

The micrOMEGAs user's manual, version 4.0

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Abstract

We give an up-to-date description of the micrOMEGAs functions. Only the routines which are available for the users are described. Examples on how to use these functions can be found in the sample main programs distributed with the code.

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1 Introduction

`micrOMEGAs` is a code to calculate the properties of cold dark matter(CDM) in a generic model of particle physics. First developed to compute the relic density of dark matter, the code also computes the rates for dark matter direct and indirect detection. `micrOMEGAs` calculates CDM properties in framework of a model of particles interaction presented in CalcHEP format [19]. It is assumed that the model is invariant under a discrete symmetry like R-parity (which is even for all standard particles and odd for some new particles including the dark matter candidate) ensures the stability of the lightest odd particle (LOP). CalcHEP package is included in `micrOMEGAs` and used for matrix elements calculations. All annihilation and coannihilation channels are included in the computation of the relic density. This manual gives an up-to-date description of all `micrOMEGAs` functions. The methods used to compute the different dark matter properties are described in references [1, 2, 3, 4, 5, 6]. These references also contain a more complete description of the code. In the following the cold dark matter candidate also called LOP or weakly-interactive massive particle (WIMP) will be denoted by χ .

`micrOMEGAs` contains both C and Fortran routines. Below we describe only the C-version of the routines, in general we use the same names and the same types of argument for both C and Fortran functions. We always use `double(real*8)` variables for float point numbers and `int(INTEGER)` for integers. In this manual we use `FD` for file descriptor variables, the file descriptors are `FILE*` in C and `channel number` in Fortran. The symbol `&` before the names of variables in C-functions stands for the address of the variable. It is used for *output parameters*. In Fortran calls there is no need for `&` since all parameters are passed via addresses. In C programs one can substitute `NULL` for any output parameter which the user chooses to ignore. In Fortran one can substitute `cNull`, `iNull`, `r8Null` for unneeded parameters of *character*, *integer* and *real*8* type respectively.

A few C-functions use pointer variables that specify an *address* in the computer memory. Because pointers do not exist in Fortran one uses any other type of variable whose length is sufficient to store a computer address, for example `INTEGER*8`.

The complete format for all functions can be found in `sources/micromegas.h` (for C) or `sources/micromegas_f.h` (for Fortran). Examples on how to use these functions are provided in the `MSSM/main.c[F]` file.

2 Discrete symmetry in micrOMEGAs.

`micrOMEGAs` exploits the fact that models of dark matter exhibit a discrete symmetry and that the fields of the model transform as $\phi \rightarrow e^{i2\pi X_\phi} \phi$ where the charge $|X_\phi| < 1$. The particles of the Standard Model are assumed to transform trivially under the discrete symmetry, $X_\phi = 0$. In the following all particles with charge $X_\phi \neq 0$ will be called *odd* and the lightest odd particle will be stable. If neutral, it can be considered as a DM candidate. Typical examples of discrete symmetries used for constructing single DM models are Z_2 and Z_3 . Multi-component DM can arise in models with larger discrete symmetries. A simple example is a model with $Z_2 \times Z'_2$ symmetry, the particles charged under $Z_2(Z'_2)$ will belong to the first (second) dark sector. The lightest particle of each sector will be stable and therefore a potential DM candidate. Another example is a model with a Z_4 symmetry. The two dark sectors contain particles with $X_\phi = \pm 1/4$ and $X_\phi = 1/2$ respectively. The

lightest particle with charge $1/4$ is always stable while the lightest particle of charge $1/2$ is stable only if its decay into two particles of charge $1/4$ is kinematically forbidden. `micrOMEGAs` assumes that all odd particles have names starting with '~', for example, `~o1` for the lightest neutralino. In versions 4.X, to distinguish the particles with different transformation properties with respect to the discrete group, that is particles belonging to different 'dark' sectors, we use the convention that the names of particles in the second 'dark' sector starts with '~~'. Note that `micrOMEGAs` does not check the symmetry of the Lagrangian, it assumes that the name convention correctly identifies all particles with the same discrete symmetry quantum numbers.

3 Downloading and compilation of micrOMEGAs.

To download micrOMEGAs, go to

<http://lapth.cnrs.fr/micromegas>

and unpack the file received, `micromegas_4.0.tgz`, with the command

```
tar -xvzf micromegas_4.0.tgz
```

This should create the directory `micromegas_4.0/` which occupies about 40 Mb of disk space. You will need more disk space after compilation of specific models and generation of matrix elements. In case of problems and questions

email: micromegas@lapth.cnrs.fr

3.1 File structure of micrOMEGAs.

<code>Makefile</code>	to compile the kernel of the package
<code>CalcHEP_src/</code>	generator of matrix elements for micrOMEGAs
<code>Packages/</code>	external codes
<code>clean</code>	to remove compiled files
<code>man/</code>	— contains the manual: description of micrOMEGAs routines
<code>newProject</code>	to create a new model directory structure
<code>sources/</code>	micrOMEGAs code
<i>MSSM model directory</i>	
<code>MSSM/</code>	
<code>Makefile</code>	to compile the code and executable for this model
<code>main.c[pp]</code> <code>main.F</code>	files with sample <i>main</i> programs
<code>lib/</code>	directory for routines specific to this model
<code>Makefile</code>	to compile the auxiliary code library <i>lib/aLib.a</i>
<code>*.c *.f</code>	source codes of auxiliary functions
<code>work/</code>	CalcHEP working directory for the generation of matrix elements
<code>Makefile</code>	to compile the library <i>work/work_aux.a</i>
<code>models/</code>	directory for files which specifies the model
<code>vars1.mdl</code>	free variables
<code>func1.mdl</code>	constrained variables
<code>prtcls1.mdl</code>	particles
<code>lgrng1.mdl</code>	Feynman rules

tmp/	auxiliary directories for CalcHEP sessions
results/	
so_generated/	storage of matrix elements generated by CalcHEP
calchep/	directory for interactive CalcHEP sessions
<i>Directories of other models which have the same structure as MSSM/</i>	
NMSSM/	Next-to-Minimal Supersymmetric Model[23, 17]
CPVMSSM/	MSSM with complex parameters[26, 16]
IDM/	Inert Doublet Model[8]
LHM/	Little Higgs Model[7]
RHNM/	Right-handed Neutrino Model[21]
SM4/	Toy model with a 4th generation of lepton and neutrino DM
Z3M/	A model with scalar DM and Z_3 discrete symmetry—[9, 42]
Z4ID/	A model with Z_4 symmetry—[9, 42]
mdlIndep/	For model independent computation of DM signals

3.2 Compilation of CalcHEP and micrOMEGAs routines.

CalcHEP and micrOMEGAs are compiled by *gmake*. Go to the micrOMEGAs directory and launch

```
gmake
```

If *gmake* is not available, then *make* should work like *gmake*. In principle micrOMEGAs defines automatically the names of *C* and *Fortran* compilers and the flags for compilation. If you meet a problem, open the file which contains the compiler specifications, `CalcHEP_src/FlagsForSh`, improve it, and launch `[g]make` again. The file is written in *sh* script format and looks like

```
# C compiler
CC="gcc"
# Flags for C compiler
CFLAGS="-g -fsigned-char"
# Disposition of header files for X11
HX11=
# Disposition of lX11
LX11="-lX11"
# Fortran compiler
FC="gfortran"
FFLAGS="-fno-automatic"
.....
```

After a successful definition of compilers and their flags, micrOMEGAs rewrites the file *FlagsForSh* into *FlagsForMake* and substitutes its contents in all *Makefiles* of the package.

`[g]make clean` deletes all generated files, but asks permission to delete *FlagsForSh*.

`[g]make flags` only generates *FlagsForSh*. It allows to check and change flags before compilation of codes.

3.3 Module structure of main programs.

Each model included in micrOMEGAs is accompanied with sample files for C and Fortran programs which call micrOMEGAs routines, the *main.c*, *main.F* files. These files consist of several modules enclosed between the instructions

```
#ifdef XXXXX
.....
#endif

Each of these blocks contains some code for a specific problem

#define MASSES_INFO           //Displays information about mass spectrum
#define CONSTRAINTS          //Displays B->sgamma, Bs->mumu, etc
#define OMEGA                 //Calculates the relic density
#define INDIRECT_DETECTION    //Signals of DM annihilation in galactic halo
#define LoopGAMMA             //Gamma-Ray lines - available only in some models
#define RESET_FORMFACTORS     //Redefinition of Form Factors and other
                             //parameters
#define CDM_NUCLEON           //Calculates amplitudes and cross-sections
                             //for DM-nucleon collisions
#define CDM_NUCLEUS           //Calculates number of events for 1kg*day
                             //and recoil energy distribution for various nuclei
#define NEUTRINO              //Calculates flux of solar neutrinos and
                             //the corresponding muon flux
#define DECAYS                //Calculates decay widths and branching ratios
#define CROSS_SECTIONS        //Calculates cross sections
#define CLEAN                 // Removes intermediate files.
#define SHOWPLOTS             //Displays graphical plots on the screen
```

Other modules which require a link to external programs can also be defined, in this case the path to the required code must be specified, for example

```
#define HIGGSBOUNDS "../Packages/HiggsBounds-4.0.0"
```

All these modules are completely independent. The user can comment or uncomment any set of *define* instructions to suit his/her need.

3.4 Compilation of codes for specific models.

After the compilation of micrOMEGAs one has to compile the executable to compute DM related observables in a specific model. To do this, go to the model directory, say MSSM, and launch

```
[g]make main=main.c
```

It should generate the executable *main*. In the same manner

```
gmake main=filename.ext
```

generates the executable *filename* based on the source file *filename.ext*. For *ext* we support 3 options: '*c*', '*F*', '*c++*' which correspond to C, FORTRAN and C++ sources. [g]make called in the model directory automatically launches [g]make in subdirectories *lib* and *work* to compile

lib/aLib.a - library of auxiliary model functions, e.g. constraints,
work/work_aux.a - library of model particles, free and dependent parameters.

3.5 Command line parameters of main programs.

The default versions of *main.c/F* programs need some arguments which have to be specified in command lines. If launched without arguments *main* explains which parameter are needed. As a rule *main* needs the name of a file containing the numerical values of the free parameters of the model. The structure of a file record should be

Name Value # comment (optional)

For instance, an Inert Doublet model (IDM) input file contains

```
Mh      125    # mass of SM Higgs
MHC     200    # mass of charged Higgs ~H+
MH3     200    # mass of odd Higgs ~H3
MHX     63.2   # mass of ~X particle
la2     0.01   # \lambda_2 coupling
laL     0.01   # 0.5*(\lambda_3+\lambda_4+\lambda_5)
```

In other cases, different inputs can be required. For example, in the MSSM with input parameters defined at the GUT scale, the parameters have to be provided in a command line. Launching `./main` will return

```
      This program needs 4 parameters:
m0      common scalar mass at GUT scale
mhf     common gaugino mass at GUT scale
a0      trilinear soft breaking parameter at GUT scale
tb      tan(beta)
Auxiliary parameters are:
sgn +/-1, sign of Higgsino mass term (default 1)
Mtp     top quark pole mass
MbMb    Mb(Mb) scale independent b-quark mass
alfSMZ  strong coupling at MZ
Example: ./main 120 500 -350 10 1 173.1
```

4 Global Parameters

The list of the global parameters and their default values are given in Tables 1, 2. The numerical value for any of these parameters can be simply reset anywhere in the code. Numerical values of scalar quark form factors can also be reset by the `calcScalarQuarkFF` routine presented below. Some physical values evaluated by `micrOMEGAs` also are presented as global variables. See Table 3.

5 Setting of model parameters, spectrum calculation, parameter display.

The independent parameters that characterize a given model are listed in the file `work/models/vars1.mdl`. Three functions can be used to set the value of these parameters:

Table 1: Global input parameters of **micrOMEGAs**

Name	default value	units	comments
deltaY	0		Difference between DM/anti-DM abundances
K_dif	0.0112	kpc ² /Myr	The normalized diffusion coefficient
L_dif	4	kpc	Vertical size of the Galaxy diffusive halo
Delta_dif	0.7		Slope of the diffusion coefficient
Tau_dif	10 ¹⁶	s	Electron energy loss time
Vc_dif	0	km/s	Convective Galactic wind
Fermi_a	0.52	fm	nuclei surface thickness
Fermi_b	-0.6	fm	parameters to set the nuclei radius with
Fermi_c	1.23	fm	$R_A = cA^{1/3} + b$
Rsun	8.5	kpc	Distance from the Sun to the center of the Galaxy
Rdisk	20	kpc	Radius of the galactic diffusion disk
rhoDM	0.3	GeV/cm ³	Dark Matter density at Rsun
Vearth	225.2	km/s	Galaxy velocity of the Earth
Vrot	220	km/s	Galaxy rotation velocity at Rsun
Vesc	600	km/s	Escape velocity at Rsun

Table 2: Global parameters of **micrOMEGAs**: nucleon quark form factors

Proton		Neutron		comments
Name	value	Name	value	
ScalarFFPd	0.0191	ScalarFFNd	0.0273	Scalar form factor
ScalarFFPu	0.0153	ScalarFFNu	0.011	
ScalarFFPs	0.0447	ScalarFFNs	0.0447	
pVectorFFPd	-0.427	pVectorFFNd	0.842	Axial-vector form factor
pVectorFFPu	0.842	pVectorFFNu	-0.427	
pVectorFFPs	-0.085	pVectorFFNs	-0.085	
SigmaFFPd	-0.23	SigmaFFNd	0.84	Tensor form factor
SigmaFFPu	0.84	SigmaFFNu	-0.23	
SigmaFFPs	-0.046	SigmaFFNs	-0.046	

Table 3: Evaluated global variables

Name	units	comments	Evaluated by
CDM1	<i>character</i>	name of first DM particle	sortOddParticles
CDM2	<i>character</i>	name of second DM particle	sortOddParticles
Mcdm1	GeV	Mass of the first Dark Matter particle	sortOddParticles
Mcdm2	GeV	Mass of the second DM particles	sortOddParticles
Mcdm	GeV	$\min(M_{cdm1}, M_{cdm2})$ if both exist	sortOddParticles
dmAsymm		Asymmetry between relic density of DM - \overline{DM}	darkOmega[FO]
fracCDM2		fraction of CDM2 in relic density.	darkOmega2

- `assignVal(name, val)`

- `assignValW(name, val)`

assign value *val* to parameter *name*. The function `assignVal` returns a non-zero value if it cannot recognize a parameter name while `assignValW` writes an error message.

- `readVar(fileName)`

reads parameters from a file. The file should contain two columns with the following format (see also Section 3.5)

```
name      value
```

`readVar` returns zero when the file has been read successfully, a negative value when the file cannot be opened for reading and a positive value corresponding to the line where a wrong file record was found.

Note that in Fortran, numerical constants should be specified as `Real*8`, for example

```
call assignValW('SW', 0.473D0)
```

A common mistake is to use `Real*4`.

The constrained parameters of the model are stored in `work/models/func1.mdl`. Some of these parameters are treated as *public* parameters. The *public* parameters include by default all particle masses and all parameters whose calculation requires external functions (except simple mathematical functions like `sin`, `cos` ..). The parameters listed above any *public* parameters in `work/models/func1.mdl` are also treated as *public*. It is possible to enlarge the list of *public* parameters. There are two ways to do this. One can type `*` before a parameter name to make it *public* or one can add a special record in `work/models/func1.mdl`

```
%Local! |
```

Then all parameters listed above this record become *public*. See example in

```
MSSM/work/models/func1.mdl
```

The calculation of the particle spectrum and of all *public* model constraints is done with:

- `sortOddParticles(txt)`

which also sorts the odd particles with increasing masses, writes the name of the lightest odd particle in `txt` and assigns the value of the mass of the lightest odd particle to the global parameter `Mcdm`. This routine returns a non zero error code for a wrong set of parameters, for example parameters for which some constraint cannot be calculated. The name of the corresponding constraint is written in `txt`. This routine has to be called after a reassignment of any input parameter. These routine was updated for the case of two DM particles. `sortOddParticles` fills text parameters `CDM1` and `CDM2` which present the name of lightest particle which names are started with one and two tildes respectively. `Mcdm1` and `Mcdm2` are masses of these particles. If we have only one kind of DM then for the absent component $M_{cdm_i} = 0$ and $CDM_i = \text{NULL}$ (in Fortran the string is filled by space symbols).

- `qNumbers(pName, &spin2,&charge3,&cdim)`

returns the quantum numbers for the particle `pName`. Here `spin2` is twice the spin of the particle; `charge3` is three times the electric charge; `cdim` is the dimension of the representation of $SU(3)_c$, it can be 1, 3, -3 or 8. The parameters `spin2`, `charge3`, `cdim` are variables of type `int`. The value returned is the PDG code. If `pName` does not correspond to any particle of the model then `qNumbers` returns zero.

- `pdg2name(nPDG)`

returns the name of the particle which PDG code is `nPDG`. If this particle does not exist in the model the return value is `NULL`. In the FORTRAN version this function is Subroutine `pdg2name(nPDG, pName)` and the character variable `pName` consists of white spaces if the particle does not exist in the model.

- `pMass(pName)`

returns the numerical value of the particle mass.

- `nextOdd(n, &pMass)`

returns the name and mass of the n^{th} odd particle assuming that particles are sorted according to increasing masses. For $n = 0$ the output specifies the name and the mass of the CDM candidate. In the FORTRAN version this function is Subroutine `nextOdd(n, pName, pMass)`

- `findVal(name,&val)`

finds the value of variable `name` and assigns it to parameter `val`. It returns a non-zero value if it cannot recognize a parameter name.

- `findValW(name)` just returns the value of variable `name` and writes an error message if it cannot recognize a parameter name. The variables accessible by these commands are all free parameters and the constrained parameters of the model (in file `model/func1.mdl`) treated as *public*.

The following routines are used to display the value of the independent and the constrained *public* parameters:

- `printVar(FD)`

prints the numerical values of all independent and *public* constrained parameters into `FD`

- `printMasses(FD, sort)`

prints the masses of 'odd' particles (those whose names started with `~`). If `sort` $\neq 0$ the masses are sorted so the mass of the CDM is given first.

- `printHiggsMasses(FD, sort)`

prints the masses and widths of 'even' scalars.

6 Relic density calculation.

6.1 Switches and auxiliary routines

- **VWdecay, VZdecay**

Switches to turn on/off processes with off-shell gauge bosons in the final state for DM annihilation and particle decays. If **VW/VZdecay=1**, the 3-body final states will be computed for annihilation processes only while if **VW/VZdecay=2** they will be included in coannihilation processes as well. By default the switches are set to (**VW/VZdecay=1**).¹ Note that **micrOMEGAs** calculates the width of each particle only once and stores the result in *Decay Table*. A second call to the function **pWidth** (whether an explicit call or within the computation of a cross section) will return the same result even if the user has changed the **VW/VZdecay** switch. We recommend to call

- **cleanDecayTable()**

after changing the switches to force **micrOMEGAs** to recalculate the widths taking into account the new value of **VW/VZdecay**. In Fortran, the subroutine

- **setVWdecay(VWdecay, VZdecay)** changes the switches and calls **cleanDecayTable()**. The **sortOddParticles** command which must be used to recompute the particle spectrum after changing the model parameters also clears the decay table.

If the particle widths were stored in the SLHA file (Susy Les Houches Accord [18]) downloaded by **micrOMEGAs**, then the SLHA value will be used by and are thus insensitive to the **VW/VZdecay** switches. To avoid downloading particle widths, one can use **slhaRead(fileName, mode=4)** to read the content of the SLHA file, see the description in Section 13.5.

The temperature dependence of the effective number of degrees of freedom can be set with

- **loadHeffGeff(char*fname)**

allows to modify the temperature dependence of the effective number of degrees of freedom by loading the file **fname** which contains a table of $h_{eff}(T), g_{eff}(T)$. A positive return value corresponds to the number of lines in the table. A negative return value indicates the line which creates a problem (e.g. wrong format), the routine returns zero when the file **fname** cannot be opened. The default file is **std_thg.tab** and is downloaded automatically if **loadHeffGeff** is not called in user's main program. Five other files are provided in the sources/data directory: **HP_A_thg.tab**, **HP_B_thg.tab**, **HP_B2_thg.tab**, **HP_B3_thg.tab**, and **HP_C_thg.tab**. They correspond to sets A, B, B2, B3, C in [10]. The user can substitute his/her own table as well, if so, the file must contain three columns containing the numerical values for T, h_{eff}, g_{eff} , the data file can also contain comments for lines starting with #.

- **improveCrossSection(p1,p2,p3,p4, Pcm, &address)**

allows to substitute a new cross-section for a given process. Here **p1, p2** are the names of particles in the initial state and **p3, p4** those in the final state. **Pcm** is the center of mass momentum and **address** ... This function is useful if for example the user wants to include her/his one-loop improved cross-section calculation in the relic density computation.

¹Including the 3-body final states can significantly increase the execution time for the relic density.

6.2 Calculation of relic density for one-component Dark Matter models.

All routines to calculate the relic density in version 3 are still available in this version. For these routines, the difference between the two dark sectors DM is ignored. These routines are intended for models with either a Z_2 or Z_3 discrete symmetry.

- **vSigma(T,Beps,fast)**

calculates the thermally averaged cross section for DM annihilation times velocity at a temperature T [GeV], see formula (2.6) in [1]. The value for σv is expressed in [pb]. The parameter *Beps* defines the criteria for including coannihilation channels as for **darkOmega** described below. The *fast* = 1/0 option switches between the *fast/accurate* calculation. The global array **vSigmaTCh** contains the contribution of different channels to **vSigma**. **vSigmaTCh[i].weight** specifies the relative weight of the i^{th} channel

vSigmaTCh[i].prtc1[j] ($j=0, 4$) defines the particles names for the i^{th} channel.

The last record in **vSigmaTCh** array has zero weight and NULL particle names. In the Fortran version, the function **vSigmaTCh(i,weight,pdg,process)** serves the same purpose. This function returns 0 if i exceeds the number of annihilation channels and 1 otherwise, $i \geq 1$. *real*8 weight* gives the relative contribution of each annihilation channel. *integer pdg(5)* contains the codes of incoming and outgoing particles in the annihilation process. *character*40 process* contains a textual description of annihilation processes.

The cross sections for semi - annihilation processes contribute to **vSigma** with a factor $\frac{1}{2}$ as described in [9]. Furthermore if an outgoing particle has a non-zero decay branching ratio to odd particles, then the annihilation cross section is reduced correspondingly.

- **darkOmega(&Xf,fast,Beps)**

calculates the dark matter relic density Ωh^2 . This routine solves the differential evolution equation using the Runge-Kutta method. $X_f = M_{cdm}/T_f$ characterizes the freeze-out temperature. The value of X_f is given for information and is also used as an input for the routine that gives the relative contribution of each channel to Ωh^2 , see **printChannels** below. The *fast* = 1 flag forces the fast calculation (for more details see Ref. [2]). This is the recommended option and gives an accuracy around 1%. The parameter *Beps* defines the criteria for including a given coannihilation channel in the computation of the thermally averaged cross-section, [2]. The recommended value is $Beps = 10^{-4} - 10^{-6}$ whereas if $Beps = 1$ only annihilation of the lightest odd particle is computed.

- **darkOmegaF0(&Xf, fast, Beps)**

calculates the dark matter relic density Ωh^2 using the freeze-out approximation. Both **darkOmega** and **darkOmegaF0** set parameter **fracCDM2** 0 or 1 depending on the name of lightest odd particle

- **printChannels(Xf,cut,Beps,prcnt,FD)**

writes into **FD** the contributions of different channels to $(\Omega h^2)^{-1}$. Here **Xf** is an input parameter which should be evaluated first in **darkOmega[F0]**. Only the channels whose relative contribution is larger than **cut** will be displayed. *Beps* plays the same role as the **darkOmega[F0]** routine. If *prcnt* $\neq 0$ the contributions are given in percent. Note that for this specific purpose we use the freeze-out approximation.

- **oneChannel(Xf,Beps,p1,p2,p3,p4)**

calculates the relative contribution of the channel $p1, p2 \rightarrow p3, p4$ to $(\Omega h^2)^{-1}$. $p1, \dots, p4$ are particle names. To sum over several channels one can write "*" instead of a particle name, e.g. "*" in place of $p1$.

- **omegaCh** is an array that contains the relative contribution and particle names for each annihilation channel. In the Fortran version one uses instead the function **omegaCh(i,weight,pdg,process)**. These array and function are similar to **vSigmaTCh** described above. The array **omegaCh** is filled after calling either **darkOmegaF0** or **printChannels**.

There is an option to calculate relic density in models with $DM - \overline{DM}$ asymmetry. In this case we assume that the number difference $DM - \overline{DM}$ is conserved in reactions. Thus initial small difference in abundances can lead to large DM asymmetry after freeze-out similar to baryonic asymmetry.

- **deltaY**

describes the difference between the DM and anti-DM abundances for the models where number of DM particles minus number of anti-DM ones is conserved in reactions of decay and collisions. In such models **deltaY** is a constant during thermal evolution of Universe. See Ref. [6].

- **dmAsymm**

is defined by equation

$$\Omega_{\pm} = \Omega \frac{1 \pm \text{dmAsymm}}{2}$$

and evaluated by **micrOMEGAs** while calculating the relic density with an initial asymmetry **deltaY**. See [6]. This parameter can also be reset after the relic density computation and will be taken into account for direct and indirect detection rates.

6.3 Calculation of relic density for two-component Dark Matter models.

- **darkOmega2(fast, Beps)**

Calculates Ωh^2 for either one- or two-components DM models. In the latter case it should give the same result as **darkOmega**. The parameters **fast** and **Besp** have the same meaning as for the **darkOmega** routine. The returned value corresponds to the sum of the contribution of the two DM components to Ωh^2 . **darkOmega2** also calculates the global parameter **fracCDM2** which represents the mass fraction of CDM2 in the total relic density

$$\Omega = \Omega_1 + \Omega_2 \tag{1}$$

$$\text{fracCDM2} = \frac{\Omega_2}{\Omega} \tag{2}$$

This parameter is then used in routines which calculate the total signal from both DM candidates in direct, indirect and neutrino telescope experiments, **nucleusRecoil**, **calcSpectrum**, and **neutrinoFlux**. The user can change the global **fracCDM2** parameter before the calculation of these observables to take into account the fact that the value of the DM fraction in the Milky Way could be different than in the early Universe.

The routines that were described in section 6.2 are not available for two-component DM models. In particular the individual channel contribution to the relic density cannot be computed and DM asymmetry is ignored. After calling **darkOmega2** the user can check the cross sections for each class of reactions (but not for individual processes) which were tabulated during the calculation of the relic density. The functions

- **vsabbF(T)**

computes the sum of cross sections for each class of reactions ($a, b = 0, 1, 2$) tabulated during the calculation of the relic density. Here **T** is the temperature in [GeV] and the

return value is $v\sigma$ in [pb]. These functions are defined in the interval $[T_{\text{start}}, T_{\text{end}}]$ where T_{start} is defined by `darkOmega2`, $T_{\text{end}}=10^{-3}\text{GeV}$. Specifically the functions available are

vs1100F vs1110F vs1120F vs1112F vs1122F vs1210F vs1211F
vs1220F vs1222F vs2200F vs2210F vs2220F vs2211F vs2221F

The temperature dependence of the equilibrium abundances can also be called by the user, the functions are named $Y_1(T)$ and $Y_2(T)$ and are defined only in the interval $T \in [T_{\text{end}}, T_{\text{start}}]$. The equilibrium abundances are accessible via the $Y_{\text{eq1}}(T)$, $Y_{\text{eq2}}(T)$ functions and the deviation from equilibrium by the functions $dY1F(T) = Y1F(T) - Y1_{\text{eq}}(T)$ and $dY2F(T) = Y2F(T) - Y2_{\text{eq}}(T)$.

7 Direct detection.

7.1 Amplitudes for elastic scattering

- `nucleonAmplitudes(CDM, qBOX, pAsi, pAsd, nAsi, nAsd)`
calculates the amplitudes for CDM-nucleon elastic scattering at zero momentum. `pAsi(nAsi)` are spin independent amplitudes for protons(neutrons) whereas `pAsd(nAsd)` are the corresponding spin dependent amplitudes. Each of these four parameters is an array of dimension 2. The zeroth (first) element of these arrays gives the χ -nucleon amplitudes whereas the second element gives $\bar{\chi}$ -nucleon amplitudes. Amplitudes are normalized such that the total cross section for either χ or $\bar{\chi}$ cross sections is²

$$\sigma_{\text{tot}} = \frac{4M_\chi^2 M_N^2}{\pi(M_\chi + M_N)^2} (|A^{SI}|^2 + 3|A^{SD}|^2) \quad (3)$$

If `qBOX=NULL` (`qBOX=NoLoop` in Fortran) tree level amplitudes are computed. In MSSM-type models with a spin 1/2 WIMP and scalar "squarks", `qBOX=FeScLoop` uses an improved tree-level calculation. `nucleonAmplitudes` returns a value different from zero only when there is an internal problem in the calculation.

`nucleonAmplitudes` depends implicitly on form factors which describe the quark contents in the nucleon. These form factors are global parameters (see Table 1 for default values)

$$TypeFFPq \quad TypeFFNq$$

where *Type* is either "Scalar", "pVector", or "Sigma", FFP and FFN denote proton and neutron and *q* specifies the quark, *d*, *u* or *s*. Heavy quark coefficients are calculated automatically.

- `calcScalarQuarkFF($m_u/m_d, m_s/m_d, \sigma_{\pi N}, \sigma_s$)`
computes the scalar coefficients for the quark content in the nucleon from the quark mass ratios $m_u/m_d, m_s/m_d$ as well as from $\sigma_{\pi N}$ and σ_s . The default values given in Table 2 are obtained for $\sigma_s = 42\text{MeV}$, $\sigma_{\pi N} = 34\text{MeV}$, $m_u/m_d = 0.56$, $m_s/m_d = 20.2$ [11]. The function `calcScalarQuarkFF(0.553, 18.9, 55., 243.5)` will reproduce the default values of the scalar quark form factors used in micrOMEGAs2.4 and earlier versions.

²All parameters are in GeV.

7.2 Scattering on nuclei

• `nucleusRecoil(f,A,Z,J,Sxx,qBOX,dNdE)`

This is the main routine of the direct detection module. The input parameters are:

- ◊ `f` - the DM velocity distribution normalized such that

$$\int_0^\infty v f(v) dv = 1$$

The units are km/s for v and s^2/km^2 for $f(v)$.

- ◊ `A` - atomic number of nucleus;
- ◊ `Z` - number of protons in the nucleus, predefined values for a wide set of isotopes are called with $Z_{\{Name\}}$;
- ◊ `J` - nucleus spin, predefined values for a wide set of isotopes are called with $J_{\{Name\}}\{atomic_number\}$.
- ◊ `Sxx` - is a routine which calculates nucleus form factors for spin-dependent interactions (`S00`, `S01`, `S11`), it depends on the momentum transfer in fm^{-1} . The available form factors are

```
SxxF19   SxxNa23   SxxNa23A  SxxAl27   SxxSi29   SxxSi29A
SxxK39   SxxGe73   SxxGe73A  SxxNb92   SxxTe125  SxxTe125A
SxxI127  SxxI127A  SxxXe129  SxxXe129A
SxxXe131 SxxXe131A SxxXe131B
```

The last character is used to distinguish different implementations of the form factor for the same isotope, see details in [4].

- ◊ `qBOX` - a parameter needed by `nucleonAmplitudes`, see the description above.

The form factors for the spin independent (SI) cross section are defined by a Fermi distribution and depend on the global parameters `Fermi_a`, `Fermi_b`, `Fermi_c`.

The returned value gives the number of events per day and per kilogram of detector material. The result depends implicitly on the global parameter `rhoDM`, the density of DM near the Earth. The distribution over recoil energy is stored in the array $dNdE$ which by default has $Nstep = 200$ elements. The value in the i^{th} element corresponds to

$$dNdE[i] = \left. \frac{dN}{dE} \right|_{E=i*keV*step}$$

in units of $(1/keV/kg/day)$. By default `step` is set to 1.

For a complex WIMP, `nucleusRecoil` averages over χ and $\bar{\chi}$. For example for ^{73}Ge , a call to this routine will be:

```
nucleusRecoil(Maxwell,73,Z_Ge,J_Ge73,SxxGe73,FeScLoop,dNdE);
```

- **setRecoilEnergyGrid(step,Nstep)**

changes the values of **step** and **Nstep** for the computation of **dNdE**.

- **Maxwell(v)**

returns

$$f(v) = \frac{c_{\text{norm}}}{v} \int_{|\vec{v}| < v_{\text{max}}} d^3\vec{v} \exp\left(-\frac{(\vec{v} - V_{\text{Earth}})^2}{\Delta v^2}\right) \delta(v - |\vec{v}|)$$

which corresponds to the isothermal model. Default values for the global parameters $\Delta v = V_{\text{rot}}$, $v_{\text{max}} = V_{\text{esc}}$, **Vearth** are listed in Table 1. c_{norm} is the normalization factor. This function is an argument of the **nucleusRecoil** function described above.

- **nucleusRecoil0(f,A,Z,J,Sp,Sn,qBOX,dNdE)**

is similar to the function **nucleusRecoil** except that the spin dependent nuclei form factors are described by Gauss functions whose values at zero momentum transfer are defined by the coefficients **Sp,Sn** [4]. Predefined values for the coefficients **Sp,Sn** are included for the nuclei listed in **nucleusrecoil** as well as ${}^3\text{He}$, ${}^{133}\text{Cs}$. Their names are

$$\begin{aligned} &Sp_{\{Nucleus\ Name\}}\{Atomic\ Number\} \\ &Sn_{\{Nucleus\ Name\}}\{Atomic\ Number\} \end{aligned}$$

One can use this routine for nuclei whose form factors are not known.

7.3 Auxiliary routines

Two auxiliary routines are provided to work with the energy spectrum computed with **nucleusRecoil** and **nucleusRecoil0**.

- **cutRecoilResult(dNdE,E1,E2)**

calculates the number of events in an energy interval defined by the values **E1,E2** in keV.

- **displayRecoilPlot(dNdE,title,E1,E2)**

plots the generated energy distribution **dNdE**. Here **title** is a character string specifying the title of the plot and **E1,E2** are minimal and maximal values for the displayed energy in keV.

8 Indirect detection

8.1 Interpolation and display of spectra

Various spectra and fluxes of particles relevant for indirect detection are stored in arrays with **NZ=250** elements. To decode and interpolate the spectrum array one can use the following functions:

- **SpectdNdE(E,spectTab)**

interpolates the tabulated spectra and returns the particle distribution **dN/dE** where **E** is the energy in GeV. For a particle number distribution the returned value is given in GeV^{-1} units while a particle flux is expressed in $(\text{sec cm}^2 \text{ sr GeV})^{-1}$.

To display the spectra as a function of energy one can use

- **displaySpectrum(spectTab,message,Emin,Emax)**

where **message** is a text string which gives a title to the plot and **Emin** and **Emax** define energy cuts.

Although structure of spectrum array actually does not need for the user, we present it below. The first (zeroth) element of array contains the maximum of spectrum energy E_{max} . As a rule E_{max} is a mass of DM particle. The i^{th} element ($1 \leq i < NZ - 1$) of the spectrum array contains the value of $E_i \frac{dN}{dE_i}$ where $E_i = E_{max} e^{Zi(i)}$,

- $Zi(i) = -7 \left(\frac{i-1}{NZ} \right)^{1.5} \log 10$.

That is the array points cover the energy interval $E_{max} \geq E > 10^{-7} E_{max}$.

8.2 Annihilation spectra

- **calcSpectrum(key,Sg,Se,Sp,Sne,Snm,Snl,&err)**

calculates the spectra of DM annihilation at rest and returns σv in cm^3/s . The calculated spectra for γ , e^+ , \bar{p} , ν_e , ν_μ , ν_τ are stored in arrays of dimension NZ as described above: **Sg**, **Se**, **Sp**, **Sne**, **Snm**, **Snl**. To remove the calculation of a given spectra, substitute **NULL** for the corresponding argument. **key** is a switch to include the polarisation of the W, Z bosons (**key=1**) or photon radiation (**key=2**). Note that final state photon radiation (FSR) is always included. When **key=2** the 3-body process $\chi\chi' \rightarrow XX + \gamma$ is computed for those subprocesses which either contain a light particle in the t-channel (of mass less than 1.2 Mcdm) or an outgoing W when Mcdm>500GeV. The FSR is then subtracted to avoid double counting. Only the electron/positron spectrum is modified with this switch. When **key=4** the contributions for each channel to total annihilation rate are written on the screen. More than one option can be switched on simultaneously by adding the corresponding values for **key**. For example both the W polarization and photon radiation effects are included if **key=3**. A problem in the spectrum calculation will produce a non zero error code, $err \neq 0$. **calcSpectrum** interpolates and sums spectra obtained by Pythia. The spectra tables are provided only for Mcdm> 2GeV. The results for a dark matter mass below 2 GeV will therefore be wrong, for example an antiproton spectrum with kinematically forbidden energies will be produced. A warning is issued for Mcdm< 2GeV.

- **spectrInfo(Xmin,spectrTab,&Ntot,&Xtot)**

provides information on the spectra generated. Here **Xmin** defines the minimum cut for the energy fraction $x=E/Mcdm$, **Ntot** and **Xtot** are calculated parameters which give on average the total number and the energy fraction of the final particles produced per collision. Note that the upper limit is **Xtot=2**.

- **vSigmaCh** is an array that contains the relative contribution and particle names for each annihilation channel. It is similar to **vSigmaTCh** described above, but list of particle has 5 positions to describe gamma radiation. For 2->2 processes **vSigmaCh[n].prtel[4]=NULL**. The array **vSigmaCh** is filled by **calcSpectrum**. In the Fortran version one uses instead the function

vSigmaCh(i,weight,pdg,process)

which also is similar to Fortran **vSigmaTCh** described above.

8.3 Distribution of Dark Matter in Galaxy.

The indirect DM detection signals depend on the DM density in our Galaxy. The DM density is given as the product of the local density at the Sun with the halo profile function

$$\rho(r) = \rho_{\odot} F_{halo}(r) \quad (4)$$

In `micrOMEGAs` ρ_{\odot} is a global parameter `rhoDM` and the Zhao profile [12]

$$F_{halo}(r) = \left(\frac{R_{\odot}}{r} \right)^{\gamma} \left(\frac{r_c^{\alpha} + R_{\odot}^{\alpha}}{r_c^{\alpha} + r^{\alpha}} \right)^{\frac{\beta-\gamma}{\alpha}} \quad (5)$$

with $\alpha = 1, \beta = 3, \gamma = 1, r_c = 20[kpc]$ is used by default. R_{\odot} , the distance from the Sun to the galactic center, is also a global parameter, `Rsun`. The parameters of the Zhao profile can be reset by

- `setProfileZhao($\alpha, \beta, \gamma, r_c$)`

The function to set another density profile is

- `setHaloProfile($F_{halo}(r)$)`

where $F_{halo}(r)$ is any function which depends on the distance from the galactic center, r , defined in [kpc] units. For instance, `setHaloProfile(hProfileEinasto)` sets Einasto profile

$$F_{halo}(r) = \exp \left[-\frac{2}{\alpha} \left(\left(\frac{r}{R_{\odot}} \right)^{\alpha} - 1 \right) \right]$$

where by default $\alpha = 0.17$, but can be changed by

- `setProfileEinasto(α)`

The command `setHaloProfile(hProfileZhao)` sets back the Zhao profile. Note that both `setProfileZhao` and `setProfileEinasto` call `setHaloProfile` to define the corresponding profile.

Dark matter annihilation in the Galaxy depends on the average of the square of the DM density, $\langle \rho^2 \rangle$. This quantity can be significantly larger than $\langle \rho \rangle^2$ when clumps of DM are present [13]. In `micrOMEGAs`, we use a simple model where f_{cl} is a constant that characterizes the fraction of the total density due to clumps and where all clumps occupy the same volume V_{cl} and have a constant density ρ_{cl} . Assuming clumps do not overlap, we get

$$\langle \rho^2 \rangle = \rho^2 + f_{cl} \rho_{cl} \rho. \quad (6)$$

This simple description allows to demonstrate the main effect of clumps: far from the Galactic center the rate of DM annihilation falls as $\rho(r)$ rather than as $\rho(r)^2$. The parameters ρ_{cl} and f_{cl} have zero default values. The routine to change these values is

- `setClumpConst(f_{cl}, ρ_{cl})`

To be more general, one could assume that ρ_{cl} and f_{cl} depend on the distance from the galactic center. The effect of clumping is then described by the equation

$$\langle \rho^2 \rangle(r) = \rho(r)(\rho(r) + \rho_{clump}^{eff}(r)), \quad (7)$$

and the function

- `setRhoClumps(ρ_{clump}^{eff})`

allows to implement a more sophisticated clump structure. To return to the default treatment of clumps call `setRhoClumps(rhoClumpsConst)` or `setClumpConst`.

8.4 Photon signal

The photon flux does not depend on the diffusion model parameters but on the angle ϕ between the line of sight and the center of the galaxy as well as on the annihilation spectrum into photons

- `gammaFluxTab(fi,dfi,sigmav,Sg,Sobs)`

multiplies the annihilation photon spectrum with the integral over the line of sight and over the opening angle to give the photon flux. `fi` is the angle between the line of sight and the center of the galaxy, `dfi` is half the cone angle which characterizes the detector resolution (the solid angle is $2\pi(1 - \cos(df_i))$), `sigmav` is the annihilation cross section, `Sg` is the DM annihilation spectra. `Sobs` is the spectra observed.

The function `gammaFluxTab` can be used for the neutrino spectra as well.

- `gammaFlux(fi,dfi,vcs)`

is the same function as `gammaFluxTab` above but corresponds to a discrete photon spectrum. `vcs` is the annihilation cross section, for instance in the \hat{M} SSM it is calculated with the `loopGamma` function. The function returns the number of photons per cm^2 of detector surface per second. Note that for $\chi\chi \rightarrow \gamma\gamma$ the result should be multiplied by a factor 2 as each annihilation leads to the production of two photons.

8.5 Propagation of charged particles.

The observed spectrum of charged particles strongly depends on their propagation in the Galactic Halo. The propagation depends on the global parameters

`K_dif, Delta_dif, L_dif, R_sun, R_disk`

as well as

`Tau_dif (positrons), Vc_dif (antiprotons)`

- `posiFluxTab(Emin,sigmav, Se, Sobs)`

computes the positron flux at the Earth. Here `sigmav` and `Se` are values obtained by `calcSpectrum`. `Sobs` is the positron spectrum after propagation. `Emin` is the energy cut to be defined by the user. Note that a low value for `Emin` increases the computation time. The format is the same as for the initial spectrum. The function `SpectrdNdE(E,Sobs)` described above can also be used for the interpolation, in this case the flux is returned in $(GeV \text{ s } cm^2 sr)^{-1}$.

- `pbarFlux(E,dSigmavdE)`

computes the antiproton flux for a given energy `E` and a differential cross section for antiproton production, `dSigmavdE`. For example, one can substitute

`dSigmavdE = σv SpectdNdE(E,SpP)`

where σv and `SpP` are obtained by `calcSpectrum`. This function does not depend on the details of the particle physics model and allows to analyse the dependence on the parameters of the propagation model.

- `pbarFluxTab(Emin,sigmav, Sp, Sobs)`

computes the antiproton flux, this function works like `posiFluxTab`,

- `solarModulation(Phi, mass, stellarTab, earthTab)`

takes into account modification of the interstellar positron/antiproton flux caused by the electro-magnetic fields in the solar system. Here `Phi` is the effective Fisk potential in

MeV, **mass** is the particle mass, **stellarTab** describes the interstellar flux, **earthTab** is the calculated particle flux in the Earth orbit.

Note that for **solarModulation** and for all ***FluxTab** routines one can use the same array for the spectrum before and after propagation.

8.6 Experimental data and backgrounds.

Background for anti-proton signal is caused by collision of protons with gas of galactic disk. It depends on proton flux and propagation parameters. But all sets of propagation parameters corresponding to the measured B/C rate should provide the same \bar{p}/p rate. It gives an opportunity of robust estimation of anti-proton background. Function

- **pBarBackgroundFlux(E)**

calculates \bar{p} background flux in $(GeV \cdot cm^2 \cdot s)^{-1}$ units using formula of [41]. There is an option to present this flux as an array of the format used by micrOMEGAs for other fluxes:

- **pBarBackgroundTab(Emax,pTab),**

where **Emax** defines an energy cut. Then one can apply to this array the **solarModulation** routine to take into account effect of solar modulation.

9 Neutrino signal from the Sun and the Earth

This module does not work yet in case of 2DM

After being captured, DM particles concentrate in the center of the Sun/Earth and then annihilate into Standard Model particles. These SM particles further decay producing neutrinos that can be observed at the Earth.

- **neutrinoFlux(f,forSun,nu, nu_bar)**

calculates muon neutrino/anti-neutrino fluxes near the surface of the Earth. Here **f** is the DM velocity distribution normalized such that $\int_0^\infty v f(v) dv = 1$. The units are km/s for **v** and s^2/km^2 for **f(v)**. At first approximation, one can use the same **Maxwell** function introduced for direct detection.

If **forSun==0** then the flux of neutrinos from the Earth is calculated, otherwise this function computes the flux of neutrinos from the Sun. The calculated fluxes are stored in **nu** and **nu_bar** arrays of dimension **NZ=250**. The neutrino fluxes are expressed in $[1/Year/km^2]$. The function **SpectdNdE(E,nu)** returns the differential flux of neutrinos in $[1/Year/km^2/GeV]$, and

displaySpectrum(nu,"nu from Sun $[1/Year/km^2/GeV]$ ",Emin,Emax,1)

allows to display the corresponding spectrum on the screen.

- **muonUpward(nu,Nu,rho, muon)**

calculates the muon flux which results from interactions of neutrinos with rocks below the detector. Here **nu** and **Nu** are input arrays containing the neutrino/anti-neutrino fluxes calculated by **neutrinoFlux**. **rho** is the Earth density $\approx 2.6g/cm^3$. **muon** is an array which stores the resulting sum of μ^+ , μ^- fluxes. **SpectdNdE(E,muon)** gives the differential muon flux in $[1/Year/km^2/GeV]$ units.

- **muonContained(nu,Nu,rho, muon)** calculates the flux of muons produced in a given detector volume. This function has the same parameters as **muonUpward** except that the outgoing array gives the differential muon flux resulting from neutrinos converted to muons

in a km^3 volume given in $[1/\text{Year}/km^3/\text{GeV}]$ units. `rho` is the density of the detector in g/cm^3 .

Two functions allow to estimate the background from atmospheric neutrinos creating muons after interaction with rocks below the detector or with water inside the detector.

- `ATMmuonUpward(cosFi,E)` calculates the sum of muon and anti-muon fluxes resulting from the interaction of atmospheric neutrinos with rocks in units of $[1/\text{Year}/km^2/\text{GeV}/\text{Sr}]$. `cosFi` is the energy between the direction of observation and the direction to the center of Earth. `E` is the muon energy in GeV .

- `ATMmuonContained(cosFi, E, rho)` calculates the muon flux caused by atmospheric neutrinos produced in a given (detector) volume. The returned value for the flux is given in $1/\text{Year}/km^3/\text{GeV}$. `rho` is the density of the detector in g/cm^3 units. `cosFi` and `E` are the same as above.

10 Cross sections and decays.

The calculation of particle widths, decay channels and branching fractions can be done by the function

- `pWidth(particleName,&address)`

returns directly the particle width. If the $1 \rightarrow 2$ decay channels are kinematically accessible then only these channels are included in the width when `VWdecay,VZdecay= 0`. If not, `pWidth` compiles all open $1 \rightarrow 3$ channels and use these for computing the width. If all $1 \rightarrow 3$ channels are kinematically forbidden, `micrOMEGAs` compiles $1 \rightarrow 4$ channels. If `VWdecay(VZdecay) ≠ 0`, then `micrOMEGAs` also computes the processes with virtual $W(Z)$ which are closed kinematically and adds these to the $1 \rightarrow 2$ decay channels. Note that $1 \rightarrow 3$ decay channels with a virtual W will be computed even if the mass of the decaying particle exceeds the threshold for $1 \rightarrow 2$ decays by several GeV 's. This is done to ensure a proper matching of $1 \rightarrow 2$ and $1 \rightarrow 3$ processes. For particles other than gauge bosons, an improved routine with 3-body processes and a matching between the $1 \rightarrow 2$ and $1 \rightarrow 3$ calculations is kept for the future. The returned parameter `address` gives an address where information about the decay channels is stored. In C, the address should be of type `txtList`. For models which read a SLHA parameter file, the values of the widths and branchings are taken from the SLHA file unless the user chooses not to read this data, see (Section 13.5) for details.

- `printTxtList(address,FD)`

lists the decays and their branching fractions and writes them in a file. `address` is the address returned by `pWidth`.

- `findBr(address,pattern)`

finds the branching fraction for a specific decay channel specified in `pattern`, a string containing the particle names in the CalcHEP notation. The names are separated by commas or spaces and can be specified in any order.

- `slhaDecayPrint(pname,FD)`

uses `pWidth` described above to calculate the width and branching ratios of particle `pname` and writes down the result in SLHA format. The return value is the PDG particles code. In case of problem, for instance wrong particle names, this function returns zero. This function first tries to calculate $1 \rightarrow 2$ decays. If such decays are kinematically forbidden

then $1 \rightarrow 3$ decay channels are computed.

- **newProcess(procName)**

compiles the codes for any $2 \rightarrow 2$ or $1 \rightarrow 2$ reaction. The result of the compilation is stored in the shared library in the directory **work/so-generated/**. The name of the library is generated automatically.

The **newProcess** routine returns the *address* of the compiled code for further usage. If the process can not be compiled, then a NULL address is returned³. Note that it is also possible to compute processes with polarized massless beams, for example for a polarized electron beam use **e%** to designate the initial particle.

- **procInfo1(address,&ntot,&nin,&nout)**

provides information about the total number of subprocesses (**ntot**) stored in the library specified by **address** as well as the number of incoming (**nin**) and outgoing (**nout**) particles for these subprocesses. Typically, for collisions (decays), $nin = 2(1)$ and $nout = 2, 3$. NULL can be substitute if this information is not needed.

- **procInfo2(address,nsub,N,M)**

fills an array of particle names **N** and an array of particle masses **M** for the subprocess **nsub** ($1 \leq nsub \leq ntot$). These arrays have size $nin + nout$ and the elements are listed in the same order as in CalcHEP starting with the initial state, see the example in **MSSM/main.c**.

- **cs22(address,nsub,P,c1,c2,&err)**

calculates the cross-section for a given $2 \rightarrow 2$ process, **nsub**, with center of mass momentum $P(\text{GeV})$. The differential cross-section is integrated within the range $c1 < \cos \theta < c2$. θ is the angle between \vec{p}_1 and \vec{p}_3 in the center-of-mass frame. Here \vec{p}_1 (\vec{p}_3) denote respectively the momentum of the first initial(final) particle. *err* contains a non zero error code if **nsub** exceeds the maximum value for the number of subprocesses (given by the argument **ntot** in the routine **procInfo1**). To set the polarization of the initial massless beam, define **Helicity[i]** where $i = 0, 1$ for the 1^{th} and 2^{nd} particles respectively. The helicity is defined as the projection of the particle spin on the direction of motion. It ranges from $[-1,1]$ for spin 1 particles and from $[-0.5,0.5]$ for spin $1/2$ particles. By definition a left handed particle has a positive helicity.

- **hCollider(Pcm,pp,pName1,pName2)** calculates the cross section for particle production at hadron colliders. Here **Pcm** is the beam energy in the center-of-mass frame. **pp** is 1(-1) for $pp(p\bar{p})$ collisions. **pName1** and **pName2** are the names of outgoing particles. The value returned is the cross section in [pb]. The QCD scale is fixed to $Q \approx (m(pName1) + m(pName2))/2$.

11 Tools for model independent analysis

A model independent calculation of the DM observables is also available. After specifying the DM mass, the cross sections for DM spin dependent and spin independent scattering on proton and neutron, the DM annihilation cross section times velocity at rest and the relative contribution of each annihilation channel to the total DM annihilation cross section, one can compute the direct detection rate on various nuclei, the fluxes for photons, neutrinos and antimatter resulting from DM annihilation in the galaxy and the neutrino/muon fluxes in neutrino telescopes.

³In Fortran the format is **call newProcess(procName, address)**

- **nucleusRecoilAux(f,A,Z,J,Sxx,csIp,csIn,csDp,csDn,dNdE)**

This function is similar to **nucleusRecoil**. The additional input parameters include **csIp(csIn)** the SI cross sections for WIMP scattering on protons(neutrons) and **csDp(csDn)** the SD cross sections on protons(neutrons). A negative value for one of these cross sections is interpreted as a destructive interference between the proton and neutron amplitudes. Note that the rate of recoil depends implicitly on the WIMP mass, the global parameter **Mcdm**. The numerical value for the global parameter has to be set before calling this function.

- **nucleusRecoil0Aux(f,A,Z,J,Sp,Sn,csIp,csIn,csDp,csDn,dNdE)** is the corresponding modification of **nucleusRecoil0**.

For indirect detection, we also provide a tool for model independent studies

- **basicSpectra(pdgN, outN,Spectr)**

computes the spectra of outgoing particles and writes the result in an array of dimension 250, **Spectr**, **pdgN** is the PDG code of the particles produced in the annihilation of a pair of WIMPs. To get the spectra generated by transverse and longitudinal W's substitute $pdgN = 24 +' T'$ and $24 +' L'$ correspondingly. In the same manner $pdgN = 23 +' T'$ and $23 +' L'$ provides the spectra produced by a polarized Z boson. **outN** specifies the outgoing particle,

$$\text{outN} = \{0, 1, 2, 3, 4, 5\} \text{ for } \{\gamma, e^+, p^-, \nu_e, \nu_\mu, \nu_\tau\}$$

This function depends implicitly on the global variable **Mcdm**. Note that the propagation routines for e^+, p^-, γ can be used after this routine as usual. Note that the result of **basicSpectra** are not valid for $Mcdm < 2\text{GeV}$ as explained in the description of **calcSpectrum**.

- **captureAux(f,forSun,csIp,csIn,csDp,csDn)**

calculates the number of DM particles captured per second assuming the cross sections for spin-independent and spin-dependent interactions with protons and neutrons **csIp**, **csIn**, **csDp**, **csDn** are given as input parameters (in [pb]). A negative value for one of the cross sections is interpreted as a destructive interference between the proton and neutron amplitudes. The first two parameters have the same meaning as in the **neutrinoFlux** routine Section 9. The result depends implicitly on the global parameters **rhoDM** and **Mcdm** in Table 1.

- **basicNuSpectra(forSun,pdgN, outN,nu)**

calculates the ν_μ and $\bar{\nu}_\mu$ spectra corresponding to DM annihilating into particles specified by the PDG code **pdgN**. Effects of interaction with Sun/Earth medium as well as neutrino oscillation are taken into account [39]. **outN** should be chosen 1 for muon neutrino and -1 for anti-neutrino. The resulting spectrum is stored in the array **nu** with $NZ=250$ elements which can be checked by the **SpectdNdE(E,nu)** function.

The files **main.c/F** in the directory **mdlIndep** contain an example of the calculation of the direct detection, indirect detection and neutrino telescope signals using the routines described in this section. The numerical input data in this sample file corresponds to 'MSSM/mssmh.dat'.

12 Additional routines for specific models

The models included in `micrOMEGAs` contain some specific routines which we describe here for the sake of completeness. The current distribution includes the following models: **MSSM**, **NMSSM**, **CPVMSSM**, **IDM** (inert doublet model), **LHM** (little Higgs model), **RHNM** (a Right-handed Neutrino model), **SM4** (toy model with 4th generation lepton), and **Z3M** (doublet and singlet model with Z_3 symmetry).

Some of these models contain a special routine for reading the input parameters:

- `readVarMSSM`, `readVarNMSSM`, `readVarCPVMSSM`, `readVarlHiggs`, `readVarRHNM`.

These routines are similar to the general `readVar` routine described in Section 5 but they write a warning when a parameter is not found in the input file and display the default values for these parameters.

The supersymmetric models contain several additional routines to calculate the spectrum and compute various constraints on the parameter space of the models. Some functions are common to the **MSSM**, **NMSSM**, **CPVMSSM** models:

- `o1Contents(FD)`

prints the neutralino LSP components as the $\tilde{B}, \tilde{W}, \tilde{h}_1, \tilde{h}_2$ fractions. For the **NMSSM** the fifth component is the singlino fraction \tilde{S} . The sum of the squares of the LSP components should add up to 1.

12.1 MSSM

The **MSSM** has a long list of low scale independent model parameters, those are specified in the SLHA file [18, 14]. They are directly implemented as parameters of the model. For **EWSB** scenarios the input parameters are the soft parameters, the names of these parameters are given in the `MSSM/mssm[1/2].par` files. The user can assign new values to these parameters by means of `assignVal` or `readVarMSSM`.

- `spectEwsbMSSM()`

calculates the masses of Higgs and supersymmetric particles in the **MSSM** including one-loop corrections starting from weak scale input parameters.

In these functions `spect` stands for one of the spectrum calculators `suspect`, `isajet`, `spheno`, or `softSusy`. The default spectrum calculator package is `SuSpect`. To work with another package one has to specify the appropriate path in `MSSM/lib/Makefile`. For this, the environment variables `ISAJET`, `SPHENO` or `SOFTSUSY` must be redefined accordingly. Note that we also provide a special interface for `ISAJET` to read a SLHA file. This means that the user must first compile the executable `isajet_slha` which sets up the SLHA interface in `ISAJET`. Specific instructions are provided in the `README` file.

For other **MSSM** scenarios, the parameters at the electroweak symmetry breaking scale are derived from an input at high scale. The same codes `suspect`, `isajet`, `spheno`, or `softSusy` are used for this. The corresponding routines are:

- `spectSUGRA(tb, MG1, MG2, MG3, A1, At, Ab, signMu, MHu, MHd, M11, M13, Mr1, Mr3, Mq1, Mq3, Mu1, Mu3, Md1, Md3)`

assumes that all input parameters except `tb` and `signMu` are defined at the GUT scale. The **SUGRA/CMSSM** scenario is a special case of this general routine.

- `spectSUGRAnuh(tb, MG1, MG2, MG3, A1, At, Ab, M11, M13, Mr1, Mr3, Mq1, Mq3, Mu1, Mu3, Md1, Md3, mu, MA)`

realizes a SUGRA scenario with non universal Higgs parameters. Here the `Mhu`, `MHd` parameters in the Higgs potential are replaced with the `mu` parameter defined at the EWSB scale and `MA`, the pole mass of the CP-odd Higgs. The `signMu` parameter is omitted because `mu` is defined explicitly.

- `spectAMSB(am0,m32,tb,sng)`.

does the same as above within the `AMSB` model.

We have an option to directly read a `SLHA` input file, this uses the function

- `lesHinput(file_name)`

which returns a non-zero number in case of problem.

The routines for computing constraints are (see details in [2]).

- `deltarho()`

calculates the $\Delta\rho$ parameter in the MSSM. It contains for example the stop/sbottom contributions, as well as the two-loop QCD corrections due to gluon exchange and the correction due to gluino exchange in the heavy gluino limit.

- `bsgnlo(&SMbsg)`

returns the value of the branching ratio for $b \rightarrow s\gamma$, see Appendix A. We have included some new contributions beyond the leading order that are especially important for high $\tan\beta$. `SMbsg` gives the SM contribution.

- `bsmumu()`

returns the value of the branching ratio $B_s \rightarrow \mu^+\mu^-$ in the MSSM. It includes the loop contributions due to chargino, sneutrino, stop and Higgs exchange. The Δm_b effect relevant for high $\tan\beta$ is taken into account.

- `btaunu()`

computes the ratio between the MSSM and SM branching fractions for $\bar{B}^+ \rightarrow \tau^+\nu_\tau$.

- `gmuon()`

returns the value of the supersymmetric contribution to the anomalous magnetic moment of the muon.

- `Rl23()`

computes the ratio of the MSSM to SM value for R_{l23} in $K^+ \rightarrow \mu\nu$ due to a charged higgs contribution, see Eq.70 in [6].

- `dtaunu(&dmmu)`

computes the branching ratio for $D_s^+ \rightarrow \tau^+\nu_\tau$. `dmmu` gives the branching ratio for $D_s^+ \rightarrow \mu^+\nu_\mu$

- `masslimits()`

returns a positive value and prints a WARNING when the choice of parameters conflicts with a direct accelerator limits on sparticle masses from LEP. The constraint on the light Higgs mass from the LHC is included.

We have added a routine for an interface with `superIso` [32]. This code is not included in `micrOMEGAs` so one has first to define the global environment variable `superIso` to specify the path to the package.

- `callSuperIsoSLHA()`

launches `superIso` and downloads the `SLHA` file which contains the results. The return value is zero when the program was executed successfully. Results for specific observables can be obtained by the command `slhaValFormat` described in section (13.5). Both `superIso` and `callSuperIsoSLHA` use a file interface to exchange data. The `delFiles` flag specifies whether to save or delete the intermediate files.

- `loopGamma(&vcs_gz,&vcs_gg)`

calculates σv for loop induced processes of neutralino annihilation into γZ and into $\gamma\gamma$. The result is given in $\frac{cm^3}{s}$. In case of a problem the function returns a non-zero value.

12.2 The NMSSM

As in the MSSM there are specific routines to compute the parameters of the model as specified in SLHA. The spectrum calculator is `NMSPEC` [23] in the `NMSSMTools_4.0` package [29].

- `nmssmEWSB(void)`

calculates the masses of Higgs and supersymmetric particles in the NMSSM starting from the weak scale input parameters. These can be downloaded by the `readVarNMSSM` routine. [24]

- `nmssmSUGRA(m0,mhf,a0,tb,sgn,Lambda,aLambda,aKappa)`

calculates the parameters of the NMSSM starting from the input parameters of the `CNMSSM`.

The routines for computing constraints are taken from `NMSSMTools` (see details in [3]).

- `bsgnlo(&M,&P)`, `bsmumu(&M,&P)`, `btaunu(&M,&P)`, `gmuon(&M,&P)`

are the same as in the MSSM case. Here the output parameters `M` and `P` give information on the lower/upper experimental limits [25]

- `deltaMd()`, `deltaMs()`

compute the supersymmetric contribution to the $B_{d,s}^0 - \overline{B}_{d,s}^0$ mass differences, ΔM_d and ΔM_s .

- `NMHwarn(FD)`

is similar to `masslimits_` except that it also checks the constraints on the Higgs masses, returns the number of warnings and writes down warnings in the file `FD`.

- `loopGamma(&vcs_gz,&vcs_gg)`

calculates σv for loop induced processes of neutralino annihilation into γZ and into $\gamma\gamma$. The result is given in $\frac{cm^3}{s}$. In case of a problem the function returns a non-zero value.

12.3 The CPVMSSM

The independent parameters of the model include, in addition to some standard model parameters, only the weak scale soft SUSY parameters. The independent parameters are listed in `CPVMSSM/work/models/vars1.mdl`. Masses, mixing matrices and parameters of the effective Higgs potential are read directly from `CPsuperH` [26, 27], together with the masses and the mixing matrices of the neutralinos, charginos and third generation sfermions. Masses of the first two generations of sfermions are evaluated (at tree-level) within `micrOMEGAs` in terms of the independent parameters of the model.

The routines for computing constraints are taken from `CPsuperH`, [28]

- `bsgnlo()`, `bsmumu()`, `btaunu()`, `gmuon()`

are the same as in the MSSM case.

- `deltaMd()`, `deltaMs()`

are the same as in the NMSSM case.

- `Bd11()`

computes the supersymmetric contribution to the branching fractions for $B_d \rightarrow \tau^+ \tau^-$ in the CPVMSSM.

- `ABsg()`

computes the supersymmetric contribution to the asymmetry for $B \rightarrow X_s \gamma$.

- `EDMe1()`, `EDMmu()`, `EDMT1()`

return the value of the electric dipole moment of the electron, d_e , the muon, d_μ , and of Thallium, d_{Tl} in units of ecm .

13 Tools for new model implementation.

It is possible to implement a new particle physics model in `micrOMEGAs`. For this the model must be specified in the CalcHEP format. `micrOMEGAs` then relies on CalcHEP to generate the libraries for all matrix elements entering DM calculations. Below we describe the main steps and tools for implementing a new model.

13.1 Main steps

- The command `./newProject MODEL` launched from the root `micrOMEGAs` directory creates the directory `MODEL`. This directory and the subdirectories contain all files needed to run `micrOMEGAs` with the exception of the new model files.
- The new model files in the CalcHEP format should then be included in the subdirectory `MODEL/work/models`. The files needed are `vars1.mdl`, `func1.mdl`, `prtcls1.mdl`, `lgrng1.mdl`, `extlib1.mdl`. For more details on the format and content of model files see [19].
- For odd particles and for the Higgs sector it is recommended to use the widths that are (automatically) calculated internally by CalcHEP/`micrOMEGAs`. For this one has to add the `'!'` symbol before the definition of the particle's width in the file `prtcls1.mdl`, for example

Full	name	P	aP	PDG	2*spin	mass	width	color
Higgs	1	h1	h1	25	0	Mh1	!wh1	1

- Some models contain external functions, if this is the case they have to be compiled and stored in the `MODEL/lib/aLib.a` library. These functions should be written in C and both functions and their arguments have to be of type `double`. The library `aLib.a` can also contain some functions which are called directly from the `main` program. The `MODEL/Makefile` automatically launches `make` in the `lib` directory and compiles the external functions provided the prototypes of these external functions are specified in `MODEL/lib/pmodel.h`. The user can of course rewrite his own `lib/Makefile` if need be.

If the new `aLib.a` library needs some other libraries, their names should be added to the `SSS` variable defined in `MODEL/Makefile`.

The *MODEL* directory contains both C and FORTRAN samples of *main* routines. In these sample main programs it is assumed that input parameters are provided in a separate file. In this case the program can be launched with the command:

```
./main data1.par
```

Note that for the direct detection module all quarks must be massive. However the cross sections do not depend significantly on the exact numerical values for the masses of light quarks.

13.2 Automatic width calculation

Automatic width calculation can be implemented by inserting the '!' symbol before the name of the particle width in the CalcHEP particle table (file `prtcls1.mdl`). In this case the width parameter should not be defined as a free or constrained parameter. Actually the `pWidth` function described in section 10 is used for width calculation in this case. We recommend to use the automatic width calculation for all particles from the 'odd' sector and for Higgs particles. For models which use SLHA parameter transfer (Section 13.5), the automatic width option will use the widths contained in the SLHA file unless the user chooses the option to ignore this data in the SLHA file, see section 13.5.

13.3 Using LanHEP for model file generation.

For models with a large number of parameters and various types of fields/particles such as the MSSM, it is more convenient to use an automatic tool to implement the model. LanHEP is a tool for Feynman rules generation. A few minor modifications to the default format of LanHEP have to be taken into account to get the model files into the `micrOMEGAs` format.

- The `lhhep` command has to be launched with the `-evl 2` flag

```
lhhep source_file -evl 2
```

Such a flag provides the correct level of optimization for the model's Feynman rules.

- The default format for the file `prtcls1.mdl` which specifies the particle content has to be modified to include a column containing the PDG code of particles. For this, first add the following command in the LanHEP source code, before specifying the particles

```
prtcformat fullname:
'Full Name ', name:' P ', aname:' aP', pdg:' number ',
spin2,mass,width, color, aux, texname: '> LaTeX(A) <',
atexname:'> LateX(A+) <' .
```

Then for each particle define the PDG code. For instance:

```
vector 'W+'/'W-': ('W boson', pdg 24, mass MW, width wW).
```

- LanHEP does not generate the file `extlib1.mdl`. `micrOMEGAs` works without this file but it is required for a `CalcHEP` interactive session. The role of this file is to provide the linker with the paths to all user's libraries needed at compilation. For example for the `lib/aLib.a` library define

```
$CALCHEP/../../MODEL/lib/aLib.a
```

For examples see the `extlib1.mdl` files in the directory of the models provided.

13.4 QCD functions

Here we describe some QCD functions which can be useful for the implementation of a new model.

- `initQCD(alfsMZ,McMc,MbMb,Mtp)`

This function initializes the parameters needed for the functions listed below. It has to be called before any of these functions. The input parameters are the QCD coupling at the Z scale, $\alpha_s(M_Z)$, the quark masses, $m_c(m_c)$, $m_b(m_b)$ and $m_t(pole)$.

- `alphaQCD(Q)`

calculates the running α_s at the scale Q in the \overline{MS} scheme. The calculation is done using the NNLO formula in [22]. Thresholds for the b-quark and t-quark are included in n_f at the scales $m_b(m_b)$ and $m_t(m_t)$ respectively.

- `MtRun(Q)`, `MbRun(Q)`, `McRun(Q)`

calculates top, bottom and charm quarks running masses evaluated at NNLO.

- `MtEff(Q)`, `MbEff(Q)`, `McEff(Q)`,

calculates effective top, bottom and charm quark masses using [22]

$$M_{eff}^2(Q) = M(Q)^2 [1 + 5.67a + (35.94 - 1.36n_f)a^2 + (164.14 - n_f(25.77 - 0.259n_f))a^3] \quad (8)$$

where $a = \alpha_s(Q)/\pi$, $M(Q)$ and $\alpha_s(Q)$ are the quark masses and running strong coupling in the \overline{MS} -scheme. In `micrOMEGAs`, we use the effective quark masses calculated at the scale $Q = 2M_{cdm}$. In some special cases one needs a precise treatment of the light quarks masses. The function

- `MqRun(M2GeV, Q)`

returns the running quark mass defined at a scale of 2 GeV. The corresponding effective mass needed for the Higgs decay width is given by

- `Mqeff(M2GeV, Q)`

13.5 SLHA reader

Very often the calculation of the particle spectra for specific models is done by some external program which writes down the particle masses, mixing angles and other model parameters in a file with the so-called **SLHA** format [18, 14]. The `micrOMEGAs` program contains routines for reading files in the SLHA format. Such routines can be very useful for the implementation of new models.

In general a SLHA file contains several pieces of information which are called blocks. A block is characterized by its name and, sometimes, by its energy scale. Each block contains the values of several physical parameters characterized by a *key*. The key consists in a sequence of integer numbers. For example:

```

BLOCK MASS      # Mass spectrum
#  PDG Code      mass              particle
      25      1.15137179E+02      # lightest neutral scalar
      37      1.48428409E+03      # charged Higgs

```

```

BLOCK NMIX      # Neutralino Mixing Matrix
  1  1      9.98499129E-01      # Zn11
  1  2     -1.54392008E-02      # Zn12

```

```

BLOCK Au Q= 4.42653237E+02      # The trilinear couplings
  1  1     -8.22783075E+02      # A_u(Q) DRbar
  2  2     -8.22783075E+02      # A_c(Q) DRbar

```

- `slhaRead(filename,mode)`

downloads all or part of the data from the file `filename`. `mode` is an integer which determines which part of the data should be read from the file, `mode= 1*m1+2*m2+4*m4` where

```

m1 = 0/1 -   overwrites all/keeps old data
m2 = 0/1 -   reads DECAY /does not read   DECAY
m4 = 0/1 -   reads BLOCK/does not read   BLOCK

```

For example `mode=2` (`m1=0,m2=1`) is an instruction to overwrite all previous data and read only the information stored in the BLOCK sections of `filename`. In the same manner `mode=3` is an instruction to add information from DECAY to the data obtained previously. `slhaRead` returns the values:

```

 0 - successful reading
-1 - can not open the file
-2 - error in spectrum calculator
-3 - no data
n>0 - wrong file format at line n

```

- `slhaValExists(BlockName, keylength, key1, key2,...)`

checks the existence of specific data in a given block. `BlockName` can be substituted with any case spelling. The `keylength` parameter defines the length of the key set `{key1,key2,...}`. For example `slhaValExists("Nmix",2,1,2)` will return 1 if the neutralino mass mixing element Zn12 is given in the file and 0 otherwise.

- `slhaVal(BlockName,Q, keylength, key1, key2,.....)`

is the main routine which allows to extract the numerical values of parameters. `BlockName` and `keylength` are defined above. The parameter `Q` defines the scale dependence. This parameter is relevant only for the blocks that contain scale dependent parameters, it will be ignored for other blocks, for example those that give the particle pole masses. In general a SLHA file can contain several blocks with the same name but different scales (the scale is specified after the name of the block). `slhaVal` uses the following algorithm to read the scale dependent parameters. If `Q` is less(greater) than all the scales used in the different blocks for a given parameter `slhaVal` returns the value corresponding to the minimum(maximum) scale contained in the file. Otherwise `slhaVal` reads the values corresponding to the two scales Q_1 and Q_2 just below and above `Q` and performs a linear interpolation with respect to $\log(Q)$ to evaluate the returned values.

Recently it was proposed to use an extension of the SLHA interface to transfer Flavour Physics data [31]. Unfortunately the structure of the new blocks is such that they cannot be read with the `slhaVal` routine. We have added two new routines for reading such data

- `slhaValFormat(BlockName, Q, format)`

where the *format* string allows to specify data which one would like to extract from the given block *BlockName*. For instance, to get the $b \rightarrow s\gamma$ branching ratio from the block

Block FOBS # Flavour observables

```
# ParentPDG type value      q  NDA ID1 ID2 ID3 ... comment
   5      1  2.95061156e-04  0    2   3  22      # BR(b->s gamma)
  521     4  8.35442304e-02  0    2 313  22      # Delta0(B->K* gamma)
  531     1  3.24270419e-09  0    2  13 -13      # BR(B_s->mu+ mu-)
...

```

one has to use the command `slhaValFormat("FOBS", 0., "5 1 %E 0 2 3 22")`. In this command the *format* string is specified in C-style. The same routine can be used to read HiggsBound SLHA output.

A block can also contain a textual information. For example, in HIGGSBOUNDS a block contains the following records,

```
Block HiggsBoundsResults
#CHANNELTYPE 1: channel with the highest statistical sensitivity
  1      1      328      # channel id number
  1      2      1      # HBresult
  1      3  0.72692779334500290  # obsratio
  1      4      1      # ncombined
  1      5 ||(p p)->h+..., h=1 where h is SM-like (CMS-PAS-HIG-12-008)|| # text description of channel

```

In particular, the last record contains the name of the channel which gives the strongest constraint on the Higgs. To extract the name of this channel one can use the new function

- `slhaSTRFormat("HiggsBoundsResults", "1 5 || %[^\n]||", channel);`

which will write the channel name in the text parameter *channel*.

- `slhaWarnings(fileName)`

writes into the file the warnings or error message stored in the SPINFO block and returns the number of warnings. If `FD=NULL` the warnings are not written in a file.

- `slhaWrite(fileName)`

writes down the information stored by `readSLHA` into the file. This function can be used for testing purposes.

SLHA also describes the format of the information about particle decay widths. Even though `micrOMEGAs` also performs the width calculation, one might choose to read the information from the SLHA file.

- `slhaDecayExists(pNum)`

checks whether information about the decay of particle *pNum* exists in the SLHA file. *pNum* is the particle PDG code. This function returns the number of decay channels. In particular zero means that the SLHA file contains information only about the total width, not on branching ratios while -1 means that even the total width is not given.

- `slhaWidth(pNum)`

returns the value of particle width.

- `slhaBranch(pNum, N, nCh)`

returns the branching ratio of particle *pNum* into the *N*-th decay channel. Here

$0 < N \leq \text{slhaDecayExists}(\text{pNum})$. The array `nCh` is an output which specifies the PDG numbers of the decay products, the list is terminated by zero.

The functions `slhaValExists`, `slhaVal`, `slhaDecayExists`, `slhaWidth` can be used directly in CalcHEP model files, see an example in `MSSM/calchep/models/func2.mdl`. Note that in this example the call to `slhaRead` is done within the function `suspectSUGRAc`.

13.5.1 Writing an SLHA input file

We have included in the `micrOMEGAS` package some routines which allow to write an SLHA input file and launch the spectrum generator via the CalcHEP *constraints* menu. This way a new model can be implemented without the use of external libraries. The routines are called from `func1.mdl`, see example below.

- `openAppend(fileName)`

deletes the input file `fileName` and stores its name. This file will then be filled with the function `aPrintF`.

- `aPrintF(format,...)`

opens the file `fileName` and writes at the end of the file the input parameters needed in the SLHA format or in any other format understood by the spectrum calculator. The arguments of `aPrintF` are similar to the arguments of the standard `printf` function.

- `System(command, ...)` generates a command line which is launched by the standard `system` C-function. The parameter *command* works here like a format string and can contain `%s`, `%d` elements. These are replaced by the next parameters of the `System` call.

For example to write directly the SLHA model file needed by `SuSpect` to compute the spectrum in a CMSSM(SUGRA) model, one needs to add the following sequence in the `func1.mdl` model file.

```
open |openAppend("suspect2_lha.in")
input1|aPrintF("Block MODSEL # Select model\n 1 1 # SUGRA\n")
input2|aPrintF("Block SMINPUTS\n 5 %E#mb(mb)\n 6 %E#mt(pole)\n",MbMb,Mtp)
input3|aPrintF("BLOCK MINPAR\n 1 %E #m0\n 2 %E #m1/2\n ",Mzero,Mhalf)
input4|aPrintF("3 %E #tb\n 4 %E #sign(mu)\n 5 %E #A0\n",tb,sgn,A0)
sys |System("./suspect2.exe")
rd |slhaRead("suspect2_lha.out",0)
```

It is possible to cancel the execution of a program launched with `System` if it runs for too long. For this we have introduced two global parameters `sysTimeLim` and `sysTimeQuant`. `sysTimeLim` sets a time limit in milliseconds for `System` execution, if `sysTimeLim==0` (the default value) the execution time is not checked. The time interval between checks of the status of the program launched with `System` is specified by the parameter `sysTimeQuant`, the default value is set to 10. Note that it is preferable not too use too large a value for `sysTimeQuant` as it defines the lower time limit for a system call. In Fortran use `call setSysTimeLim(sysTimeLim,sysTimeQuant)` to reset the default time control parameters.

The function prototypes are available in `CalcHEP_src/c_source/SLHAplus/include/SLHAplus.h`

13.6 Routines for diagonalisation.

Very often in a new model one has to diagonalize mass matrices. Here we present some numerical routines for diagonalizing matrices. Our code is based on the `jacobi` routine provided in [30]. To use the same routine for a matrix of arbitrary size, we use a C option that allows to write routines with an arbitrary number of arguments.

- `initDiagonal()` should be called once before any other `rDiagonal(A)` routine described below. `initDiagonal()` assigns zero value to the internal counter of eigenvalues and rotation matrices. Returns zero.

- `rDiagonal(d,M11,M12,...M1d,M22,M23...Mdd)`

diagonalizes a symmetric matrix of dimension `d`. The $d(d+1)/2$ matrix elements, `Mij` ($i \leq j$), are given as arguments. The function returns an integer number `id` which serves as an identifier of eigenvalues vector and rotation matrix.

- `MassArray(id, i)`

returns the eigenvalues m_i ordered according to their absolute values.

- `MixMatrix(id,i,j)`

returns the rotation matrix R_{ij} where

$$M_{ij} = \sum_k R_{ki} m_k R_{kj}$$

A non-symmetric matrix, for example the chargino mass matrix in the MSSM, is diagonalized by two rotation matrices,

$$M_{ij} = \sum_k U_{ki} m_k V_{kj}.$$

- `rDiagonalA(d,M11,M12...M1d,M21,M22...Mdd)`

diagonalizes a non-symmetric matrix, the d^2 matrix elements, `Mij`, are given as arguments. The eigenvalues and the V rotation matrix are calculated as above with `MassArray` and `MixMatrix`.

- `MixMatrixU(id,i,j)`

returns the rotation matrix U_{ij} .

The function prototypes can be found in

`CalcHEP_src/c_source/SLHAplus/include/SLHAplus.h`

14 Mathematical tools.

Some mathematical tools used by `micrOMEGAs` are available only in C format. Prototypes of these functions can be found in

`sources/micromegas_aux.h`

- `simpson(F, x1, x2, eps)`

numerical integration of the function $F(x)$ in the interval $[x1, x2]$ with relative precision eps . `simpson` uses an adaptive algorithm for integrand evaluation and increases the number of function calls in the regions where the integrand has peaks.

- `gauss(F,x1,x2,N)`

performs Gauss N-point integration for $N < 8$.

- **odeint**(Y, Dim, x1, x2, eps, h1, deriv)

solves a system of *Dim* differential equations in the interval $[x1, x2]$. The array *Y* contains the starting variables at *x1* as an input and is replaced by the resulting values at *x2* as an output. *eps* determines the precision of the calculation and *h1* gives an estimation of step of calculation. The function **deriv** calculates $Y'_i = dY_i/dx$ with the call *deriv*(*x*, *Y*, *Y'*). The Runge-Kutta method is used, see details in [30].

- **buildInterpolation**(F, x1, x2, eps, &Dim, &X, &Y)

constructs a cubic interpolation of the function *F* in the interval $[x1, x2]$. *eps* controls the precision of interpolation. If *eps* < 0 the absolute precision is fixed, otherwise a relative precision is required. The function checks that after removing any grid point, the function at that point can be reproduced with a precision *eps* using only the other points. It means that the expected precision of interpolation is about *eps*/16. *Dim* gives the number of points in the constructed grid. *X* and *Y* are variables of the **double*** type. The function allocates memory for *Dim* array for each of these parameters. *X*[*i*] contains the x-grid while *Y*[*i*] = *F*(*X*[*i*]).

- **polint4**(x, Dim, X, Y)

performs cubic interpolation for *Dim*-dimension arrays *X*, *Y*. A similar function, **polint3**, performs quadratic interpolation.

- **bessk0**(x)

The Bessel function of zero order of the second kind.

- **displayFunc**(F, x1, x2, title)

displays a plot of function *F*(*x*) in the $[x1, x2]$ interval. *title* is a text which appears as the title of the plot.

- **displayFunc10**(F, x1, x2, title)

displays *F*(10^{*x*}) in the $[x1, x2]$ interval.

A An updated routine for $b \rightarrow s\gamma$ in the MSSM

The calculation of $b \rightarrow s\gamma$ was described in micromegas1.3 [2]. The branching fraction reads

$$B(\bar{B} \rightarrow X_s \gamma) = B(\bar{B} \rightarrow X_c e \bar{\nu}) \left| \frac{V_{ts}^* V_{tb}}{V_{cb}} \right|^2 \frac{6\alpha_{em}}{\pi f(z_0)} K_{NLO}(\delta) \quad (9)$$

where $\alpha_{em} = 1/137.036$, the factor K_{NLO} involves the photon energy cut-off parameter δ and $f(z_0) = 0.542 - 2.23(\sqrt{z_0} - 0.29)$ depends on $z_0 = (m_c/m_b)^2$ defined in terms of pole masses. In the code the standard model and Higgs contribution at NLO were included as well as the leading order SUSY contributions. However in the last few years the NNLO standard model contribution has been computed [33] and shown to lead to large corrections, shifting the standard model value by over 10%. It was also argued that the NNLO SM result could be reproduced from the NLO calculation by appropriately choosing the scale for the c-quark mass [34, 35].

In this improved version of the **bsgnlo** routine, we have changed the default value for the parameter $z_1 = (m_c/m_b)^2$ where m_c is the \overline{MS} running charm mass $m_c(m_b)$. Taking $z_1 = 0.29$ allows to reproduce the NNLO result. It is therefore no longer necessary to apply a shift to the micromegas output of $b \rightarrow s\gamma$ to reproduce the SM value.

We have also updated the default values for the experimentally determined quantities

$B(\bar{B} \rightarrow X_c e \bar{\nu})$	0.1064 [37]
C_{sl}	0.546 [35]
$ V_{ts}^* V_{tb}/V_{cb} ^2$	0.9613 [37]
A	0.808
λ	0.2253
$\bar{\rho}$	0.132
$\bar{\eta}$	0.341
m_b/m_s	50
$\lambda_2 \approx \frac{1}{4}(m_{B^*}^2 - m_B^2)$	0.12 GeV ² [36]
$\alpha_s(M_Z)$	0.1189

Table 4: Default values in micrOMEGAs

in Eq. 9, see Table A, and we have replaced the factor $f(z_0)$ by C_{sl} where

$$C_{sl} = \left| \frac{V_{ub}}{V_{cb}} \right|^2 \frac{\Gamma(\bar{B} \rightarrow X_c e \bar{\nu})}{\Gamma(\bar{B} \rightarrow X_u e \bar{\nu})} \quad (10)$$

accounts for the m_c dependence in $\bar{B} \rightarrow X_c e \bar{\nu}$.

The CKM matrix elements in the Wolfenstein parametrisation given in Table A are used to compute the central value of $ckmf$ at order λ^4 ,

$$ckmf = \left| \frac{V_{ts}^* V_{tb}}{V_{cb}} \right|^2 = 1 + \lambda^2(2\bar{\rho} - 1) + \lambda^4(\bar{\rho}^2 + \bar{\eta}^2 - A^2) \quad (11)$$

With these default values the NLO- improved SM contribution is $B(\bar{B} \rightarrow X_s \gamma)|_{\text{SM}} = 3.27 \times 10^{-4}$ which corresponds to the result of Gambino and Giordano [35] after correcting for the slightly different CKM parameter used ($ckmf = 0.963$).

We have performed a comparison with superIso3.1 which includes the NNLO SM calculation for 10^5 randomly generated MSSM scenarios. The results are presented in Fig. A after applying a correction factor in superISO to account for the different value for the overall factor $F = B(\bar{B} \rightarrow X_c e \bar{\nu}) \left| \frac{V_{ts}^* V_{tb}}{V_{cb}} \right|^2 / C_{sl}$. The ratio of $F_{\text{micro}}/F_{\text{ISO}} = 0.942$. The two codes agree within 5% most of the time.

B Solution of equation for relic density in case of 2 DM components.

In terms of $\Delta Y_i = Y_i - \bar{Y}_i$, where \bar{Y}_i denotes equilibrium abundance, evolution equation takes a form

$$3H \frac{\Delta Y_i}{ds} = -C_i + A_{ij}(T) \Delta Y_j + Q_{ijk}(T) \Delta Y_j \Delta Y_k, \quad (12)$$

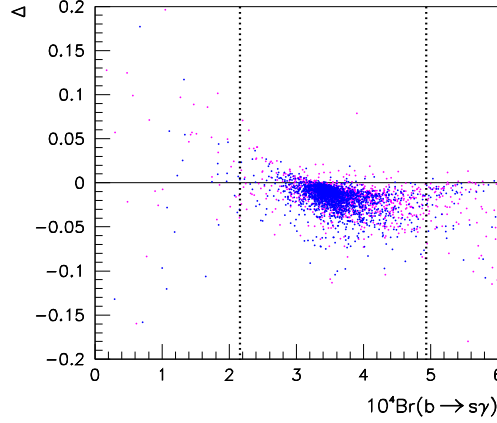


Figure 1: Relative difference for $B(\bar{B} \rightarrow s\gamma)$ between micromegas2.4 and superIso3.1. the vertical lines show the 3σ experimentally measured value.

where

$$C_i = 3H \frac{d\bar{Y}_i}{ds}, \quad (13)$$

$$A = \begin{pmatrix} 2(\sigma_v^{1100} + \sigma_v^{1122} + \sigma_v^{1120})\bar{Y}_1 & -(\sigma_v^{1120} + 2\sigma_v^{1122})\frac{\bar{Y}_1^2}{\bar{Y}_2} \\ -\sigma_v^{1120}\bar{Y}_1 - 2\sigma_v^{1122}\bar{Y}_1 & 2(\sigma_v^{2200} + \sigma_v^{2211})\bar{Y}_2 + 0.5(\sigma_v^{1210} + \sigma_v^{1120}\frac{\bar{Y}_1}{\bar{Y}_2})\bar{Y}_1 \end{pmatrix} \quad (14)$$

$$Q_1 = \begin{pmatrix} \sigma_v^{1100} + \sigma_v^{1122} + \sigma_v^{1120} & 0 \\ 0 & -\sigma_v^{2211} \end{pmatrix}, \quad (15)$$

$$Q_2 = \begin{pmatrix} -\sigma_v^{1120} - \sigma_v^{1122} & \frac{1}{2}\sigma_v^{1210} \\ 0 & \sigma_v^{2200} + \sigma_v^{2211} \end{pmatrix}. \quad (16)$$

$$(17)$$

At large temperatures we expect the densities of both DM components to be close to their equilibrium values. Here solution can be presented in the form

$$\Delta Y_i(s) = A_{ij}^{-1}(s)C_j(s). \quad (18)$$

where exponential factors in A and C are cancel one other, thus $\Delta Y_i(s)$ slow depends on temperature that retroactively justifies neglecting of left part of Eq.12. Solution (18) can be used as a stating point for complete Eq.12.

In general in micrOMEGAs [1] the equation for the abundance is solved numerically starting from large temperatures. However, this procedure poses a problem for Eq. (12). The step of the numerical solution is inversely proportional to $A(T)$ and as long as $A(T)$ is not suppressed by the Boltzmann factor included in \bar{Y} , the step is too small and the numerical method fails.

To avoid this problem, we use the fact that at large temperatures one can neglect the Q term in Eq. (12) and write the explicit solution for the linearised equation. The

approximate solution in the case of large A is One can use Eq. (18) to find the lowest temperature where $\Delta Y_i \approx 0.05 Y_i$ and start solving numerically Eq. (12) from this temperature. In the general case it gives a reasonable step for the numerical solution $\delta s/s \approx 0.1$, where s is the variable of integration. This method can, however, lead to some numerical problems if the masses of the two dark matter particles are very different. Let us call the light particle ℓ and the heavy particle h . We have to start the numerical solution at a temperature T above the freeze-out temperature of the heaviest DM,

$$T_{\text{fo}h} \approx M_h/25. \quad (19)$$

At this temperature,

$$\frac{Y_\ell}{Y_h} \approx \exp \frac{M_h - M_\ell}{T_{\text{fo}h}}, \quad (20)$$

and the step in the numerical solution of the two component equations will be suppressed by a factor $\exp(-M_h - M_\ell)/T_{\text{fo}h}$. This small step size is problematic when solving numerically the equation with the Runge-Kutta method. This occurs when $M_h/M_\ell > 2$. In this case the equation for the heavy component must be solved independently assuming that the light component has reached its equilibrium density. If $M_h/M_\ell < 2$, the Runge-Kutta procedure can be used to successfully solve the thermal evolution equations (12).

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