3  | admissibleSub              | 554 |



|           |                            |     |
|-----------|----------------------------|-----|
| 7.5.19.4  | isUpperTriangular          | 555 |
| 7.5.19.5  | appendWeight2Ord           | 556 |
| 7.5.19.6  | elimWeight                 | 557 |
| 7.5.19.7  | extendedTensor             | 557 |
| 7.5.20    | nctools_lib                | 559 |
| 7.5.20.1  | Gweights                   | 559 |
| 7.5.20.2  | weightedRing               | 560 |
| 7.5.20.3  | ndcond                     | 561 |
| 7.5.20.4  | Weyl                       | 562 |
| 7.5.20.5  | makeWeyl                   | 562 |
| 7.5.20.6  | makeHeisenberg             | 563 |
| 7.5.20.7  | Exterior                   | 564 |
| 7.5.20.8  | findimAlgebra              | 564 |
| 7.5.20.9  | superCommutative           | 565 |
| 7.5.20.10 | rightStd                   | 567 |
| 7.5.20.11 | rightNF                    | 568 |
| 7.5.20.12 | rightModulo                | 568 |
| 7.5.20.13 | moduloSlim                 | 569 |
| 7.5.20.14 | ncRelations                | 570 |
| 7.5.20.15 | isCentral                  | 570 |
| 7.5.20.16 | isNC                       | 571 |
| 7.5.20.17 | isCommutative              | 572 |
| 7.5.20.18 | isWeyl                     | 572 |
| 7.5.20.19 | UpOneMatrix                | 572 |
| 7.5.20.20 | AltVarStart                | 573 |
| 7.5.20.21 | AltVarEnd                  | 574 |
| 7.5.20.22 | IsSCA                      | 575 |
| 7.5.20.23 | makeModElimRing            | 577 |
| 7.5.20.24 | embedMat                   | 577 |
| 7.5.21    | olga_lib                   | 578 |
| 7.5.21.1  | locStatus                  | 578 |
| 7.5.21.2  | testLocData                | 581 |
| 7.5.21.3  | isInS                      | 581 |
| 7.5.21.4  | fracStatus                 | 582 |
| 7.5.21.5  | testFraction               | 583 |
| 7.5.21.6  | leftOre                    | 584 |
| 7.5.21.7  | rightOre                   | 585 |
| 7.5.21.8  | convertRightToLeftFraction | 586 |
| 7.5.21.9  | convertLeftToRightFraction | 587 |
| 7.5.21.10 | addLeftFractions           | 589 |
| 7.5.21.11 | multiplyLeftFractions      | 590 |
| 7.5.21.12 | areEqualLeftFractions      | 591 |
| 7.5.21.13 | isInvertibleLeftFraction   | 592 |
| 7.5.21.14 | invertLeftFraction         | 592 |
| 7.5.21.15 | isZeroFraction             | 593 |
| 7.5.21.16 | isOneFraction              | 594 |
| 7.5.21.17 | normalizeMonoidal          | 594 |
| 7.5.21.18 | normalizeRational          | 595 |
| 7.5.21.19 | testOlga                   | 595 |
| 7.5.21.20 | testOlgaExamples           | 595 |
| 7.5.22    | perron_lib                 | 596 |
| 7.5.22.1  | perron                     | 596 |
| 7.5.23    | purityfiltration_lib       | 597 |

|           |                                                                  |     |
|-----------|------------------------------------------------------------------|-----|
| 7.5.23.1  | projectiveDimension                                              | 597 |
| 7.5.23.2  | purityFiltration                                                 | 598 |
| 7.5.23.3  | purityTriang                                                     | 599 |
| 7.5.23.4  | gradeNumber                                                      | 600 |
| 7.5.23.5  | showgrades                                                       | 601 |
| 7.5.23.6  | allExtOfLeft                                                     | 601 |
| 7.5.23.7  | allExtOfRight                                                    | 602 |
| 7.5.23.8  | doubleExt                                                        | 603 |
| 7.5.23.9  | allDoubleExt                                                     | 603 |
| 7.5.23.10 | is_pure                                                          | 605 |
| 7.5.23.11 | purelist                                                         | 605 |
| 7.5.24    | qmatrix.lib                                                      | 606 |
| 7.5.24.1  | quantMat                                                         | 606 |
| 7.5.24.2  | qminor                                                           | 607 |
| 7.5.24.3  | SymGroup                                                         | 608 |
| 7.5.24.4  | LengthSymElement                                                 | 608 |
| 7.5.24.5  | LengthSym                                                        | 608 |
| 7.5.25    | ratgb.lib                                                        | 609 |
| 7.5.25.1  | ratstd                                                           | 609 |
| 7.6       | Graded commutative algebras (SCA)                                | 611 |
| 7.7       | LETTERPLACE                                                      | 613 |
| 7.7.1     | Examples of use of LETTERPLACE                                   | 613 |
| 7.7.2     | Example of use of LETTERPLACE over $\mathbb{Z}$                  | 614 |
| 7.7.3     | Functionality and release notes of LETTERPLACE                   | 617 |
| 7.7.4     | References and history of LETTERPLACE                            | 618 |
| 7.8       | Functions (letterplace)                                          | 619 |
| 7.8.1     | freeAlgebra (letterplace)                                        | 619 |
| 7.8.2     | lift (letterplace)                                               | 620 |
| 7.8.3     | liftstd (letterplace)                                            | 621 |
| 7.8.4     | ncgen                                                            | 621 |
| 7.8.5     | reduce (letterplace)                                             | 622 |
| 7.8.6     | rightstd (letterplace)                                           | 623 |
| 7.8.7     | std (letterplace)                                                | 623 |
| 7.8.8     | syz (letterplace)                                                | 623 |
| 7.8.9     | twostd (letterplace)                                             | 624 |
| 7.9       | Mathematical background (letterplace)                            | 625 |
| 7.9.1     | Free associative algebras                                        | 625 |
| 7.9.2     | Monomial orderings on free algebras                              | 625 |
| 7.9.3     | Groebner bases for two-sided ideals in free associative algebras | 627 |
| 7.9.4     | Bimodules and syzygies and lifts                                 | 628 |
| 7.9.5     | Letterplace correspondence                                       | 628 |
| 7.10      | LETTERPLACE libraries                                            | 629 |
| 7.10.1    | fpadim.lib                                                       | 629 |
| 7.10.1.1  | lpKDimCheck                                                      | 630 |
| 7.10.1.2  | lpKDim                                                           | 631 |
| 7.10.1.3  | lpMonomialBasis                                                  | 631 |
| 7.10.1.4  | lpHilbert                                                        | 632 |
| 7.10.1.5  | lpSickleDim                                                      | 633 |
| 7.10.2    | fpalgebras.lib                                                   | 634 |
| 7.10.2.1  | operatorAlgebra                                                  | 634 |
| 7.10.2.2  | serreRelations                                                   | 635 |
| 7.10.2.3  | fullSerreRelations                                               | 635 |

|           |                           |     |
|-----------|---------------------------|-----|
| 7.10.2.4  | ademRelations             | 636 |
| 7.10.2.5  | baumslagSolitar           | 636 |
| 7.10.2.6  | baumslagGroup             | 637 |
| 7.10.2.7  | crystallographicGroupP1   | 637 |
| 7.10.2.8  | crystallographicGroupPM   | 638 |
| 7.10.2.9  | crystallographicGroupPG   | 638 |
| 7.10.2.10 | crystallographicGroupP2MM | 639 |
| 7.10.2.11 | crystallographicGroupP2   | 639 |
| 7.10.2.12 | crystallographicGroupP2GG | 640 |
| 7.10.2.13 | crystallographicGroupCM   | 640 |
| 7.10.2.14 | crystallographicGroupC2MM | 641 |
| 7.10.2.15 | crystallographicGroupP4   | 641 |
| 7.10.2.16 | crystallographicGroupP4MM | 642 |
| 7.10.2.17 | crystallographicGroupP4GM | 642 |
| 7.10.2.18 | crystallographicGroupP3   | 643 |
| 7.10.2.19 | crystallographicGroupP31M | 643 |
| 7.10.2.20 | crystallographicGroupP3M1 | 644 |
| 7.10.2.21 | crystallographicGroupP6   | 645 |
| 7.10.2.22 | crystallographicGroupP6MM | 645 |
| 7.10.2.23 | dyckGroup1                | 646 |
| 7.10.2.24 | dyckGroup2                | 646 |
| 7.10.2.25 | dyckGroup3                | 647 |
| 7.10.2.26 | fibonacciGroup            | 647 |
| 7.10.2.27 | tetrahedronGroup          | 648 |
| 7.10.2.28 | triangularGroup           | 648 |
| 7.10.3    | fpaprops_lib              | 649 |
| 7.10.3.1  | lpNoetherian              | 649 |
| 7.10.3.2  | lpIsSemiPrime             | 650 |
| 7.10.3.3  | lpIsPrime                 | 650 |
| 7.10.3.4  | lpGlDimBound              | 651 |
| 7.10.3.5  | lpSubstitute              | 651 |
| 7.10.3.6  | lpCalcSubstDegBound       | 652 |
| 7.10.4    | freegb_lib                | 652 |
| 7.10.4.1  | isFreeAlgebra             | 653 |
| 7.10.4.2  | lpDegBound                | 653 |
| 7.10.4.3  | lpVarBlockSize            | 653 |
| 7.10.4.4  | lpNcgenCount              | 654 |
| 7.10.4.5  | lpDivision                | 654 |
| 7.10.4.6  | lpGBPres2Poly             | 655 |
| 7.10.4.7  | isOrderingShiftInvariant  | 655 |
| 7.10.4.8  | makeLetterplaceRing       | 656 |
| 7.10.4.9  | letplaceGBasis            | 657 |
| 7.10.4.10 | lieBracket                | 658 |
| 7.10.4.11 | setLetterplaceAttributes  | 658 |
| 7.10.4.12 | testLift                  | 659 |
| 7.10.4.13 | testSyz                   | 662 |
| 7.10.5    | ncHilb_lib                | 665 |
| 7.10.5.1  | nchilb                    | 665 |
| 7.10.5.2  | rcolon                    | 668 |
| 7.10.6    | ncrat_lib                 | 668 |
| 7.10.6.1  | ncInit                    | 669 |
| 7.10.6.2  | ncVarsGet                 | 669 |
| 7.10.6.3  | ncVarsAdd                 | 669 |

|           |                             |     |
|-----------|-----------------------------|-----|
| 7.10.6.4  | ncratDefine                 | 670 |
| 7.10.6.5  | ncratAdd                    | 670 |
| 7.10.6.6  | ncratSubtract               | 671 |
| 7.10.6.7  | ncratMultiply               | 671 |
| 7.10.6.8  | ncratInvert                 | 672 |
| 7.10.6.9  | ncratSPrint                 | 672 |
| 7.10.6.10 | ncratPrint                  | 673 |
| 7.10.6.11 | ncratFromString             | 673 |
| 7.10.6.12 | ncratFromPoly               | 673 |
| 7.10.6.13 | ncratPower                  | 674 |
| 7.10.6.14 | ncratEvaluateAt             | 674 |
| 7.10.6.15 | ncrepGet                    | 674 |
| 7.10.6.16 | ncrepAdd                    | 675 |
| 7.10.6.17 | ncrepSubtract               | 676 |
| 7.10.6.18 | ncrepMultiply               | 677 |
| 7.10.6.19 | ncrepInvert                 | 678 |
| 7.10.6.20 | ncrepPrint                  | 679 |
| 7.10.6.21 | ncrepDim                    | 680 |
| 7.10.6.22 | ncrepSubstitute             | 680 |
| 7.10.6.23 | ncrepEvaluate               | 681 |
| 7.10.6.24 | ncrepEvaluateAt             | 682 |
| 7.10.6.25 | ncrepIsDefinedDim           | 682 |
| 7.10.6.26 | ncrepIsDefined              | 683 |
| 7.10.6.27 | ncrepIsRegular              | 684 |
| 7.10.6.28 | ncrepRegularZeroMinimize    | 685 |
| 7.10.6.29 | ncrepRegularMinimize        | 686 |
| 7.10.6.30 | ncrepGetRegularZeroMinimal  | 686 |
| 7.10.6.31 | ncrepGetRegularMinimal      | 687 |
| 7.10.6.32 | ncrepPencilGet              | 688 |
| 7.10.6.33 | ncrepPencilCombine          | 689 |
| 7.11      | Release Notes (letterplace) | 689 |

## Appendix A Examples . . . . . 691

|       |                                        |     |
|-------|----------------------------------------|-----|
| A.1   | Programming                            | 691 |
| A.1.1 | Basic programming                      | 691 |
| A.1.2 | Writing procedures and libraries       | 692 |
| A.1.3 | Rings associated to monomial orderings | 695 |
| A.1.4 | Long coefficients                      | 696 |
| A.1.5 | Parameters                             | 698 |
| A.1.6 | Formatting output                      | 699 |
| A.1.7 | Cyclic roots                           | 699 |
| A.1.8 | Parallelization with ssi links         | 700 |
| A.1.9 | Dynamic modules                        | 701 |
| A.2   | Computing Groebner and Standard Bases  | 702 |
| A.2.1 | groebner and std                       | 702 |
| A.2.2 | Groebner basis conversion              | 704 |
| A.2.3 | slim Groebner bases                    | 706 |
| A.3   | Commutative Algebra                    | 706 |
| A.3.1 | Saturation                             | 707 |
| A.3.2 | Finite fields                          | 707 |
| A.3.3 | Elimination                            | 709 |
| A.3.4 | Free resolution                        | 712 |

|        |                                              |     |
|--------|----------------------------------------------|-----|
| A.3.5  | Handling graded modules                      | 715 |
| A.3.6  | Computation of Ext                           | 716 |
| A.3.7  | Depth                                        | 719 |
| A.3.8  | Factorization                                | 720 |
| A.3.9  | Primary decomposition                        | 721 |
| A.3.10 | Normalization                                | 723 |
| A.3.11 | Kernel of module homomorphisms               | 724 |
| A.3.12 | Algebraic dependence                         | 725 |
| A.4    | Singularity Theory                           | 726 |
| A.4.1  | Milnor and Tjurina number                    | 726 |
| A.4.2  | Critical points                              | 728 |
| A.4.3  | Polar curves                                 | 729 |
| A.4.4  | T1 and T2                                    | 731 |
| A.4.5  | Deformations                                 | 733 |
| A.4.6  | Invariants of plane curve singularities      | 736 |
| A.4.7  | Branches of space curve singularities        | 738 |
| A.4.8  | Classification of hypersurface singularities | 741 |
| A.4.9  | Resolution of singularities                  | 743 |
| A.5    | Invariant Theory                             | 744 |
| A.5.1  | G <sub>a</sub> -Invariants                   | 744 |
| A.5.2  | Invariants of a finite group                 | 745 |
| A.6    | Geometric Invariant Theory                   | 746 |
| A.6.1  | GIT-Fans                                     | 746 |
| A.7    | Non-commutative Algebra                      | 748 |
| A.7.1  | Left and two-sided Groebner bases            | 748 |
| A.7.2  | Right Groebner bases and syzygies            | 750 |
| A.8    | Applications                                 | 752 |
| A.8.1  | Solving systems of polynomial equations      | 752 |
| A.8.2  | AG codes                                     | 756 |

## **Appendix B Polynomial data . . . . . 759**

|       |                                        |     |
|-------|----------------------------------------|-----|
| B.1   | Representation of mathematical objects | 759 |
| B.2   | Monomial orderings                     | 760 |
| B.2.1 | Introduction to orderings              | 760 |
| B.2.2 | General definitions for orderings      | 760 |
| B.2.3 | Global orderings                       | 761 |
| B.2.4 | Local orderings                        | 761 |
| B.2.5 | Module orderings                       | 761 |
| B.2.6 | Matrix orderings                       | 762 |
| B.2.7 | Product orderings                      | 764 |
| B.2.8 | Extra weight vector                    | 764 |
| B.2.9 | Pseudo ordering L                      | 765 |

## Appendix C Mathematical background . . . . . 766

|     |                                                                   |     |
|-----|-------------------------------------------------------------------|-----|
| C.1 | Standard bases . . . . .                                          | 766 |
| C.2 | Hilbert function . . . . .                                        | 766 |
| C.3 | Syzygies and resolutions . . . . .                                | 767 |
| C.4 | Characteristic sets . . . . .                                     | 768 |
| C.5 | Gauss-Manin connection . . . . .                                  | 769 |
| C.6 | Toric ideals and integer programming . . . . .                    | 773 |
|     | C.6.1 Toric ideals . . . . .                                      | 773 |
|     | C.6.2 Algorithms . . . . .                                        | 773 |
|     | C.6.2.1 The algorithm of Conti and Traverso . . . . .             | 773 |
|     | C.6.2.2 The algorithm of Pottier . . . . .                        | 774 |
|     | C.6.2.3 The algorithm of Hosten and Sturmfels . . . . .           | 774 |
|     | C.6.2.4 The algorithm of Di Biase and Urbanke . . . . .           | 774 |
|     | C.6.2.5 The algorithm of Bigatti, La Scala and Robbiano . . . . . | 775 |
|     | C.6.3 The Buchberger algorithm for toric ideals . . . . .         | 775 |
|     | C.6.4 Integer programming . . . . .                               | 775 |
|     | C.6.5 Relevant References . . . . .                               | 776 |
| C.7 | Non-commutative algebra . . . . .                                 | 776 |
| C.8 | Decoding codes with Groebner bases . . . . .                      | 776 |
|     | C.8.1 Codes and the decoding problem . . . . .                    | 777 |
|     | C.8.2 Cooper philosophy . . . . .                                 | 778 |
|     | C.8.3 Generalized Newton identities . . . . .                     | 779 |
|     | C.8.4 Fitzgerald-Lax method . . . . .                             | 780 |
|     | C.8.5 Decoding method based on quadratic equations . . . . .      | 781 |
|     | C.8.6 References for decoding with Groebner bases . . . . .       | 782 |
| C.9 | References . . . . .                                              | 783 |

## Appendix D SINGULAR libraries . . . . . 785

|     |                                |     |
|-----|--------------------------------|-----|
| D.1 | standard_lib . . . . .         | 785 |
|     | D.1.1 qslimlib . . . . .       | 786 |
|     | D.1.2 par2varRing . . . . .    | 786 |
| D.2 | General purpose . . . . .      | 787 |
|     | D.2.1 all_lib . . . . .        | 787 |
|     | D.2.2 compregb_lib . . . . .   | 790 |
|     | D.2.3 general_lib . . . . .    | 790 |
|     | D.2.4 grobcov_lib . . . . .    | 791 |
|     | D.2.5 inout_lib . . . . .      | 796 |
|     | D.2.6 modular_lib . . . . .    | 796 |
|     | D.2.7 parallel_lib . . . . .   | 797 |
|     | D.2.8 polylib_lib . . . . .    | 798 |
|     | D.2.9 redcgs_lib . . . . .     | 799 |
|     | D.2.10 random_lib . . . . .    | 802 |
|     | D.2.11 resources_lib . . . . . | 802 |
|     | D.2.12 ring_lib . . . . .      | 803 |
|     | D.2.13 tasks_lib . . . . .     | 804 |
| D.3 | Linear algebra . . . . .       | 805 |
|     | D.3.1 matrix_lib . . . . .     | 806 |
|     | D.3.2 linalg_lib . . . . .     | 807 |
| D.4 | Commutative algebra . . . . .  | 809 |
|     | D.4.1 absfact_lib . . . . .    | 809 |
|     | D.4.2 algebra_lib . . . . .    | 809 |

|        |                      |     |
|--------|----------------------|-----|
| D.4.3  | assprimeszerodim_lib | 810 |
| D.4.4  | cisimplicial_lib     | 810 |
| D.4.5  | curveInv_lib         | 811 |
| D.4.6  | elim_lib             | 812 |
| D.4.7  | ellipticcovers_lib   | 812 |
| D.4.8  | ffmodstd_lib         | 813 |
| D.4.9  | grwalk_lib           | 815 |
| D.4.10 | homolog_lib          | 815 |
| D.4.11 | integralbasis_lib    | 816 |
| D.4.12 | intprog_lib          | 817 |
| D.4.13 | locnormal_lib        | 817 |
| D.4.14 | moddiq_lib           | 817 |
| D.4.15 | modnormal_lib        | 818 |
| D.4.16 | modstd_lib           | 818 |
| D.4.17 | monomialideal_lib    | 819 |
| D.4.18 | mprimdec_lib         | 820 |
| D.4.19 | mregular_lib         | 821 |
| D.4.20 | nfmodstd_lib         | 822 |
| D.4.21 | nfmodsyz_lib         | 823 |
| D.4.22 | noether_lib          | 823 |
| D.4.23 | normal_lib           | 824 |
| D.4.24 | normaliz_lib         | 825 |
| D.4.25 | pointid_lib          | 827 |
| D.4.26 | primdec_lib          | 828 |
| D.4.27 | primdecint_lib       | 830 |
| D.4.28 | primitiv_lib         | 830 |
| D.4.29 | realrad_lib          | 831 |
| D.4.30 | reesclos_lib         | 831 |
| D.4.31 | rstandard_lib        | 831 |
| D.4.32 | sagbi_lib            | 832 |
| D.4.33 | sing4ti2_lib         | 833 |
| D.4.34 | symodstd_lib         | 833 |
| D.4.35 | toric_lib            | 834 |
| D.5    | Algebraic geometry   | 834 |
| D.5.1  | brillnoether_lib     | 834 |
| D.5.2  | chern_lib            | 834 |
| D.5.3  | deRham_lib           | 838 |
| D.5.4  | divisors_lib         | 838 |
| D.5.5  | hess_lib             | 840 |
| D.5.6  | numerAlg_lib         | 840 |
| D.5.7  | numerDecom_lib       | 841 |
| D.5.8  | orbitparam_lib       | 842 |
| D.5.9  | paraplanecurves_lib  | 843 |
| D.5.10 | resbinomial_lib      | 844 |
| D.5.11 | resgraph_lib         | 846 |
| D.5.12 | resjung_lib          | 847 |
| D.5.13 | resolve_lib          | 847 |
| D.5.14 | reszeta_lib          | 848 |
| D.5.15 | schubert_lib         | 849 |
| D.5.16 | sheafcoh_lib         | 852 |
| D.5.17 | JMBTest_lib          | 852 |
| D.5.18 | JMSConst_lib         | 853 |
| D.6    | Singularities        | 854 |

|        |                            |     |
|--------|----------------------------|-----|
| D.6.1  | alexpoly_lib               | 854 |
| D.6.2  | arcpoint_lib               | 854 |
| D.6.3  | arnoldclassify_lib         | 855 |
| D.6.4  | classify_lib               | 856 |
| D.6.5  | classify2_lib              | 857 |
| D.6.6  | classify_aeq_lib           | 858 |
| D.6.7  | classifyceq_lib            | 859 |
| D.6.8  | classifyci_lib             | 859 |
| D.6.9  | classifyMapGerms_lib       | 860 |
| D.6.10 | curvepar_lib               | 861 |
| D.6.11 | deform_lib                 | 861 |
| D.6.12 | equising_lib               | 862 |
| D.6.13 | gmssing_lib                | 862 |
| D.6.14 | gmspoly_lib                | 864 |
| D.6.15 | hnoether_lib               | 864 |
| D.6.16 | kskernel_lib               | 866 |
| D.6.17 | mondromy_lib               | 866 |
| D.6.18 | qhmoduli_lib               | 867 |
| D.6.19 | realclassify_lib           | 867 |
| D.6.20 | sing_lib                   | 868 |
| D.6.21 | spcurve_lib                | 869 |
| D.6.22 | spectrum_lib               | 870 |
| D.6.23 | surfacesignature_lib       | 870 |
| D.7    | Invariant theory           | 871 |
| D.7.1  | finvar_lib                 | 871 |
| D.7.2  | ainvar_lib                 | 873 |
| D.7.3  | rinvar_lib                 | 873 |
| D.7.4  | invar_lib                  | 874 |
| D.7.5  | stratify_lib               | 875 |
| D.8    | Symbolic-numerical solving | 875 |
| D.8.1  | ffsolve_lib                | 875 |
| D.8.2  | interval_lib               | 876 |
| D.8.3  | presolve_lib               | 877 |
| D.8.4  | solve_lib                  | 878 |
| D.8.5  | triang_lib                 | 878 |
| D.8.6  | ntsolve_lib                | 879 |
| D.8.7  | recover_lib                | 879 |
| D.8.8  | rootisolation_lib          | 880 |
| D.8.9  | signcond_lib               | 882 |
| D.8.10 | zeroset_lib                | 882 |
| D.9    | Visualization              | 883 |
| D.9.1  | graphics_lib               | 883 |
| D.9.2  | latex_lib                  | 884 |
| D.9.3  | surf_lib                   | 885 |
| D.9.4  | surfex_lib                 | 885 |
| D.10   | Coding theory              | 886 |
| D.10.1 | brnoeth_lib                | 886 |
| D.10.2 | decodegb_lib               | 887 |
| D.11   | System and Control theory  | 888 |
| D.11.1 | Control theory background  | 888 |
| D.11.2 | control_lib                | 888 |
| D.11.3 | jacobson_lib               | 889 |
| D.11.4 | findifs_lib                | 890 |



|      |                                |     |
|------|--------------------------------|-----|
| D.12 | Teaching                       | 891 |
|      | D.12.1 aksaka_lib              | 891 |
|      | D.12.2 atkins_lib              | 892 |
|      | D.12.3 crypto_lib              | 893 |
|      | D.12.4 hyperel_lib             | 896 |
|      | D.12.5 teachstd_lib            | 897 |
|      | D.12.6 weierstr_lib            | 898 |
|      | D.12.7 rootsmr_lib             | 898 |
|      | D.12.8 rootsur_lib             | 899 |
| D.13 | Tropical Geometry              | 900 |
|      | D.13.1 cimonom_lib             | 900 |
|      | D.13.2 gfan_lib                | 901 |
|      | D.13.3 gitfan_lib              | 904 |
|      | D.13.4 polymake_lib            | 906 |
|      | D.13.5 realizationMatroids_lib | 908 |
|      | D.13.6 tropical_lib            | 909 |
|      | D.13.7 tropicalNewton_lib      | 913 |
| D.14 | Miscellaneous libraries        | 914 |
|      | D.14.1 arr_lib                 | 914 |
|      | D.14.2 customstd_lib           | 917 |
|      | D.14.3 phindex_lib             | 917 |
|      | D.14.4 polybori_lib            | 918 |
| D.15 | Experimental libraries         | 920 |
|      | D.15.1 autgradalg_lib          | 920 |
|      | D.15.2 combinat_lib            | 921 |
|      | D.15.3 difform_lib             | 921 |
|      | D.15.4 finitediff_lib          | 924 |
|      | D.15.5 goettsche_lib           | 925 |
|      | D.15.6 graal_lib               | 926 |
|      | D.15.7 GND_lib                 | 927 |
|      | D.15.8 gradedModules_lib       | 927 |
|      | D.15.9 maxlike_lib             | 929 |
|      | D.15.10 methods_lib            | 930 |
|      | D.15.11 modules_lib            | 930 |
|      | D.15.12 modwalk_lib            | 933 |
|      | D.15.13 multigrading_lib       | 934 |
|      | D.15.14 nets_lib               | 937 |
|      | D.15.15 pfd_lib                | 938 |
|      | D.15.16 polyclass_lib          | 939 |
|      | D.15.17 ringgb_lib             | 940 |
|      | D.15.18 rwalk_lib              | 940 |
|      | D.15.19 sets_lib               | 940 |
|      | D.15.20 stanleyreisner_lib     | 941 |
|      | D.15.21 swalk_lib              | 942 |
|      | D.15.22 systhreads_lib         | 942 |
|      | D.15.23 VecField_lib           | 943 |

|          |                                           |            |
|----------|-------------------------------------------|------------|
| <b>8</b> | <b>Release Notes .....</b>                | <b>944</b> |
| 8.1      | News and changes .....                    | 944        |
| 8.2      | Singular 3 and Singular 4 .....           | 950        |
| 8.2.1    | Version schema for Singular .....         | 950        |
| 8.2.2    | Notes for Singular users .....            | 950        |
| 8.2.3    | Notes for developers .....                | 952        |
| 8.2.4    | Building Singular .....                   | 952        |
| 8.2.5    | Side-by-side installation .....           | 952        |
| 8.3      | libSingular .....                         | 953        |
| 8.4      | Download instructions .....               | 953        |
| 8.5      | Used environment variables .....          | 953        |
| 8.6      | Unix installation instructions .....      | 954        |
| 8.7      | Windows installation instructions .....   | 955        |
| 8.8      | Macintosh installation instructions ..... | 955        |
| <b>9</b> | <b>Index .....</b>                        | <b>956</b> |