

# WHIZARD 2.0

## A generic Monte-Carlo integration and event generation package for multi-particle processes

### MANUAL<sup>1</sup>

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## ABSTRACT

**WHIZARD** is a program system designed for the efficient calculation of multi-particle scattering cross sections and simulated event samples. The generated events can be written to file in various formats (including HepMC, LHEF, STDHEP, and ASCII) or analyzed directly on the parton level using a built-in **L<sup>A</sup>T<sub>E</sub>X**-compatible graphics package.

Complete tree-level matrix elements are generated automatically for arbitrary partonic multi-particle processes by calling the built-in matrix-element generator **O'Mega**. Beyond hard matrix elements, **WHIZARD** can generate (cascade) decays with complete spin correlations. Various models beyond the SM are implemented, in particular, the MSSM is supported with an interface to the SUSY Les Houches Accord input format. Matrix elements obtained by alternative methods (e.g., including loop corrections) may be interfaced as well.

The program uses an adaptive multi-channel method for phase space integration, which allows to calculate numerically stable signal and background cross sections and generate unweighted event samples with reasonable efficiency for processes with up to eight and more final-state particles. Polarization is treated exactly for both the initial and final states. Quark or lepton flavors can be summed over automatically where needed.

For hadron collider physics, an interface to the LHAPDF library is provided.

For showering, fragmenting and hadronizing the final state, a PYTHIA and HERWIG interface are provided which follow the Les Houches Accord.

The **WHIZARD** distribution is available at

<http://whizard.event-generator.org>

or at

<http://projects.hepforge.org/whizard>

where also the **svn** repository is located.

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# Chapter 1

## Introduction

### 1.1 Disclaimer

*This is a very preliminary version of the WHIZARD manual. Many parts are still missing or incomplete, and some parts will be rewritten and improved soon. To find updated versions of the manual, visit the WHIZARD website*

<http://whizard.event-generator.org>

*or consult the current version in the `svn` repository on <http://projects.hepforge.org/whizard> directly.*

*For information that is not (yet) written in the manual, please consult the examples in the WHIZARD distribution. You will find these in the subdirectory `share/examples` of the main directory where WHIZARD is installed.*

## 1.2 Overview

## 1.3 About examples in this manual

Although **WHIZARD** has been designed as a Monte Carlo event generator for LHC physics, several elementary steps and aspects of its usage throughout the manual will be demonstrated with the famous textbook example of  $e^+e^- \rightarrow \mu^+\mu^-$ . This is the same process, the textbook by Peskin/Schroeder [14] uses as a prime example to teach the basics of quantum field theory. We use this example not because it is very special for **WHIZARD** or at the time being a relevant physics case, but simply because it is the easiest fundamental field theoretic process without the complications of structured beams (which can nevertheless be switched on like for ISR and beamstrahlung!), the need for jet definitions/algorithms and flavor sums; furthermore, it easily accomplishes a demonstration of polarized beams. After the basics of **WHIZARD** usage have been explained, we move on to actual physics cases from Tevatron or LHC.

# Chapter 2

## Installation

### 2.1 Prerequisites and Installation

The concept of the WHIZARD installation has been changed from version 1 to version 2. Now WHIZARD is centrally installed on a computer, e.g. in the `/usr/local`, and then the user has a working space which is completely separated from the WHIZARD installation directory. The WHIZARD tarball can be downloaded either from the WHIZARD webpage, <http://whizard.event-generator.org>, or the corresponding HepForge webpage, <http://projects.hepforge.org/whizard>. On the WHIZARD webpage, one can either download the tarball of the most recent version (or older versions), or one can check out the latest version from the subversion (svn) repository. The latter is only recommended for developers and users willing to accept that maybe not all newly installed features are already working. The check-out from the svn repository is done with the following command:

```
svn checkout http://svn.hepforge.org/whizard/trunk/ SomeLocalDir
```

Note again, that the subversion contains the latest developer version. In order to be able, to compile this, one has to first generate the `configure` script out of the file `configure.ac` by running `autoreconf` (NOT `autoconf`) which is part of the `autoconf/automake` (<http://www.gnu.org/software/autoconf/> and <http://www.gnu.org/software/automake>) package. Furthermore, the development version also needs the `noweb` tools to be installed on the system in order to extract the source codes and documentation from several so called `.nw` files. The `noweb` package can be downloaded and installed from <http://www.cs.tufts.edu/~nr/noweb/>.

The general prerequisites for the installation (i.e. also from the tarball, not only from the svn) are standard tools for software development like `make` etc., and two different compilers, a `FORTRAN2003` for the WHIZARD core and its corresponding libraries as well as an `O'Caml` compiler for the `O'Mega` matrix element generator.

Unpack the tarball, go to the WHIZARD directory, create a new directory and go to it. In that directory, perform a `../configureFC=<yourFORTRANcompiler>--prefix=/usr/local`. Note that this is because the source and compile directories should be different to avoid any problems during compilation and installation. `../configure--help` shows you the options for the `configure` process you have. The `FC` environment variable allows you to specify your

**FORTRAN** compiler of choice. Note that **WHIZARD** 2 has been written in **FORTRAN2003** in a fully object-oriented way. We highly recommend usage of the standard **gfortran** compiler from version 4.5.0 on. You can access the help menu of configure by `../configure --help`. `./configure -V` shows you the actual version of your downloaded **WHIZARD** distribution. The possible environment variables are:

CC	C compiler command
CFLAGS	C compiler flags
LDLFLAGS	linker flags, e.g. <code>-L&lt;lib dir&gt;</code> if you have libraries in a nonstandard directory <code>&lt;lib dir&gt;</code>
LIBS	libraries to pass to the linker, e.g. <code>-l&lt;library&gt;</code>
CPPFLAGS	C/C++/Objective C preprocessor flags, e.g. <code>-I&lt;include dir&gt;</code> if you have headers in a nonstandard directory <code>&lt;include dir&gt;</code>
CPP	C preprocessor
FC	Fortran compiler command
FCFLAGS	Fortran compiler flags
CXX	C++ compiler command
CXXFLAGS	C++ compiler flags
CXXCPP	C++ preprocessor

For most of these there is no need to be set during installation.

The configure process checks for the build and host system type; only if this is not detected automatically, the user would have to specify this by himself. After that system-dependent files are searched for, LaTeX and Acroread for documentation and plots, the **FORTRAN** compiler is checked, and finally the O’Caml compiler. The next step is the checks for external programs like LHAPDF and HepMC. Finally, all the Makefiles are being built.

The compilation is done by invoking `make` and finally `make install`. You could also do a `make check` in order to test whether the compilation has produced sane files on your system. This is highly recommended.

Be aware that there be problems for the installation if the install path or a user’s home directory is part of an AFS file system. Several times problems were encountered connected with conflicts with permissions inside the OS permission environment variables and the AFS permission flags which triggered errors during the `make install` procedure.

It is possible to compile **WHIZARD** without the O’Caml parts of O’Mega, namely by using the `--disable-omega` option of the configure. This will result in a built of **WHIZARD** with the O’Mega Fortran library, but without the binaries for the matrix element generation. All selftests (cf. 2.1.2) requiring O’Mega matrix elements are thereby switched off. Note that you can install such a built (e.g. on a batch system without O’Caml installation), but the try to build a distribution (all `make distxxx` targets) will fail.

### 2.1.1 Installation of optional external programs

There are several optional external programs that can be installed and linked to **WHIZARD**. The probably most important is LHAPDF [16] for using parton distribution functions (PDFs) for

hadron colliders. Other external programs or libraries are HepMC [17] and StdHEP [?] for the corresponding event formats. As LHAPDF has become a widely accepted tool and is easily available from the Hepforge server, we decided not to ship WHIZARD with a standard PDF. Note that because we believe this code is outdated by now, from WHIZARD 2.0 on we do no longer support the PDFLIB interface from the CERNLIB library.

As the LHAPDF homepage states, it provides a unified and easy to use interface to modern PDF sets. The LHAPDF package is available from the Hepforge address: <http://projects.hepforge.org/lhapdf/>. A comprehensive manual and description on how to install it. The basic procedure is the same as for WHIZARD itself, namely unpack it, configure it with a flag `FC=<your compiler>` for the Fortran 95 compiler and a flag `--prefix=<install dir.>` for the install directory, and then do a `make`. After that you can do an optional `make check`. Finally, install LHAPDF by doing `make install`. It is not mandatory to compile LHAPDF with the same Fortran compiler as WHIZARD, but of course desirable. In the worst case, when configuring WHIZARD you have to specify the run-time library for the Fortran compiler for LHAPDF by `LIBS=-L<Fortran run time>`. If this library is in a system-accessible library path, this is not necessary. When configuring WHIZARD, WHIZARD looks for the binary `lhapdf-config` (which is present since LHAPDF version 4.1.0): if this file is in an executable path, the environment variables for LHAPDF are automatically recognized by WHIZARD, as well as the version number. This should look like this in the `configure` output:

```
configure: -----
configure: --- LHAPDF ---
configure:
checking for lhapdf-config... /usr/local/bin/lhapdf-config
checking the LHAPDF version... 5.8.2
checking the LHAPDF pdfsets path... /usr/local/share/lhapdf/PDFsets
checking for initpdfsetm in -lLHAPDF... yes
configure: -----
```

If you want to use a different LHAPDF (e.g. because the one installed on your system by default is an older one), the preferred way to do so is to put the `lhapf-config` in an executable path that is checked before the system paths, e.g. `<home>/bin`.

A possible error could arise if LHAPDF had been compiled with a different Fortran compiler than WHIZARD, and if the run-time library of that Fortran compiler had not been included in the WHIZARD configure process. The output then looks like this:

```
configure: -----
configure: --- LHAPDF ---
configure:
checking for lhapdf-config... /usr/local/bin/lhapdf-config
checking the LHAPDF version... 5.8.2
checking the LHAPDF pdfsets path... /usr/local/share/lhapdf/PDFsets
checking for initpdfsetm in -lLHAPDF... no
configure: -----
```

So, the WHIZARD configure found the LHAPDF distribution, but could not link because it could not resolve the symbols inside the library. In case of failure, for more details confer the `config.log`.

The HepMC package [17] is an object oriented event record written in C++ for High Energy Physics Monte Carlo Generators. Many extensions from HEPEVT, the Fortran HEP standard, are supported: the number of entries is unlimited, spin density matrices can be stored with each vertex, flow patterns (such as color) can be stored and traced, integers representing random number generator states can be stored, and an arbitrary number of event weights can be included. The HepMC webpage is: <https://savannah.cern.ch/projects/hepmc/>, and the package can be downloaded from <http://lcgapp.cern.ch/project/simu/HepMC/download/>. Detailed information on the installation and usage can be found there. We give here only some brief details relevant for the usage with WHIZARD: For the compilation of HepMC one needs a C++ compiler. Then the procedure is the same as for the WHIZARD package, namely configure HepMC: `configure --with-momentum=GEV --with-length=MM --prefix=<install dir>`. Note that the particle momentum and decay length flags are mandatory, and we highly recommend to set them to the values GEV and MM, respectively. After configuration, do `make`, an optional `make check` (which might sometimes fail for non-standard values of momentum and length), and finally `make install`.

A WHIZARD configuration for HepMC is a bit lengthier as the C++ details have to be checked first:

```
configure: -----
configure: --- HepMC ---
configure:
checking for g++... g++
checking whether we are using the GNU C++ compiler... yes
checking whether g++ accepts -g... yes
checking dependency style of g++... gcc3
checking whether we are using the GNU C++ compiler... (cached) yes
checking whether g++ accepts -g... (cached) yes
checking dependency style of g++... (cached) gcc3
checking how to run the C++ preprocessor... g++ -E
checking for ld used by g++... /usr/bin/ld
checking if the linker (/usr/bin/ld) is GNU ld... yes
checking whether the g++ linker (/usr/bin/ld) supports shared libraries... yes
checking for g++ option to produce PIC... -fPIC -DPIC
checking if g++ PIC flag -fPIC -DPIC works... yes
checking if g++ static flag -static works... yes
checking if g++ supports -c -o file.o... yes
checking if g++ supports -c -o file.o... (cached) yes
checking whether the g++ linker (/usr/bin/ld) supports shared libraries... yes
checking dynamic linker characteristics... GNU/Linux ld.so
checking how to hardcode library paths into programs... immediate
checking the HepMC version... 2.05.01
checking for LDFLAGS_STATIC: host system is linux-gnu: static flag...
checking for GenEvent class in -lHepMC... yes
checking whether we are using the GNU Fortran compiler... (cached) yes
checking whether /usr/bin/gfortran accepts -g... (cached) yes
configure: -----
```

If WHIZARD does not automatically find the HepMC distribution (because it is installed in a non-standard path), or you want to use a different version than installed on your system, then

set the environment variable `HEPMC_DIR` during the `WHIZARD` configuration to the corresponding HepMC installation directory:

```
./configure HEPMC_DIR=<HEPMC install dir.> CXXFLAGS=<C++ flags>
```

The environment variable `CXXFLAGS` allows you to set specific C/C++ preprocessor flags, e.g. non-standard include paths for header files.

Besides the fact that StdHEP contains a set of translation routines which convert Herwig, Jetset, Isajet, or QQ events to and from the standard HEP event format, it also contains utility routines to work with the HEPEVT common block and a set of I/O routines. The second point is the interesting one for the usage with `WHIZARD`, as StdHEP provides a possibility to write machine-independent binary event files, using either the HEPEVT or the HEPRUP/HEPEUP common block. The StdHEP webpage is <http://cepa.fnal.gov/psm/stdhep/> and the package can be downloaded from <http://cepa.fnal.gov/psm/stdhep/getStdHep.shtml>. StdHEP is written in Fortran77. Although not really necessary, we strongly advice to compile StdHEP with the same compiler as `WHIZARD`. Otherwise, one has to add the corresponding Fortran77 run-time libraries to the configure command for `WHIZARD`. In order to compile StdHEP with a modern Fortran90/95/03 compiler, add the line `F77 = <your Fortran compiler>` below the `MAKE=make` statement in the GNUmakefile of the StdHEP distribution after you extracted the tarball (Note that there might be some difficulties that some modern compilers do not understand the D debugging precompiler statements in some of the files. In that case just replace them by comment characters, C. Also, some of the hard-coded compiler flags are tailor-made for old-fashioned g77). After that just do `make`. Copy the libraries created in the `lib` directory of your StdHEP distribution to a directory which is in the `LD_LIBRARY_PATH` of your computer.

The `WHIZARD` configure script will search for the two libraries `libFmcfio.a` and `libstdhep.a`. When `WHIZARD` does not find the StdHEP library, you have to set the location of the two libraries explicitly:

```
./configure ... ... ... STDHEP=<stdhep path>/libstdhep.a
FMCFIO=<fmcfio path>/libFmcfio.a
```

The corresponding configure output will look like this:

```
configure: -----
configure: --- STDHEP ---
configure:
checking for libFmcfio.a... /usr/local/lib/libFmcfio.a
checking for libstdhep.a... /usr/local/lib/libstdhep.a
checking for stdxwinit in -lstdhep -lFmcfio... yes
configure: -----
```

In the last line, `WHIZARD` checks whether it can correctly access functions from the library. If some symbols could not be resolved, it will put a “no” in the last entry. Then the `config.log` will tell you more about what went wrong in detail.

### 2.1.2 WHIZARD self tests/checks

WHIZARD has a number of self-consistency checks and test which assure that most of its features are running in the intended way. The standard procedure to invoke these self tests is to perform a `make check` from the `build` directory. If `src` and `build` directories are the same, all relevant files for these self-tests reside in the `test` subdirectory of the main WHIZARD directory. In that case, one could in principle just call the scripts individually from the command line. Note, that if `src` and `build` directory are different as recommended, then the input files will have been installed in `prefix/share/whizard/test`, while the corresponding test shell scripts remain in the `srcdir/test` directory. As the main shell script `run_whizard.sh` has been built in the `build` directory, one now has to copy the files over by and set the correct paths by hand, if one wishes to run the test scripts individually. `make check` still correctly performs all WHIZARD self-consistency tests.

There are additional, quite extensiv numerical tests for validation and backwards compatibility checks for SM and MSSM processes. As a standard, these extended self tests are not invoked. However, they can be enabled by setting the configure option `--enable-extrnum-checks`. On the other hand, the standard self-consistency checks can be completely disabled with the option `--disable-default-checks`.

## 2.2 Setting up a user work space

When WHIZARD is installed on a system it can be used by any user in a multi-user environment.

# Chapter 3

## Getting Started

WHIZARD can run as a stand-alone program. You (the user) can steer WHIZARD either interactively or by a script file. We will first describe the latter method, since it will be the most common way to interact with the WHIZARD system.

### 3.1 Hello World

The script is written in SINDARIN. This is a DSL – a domain-specific scripting language that is designed for the single purpose of steering WHIZARD.

Previous versions of the program, similar to most high-energy physics programs, relied on a bunch of input files that the user had to provide in some obfuscated format. This approach is sufficient for straightforward applications. However, once you get experienced with a program, you start thinking about uses that the program’s authors did not foresee. In case of a Monte Carlo package, typical abuses are parameter scans, complex patterns of cuts and reweighting factors, or data analysis without recourse to external packages. This requires more flexibility.

Instead of transferring control over data input to some generic scripting language like PERL or PYTHON (or even C++), which come with their own peculiarities and learning curves, we decided to unify data input and scripting in a dedicated steering language that is particularly adapted to the needs of Monte-Carlo integration, simulation, and simple analysis of the results. Thus we discovered what everybody knew anyway: that W(h)izards communicate in SINDARIN, Scripting INtegration, Data Analysis, Results display and INterfaces.

Now since SINDARIN *is* a programming language, we honor the old tradition of starting with the famous Hello World program. In SINDARIN this reads

```
printf "Hello World!"
```

Open your favorite editor, type this text, and save it into a file named `hello.sin`.

Now we assume that you – or your kind system administrator – has installed WHIZARD in your executable path. Then you should open a command shell and execute

```
> whizard -r hello.sin
```

and if everything works well, you get the output

```
| Writing log to 'whizard.log'
|=====
|                               WHIZARD 2.0.0_rc1
|=====
| Initializing process library 'processes'
| Reading model file 'SM.mdl'
| Using model: SM
| Reading commands from file 'hello.sin'
Hello World!
| WHIZARD run finished.
|=====
```

## 3.2 A Simple Calculation

You may object that **WHIZARD** is not exactly designed for printing out plain text. So let us demonstrate a more useful example.

Looking at the Hello World output, we first observe that the program writes a log file named (by default) `whizard.log`. This file receives all screen output, except for the output of external programs that are called by **WHIZARD**. You don't have to cache **WHIZARD**'s screen output yourself.

After the welcome banner, **WHIZARD** tells you that it initializes a *process library*, and it reads a physics *model*. The process library is initially empty. It is ready for receiving definitions of elementary high-energy physics processes (scattering or decay) that you provide. The processes are set in the context of a definite model of high-energy physics. By default this is the Standard Model, dubbed **SM**.

Here is the SINDARIN code for defining a SM physics process, computing its cross section, and generating a simulated event sample:

```
process ee = e1, E1 => e2, E2
compile

sqrtS = 360 GeV
integrate (ee)

n_events = 10
sample_format = lhef
$sample = "ee"
simulate (ee)
```

As before, you save this text in a file (named, e.g., `ee.sin`) which is run by

```
> whizard -r ee.sin
```

(We will come to the meaning of the `-r` option later.) This produces a lot of output. We break it down into pieces.

The startup is as before:

```

| Writing log to 'whizard.log'
|=====
|                               WHIZARD 2.0.0_rc1
|=====
| Initializing process library 'processes'
| Reading model file 'SM.mdl'
| Using model: SM
| Reading commands from file 'ee.sin'
| Added process to library 'processes':
|   [0] ee = e-, e+ => mu-, mu+
|
| Generating code for process library 'processes'
| Calling OMega for process 'ee'
| command:
| /home/kilian/whizard/build/nagfor/src/omega/bin/omega_SM.opt -o
| ee.f90 -target:whizard -target:parameter_module parameters_SM
| -target:module ee -target:md5sum 6ABA33BC2927925D0F073B1C1170780A
| -fusion:progress -scatter 'e- e+ => mu- mu+'
[1/1] e- e+ => mu- mu+ ... done. [time: 0.03 secs, total: 0.03 secs, remaining: 0.00 secs]
all processes done. [total time: 0.03 secs]
SUMMARY: 6 fusions, 2 propagators, 2 diagrams
| Writing interface code for process library 'processes'
| Compiling process library 'processes'

| Loading process library 'processes'
| Process 'ee': updating previous configuration
sqrtS = 3.600000000000000E+02
| Integrating process 'ee'
| Generating phase space, writing file 'ee.phs' (this may take a while)
| Found 2 phase space channels.
Warning: No cuts have been defined.

| Using partonic energy as event scale.
| iterations = 3:1000, 3:10000
| Creating VAMP integration grids:
| Using phase-space channel equivalences.
| 1000 calls, 2 channels, 2 dimensions, 20 bins, stratified = T
|=====
| It      Calls  Integral[fb]  Error[fb]  Err[%]    Acc  Eff[%]  Chi2 N[It] |
|=====
|   1      1000  8.3366006E+02  1.47E+00  0.18    0.06*  40.12
|   2      1000  8.3357740E+02  8.16E-01  0.10    0.03*  40.11
|   3      1000  8.3214263E+02  1.01E+00  0.12    0.04   57.40
| -----
|   3      3000  8.3311382E+02  5.83E-01  0.07    0.04   57.40  0.69  3
| -----
|   4      10000 8.3325834E+02  1.10E-01  0.01    0.01*  57.02
|   5      10000 8.3333796E+02  1.11E-01  0.01    0.01   57.03
|   6      10000 8.3323772E+02  1.11E-01  0.01    0.01   57.03
| -----
|   6      30000 8.3327798E+02  6.41E-02  0.01    0.01   57.03  0.23  3
| =====

```

```
n_events = 10
$sample => "ee"
| Initializing simulation for processes ee:
| Simulation mode = unweighted, event_normalization = '1'
| No analysis setup has been provided.
| Writing events in LHEF format to file 'ee.lhef'
| Generating 10 events ...
| Writing events in internal format to file 'ee.evx'
| Event sample corresponds to luminosity [fb-1] = 0.1200E-01
| ... done
| Simulation finished.
| There were no errors and 1 warning(s).
| WHIZARD run finished.
=====|
```

# Chapter 4

## SINDARIN: The WHIZARD command language

### 4.1 Why a specialized language?

A conventional physics application program gets its data from a set of input files. Alternatively, it is called as a library, so the user has to write his own code to interface ist, or it combines these two approaches. WHIZARD 1 was built in this way: there were some input files which were written by the user, and it could be called both stand-alone or as an external library.

WHIZARD 2 is also a stand-alone program. It comes with its own full-fledged script language, called SINDARIN. All interaction between the user and the program is done in SINDARIN expressions, commands, and scripts. Two main reasons led us to this choice:

- In any nontrivial physics study, cuts and (parton- or hadron-level) analysis are of central importance. The task of specifying appropriate kinematics and particle selection for a given process is well defined, but it is impossible to cover all possibilities in a simple format like the cut files of WHIZARD 1.

The usual way of dealing with this problem is to write analysis driver code (often in C++), using external libraries for Lorentz algebra etc. However, the overhead of writing correct C++ or Fortran greatly blows up problems that could be formulated in a few lines of text.

- While many problems lead to a repetitive workflow (process definition, integration, simulation), there are more involved tasks that involve parameter scans, comparisons of different processes, conditional execution, or writing output in widely different formats. This is easily done by a steering script, which should be formulated in a complete language.

The SINDARIN language is built specifically around event analysis, suitably extended to support steering, including data types, loops, conditionals, and I/O.

It would have been possible to use an established general-purpose language for these tasks. For instance, O'Caml which is a functional language would be a suitable candidate, and the

matrix-element generator is written in that language. Another candidate would be a popular scripting language such as PYTHON.

We do plan to support interfaces for commonly used languages in the future. However, introducing a speal-purpose language has the three distinct advantages: First, it is compiled and executed by the very Fortran code that handles data and thus accesses it without interfaces. Second, it can be designed with a syntax especially suited to the task of event handling and Monte-Carlo steering, and third, the user is not forced to learn all those features of a generic language that are of no relevance to the application he is interested in.

## 4.2 Basic concepts

### 4.2.1 SINDARIN scripts

SINDARIN scripts are contained in files. The user can create them using any editor of his choice. By convention, the files have the extension ‘.sin’. WHIZARD executes a script if the filename is given as an argument to the program:

```
$ whizard script.sin
```

Alternatively, scripts can be executed line by line interactively; we describe this below in Sec.4.4.

#### Commands

A SINDARIN script as a whole is a sequence of *commands*, similar to the commands in any imperative language such as Fortran or C. Examples of commands are `integrate` or `simulate`.

The script is free-form, i.e., indentation, extra whitespace and newlines are syntactically insignificant. In contrast to most languages, there is no command separator. Commands simply follow each other, just separated by whitespace.

Nevertheless, we recommend to use some line-oriented format and meaningful indentation, so the logical structure of a script is made explicit. How this is done in detail, is up to the script writer.

A command may consist of a *keyword*, a list of *arguments* in brackets (...), and an *option* script. In some cases, there is a zeroth (*suffix*) argument without brackets, immediately following the keyword.

Arguments enclosed in square brackets [] also exist. They have a special meaning, they denote subevents (collections of momenta) in event analysis.

The option script, if any, is enclosed in braces {...}. It is also a sequence of commands (possibly with their own options). Usually, it has the purpose of setting specific parameters in a context local to the command. The braces indicate a scoping unit; most parameters will be restored their previous values when the execution of that command is completed.

### 4.2.2 Data types and expressions

SINDARIN data are classified by their types. The language supports the classical numeric types

- **int** for integer: machine-default, usually 32 bit;
- **real**, usually *double precision* or 64 bit;
- **complex**, consisting of real and imaginary part equivalent to a **real** each.

SINDARIN contains arithmetic expressions and functions very much similar to conventional languages. In arithmetic expressions, the three numeric types can be mixed as appropriate. The computation essentially follows the rules for mixed arithmetic in Fortran or C.

The names of numerical variables consist of alphanumeric characters and underscores. The first character must not be a digit. Character case does matter. In this manual we follow the convention that variable names consist of lower-case letters, digits, and underscores only.

Exclusively in the context of particle selections (event analysis), there are *observables* as special numeric objects. They are used like numeric variables, but they are never declared or assigned. They get their value assigned dynamically, computed from the particle momentum configuration. Observable names begin with a capital letter.

The language also has the following standard types:

- **logical** (a.k.a. boolean). Logical variable names are prefixed by a ? (question mark) sign, otherwise the same rules as for numerical variables apply.
- **string** (arbitrary length). String variable names have a \$ (dollar) sign as prefix.

There are comparisons, logical operations, string concatenation, and a mechanism for formatting objects as strings for output.

Furthermore, SINDARIN supports two data types tailored specifically for Monte Carlo:

- **alias** objects denote a set of particle species. Alias names have no prefix, they are distinguished from numerics by context.
- **subevt** objects denote a collection of particle momenta within an event. Their names are prefixed by a @ (at) sign.

Each variable or object has a well-defined type. Some variables are predefined by the system (*intrinsic variables*). They are further separated into *independent* parameters that can be reset by the user, and *derived* parameters that are automatically computed by the program. On top of that, the user is free to introduce his own variables (*user variables*).

The script can assign new values to intrinsic variables, no value is hard-coded. User variables must be declared with any of the basic types when they are introduced. They retain this type until the end of program execution.

In the current implementation, SINDARIN has no container data types such as lists, arrays, or hashes. They may be introduced in a future language extension.

### 4.2.3 Special objects

In addition to the basic data types, SINDARIN contains objects that serve special purposes. Most of them do not occur in expressions, but there are assignment statements and operations acting on them with specific rules. Examples are physics models, processes, cut expressions, iteration specifiers, histograms, and more. These objects are intrinsic.

### 4.2.4 Control structures

Any computer language has some concept of conditionals and loops. In SINDARIN, conditionals are represented by the usual `if-then-elsif-else-endif` sequence. Loops are labeled according to their main purpose in Monte Carlo context, parameter scans. The loop syntax is therefore `scan variable = (values) {...}`.

## 4.3 Basic data and expressions

### 4.3.1 Real-valued objects

Real literals have their usual form, mantissa and, optionally, exponent:

```
0. 3.14 -.5 2.345e-3 .890E-023
```

Internally, real values are treated as double precision. The values are read by the Fortran library, so details depend on its implementation.

A special feature of SINDARIN is that numerics (real and integer) can be immediately followed by a physical unit. The supported units are presently hard-coded, they are

meV	eV	keV	MeV	GeV	TeV
nbar	pbar	fbarn	abarn		
rad	mrad	degree			
	%				

If a number is followed by a unit, it is automatically normalized to the corresponding default unit: `14.Tev` is transformed into the real number 14000. Default units are `GeV`, `fbarn`, and `rad`. The `%` sign after a number has the effect that the number is multiplied by 0.01. Note that no checks for consistency of units are done, so you can add `1 meV + 3 abarn` if you absolutely wish to. Omitting units is always allowed, in that case, the default unit is assumed.

Units are not treated as variables. In particular, you can't write `theta / degree`, the correct form is `theta / 1 degree`.

There is a single predefined real constant, namely  $\pi$  which is referred to by the keyword `pi`. The arithmetic operators are

```
+ - * / ^
```

with their obvious meaning and the usual precedence rules. Grouping is done using ordinary brackets () .

SINDARIN supports a bunch of standard functions, mostly equivalent to their Fortran counterparts:

```
abs  sgn  mod  modulo
sqrt  exp  log  log10
sin  cos  tan  asin  acos  atan
          sinh  cosh  tanh
```

(Unlike Fortran, the `sgn` function takes only one argument and returns 1., 0., or -1.) The function argument is enclosed in brackets: `sqrt (2.)`, `tan (11.5 degree)`.

There are two functions with two real arguments:

```
max  min
```

with `min (mZ, mH)` as an example.

The following functions of a real convert to integer:

```
int  nint  floor  ceiling
```

and this converts to complex type:

```
complex
```

Real values can be compared by the following operators, the result is a logical value:

```
==  <>
>  <  >=  <=
```

In SINDARIN, it is possible to have more than two operands in a logical expressions. The comparisons are done from left to right. Hence,

```
115 GeV < mH < 180 GeV
```

is valid SINDARIN code and evaluates to `true` if the Higgs mass is in the given range.

Tests for equality and inequality with machine-precision real numbers are notoriously unreliable and should be avoided altogether. To deal with this problem, SINDARIN has the “fuzzy” comparison operators

```
==~  <>~
```

which should be read as “equal (unequal) up to a tolerance”, where the tolerance is given by the real-valued intrinsic variable `tolerance`. This variable is initially zero, but can be set to any value (for instance, `tolerance = 1.e-13` by the user. Note that these operators, in contrast to `==` vs. `<>`, are not mutually exclusive.

### 4.3.2 Integer-valued objects

Integer literals are obvious:

```
1 -98765 0123
```

Integers are always signed. Their range is the default-integer range as determined by the Fortran compiler.

Like real values, integer values can be followed by a physical unit: `1 TeV`, `30 degree`. This actually transforms the integer into a real.

Standard arithmetics is supported:

```
+ - * / ^
```

It is important to note that there is no fraction datatype, and pure integer arithmetics does not convert to real. Hence `3/4` evaluates to 0.

Since all arithmetics is handled by the underlying Fortran library, integer overflow is not detected. If in doubt, do real arithmetics.

Integer functions are more restricted than real functions. We support the following:

```
abs  sgn  mod  modulo
      max  min
```

and the conversion functions

```
real  complex
```

Comparisons of integers among themselves and with reals are possible using the same set of comparison operators as real values. This includes the operators `==~` and `<>~`.

### 4.3.3 Complex-valued objects

Complex values are currently not yet used by the physics models implemented in `WHIZARD`, so they play no role in most applications.

There is no form for complex literals. Complex values must be created via an arithmetic expression,

```
complex c = 1 + 2 * I
```

where `I` is a predefined constant.

The standard arithmetic operations are supported (also mixed with real and integer). Support for functions is incomplete, among the supported functions there is `sqr`, `log`, `exp`.

### 4.3.4 Logical-valued objects

There are two predefined logical constants, `true` and `false`. Logicals are *not* equivalent to integers (like in C) or to strings (like in PERL), but they make up a type of their own. Only in `printf` output, they are treated as strings, that is, they require the `%s` conversion specifier.

The names of logical variables begin with a question mark ?:

```
?higgs_decays_into_tt = mH > 2 * mtop
```

Logical expressions use the standard boolean operations

or and not

Furthermore, the results of comparisons (see above) are logicals.

A primary use for intrinsic logicals are flags that change the behavior of commands. For instance, `?unweighted = true` and `?unweighted = false` switch the unweighting of a simulated event samples on and off.

Logical expressions are furthermore used for specifying cut and analysis expressions.

### 4.3.5 String-valued objects and string operations

String literals are enclosed in double quotes: "This is a string." The empty string is "".

String variables begin with \$. There is only one string operation, concatenation

```
$string = "abc" // "def"
```

However, it is possible to variables and values to a string using the `sprintf` function. This function is an interface to the system's C function `sprintf` with some restrictions and modifications. The allowed conversion specifiers are

%d %i (integer)
%e %f %g %E %F %G (real)
%s (string and logical)

The conversions can use flag parameter, field width, and precision, but length modifiers are not supported since they have no meaning for the application.

The `sprint` function has the syntax

```
sprintf format-string (arg-list)
```

where the format string contains the mentioned conversion specifiers. The argument list is optional. The arguments are separated by commas. Allowed arguments are integer, real, logical, and string variables, and numeric expressions. Logical and string expressions can also be printed, but they have to be dressed as *anonymous variables*. A logical anonymous variable has the form `?(logical-expr)` (example: `?(mH > 115 GeV)`). A string anonymous variable has the form `$(string-expr)`.

Example:

```
string $unit = "GeV"
string $str = sprintf "mW = %f %s" (mW, $unit)
```

The related `printf` command with the same syntax prints the formatted string to standard output. There is also a `sprint` function and a `print` command; they have no format string but typeset their arguments in a default format.

### 4.3.6 Particle aliases

A particle species is denoted by its name as a string: `W+`. Alternatively, it can be addressed by an `alias`. The model files predefine aliases for all particles. For instance, the  $W^+$  has the alias `Wp`. Aliases are used like variables, but in contexts where a particle species is expected.

In particular, aliases can specify *alternatives* for particle species. This is done by concatenating particle names or aliases by colons `:`. Example:

```
alias quark = u:d:s
```

Aliases are used for defining processes with summation over flavors, and for defining classes of particles for analysis.

Each model predefines the multi-species alias `colored` and `charged`.

## 4.4 WHISH – The WHIZARD Shell/Interactive mode

WHIZARD can be also run in the interactive mode using its own shell environment. This is called the WHIZARD Shell (WHISH). For this purpose, one starts with the command

```
/home/user$ whizard --interactive
```

or

```
/home/user$ whizard -i
```

The WHISH can be closed by the `quit` command:

```
whish? quit
```

## 4.5 Using processes from several different models

When using two different models which need an SLHA input file, these *have* to be provided for both models. Otherwise WHIZARD will not be performing the phase-space setup for the second process.

Note that when using more than one models, the setting of parameters *after* the last model and process declarations only affects the active – i.e. the last – model. If one wants to set a parameter for all models in the input file, one has to repeat the model setting for every defined model. Although this might seem cumbersome at first, it is nevertheless a sensible procedure since the parameters defined by the user might anyhow not be defined or available for all chosen models.

# Chapter 5

## Examples

In this chapter we discuss the running and steering of WHIZARD with the help of several examples. These examples can be found in the `share/examples` directory of your installation.



# **Chapter 6**

## **Implemented physics**

**6.1 The Monte-Carlo integration routine: VAMP**

**6.2 The Phase-Space Setup**

**6.3 The hard interaction models**

**6.3.1 The Standard Model and friends**

**6.3.2 Beyond the Standard Model**

MODEL TYPE	with CKM matrix	trivial CKM
Yukawa test model	---	Test
QED with $e, \mu, \tau, \gamma$	---	QED
QCD with $d, u, s, c, b, t, g$	---	QCD
Standard Model	SM_CKM	SM
SM with anomalous gauge couplings	SM_ac_CKM	SM_ac
SM with anomalous top couplings	---	SM_top
SM with K matrix	---	SM_KM
MSSM	MSSM_CKM	MSSM
MSSM with gravitinos	---	MSSM_Grav
NMSSM	NMSSM_CKM	NMSSM
extended SUSY models	---	PSSSM
Littlest Higgs	---	Littlest
Littlest Higgs with ungauged $U(1)$	---	Littlest_Eta
Littlest Higgs with $T$ parity	---	Littlest_Tpar
Simplest Little Higgs (anomaly-free)	---	Simplest
Simplest Little Higgs (universal)	---	Simplest_univ
SM with graviton	---	Xdim
UED	---	UED
SM with $Z'$	---	Zprime
“SQED” with gravitino	---	GravTest
Augmentable SM template	---	Template

Table 6.1: *List of models available in WHIZARD.* There are pure test models or models implemented for theoretical investigations, a long list of SM variants as well as a large number of BSM models.

# Chapter 7

## Event generation and analysis

In order to perform a physics analysis with **WHIZARD** one has to generate events. This seems to be a trivial statement, but as there have been any questions like "My **WHIZARD** does not produce plots – what has gone wrong?" we believe that repeating that rule is worthwhile. Of course, it is not mandatory to use **WHIZARD**'s own analysis set-up, the user can always choose to just generate events and use his/her own analysis package like ROOT, or TopDrawer, or you name it for the analysis.

Accordingly, we first start to describe how to generate events and what options there are – different event formats, renaming output files, using weighted or unweighted events with different normalizations. How to re-use and manipulate already generated event samples, how to limit the number of events per file, etc. etc.

### 7.1 Event generation

To explain how event generation works, we again take our favourite example,  $e^+e^- \rightarrow \mu^+\mu^-$ ,

```
process eemm = e1, E1 => e2, E2  
compile
```

The command to trigger generation of events is `simulate (<proc_name>) { <options> }`, so in our case – neglecting any options for now – simply:

```
simulate (eemm)
```

When you run this SINDARIN file you will experience a fatal error: **FATAL ERROR: Process 'eemm' must be integrated before simulation..** This is because you have to provide **WHIZARD** with the information of the corresponding cross section, phase space parameterization and grids, i.e. you have to integrate a process before you could generate events. A corresponding `integrate` command like

```
sqrtS = 500 GeV  
integrate (eemm) { iterations = 3:10000 }
```

obviously has to appear *before* the corresponding `simulate` command (otherwise you would be punished by the same error message as before). Putting things in the correct order results in an output like:

```
| Loading process library 'processes'
| Process 'eemm': updating configuration
sqrt = 500.0000000000000000
| Integrating process 'eemm'
| Generating phase space configuration ...
| ... found 2 phase space channels, collected in 2 groves.
| Phase space: found 2 equivalences between channels.
| Wrote phase-space configuration file 'eemm.phs'.
Warning: No cuts have been defined.
| Using partonic energy as event scale.
| iterations = 3:10000
| Creating VAMP integration grids:
| Using phase-space channel equivalences.
| 10000 calls, 2 channels, 2 dimensions, 20 bins, stratified = T
=====
| It      Calls  Integral[fb]  Error[fb]   Err[%]    Acc  Eff[%]  Chi2 N[It] |
=====|
  1      10000  4.2823916E+02  6.75E-02   0.02    0.02*  40.29
  2      10000  4.2823862E+02  4.37E-02   0.01    0.01*  40.29
  3      10000  4.2824459E+02  3.38E-02   0.01    0.01*  40.29
=====
| 3      30000  4.2824192E+02  2.48E-02   0.01    0.01  40.29   0.01  3
=====
| Process 'eemm':
|   time estimate for 10000 unweighted events = 0h 00m 00.469s
| -----
| Initializing simulation for processes eemm:
| Simulation mode = unweighted, event_normalization = '1'
| No analysis setup has been provided.
| Simulation finished.
| There were no errors and 1 warning(s).
| WHIZARD run finished.
=====|
```

So, `WHIZARD` tells you that it has entered simulation mode, but besides this, it has not done anything. The next step is that you have to demand event generation – there are two ways to do this: you could either specify a certain number, say 42, of events you want to have generated by `WHIZARD`, or you could provide a number for an integrated luminosity of some experiment. (Note, that if you choose to take both options, `WHIZARD` will take the one which gives the larger event sample. This, of course, depends on the given process(es) – as well as cuts – and its corresponding cross section(s).) The first of these options is set with the command: `n_events = <number>`, the second with `luminosity = <number> <opt. unit>`.

Another important point already stated several times in the manual is that `WHIZARD` follows the commands in the steering SINDARIN file in a chronological order. Hence, a given number of events or luminosity *after* a `simulate` command will be ignored – or are relevant only for any `simulate` command potentially following further down in the SINDARIN file. So, in our

case, try:

```
n_events = 500
luminosity = 10
simulate (eemm)
```

Per default, numbers for integrated luminosity are understood as inverse femtobarn. So, for the cross section above this would correspond to 4282 events, clearly superseding the demand for 500 events. After reducing the luminosity number from ten to one inverse femtobarn, 500 is the larger number of events taken by **WHIZARD** for event generation. Now **WHIZARD** tells you:

```
| No analysis setup has been provided.
| Generating 500 events ...
| Writing events in internal format to file 'whizard.evx'
| Event sample corresponds to luminosity [fb-1] = 1.167
```

I.e., it evaluates the luminosity to which the sample of 500 events would correspond to, which is now, of course, bigger than the  $1\text{fb}^{-1}$  explicitly given for the luminosity. Furthermore, you can read off that a file **whizard.evx** has been generated, containing the demanded 500 events. Files with the suffix **.evx** are binary format event files, using a machine-independent **WHIZARD**-specific event file format. Before we list the event formats supported by **WHIZARD**, the next two section tell you more about unweighted and weighted events as well as different possibilities to normalize events in **WHIZARD**.

As already explained for the libraries, as well as the phase space and grid files, **WHIZARD** is trying to re-use as much information as possible. The same holds for the event files. There are special MD5 check sums testing the integrity and compatibility of the event files. If you demand for a process with an already existing event file less or equally many events as generated before, **WHIZARD** will not generate again but re-use the existing events (as will be explained below, the events are stored in a **WHIZARD**-own binary event format, i.e. in a so-called **.evx** file. If you suppress generation of that file, as will be described in subsection 7.1.3 then **WHIZARD** has to generate events all the time). Re-using event files is very practical for doing several different analyses with the same data, especially if there are many and big data samples. Consider the case, there is an event file with 200 events, and you now ask **WHIZARD** to generate 300 events, then it will re-use the 200 events (if MD5 check sums are OK!), generate the remaining 100 events and append them to the existing file. If the user for some reason, however, wants to regenerate events (i.e. ignoring possibly existing events), there is the command option **whizard --rebuild-events**.

### 7.1.1 Unweighted and weighted events

**WHIZARD** is able to generate unweighted events, i.e. events that are distributed uniformly and each contribute with the same event weight to the whole sample. This is done by mapping out the phase space of the process under consideration according to its different phase space channels (which each get their own weights), and then unweighting the sample of weighted events. Only a sample of unweighted events could in principle be compared to a real data sample from some

experiment. The seventh column in the **WHIZARD** iteration/adaptation procedure tells you about the efficiency of the grids, i.e. how well the phase space is mapped to a flat function. The better this is achieved, the higher the efficiency becomes, and the closer the weights of the different phase space channels are to uniformity. This means, for higher efficiency less weighted events ("calls") are needed to generate a single unweighted event. An efficiency of 10 % means that ten weighted events are needed to generate one single unweighted event. After the integration is done, **WHIZARD** uses the duration of calls during the adaptation to estimate a time interval needed to generate 10,000 unweighted events. The ability of the adaptive mult-channel Monte Carlo decreases with the number of integrations, i.e. with the number of final state particles. Adding more and more final state particles in general also increases the complexity of phase space, especially its singularity structure. For a  $2 \rightarrow 2$  process the efficiency is roughly of the order of several tens of per cent. As a rule of thumb, one can say that with every additional pair of final state particle the average efficiency one can achieve decreases by a factor of five to ten.

The default of **WHIZARD** is to generate *unweighted* events. One can use the logical variable `?unweighted = false` to disable unweighting and generate weighted events. (The command `?unweighted = true` is a tautology, because `true` is the default for this variable.) Note that again this command has to appear *before* the corresponding **simulate** command, otherwise it will be ignored or effective only for any **simulate** command appearing later in the SINDARIN file.

**Excess events** to be done...

### 7.1.2 Choice on event normalizations

There are basically four different choices to normalize event weights (... denotes the average) :

1.  $\langle w_i \rangle = 1, \quad \langle \sum_i w_i \rangle = N$
2.  $\langle w_i \rangle = \sigma, \quad \langle \sum_i w_i \rangle = N \times \sigma$
3.  $\langle w_i \rangle = 1/N, \quad \langle \sum_i w_i \rangle = 1$
4.  $\langle w_i \rangle = \sigma/N, \quad \langle \sum_i w_i \rangle = \sigma$

So the four options are to have the average weight equal to unity, to the cross section of the corresponding process, to one over the number of events, or the cross section over the event calls. In these four cases, the event weights sum up to the event number, the event number times the cross section, to unity, and to the cross section, respectively. Note that neither of these really guarantees that all event weight individually lie in the interval  $0 \leq w_i \leq 1$ .

The user can steer the normalization of events by using in SINDARIN input files the string variable `$event_normalization`. The default is `$event_normalization = "auto"`, which uses option 1 for unweighted and 2 for weighted events, respectively. Note that this is also what the Les Houches Event Format (LHEF) demands for both types of events. This is **WHIZARD**'s preferred mode, also for the reason, that event normalizations are independent from the number of events. Hence, event samples can be cut or expanded without further need to adjust

Format	Type	remark	ext.
Athena	ASCII	variant of HEPEVT	no
debug	ASCII	most verbose <b>WHIZARD</b> format	no
default	ASCII	<b>WHIZARD</b> verbose format	no
evx	binary	<b>WHIZARD</b> 's home-brew	no
HepMC	ASCII	HepMC format	yes
HEPEVT	ASCII	<b>WHIZARD</b> 1 style	no
LHA	ASCII	<b>WHIZARD</b> 1/old Madgraph style	no
LHEF	ASCII	Les Houches accord compliant	no
long	ASCII	variant of HEPEVT	no
StdHEP (HEPEVT)	binary	based on HEPEVT common block	yes
StdHEP (HEPRUP/EUP)	binary	based on HEPRUP/EUP common block	yes

Table 7.1: *Event formats supported by **WHIZARD**, classified according to ASCII/binary formats and whether an external program or library is needed to generate a file of this format.*

the normalization. The unit normalization (option 1) can be switched on also for weighted events by setting the event normalization variable equal to "1" or "unity". Option 2 can be demanded by setting `event_normalization = "sigma"`. Options 3 and 4 can be set by "`1/n`" and "`sigma/n`", respectively. **WHIZARD** accepts small and capital letter for these expressions.

In the following section we show some examples when discussing the different event formats available in **WHIZARD**.

### 7.1.3 Supported event formats

Event formats can either be distinguished whether they are plain text (i.e. ASCII) formats or binary formats. Besides this, one can classify event formats according to whether they are natively supported by **WHIZARD** or need some external program or library to be linked. Table 7.1 gives a complete list of all event formats available in **WHIZARD**. The second column shows whether these are ASCII or binary formats, the third column contains brief remarks about the corresponding format, while the last column tells whether external programs or libraries are needed (which is the case only for StdHEP and HepMC formats).

The ".evx" is **WHIZARD**'s native binary event format. If you demand event generation and do not specify anything further, **WHIZARD** will write out its events exclusively in this binary format. So in the examples discussed in the previous sections (where we omitted all details about event formats), in all cases this and only this internal binary format has been generated. The generation of this raw format can be suppressed (e.g. if you want to have only one specific event file type) by setting the variable `$write_raw = false`. However, if the raw event file is not present, **WHIZARD** is not able to re-use existing events (e.g. from an ASCII file) and will regenerate events for a given process.

Other event formats can be written out by setting the variable `sample_format = <format>`, where `<format>` can be any of the following supported variables:

- **ascii**: a quite verbose ASCII format which contains lots of information (an example is shown in the appendix).  
Standard suffix: `.evt`
- **debug**: an even more verbose ASCII format intended for debugging which prints out also information about the internal data structures  
Standard suffix: `.debug`
- **hepevt**: ASCII format that writes out a specific incarnation of the HEPEVT common block (**WHIZARD 1** back-compatibility)  
Standard suffix: `.hepevt`
- **short**: abbreviated variant of the previous HEPEVT (**WHIZARD 1** back-compatibility)  
Standard suffix: `.short.evt`
- **long**: HEPEVT variant that contains a little bit more information than the short format but less than HEPEVT (**WHIZARD 1** back-compatibility)  
Standard suffix: `.long.evt`
- **athena**: HEPEVT variant suitable for read-out in the ATLAS ATHENA software environment (**WHIZARD 1** back-compatibility)  
Standard suffix: `.athena.evt`
- **lha**: Implementation of the Les Houches Accord as it was in the old MadEvent and **WHIZARD 1**  
Standard suffix: `.lha`
- **lhef**: Formatted Les Houches Accord implementation that contains the XML headers  
Standard suffix: `.lhef`
- **hepmc**: HepMC ASCII format (only available if HepMC is installed and correctly linked)  
Standard suffix: `.hepmc`
- **stdhep**: StdHEP binary format based on the HEPEVT common block (only available if StdHEP is installed and correctly linked)  
Standard suffix: `.stdhep`
- **stdhep\_up**: StdHEP binary format based on the HEPUP/HEPEUP common blocks (only available if StdHEP is installed and correctly linked)  
Standard suffix: `.up.stdhep`

Of course, the variable `sample_format` can contain more than one of the above identifiers, in which case more than one different event file format is generated. The list above also shows the standard suffixes for these event formats (remember, that the native binary format of **WHIZARD** does have the suffix `.evx`). (The suffix of the different event format can even be changed by the user by setting the corresponding variable `$extension_lhef = "foo"` or `$extension_ascii_short = "bread"`. The dot is automatically included.)

The name of the corresponding event sample is taken to be the string of the name of the first process in the `simulate` statement. Remember, that conventionally the events for all processes in one `simulate` statement will be written into one single event file. So `simulate (proc1, proc2)` will write events for the two processes `proc1` and `proc2` into one single event file with name `proc1.evx`. The name can be changed by the user with the command `$sample = "<name>"`.

The commands `$sample` and `sample_format` are both accepted as optional arguments of a `simulate` command, so e.g. `simulate (proc) { $sample = "foo" sample_format = hepmc }` generates an event sample in the HepMC format for the process `proc` in the file `foo.hepmc`.

Examples for event formats (in the sequel, we gave the numbers out as single precision for better readability), for specifications of the event formats correspond the different accords and publications:

### HEPEVT:

The HEPEVT is an ASCII event format that does not contain an event file header. There is a one-line header for each single event, containing four entries. The number of particles in the event (`ISTHEP`), which is four for our example process  $e^+e^- \rightarrow \mu^+\mu^-$ , but could be larger if e.g. beam remnants are demanded to be included in the event. The second entry and third entry are the number of outgoing particles and beam remnants, respectively. The event weight is the last entry. For each particle in the event there are three lines: the first one is the status according to the HEPEVT format, `ISTHEP`, the second one the PDG code, `IDHEP`, then there are the one or two possible mother particle, `JMOHEP`, the first and last possible daughter particle, `JDAHEP`, and the polarization. The second line contains the three momentum components,  $p_x$ ,  $p_y$ ,  $p_z$ , the particle energy  $E$ , and its mass,  $m$ . The last line contains the position of the vertex in the event reconstruction.

4	2	0	1.00000000
2	11	0	0
0.00000000	0.00000000	249.999999	250.000000
0.00000000	0.00000000	0.00000000	0.00000000
2	-11	0	0
0.00000000	0.00000000	-249.999999	250.000000
0.00000000	0.00000000	0.00000000	0.00000000
1	13	1	2
225.985918	-80.1076510	70.8033735	250.000000
0.00000000	0.00000000	0.00000000	0.00000000
1	-13	1	2
-225.985918	80.1076510	-70.8033735	250.000000
0.00000000	0.00000000	0.00000000	0.00000000

### ASCII SHORT:

This is basically the same as the HEPEVT standard, but very much abbreviated. The header line for each event is identical, but first line per particle does only contain the PDG and the polarization, while the vertex information line is omitted.

4	2	0	1.00000000
11	0		
0.00000000	0.00000000	249.999999	250.000000
-11	0		
0.00000000	0.00000000	-249.999999	250.000000

13	0			
225.985918	-80.1076510	70.8033735	250.000000	0.10565838
-13	0			
-225.985918	80.1076510	-70.8033735	250.000000	0.10565838

**ASCII LONG:**

Identical to the ASCII short format, but after each event there is a line containing two values: the value of the sample function to be integrated over phase space, so basically the squared matrix element including all normalization factors, flux factor, structure functions etc.

4	2	0	1.00000000	
11	0			
0.00000000	0.00000000	249.999999	250.000000	5.11003380E-004
-11	0			
0.00000000	0.00000000	-249.999999	250.000000	5.11003380E-004
13	0			
225.985918	-80.1076510	70.8033735	250.000000	0.10565838
-13	0			
-225.985918	80.1076510	-70.8033735	250.000000	0.10565838
435.480971	1.00000000			

**ATHENA:**

Quite similar to the HEPEVT ASCII format. The header line, however, does contain only two numbers: an event counter, and the number of particles in the event. The first line for each particle lacks the polarization information (irrelevant for the ATHENA environment), but has as leading entry an ordering number counting the particles in the event. The vertex information line has only the four relevant position entries.

1	4			
1	2	11	0	3
0.00000000	0.00000000	249.999999	250.000000	5.11003380E-004
0.00000000	0.00000000	0.00000000	0.00000000	
2	2	-11	0	3
0.00000000	0.00000000	-249.999999	250.000000	5.11003380E-004
0.00000000	0.00000000	0.00000000	0.00000000	
3	1	13	1	2
225.985918	-80.1076510	70.8033735	250.000000	0.10565838
0.00000000	0.00000000	0.00000000	0.00000000	
4	1	-13	1	2
-225.985918	80.1076510	-70.8033735	250.000000	0.10565838
0.00000000	0.00000000	0.00000000	0.00000000	

**LHA:**

This is the implementation of the Les Houches Accord, as it was used in WHIZARD 1 and the old MadEvent. There is a first line containing six entries: 1. the number of particles in the event, NUP, 2. the subprocess identification index, IDPRUP, 3. the event weight, XWGTUP, 4. the scale of the process, SCALUP, 5. the value or status of  $\alpha_{QED}$ , AQEDUP, 6. the value for  $\alpha_s$ , AQCDUP. The next seven lines contain as many entries as there are particles in the event: the first one has the PDG codes, IDUP, the next two the first and second mother of the particles, MOTHUP, the fourth and fifth line the two color indices, ICOLUP, the next one the status of the particle, ISTUP, and the last line the polarization information, ISPINUP. At the end of the event there are as lines for each particles with the counter in the event and the four-vector of the particle. For more information on this event format confer [9].

```

4      1      1.0000000000  500.000000  -1.000000   0.117800
11     -11     13    -13
0       0      1       1
0       0      2       2
0       0      0       0
0       0      0       0
-1      -1      1       1
9       9      9       9
1  250.00000000000  0.0000000000  0.0000000000  249.999999995
2  250.00000000000  0.0000000000  0.0000000000 -249.999999995
3  250.00000000000  223.6404152843 -102.7925182666  43.8024162280
4  250.00000000000 -223.6404152843  102.7925182666 -43.8024162280

```

**LHEF:**

This is the modern version of the Les Houches accord event format (LHEF), for the details confer the corresponding publication [11].

```

<LesHouchesEvents version="1.0">
<header>
  <generator_name>WHIZARD</generator_name>
  <generator_version>2.0.0</generator_version>
</header>
<init>
  11      -11  250.000000  250.000000          -1          -1          -1          -1          3
  0.347536454  1.413672505E-004  1.00000000          1
</init>
<event>
  4      1  1.00000000  500.000000  -1.00000000  0.117800000
  11     -1      0      0      0      0  0.00000000  0.00000000  249.999999
  -11     -1      0      0      0      0  0.00000000  0.00000000 -249.999995
  13      1      1      2      0      0  223.640415 -102.792518  43.8024162
  -13     1      1      2      0      0 -223.640415  102.792518 -43.8024162
</event>
</LesHouchesEvents>

```

Sample files for the default ASCII format as well as for the debug event format are shown in the appendix.

#### 7.1.4 Negative weight events



# Chapter 8

## Technical details – Advanced Spells

### 8.0.5 Efficiency and tuning

Since massless fermions and vector bosons (or almost massless states in a certain approximation) lead to restrictive selection rules for allowed helicity combinations in the initial and final state. To make use of this fact for the efficiency of the **WHIZARD** program, we are applying some sort of heuristics: **WHIZARD** dices events into all combinatorially possible helicity configuration during a warm-up phase. The user can specify a helicity threshold which sets the number of zeros **WHIZARD** should have got back from a specific helicity combination in order to ignore that combination from now on. By that mechanism, typically half up to more than three quarters of all helicity combinations are discarded (and hence the corresponding number of matrix element calls). This reduces calculation time up to more than one order of magnitude. **WHIZARD** shows at the end of the integration those helicity combinations which finally contributed to the process matrix element.

Note that this list – due to the numerical heuristics – might very well depend on the number of calls for the matrix elements per iteration, and also on the corresponding random number seed.



# Appendix A

## SINDARIN Reference

*This appendix is out-of-date and needs revision.*

In the SINDARIN language, there are certain pre-defined constructors or commands that cannot be used in different context by the user, which are – in alphabetical order – `alias`, `all`, `$analysis_filename`, `and`, `as`, `any`, `beams`, `cplx`, `combine`, `compile`, `cuts`, `$description`, `echo`, `else`, `exec`, `expect`, `false`, `?fatal_beam_decay`, `if`, `include`, `int`, `integrate`, `iterations`, `$label`, `lhapdf`, `library`, `load`, `luminosity`, `model`, `n_events`, `no`, `observable`, `or`, `$physical_unit`, `plot`, `process`, `read_slha`, `real`, `?rebuild`, `?recompile`, `record`, `$restrictions`, `results`, `$sample`, `sample_format`, `scan`, `seed`, `show`, `simulate`, `sqrts`, `then`, `$title`, `tolerance`, `true`, `unstable`, `write_analysis`, `write_slha`, `$ xlabel`, and `$ ylabel`. Also units are fixed, like `degree`, `eV`, `keV`, `q MeV`, `GeV`, and `TeV`. Again, these tags are locked and not user-redefinable. There functionality will be listed in detail below. Furthermore, a variable with a preceding question mark, ?, is a logical, while a preceding hash, #, denotes a character string variable. Also, a lot of unary and binary operators exist, `+ - \ , = : => < > <= >= ^ () [] {} ~~~, as well as quotation marks, ”. Note that the different parentheses and brackets fulfill different purposes, which will be explained below. Comments in a line can be marked by a hash, #, or an exclamation mark, !.`

- `alias`

This allows to define a collective expression for a class of particles, e.g. to define a generic expression for leptons, neutrinos or a jet as `alias lepton = e1:e2:e3:E1:E2:E3`, `alias neutrino = n1:n2:n3:N1:N2:N3`, and `alias jet = u:d:s:c:U:D:S:C:g`, respectively.

- `all`

`all` is a function that works on a logical expression and a list, `all <log_expr> [<list>]`, and returns `true` if and only if `log_expr` is fulfilled for *all* entries in `list`, and `false` otherwise. Examples: `all Pt > 100 GeV [lepton]` checks whether all leptons are harder than 100 GeV, `all Dist > 2 [u:U, d:D]` checks whether all pairs of corresponding quarks are separated in *R* space by more than 2. Logical expressions with `all` can be logically combined with `and` and `or`. (cf. also `any`, `and`, `no`, and `or`)

- `$analysis_filename`

This character variable allows to create a L<sup>A</sup>T<sub>E</sub>Xfile for the user analysis, and to specify its name. If this variable is not set, the analysis will be directed to the screen output. (cf. also `write_analysis`)

- **and**

This is the standard two-place logical connective that has the value true if both of its operands are true, otherwise a value of false. It is applied to logical values, e.g. cut expressions. (cf. also `or`).

- **as**

cf. `compile`

- **ascii**

Specifier for the `sample_format` command to demand the generation of the standard WHIZARD verbose ASCII event files. (cf. also `$sample`, `sample_format`)

- **any**

`any` is a function that works on a logical expression and a list, `any <log_expr> [<list>]`, and returns `true` if `log_expr` is fulfilled for any entry in `list`, and `false` otherwise. Examples: `any PDG == 13 [lepton]` checks whether any lepton is a muon, `any E > 2 * mW [jet]` checks whether any jet has an energy of twice the  $W$  mass. Logical expressions with `any` can be logically combined with `and` and `or`. (cf. also `all`, `and`, `no`, and `or`)

- **athena**

Specifier for the `sample_format` command to demand the generation of the ATHENA variant for HEPEVT ASCII event files. (cf. also `$sample`, `sample_format`)

- **beams**

This specifies the contents and structure of the beams. If this command is absent in the input file, WHIZARD automatically takes the two incoming partons (or one for decays) of the corresponding process as beam particles and no structure functions are applied. Protons and antiprotons as beam particles are predefined as `p` and `pbar`, respectively. A structure function, like `lhapdf`, `ISR`, `EPA` and so on are switched on as e.g. `beams = p, p => lhapdf`. (cf. also `circe`, `circe2`, `lhapdf`).

- **cmplx**

Defines a complex variable. (to be finalized still)

- **combine**

The `combine [<list1>, <list2>]` operation makes a particle list whose entries are the result of adding (the momenta of) each pair of particles in the two input lists `list1`, `list2`. For example, `combine [incoming lepton, lepton]` constructs all mutual pairings of an incoming lepton with an outgoing lepton (an alias for the leptons has to be defined, of course).

- **compile**

The `compile` command is mandatory, it invokes the compilation of the process(es) (i.e. the

matrix element file(s)) to be compiled as a shared library. This shared object file has the standard name `processes.so` and resides in the `.libs` subdirectory of the corresponding user workspace. If the user has defined a different library name `lib_name` with the `library` command, then WHIZARD compiles this as the shared object `.libs/lib_name.so`. (This allows to split process classes and to avoid too large libraries.) Another possibility is to use the command `compile as "static_name"`. This will compile and link the process library in a static way and create the static executable `static_name` in the user workspace. (cf. also `library`, `load`)

- **`cuts`**

This command defines the cuts to be applied to certain processes. The syntax is: `cuts = <log_class> <log_expr> [<unary or binary particle (list) arg>]`, where the cut expression must be initialized with a logical classifier `log_class` like `all`, `any`, `no`. The logical expression `log_expr` contains the cut to be evaluated. Note that this need not only be a kinematical cut expression like `E > 10 GeV` or `5 degree < Theta < 175 degree`, but can also be some sort of trigger expression or event selection, e.g. `PDG == 15` would select a tau lepton. Whether the expression is evaluated on particles or pairs of particles depends on whether the discriminating variable is unary or binary, `Dist` being obviously binary, `Pt` being unary. Note that some variables are both unary and binary, e.g. the invariant mass `M`. Cut expressions can be connected by the logical connectives `and` and `or`. The `cuts` statement acts on all subsequent process integrations and analyses until a new `cuts` statement appears. (cf. also `all`, `any`, `Dist`, `E`, `M`, `no`, `Pt`).

- **`debug`**

Specifier for the `sample_format` command to demand the generation of the very verbose WHIZARD ASCII event file format intended for debugging. (cf. also `$sample`, `sample_format`)

- **`degree`**

Expression specifying the physical unit of degree for angular variables, e.g. the cut expression function `Theta`. (if no unit is specified for angular variables, radians are used).

- **`$description`**

String variable that allows to specify a description text for the analysis, `$description = "analysis description text"`. This line appears below the title of a corresponding analysis, on top of the respective plot. (cf. `analysis`, `$title`)

- **`echo`**

Allows to put verbose information on the screen during execution, e.g. `echo ("Hello, world!")`. (cf. also `show`)

- **`else`**

cf. `if`

- **`eV`**

Physical unit, stating that the corresponding number is in electron volt.

- **exec**

Constructor `exec ("<cmd_name>")` that demands WHIZARD to execute/run the command `cmd_name`. For this to work that specific command must be present either in the path of the operating system or as a command in the user workspace.

- **expect**

The binary function `expect` compares two numerical expressions whether they are fulfill a certain ordering condition or are equal up to a specific uncertainty or tolerance which can be set by the specifier `tolerance`, i.e. in principle it checks whether a logical expression is true. The `expect` function does actually not just check a value for correctness, but also records its result. If failures are present when the program terminates, the exit code is nonzero. The syntax is `expect (<num1> <log_comp> <num2>)`, where `num1` and `num2` are two numerical values (or corresponding variables) and `log_comp` is one of the following logical comparators: `<`, `>`, `<=`, `<>=`,  `$\approx$` ,  `$\sim\sim$` ,  `$\sim$` . (cf. also `<`, `>`, `<=`, `<>=`,  `$\approx$` ,  `$\sim\sim$` ,  `$\sim$` , `tolerance`).

- **\$extension\_ascii**

String variable that allows via `$extension_ascii = "<suffix>"` to specify the suffix for the file `name.suffix` to which events in the standard WHIZARD verbose ASCII format are written. If not set, the default file name and suffix is `<process_name>.evt`. (cf. also `sample_format`, `$sample`)

- **\$extension\_ascii\_long**

String variable that allows via `$extension_ascii_long = "<suffix>"` to specify the suffix for the file `name.suffix` to which events in the so called long variant of the WHIZARD 1 style HEPEVT ASCII format are written. If not set, the default file name and suffix is `<process_name>.long.evt`. (cf. also `sample_format`, `$sample`)

- **\$extension\_ascii\_short**

String variable that allows via `$extension_ascii_short = "<suffix>"` to specify the suffix for the file `name.suffix` to which events in the so called short variant of the WHIZARD 1 style HEPEVT ASCII format are written. If not set, the default file name and suffix is `<process_name>.short.evt`. (cf. also `sample_format`, `$sample`)

- **\$extension\_debug**

String variable that allows via `$extension_debug = "<suffix>"` to specify the suffix for the file `name.suffix` to which events in a long verbose format with debugging information are written. If not set, the default file name and suffix is `<process_name>.debug`. (cf. also `sample_format`, `$sample`)

- **\$extension\_hepevt**

String variable that allows via `$extension_hepevt = "<suffix>"` to specify the suffix for the file `name.suffix` to which events in the WHIZARD 1 style HEPEVT ASCII format are written. If not set, the default file name and suffix is `<process_name>.hepevt`. (cf. also `sample_format`, `$sample`)

- **\$extension\_hepmc**

String variable that allows via `$extension_hepmc = "<suffix>"` to specify the suffix for the file `name.suffix` to which events in the HepMC format are written. If not set, the default file name and suffix is `<process_name>.hepmc`. (cf. also `sample_format`, `$sample`)

- **\$extension\_lhef**

String variable that allows via `$extension_lhef = "<suffix>"` to specify the suffix for the file `name.suffix` to which events in the LHEF format are written. If not set, the default file name and suffix is `<process_name>.lhef`. (cf. also `sample_format`, `$sample`)

- **false**

Constructor stating that a logical expression or variable is false, e.g. `?<log_var> = false`. (cf. also `true`).

- **?fatal\_beam\_decay**

Logical variable that let the user decide whether the possibility of a beam decay is treated as a fatal error or only as a warning. An example is a process  $bt \rightarrow X$ , where the bottom quark as an initial state particle appears as a possible decay product of the second incoming particle, the top quark. This might trigger inconsistencies or instabilities in the phase space set-up.

- **GeV**

Physical unit, energies in  $10^9$  electron volt. This is the default energy unit of WHIZARD.

- **hepevt**

Specifier for the `sample_format` command to demand the generation of HEPEVT ASCII event files. (cf. also `$sample`, `sample_format`)

- **hepmc**

Specifier for the `sample_format` command to demand the generation of HepMC ASCII event files. Note that this is only available if the HepMC package is installed and correctly linked. (cf. also `$sample`, `sample_format`)

- **if**

Conditional clause with the construction `if <log_expr> then <expr> else <expr>`. Note that there must be an `end if` statement. For more complicated expressions it is better to use expressions in parentheses: `if (<log_expr>) then {<expr>} else {<expr>}`. Examples are a selection of up quarks over down quarks depending on a logical variable: `if ?ok then u else d`, or the setting of an integer variable depending on the rapidity of some particle: `if (eta > 0) then { a = +1} else { a = -1}`. The `then` constructor is not mandatory and can be omitted.

- **include**

The `include` statement, `include ("file.sin")` allows to include external SINDARIN

files `file.sin` into the main WHIZARD input file. A standard example is the inclusion of the standard cut file `default_cuts.sin`.

- **int**

This is a constructor to specify integer constants in the input file. Strictly speaking, it is a unary function setting the value `int_val` of the integer variable `int_var`: `int <int_var> = <int_val>`. (cf. also `real` and `cmplx`)

- **integrate**

The `integrate (<proc_name>) { <integrate_options> }` command invokes the integration (phase-space grid generation and Monte-Carlo sampling of the process `proc_name` (which can also be a list of processes) with the integration options `<integrate_options>`. Right now the only option is to specify the number of iterations and calls per integration during the Monte-Carlo phase-space integration via `iterations = <n_iterations>:<n_calls>`. Note that this can be list, separated by colons, which breaks up the integration process into units of the specified number of integrations and calls each.

- **iterations**

Option to set the number of iterations and calls per iteration during the Monte-Carlo phase-space integration process, cf. `integrate`.

- **keV**

Physical unit, energies in  $10^3$  electron volt.

- **\$label**

This is a string variable, `$label = "label_name"` that allows to specify a label `label_name` for analysis plots on the  $x$  axis. It is only taken into account if the variable `$xlabel` has not been set, in which case it is overwritten by the string value of that variable. (cf. also `xlabel`, `ylabel`).

- **lha**

Specifier for the `sample_format` command to demand the generation of the WHIZARD 1 LHA ASCII event format files. (cf. also `$sample`, `sample_format`)

- **lhapdf**

This is a specifier to demand calling LHAPDF parton densities to integrate processes in hadron collisions. (cf. `beams`)

- **lhef**

Specifier for the `sample_format` command to demand the generation of the Les Houches Accord (LHEF) event format files, with the XML headers. (cf. also `$sample`, `sample_format`)

- **library**

The command `library = "<lib_name>"` allows to specify a separate shared object library archive `lib_name.so`, not using the standard library `processes.so`. Those libraries (when using shared libraries) are located in the `.libs` subdirectory of the user workspace.

Specifying a separate library is useful for splitting up large lists of processes, or to restrict a larger number of different loaded model files to one specific process library. (cf. also `compile`, `load`)

- **load**

The `load` command allows to load again a library if some details have been changed (processes added, redefined or maybe changed. (cf. also `compile`, `library`)

- **long**

Specifier for the `sample_format` command to demand the generation of the long variant of HEPEVT ASCII event files. (cf. also `$sample`, `sample_format`)

- **luminosity** This specifier `luminosity = <num>` sets the integrated luminosity for the event generation of the processes in the SINDARIN input files. Note that WHIZARD itself chooses the number from the `luminosity` or from the `n_events` specifier, whichever would give the larger number of events. As this depends on the cross section under consideration, it might be different for different processes in the process list. Furthermore, the `luminosity` or `n_events` command has to be invoked *after* the corresponding logical variable which tells WHIZARD to write an event file in a specific format. (cf. `n_events`, `$sample`, `sample_format`)

- **MeV**

Physical unit, energies in  $10^6$  electron volt.

- **model**

With this specifier, `model = <MODEL_NAME>`, one sets the hard interaction physics model for the processes defined after this model specification. The list of available models can be found in Table 6.1. Note that the model specification can appear arbitrarily often in a SINDARIN input file, e.g. for compiling and running processes defined in different physics models.

- **no**

`no` is a function that works on a logical expression and a list, `no <log_expr> [<list>]`, and returns `true` if and only if `log_expr` is fulfilled for *none* of the entries in `list`, and `false` otherwise. Examples: `no Pt < 100 GeV [lepton]` checks whether no lepton is softer than 100 GeV. It is the logical opposite of the function `all`. Logical expressions with `no` can be logically combined with `and` and `or`. (cf. also `all`, `any`, `and`, and `or`)

- **n\_events**

This specifier `n_events = <num>` sets the number of events for the event generation of the processes in the SINDARIN input files. Note that WHIZARD itself chooses the number from the `n_events` or from the `luminosity` specifier, whichever would give the larger number of events. As this depends on the cross section under consideration, it might be different for different processes in the process list. Furthermore, the `n_events` or `luminosity` command has to be invoked *after* the corresponding logical variable which

tells WHIZARD to write an event file in a specific format. (cf. `luminosity`, `$sample`, `sample_format`)

- **observable**

With this, `observable = <obs_spec>`, the user is able to define a variable specifier `obs_spec` for observables. These can be reused in the analysis, e.g. as a `record`, as functions of the fundamental kinematical variables of the processes. (cf. `analysis`, `record`)

- **or**

This is the standard two-place logical connective that has the value true if one of its operands is true, otherwise a value of false. It is applied to logical values, e.g. cut expressions. (cf. also `and`).

- **\$physical\_unit**

This is a string variable, `$physical_unit = "<unit_name>"`, that allows to set a L<sup>A</sup>T<sub>E</sub>Xname `unit_name` for the physical unit of a label of an analysis plot. This unit is then also used for calculations within the analysis set-up.

- **plot**

(cf. `record`)

- **process**

Allows to set a hard interaction process, either for a decay process `decay_proc` as `process <decay_proc> = <mother> => <daughter1>, <daughter2>, ...`, or for a scattering process `scat_proc` as `<incoming1>, <incoming2> => <outgoing1>, <outgoing2>, ...`. Note that there can be arbitrarily many processes to be defined in a SINDARIN input file. (cf. also `restrictions`)

- **read\_slha**

Tells WHIZARD to read in an input file in the SUSY Les Houches accord (SLHA), as `read_slha ("slha_file.slha")`. Note that the files for the use in WHIZARD should have the suffix `.slha`. (cf. also `write_slha`)

- **real**

This is a constructor to specify real constants in the input file. Strictly speaking, it is a unary function setting the value `real_val` of the integer variable `real_var`: `real <real_var> = <real_val>`. (cf. also `int` and `cplx`)

- **?rebuild**

The logical variable `?rebuild = true/false` specifies whether the matrix element code for processes is re-generated by the matrix element generator O'Mega (e.g. if the process has been changed, but not its name). This can also be set as a command-line option `whizard --rebuild`. The default is `false`, i.e. code is never re-generated if it is present and the MD5 checksum is valid. (cf. also `recompile`).

- **?recompile**

The logical variable `?recompile = true/false` specifies whether the matrix element code for processes is re-compiled (e.g. if the process code has been manually modified by the user). This can also be set as a command-line option `whizard --recompile`. The default is `false`, i.e. code is never re-compiled if its corresponding object file is present. (cf. also `rebuild`)

- **record**

The `record` constructor provides an internal data structure in SINDARIN input files. Its syntax is in general `record <record_name> (<cmd_expr>)`. The `<cmd_expr>` could be the definition of a tuple of points for a histogram or an `eval` constructor that tells WHIZARD e.g. by which rule to calculate an observable to be stored in the record `record_name`. (cf. also `eval`)

- **\$restrictions**

This is an optional argument for process definitions. It defines a string variable, `process <process_name> = <particle1>, <particle2> => <particle3>, <particle4>, ... { $restrictions = "<restriction_def>" }`. The string argument `restriction_def` is directly transferred during the code generation to the matrix element generator O'Mega. It has to be of the form `n1 + n2 + ... ~ <particle (list)>`, where `n1` and so on are the numbers of the particles above in the process definition. The tilde specifies a certain intermediate state to be equal to the particle(s) in `particle (list)`. An example is `process eemm_z = e1, E1 => e2, E2 { $restrictions = "1+2 ~ Z" }` restricts the code to be generated for the process  $e^-e^+ \rightarrow \mu^-\mu^+$  to the *s*-channel  $Z$ -boson exchange. (cf. also `process`)

- **results**

Only used in the combination `show(results)`. Forces WHIZARD to print out a results summary for the integrated processes. (cf. also `show`)

- **\$sample**

String variable to set the (base) name of the event output format, e.g. `$sample = "foo"` will result in an intrinsic binary format event file `foo.evx`. (cf. also `sample_format`, `simulate`, `hepevt`, `ascii`, `athena`, `debug`, `long`, `short`, `hepmc`, `lhef`, `lha`, `stdhep`, `stdhep_up`)

- **sample\_format**

Variable that allows the user to specify additional event formats beyond the WHIZARD native binary event format. Its syntax is `sample_format = <format>`, where `<format>` can be any of the following specifiers: `hepevt`, `ascii`, `athena`, `debug`, `long`, `short`, `hepmc`, `lhef`, `lha`, `stdhep`, `stdhep_up`. (cf. also `$sample`, `simulate`, `hepevt`, `ascii`, `athena`, `debug`, `long`, `short`, `hepmc`, `lhef`, `lha`, `stdhep`, `stdhep_up`)

- **scan**

Constructor to perform loops over variables or scan over processes in the integration procedure. The syntax is `scan <var> <var_name> (<value list> or <value_init> =>`

`<value_fin> /<incrementor> <increment>) { <scan_cmd> }`. The variable `var` can be specified if it is not a real, e.g. an integer. `var_name` is the name of the variable which is also allowed to be a predefined one like `seed`. For the scan, one can either specify an explicit list of values `value list`, or use an initial and final value and a rule to increment. The `scan_cmd` can either be just a `show` to print out the scanned variable or the integration of a process. Examples are: `scan seed (32 => 1 / / 2) { show (seed_value) }`, which runs the seed down in steps 32, 16, 8, 4, 2, 1 (division by two). `scan mW (75 GeV, 80 GeV => 82 GeV /+ 0.5 GeV, 83 GeV => 90 GeV /* 1.2) { show (sw) }` scans over the  $W$  mass for the values 75, 80, 80.5, 81, 81.5, 82, 83 GeV, namely one discrete value, steps by adding 0.5 GeV, and increase by 20 % (the latter having no effect as it already exceeds the final value). It prints out the corresponding value of the effective mixing angle which is defined as a dependent variable in the model input file(s). `scan sqrts (500 GeV => 600 GeV /+ 10 GeV) { integrate (proc) }` integrates the process `proc` in eleven increasing 10 GeV steps in center-of-mass energy from 500 to 600 GeV.

- **seed**

Integer variable `seed = <num>` that allows to set a specific random seed `num`. If not set, WHIZARD takes the time from the system clock to determine the random seed.

- **short**

Specifier for the `sample_format` command to demand the generation of the short variant of HEPEVT ASCII event files. (cf. also `$sample`, `sample_format`)

- **show**

This is a unary function that is operating on specific constructors in order to print them out in the WHIZARD screen output as well as the log file `whizard.log`. Examples are `show(<parameter_name>)` to issue a specific parameter from a model or a constant defined in a SINDARIN input file, `show(integral(<proc_name>))`, `show(library)`, `show(results)`, or `show(jvari)` for any arbitrary variable. (cf. also `echo`, `library`, `results`)

- **simulate**

This command invokes the generation of events for the process `proc` by means of `simulate (<proc>)`. It accepts both `$sample` and `sample_format` as optional arguments. (cf. also `integrate`, `luminosity`, `n_events`, `$sample`, `sample_format`)

- **sqrts**

Real variable in order to set the center-of-mass energy for the collisions (collider energy  $\sqrt{s}$ , not hard interaction energy  $\sqrt{s_{\text{NN}}}$ ): `sqrts = <num> <phys_unit>`. The physical unit can be one of the following `eV`, `keV`, `MeV`, `GeV`, and `TeV`. If absent, WHIZARD takes `GeV` as its standard unit.

- **stable**

This constructor allows particles in the final states of processes in decay cascade set-up

to be set as stable, and not letting them decay. The syntax is `stable <particle_name>`. (cf. also `unstable`)

- **`stdhep`**

Specifier for the `sample_format` command to demand the generation of binary StdHEP event files based on the HEPEVT common block. Note that this is only available if the StdHEP package is installed and correctly linked. (cf. also `$sample`, `sample_format`)

- **`stdhep_up`**

Specifier for the `sample_format` command to demand the generation of binary StdHEP event files based on the HEPRUP/HEPEUP common blocks. Note that this is only available if the StdHEP package is installed and correctly linked. (cf. also `$sample`, `sample_format`)

- **`TeV`**

Physical unit, for energies in  $10^{12}$  electron volt.

- **`then`**

Alternative option inside a conditional clause, not mandatory, hence maybe be omitted, cf. `if`.

- **`$title`**

This string variable sets the title of a plot in a WHIZARD analysis setup, e.g. a histogram or an observable. The syntax is `$title = "<your title>"`. This title appears as a section header in the analysis file, but not in the screen output of the analysis. (cf. also `$description`, `$label`, `$xlabel`, `$ylabel`).

- **`tolerance`**

Real variable that defines the tolerance with which the (logical) function `expect` accepts equality or inequality: `tolerance = <num>`. This can e.g. be used for cross-section tests and backwards compatibility checks. (cf. also `expect`)

- **`true`**

Constructor stating that a logical expression or variable is true, e.g. `?<log_var> = true`. (cf. also `false`).

- **`unstable`**

This constructor allows to let final state particles of the hard interaction undergo a subsequent (cascade) decay (in the on-shell approximation). For this the user has to define the list of desired Decay channels as `unstable <mother> (<decay1>, <decay2>, ...)`, where `mother` is the mother particle, and the argument is a list of decay channels. Note that these have to be provided by the user as in the example in Fig. A.1. First, the  $Z$  decays to electrons and up quarks are generated, then  $ZZ$  production at a 500 GeV ILC is called, and then both  $Z$ s are decayed according to the probability distribution of the two generated decay matrix elements. This obviously allows also for inclusive decays. (cf. also `stable`)

```

process zee = Z => e1, E1
process zuu = Z => u, U
process zz = e1, E1 => Z, Z
compile
integrate (zee) { iterations = 1:100 }
integrate (zuu) { iterations = 1:100 }
sqrtS = 500 GeV
integrate (zz) { iterations = 3:5000, 2:5000 }
unstable Z (zee, zuu)

```

Figure A.1: *SINDARIN input file for unstable particles and inclusive decays.*

- **write\_analysis**

The `write_analysis` statement tells WHIZARD to write the analysis setup by the user for the SINDARIN input file under consideration. If no `$analysis_filename` is provided, the analysis (including the histograms) are printed out on the screen, otherwise they are written to a file defined by that specific string variable. (cf. also `$analysis_filename`)

- **write\_slha**

Demands WHIZARD to write out a file in the SUSY Les Houches accord (SLHA). (cf. also `read_slha`)

- **\$xlabel**

String variable, `$xlabel = "<LaTeX code>"`, that sets the *x* axis label in a plot or histogram in a WHIZARD analysis. (cf. also `label` and `$ylabel`)

- **\$ylabel**

String variable, `$ylabel = "<LaTeX code>"`, that sets the *y* axis label in a plot or histogram in a WHIZARD analysis. (cf. also `label` and `$xlabel`)

# Appendix B

## Example files

### B.1 Extended internal event formats

WHIZARD DEFAULT, .evt:

```
=====
Event record:
=====
=====
|=====
Decay tree:
  tries =           1
  acceptance probability = 1.0000000000000000
|=====
Mother process = 'qedtest_eemm'
|=====
Decay chain:
|=====
Decay configuration: [undefined]
|=====
Evaluator: Color-summed including all decays
Evaluator:
Interaction:     8
Virtual:
Particle 1
E = 250.0000000000000
P = 0.0000000000000000      0.0000000000000000      249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)1
Particle 2
E = 250.0000000000000
P = 0.0000000000000000      0.0000000000000000      -249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)2
Outgoing:
Particle 3
E = 250.0000000000000
P = 223.64041528426193      -102.79251826657215      43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (5)3
Particle 4
E = 250.0000000000000
```

```

P = -223.64041528426193          102.79251826657215          -43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (5)4
State matrix: norm = 8.67160247171198766E-003
[f(11)]
 [f(-11)]
 [f(13)]
 [f(-13)] => ME(1) = 1.000000000000
Matrix-element multiplication
Input interaction 1: 2
Input interaction 2:      5
ME(1) =
+ ME1(1) x ME2(1)
|=====
Evaluator: Color flow components including all decays
Evaluator:
Interaction:         9
Virtual:
Particle 1
E = 250.0000000000000000
P = 0.0000000000000000      0.0000000000000000      249.99999999947775
mask [fch] = [FFgT]
internal links: X => 3 4
source: (7)1
Particle 2
E = 250.0000000000000000
P = 0.0000000000000000      0.0000000000000000      -249.99999999947775
mask [fch] = [FFgT]
internal links: X => 3 4
source: (7)2
Outgoing:
Particle 3
E = 250.0000000000000000
P = 223.64041528426193      -102.79251826657215      43.802416227981048
mask [fch] = [FFgT]
internal links: 1 2 => X
source: (6)3
Particle 4
E = 250.0000000000000000
P = -223.64041528426193     102.79251826657215      -43.802416227981048
mask [fch] = [FFgT]
internal links: 1 2 => X
source: (6)4
State matrix: norm = 1.0000000000000000
[f(11)]
 [f(-11)]
 [f(13)]
 [f(-13)] => ME(1) = 8.671602471712E-03
Matrix-element multiplication
Input interaction 1: 7
Input interaction 2:      6
ME(1) =
+ ME1(1) x ME2(1)
|=====
[Process: qedtest_eemm]

Particle set:
Particle 1 [i] f(11)
Momentum:
E = 250.0000000000000000
P = 0.0000000000000000      0.0000000000000000      249.99999999947775
T = 2.61124455391836818E-007
Children: 3 4
Particle 2 [i] f(-11)

```

```

Momentum:
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 -249.9999999947775
T = 2.61124455391836818E-007
Children: 3 4
Particle 3 [o] f(13)
Momentum:
E = 250.00000000000000
P = 223.64041528426193 -102.79251826657215 43.802416227981048
T = 1.11724899955660817E-002
Parents: 1 2
Particle 4 [o] f(-13)
Momentum:
E = 250.00000000000000
P = -223.64041528426193 102.79251826657215 -43.802416227981048
T = 1.11724899955660817E-002
Parents: 1 2

Event weight = 1.0000000000000000
Excess weight = 0.0000000000000000

```

## DEBUG FORMAT:

```

=====
Event record:
=====
Process data: 1 (qedtest_eemm)
[scattering]
use separate beam setup = T
Beam data (collision):
e- (mass = 5.1100000E-04 GeV)
Polarization: [spin_type = 2, mult = 2]
State matrix: norm = 1.0000000000000000
[] => ME(1) = 0.5000000000000000
e+ (mass = 5.1100000E-04 GeV)
Polarization: [spin_type = 2, mult = 2]
State matrix: norm = 1.0000000000000000
[] => ME(1) = 0.5000000000000000
sqrt(s) = 500.0000000000000 GeV
C.m. momenta:
E = 250.00000000000000 0.0000000000000000 249.9999999947775
P = 0.0000000000000000 0.0000000000000000 -249.9999999947775
E = 250.00000000000000 0.0000000000000000 -249.9999999947775
P = 0.0000000000000000 0.0000000000000000
Lab momenta:
E = 250.00000000000000 0.0000000000000000 249.9999999947775
P = 0.0000000000000000 0.0000000000000000 -249.9999999947775
E = 250.00000000000000 0.0000000000000000 -249.9999999947775
P = 0.0000000000000000 0.0000000000000000
MD5 sum: 62BCF22F9E526C4368F761805CB3F4FA
number of structure functions = 0
number of strfun parameters = 0
number of process parameters = 2
number of parameters total = 2
number of integration channels = 1
number of bins per channel = 20
c.m. energy (sqrt(s)) = 500.0000000000000
c.m. energy (sqrt(s_hat)) = 500.0000000000000
Colliding partons = f(11) f(-11)
Incoming parton masses = 5.109999999999953E-004 5.109999999999953E-004
In-state flux factor = 1213731799.9298604
Strfun mapping factor = 1.0000000000000000
VAMP phs factor = 0.99999991062007598
Phase-space volume = 2.55294034613545786E-005

```

```

Squared matrix element = 8.67160247171198766E-003
Reweighting factor = 1.0000000000000000
Sample-function value = 268.69693914993798
-----
Structure function parameters =
Process energy scale = 500.00000000000000
-----
QCD coupling parameters =
alpha-s is fixed = T
-----
Phase-space integration parameters (input) =
0.93143035192042589 0.58760484028607607
Integration channel = 1
Phase-space integration parameters (complete) =
0.93143035192042589 0.58760484028607607
Channels: phase-space factors =
0.99999991062007598
-----
Structure function chain:
Colliding beams:
Interaction: 2
Outgoing:
Particle 1
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 249.99999999947775
mask [fch] = [FFgT]
Particle 2
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 -249.99999999947775
mask [fch] = [FFgT]
State matrix: norm = 1.0000000000000000
[f(11)]
[f(-11)] => ME(1) = 0.250000000000
-----
Evaluators:
Total number of parameters = 0
Last structure function (index) = 0 0
Outgoing particles (index) = 1 2
Colliding particles (index) = 1 2
-----
Incoming beams with all color contractions
Evaluator:
Interaction: 7
Outgoing:
Particle 1
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 249.99999999947775
mask [fch] = [FFgT]
source: (2)1
Particle 2
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 -249.99999999947775
mask [fch] = [FFgT]
source: (2)2
State matrix: norm = 1.0000000000000000
[f(11)]
[f(-11)] => ME(1) = 0.250000000000
Matrix-element multiplication
Input interaction 1: 2
Input interaction 2: [undefined]
ME(1) =
+ ME1(1)

```

```
-----
Hard interaction:
Process 'qedtest_eemm'
n_tot =          4
n_in  =          2
n_out =          2
n_flv =          1
n_hel =         16
n_col =          1
n_cin =          2
n_cff =          1
Model parameters:
 1 0.30286000000000002
 2 5.109999999999953E-004
 3 0.10570000000000000
 4 1.776999999999999
Flavor states:
 1 :      11     -11      13     -13
Helicity states:
 1 :      -1      -1      -1      -1
 2 :      -1      -1      -1       1
 3 :      -1      -1       1      -1
 4 :      -1      -1       1       1
 5 :      -1       1      -1      -1
 6 :      -1       1      -1       1
 7 :      -1       1       1      -1
 8 :      -1       1       1       1
 9 :       1      -1      -1      -1
10 :       1      -1      -1       1
11 :       1      -1       1      -1
12 :       1      -1       1       1
13 :       1       1      -1      -1
14 :       1       1      -1       1
15 :       1       1       1      -1
16 :       1       1       1       1
Color states:
 1://///
Color factors:
 1: 1   1  ( 1.0000000000000000 , 0.0000000000000000 )
Interaction:           1
Incoming:
Particle 1
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 249.9999999947775
mask [fch] = [FFgF]
internal links: X => 3 4
source: (2)1
Particle 2
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 -249.9999999947775
mask [fch] = [FFgF]
internal links: X => 3 4
source: (2)2
Outgoing:
Particle 3
E = 250.00000000000000
P = 223.64041528426193 -102.79251826657215 43.802416227981048
mask [fch] = [FFgF]
internal links: 1 2 => X
Particle 4
E = 250.00000000000000
P = -223.64041528426193 102.79251826657215 -43.802416227981048
mask [fch] = [FFgF]
internal links: 1 2 => X
```

```

State matrix: norm = 1.0000000000000000
[f(11) h(-1)]
[f(-11) h(-1)]
[f(13) h(-1)]
[f(-13) h(-1)] => ME(1) = 0.00000000000
[f(-13) h(1)] => ME(2) = 1.845852774013E-07, -2.960943317340E-24
[f(13) h(1)]
[f(-13) h(-1)] => ME(3) = 1.845852774013E-07, 7.311291744600E-25
[f(-13) h(1)] => ME(4) = 0.00000000000
[f(-11) h(1)]
[f(13) h(-1)]
[f(-13) h(-1)] => ME(5) = -3.818108556076E-05
[f(-13) h(1)] => ME(6) = 9.794450119376E-02, 4.501852634852E-02
[f(13) h(1)]
[f(-13) h(-1)] => ME(7) = -6.873979831962E-02, -3.159508072559E-02
[f(-13) h(1)] => ME(8) = 2.486236753832E-05, 2.897685239959E-05
[f(11) h(1)]
[f(-11) h(-1)]
[f(13) h(-1)]
[f(-13) h(-1)] => ME(9) = -2.486236753832E-05, 2.897685239959E-05
[f(-13) h(1)] => ME(10) = -6.873979831962E-02, 3.159508072559E-02
[f(13) h(1)]
[f(-13) h(-1)] => ME(11) = 9.794450119376E-02, -4.501852634852E-02
[f(-13) h(1)] => ME(12) = 3.818108556076E-05
[f(-11) h(1)]
[f(13) h(-1)]
[f(-13) h(-1)] => ME(13) = 0.00000000000
[f(-13) h(1)] => ME(14) = -1.845852774013E-07, 2.960943317340E-24
[f(13) h(1)]
[f(-13) h(-1)] => ME(15) = -1.845852774013E-07, -7.311291744600E-25
[f(-13) h(1)] => ME(16) = 0.00000000000
-----  

Trace including color factors (hard interaction)
Evaluator:
Interaction: 3
Incoming:
Particle 1
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)1
Particle 2
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 -249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)2
Outgoing:
Particle 3
E = 250.00000000000000
P = 223.64041528426193 -102.79251826657215 43.802416227981048
mask [fch] = [TTT]
internal links: 1 2 => X
source: (1)3
Particle 4
E = 250.00000000000000
P = -223.64041528426193 102.79251826657215 -43.802416227981048
mask [fch] = [TTT]
internal links: 1 2 => X
source: (1)4
State matrix: norm = 1.0000000000000000
[f(11)]
[f(-11)]
[]

```

```

[] => ME(1) = 3.468640988685E-02
Matrix-element multiplication
Input interaction 1: 1
Input interaction 2: [undefined]
ME(1) =
+ ME1(1)* x ME2(1) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(2)* x ME2(2) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(3)* x ME2(3) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(4)* x ME2(4) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(5)* x ME2(5) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(6)* x ME2(6) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(7)* x ME2(7) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(8)* x ME2(8) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(9)* x ME2(9) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(10)* x ME2(10) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(11)* x ME2(11) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(12)* x ME2(12) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(13)* x ME2(13) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(14)* x ME2(14) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(15)* x ME2(15) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(16)* x ME2(16) x ( 1.0000000000000000 , 0.0000000000000000 )

-----  

Exclusive sqme including color factors (hard interaction)
Evaluator:
Interaction:      5
Incoming:
Particle 1
E =    250.00000000000000
P =    0.0000000000000000      0.0000000000000000      249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)1
Particle 2
E =    250.00000000000000
P =    0.0000000000000000      0.0000000000000000      -249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)2
Outgoing:
Particle 3
E =    250.00000000000000
P =    223.64041528426193      -102.79251826657215      43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (1)3
Particle 4
E =    250.00000000000000
P =   -223.64041528426193      102.79251826657215      -43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (1)4
State matrix: norm = 1.0000000000000000
[f(11)]
[f(-11)]
[f(13)]
[f(-13)] => ME(1) = 3.468640988685E-02
Matrix-element multiplication
Input interaction 1: 1
Input interaction 2: [undefined]
ME(1) =
+ ME1(1)* x ME2(1) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(2)* x ME2(2) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(3)* x ME2(3) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(4)* x ME2(4) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(5)* x ME2(5) x ( 1.0000000000000000 , 0.0000000000000000 )

```

```

+ ME1(6)* x ME2(6) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(7)* x ME2(7) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(8)* x ME2(8) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(9)* x ME2(9) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(10)* x ME2(10) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(11)* x ME2(11) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(12)* x ME2(12) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(13)* x ME2(13) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(14)* x ME2(14) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(15)* x ME2(15) x ( 1.0000000000000000 , 0.0000000000000000 )
+ ME1(16)* x ME2(16) x ( 1.0000000000000000 , 0.0000000000000000 )

-----  

Color flow coefficients (hard interaction)
Evaluator:  

Interaction: 6  

Incoming:  

Particle 1  

E = 250.00000000000000  

P = 0.0000000000000000 0.0000000000000000 249.9999999947775  

mask [fch] = [FFgT]  

internal links: X => 3 4  

source: (7)1  

Particle 2  

E = 250.00000000000000  

P = 0.0000000000000000 0.0000000000000000 -249.9999999947775  

mask [fch] = [FFgT]  

internal links: X => 3 4  

source: (7)2  

Outgoing:  

Particle 3  

E = 250.00000000000000  

P = 223.64041528426193 -102.79251826657215 43.802416227981048  

mask [fch] = [FFgT]  

internal links: 1 2 => X  

source: (1)3  

Particle 4  

E = 250.00000000000000  

P = -223.64041528426193 102.79251826657215 -43.802416227981048  

mask [fch] = [FFgT]  

internal links: 1 2 => X  

source: (1)4  

State matrix: norm = 1.0000000000000000  

[f(11)]  

[f(-11)]  

[f(13)]  

[f(-13)] => ME(1) = 3.468640988685E-02  

Matrix-element multiplication  

Input interaction 1: 1  

Input interaction 2: [undefined]  

ME(1) =
+ |ME1(1)|^2
+ |ME1(2)|^2
+ |ME1(3)|^2
+ |ME1(4)|^2
+ |ME1(5)|^2
+ |ME1(6)|^2
+ |ME1(7)|^2
+ |ME1(8)|^2
+ |ME1(9)|^2
+ |ME1(10)|^2
+ |ME1(11)|^2
+ |ME1(12)|^2
+ |ME1(13)|^2
+ |ME1(14)|^2
+ |ME1(15)|^2

```

```

+ |ME1(16)|^2
-----
Trace including color factors (beams + strfun + hard interaction)
Evaluator:
Interaction:          4
Virtual:
Particle 1
E =      250.00000000000000
P =      0.0000000000000000      0.0000000000000000      249.9999999947775
mask [fch] = [TTT]
internal links: X => 3 4
source: (2)1
Particle 2
E =      250.00000000000000
P =      0.0000000000000000      0.0000000000000000      -249.9999999947775
mask [fch] = [TTT]
internal links: X => 3 4
source: (2)2
Outgoing:
Particle 3
E =      250.00000000000000
P =      223.64041528426193      -102.79251826657215      43.802416227981048
mask [fch] = [TTT]
internal links: 1 2 => X
source: (3)3
Particle 4
E =      250.00000000000000
P =      -223.64041528426193     102.79251826657215      -43.802416227981048
mask [fch] = [TTT]
internal links: 1 2 => X
source: (3)4
State matrix: norm = 1.0000000000000000
[]
[]
[]
[]
[] => ME(1) = 8.671602471712E-03
Matrix-element multiplication
Input interaction 1: 2
Input interaction 2:          3
ME(1) =
+ ME1(1) x ME2(1)
-----
Exclusive sqme including color factors (beams + strfun + hard interaction)
Evaluator:
Interaction:          8
Virtual:
Particle 1
E =      250.00000000000000
P =      0.0000000000000000      0.0000000000000000      249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)1
Particle 2
E =      250.00000000000000
P =      0.0000000000000000      0.0000000000000000      -249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)2
Outgoing:
Particle 3
E =      250.00000000000000
P =      223.64041528426193      -102.79251826657215      43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (5)3

```

```

Particle 4
E = 250.0000000000000
P = -223.64041528426193      102.79251826657215      -43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (5)4
State matrix: norm = 8.67160247171198766E-003
[f(11)]
[f(-11)]
[f(13)]
[f(-13)] => ME(1) = 1.00000000000
Matrix-element multiplication
Input interaction 1: 2
Input interaction 2:           5
ME(1) =
+ ME1(1) x ME2(1)
-----
Color flow coefficients (beams + strfun + hard interaction)
Evaluator:
Interaction:         9
Virtual:
Particle 1
E = 250.0000000000000
P = 0.000000000000000      0.000000000000000      249.9999999947775
mask [fch] = [FFgT]
internal links: X => 3 4
source: (7)1
Particle 2
E = 250.0000000000000
P = 0.000000000000000      0.000000000000000      -249.9999999947775
mask [fch] = [FFgT]
internal links: X => 3 4
source: (7)2
Outgoing:
Particle 3
E = 250.0000000000000
P = 223.64041528426193     -102.79251826657215      43.802416227981048
mask [fch] = [FFgT]
internal links: 1 2 => X
source: (6)3
Particle 4
E = 250.0000000000000
P = -223.64041528426193    102.79251826657215      -43.802416227981048
mask [fch] = [FFgT]
internal links: 1 2 => X
source: (6)4
State matrix: norm = 1.000000000000000
[f(11)]
[f(-11)]
[f(13)]
[f(-13)] => ME(1) = 8.671602471712E-03
Matrix-element multiplication
Input interaction 1: 7
Input interaction 2:           6
ME(1) =
+ ME1(1) x ME2(1)
-----
Phase space forest:
n_in =          2
n_out =         2
n_tot =         4
n_masses =       0
n_angles =       2
n_dim =          2
n_trees =        1

```

```

n_equiv =           1
flavors =      11   -11    13   -13
Groves and trees:
Grove     1
Tree      1
External: 4      Mask: 15
Incoming: 2      Mask: 12
Branches: 5
          8
          4
* 3       Daughters:  1   2   + (axis +)
  2
  1
    1 mass_sum = 0.1057000000000000
    2 mass_sum = 0.1057000000000000
    3 mass_sum = 0.2114000000000000
Equivalence list:
Equivalence: 1   1   Final state permutation: 1 2
               msq permutation:
               angle permutation: 1 2
                           ++
Total number of equivalences:           1

Incoming particles:
Particle 1
E = 250.0000000000000
P = 0.000000000000000      0.000000000000000      249.9999999947775
M2 = 2.6112099999999973E-007
Particle 2
E = 250.0000000000000
P = 0.000000000000000      0.000000000000000      -249.9999999947775
M2 = 2.6112099999999973E-007

Outgoing particles:
Particle 1
E = 250.0000000000000
P = 223.64041528426193      -102.79251826657215      43.802416227981048
M2 = 1.1172490000000002E-002
Particle 2
E = 250.0000000000000
P = -223.64041528426193      102.79251826657215      -43.802416227981048
M2 = 1.1172490000000002E-002

Tree particles:
Particle 1
E = 250.0000000000000
P = 223.64041528426193      -102.79251826657215      43.802416227981048
M2 = 1.1172490000000002E-002
Particle 2
E = 250.0000000000000
P = -223.64041528426193      102.79251826657215      -43.802416227981048
M2 = 1.1172490000000002E-002
Particle 3
E = 500.0000000000000
P = 0.000000000000000      0.000000000000000      0.000000000000000
M2 = 250000.00000000000
Particle 4
E = 250.0000000000000
P = 0.000000000000000      0.000000000000000      -249.9999999947775
M2 = 2.6112099999999973E-007
Particle 8
E = 250.0000000000000
P = 0.000000000000000      0.000000000000000      249.9999999947775
M2 = 2.6112099999999973E-007
-----
```

```

Inequivalent channels:
  Channel      1 :   Mult. =           1   Symm. =           1   Invar.: T F
Equivalence list:
Equivalent channels:  1   1
  Permutation: 1   2
  Mode:        3   0
-----
Particle list:
1 prt(i:11|-2.50000E+02;-0.00000E+00,-0.00000E+00,-2.50000E+02| 2.61121E-07| 1)
2 prt(i:-11|-2.50000E+02;-0.00000E+00,-0.00000E+00, 2.50000E+02| 2.61121E-07| 2)
3 prt(o:13| 2.50000E+02; 2.23640E+02,-1.02793E+02, 4.38024E+01| 1.11725E-02| 3)
4 prt(o:-13| 2.50000E+02;-2.23640E+02, 1.02793E+02,-4.38024E+01| 1.11725E-02| 4)
-----
n_in =          2
n_out =         2
n_tot =         4
sqrt_s =>    500.0000000000000000
sqrt_s_hat => 500.0000000000000000
v = [unknown real]
cw = [unknown real]
sw = [unknown real]
ee = 0.30286000000000002
alpha_em_i = [unknown real]
$fc = "/opt/gcc-4.5/bin/gfortran"
$fcflags = "-g -O2"
seed_value = 1582518999
sqrt_s = 500.0000000000000000
$model_name = "QED"
$restrictions = ""
$method = "omega"
?read_color_factors = true
?ssha_read_input = true
?ssha_read_spectrum = true
?ssha_read_decays = false
$library_name = "qedtest"
cm_momentum = 0.0000000000000000
cm_theta = 0.0000000000000000
cm_phi = 0.0000000000000000
luminosity = 0.0000000000000000
$lhapdf_file = [unknown string]
lhapdf_member = 0
lhapdf_photon_scheme = 0
isr_alpha = 0.0000000000000000
isr_q_max = 0.0000000000000000
isr_mass = 0.0000000000000000
isr_order = 3
epa_alpha = 0.0000000000000000
epa_x_min = 0.0000000000000000
epa_q_min = 0.0000000000000000
epa_e_max = 0.0000000000000000
epa_mass = 0.0000000000000000
ewa_x_min = 0.0000000000000000
ewa_q_min = 0.0000000000000000
ewa_pt_max = 0.0000000000000000
ewa_mass = 0.0000000000000000
[user variable] ?ewa_keep_momentum = false
[user variable] ?ewa_keep_energy = false
?alpha_s_is_fixed = true
?alpha_s_from_lhapdf = false
alpha_s_order = 0
alpha_s_nf = 5
?alpha_s_from_mz = true
lambda_qcd = 0.20000000000000001
?fatal_beam_decay = true
?helicity_selection_active = true

```

```

helicity_selection_threshold =      100000000000.000000
helicity_selection_cutoff =         1000
threshold_calls =                 0
min_calls_per_channel =           10
min_calls_per_bin =               10
min_bins =                        3
max_bins =                        20
?stratified = true
?use_vamp_equivalences = true
channel_weights_power =          0.25000000000000000000000000000000
$phs_file = [unknown string]
?phs_only = false
phs_threshold_s =                50.000000000000000000000000000000
phs_threshold_t =                100.000000000000000000000000000000
phs_off_shell =                  1
phs_t_channel =                  2
phs_e_scale =                     10.000000000000000000000000000000
phs_m_scale =                     10.000000000000000000000000000000
phs_q_scale =                     10.000000000000000000000000000000
?adapt_final_grids = true
?adapt_final_weights = false
?isotropic_decay = false
?diagonal_decay = false
n_events =                         0
?unweighted = true
$event_normalization = "auto"
?negative_weights = false
?keep_beams = false
$sample = ""
?read_raw = true
?write_raw = true
$extension_raw = ".evx"
$extension_default = ".evt"
$extension_debug = ".debug"
$extension_hepevt = ".hepevt"
$extension_ascii_short = ".short.evt"
$extension_ascii_long = ".long.evt"
$extension_athena = ".athena.evt"
$extension_lhef = ".lhef"
$extension_lha = ".lha"
$extension_hepmc = ".hepmc"
$extension_stdhep = ".stdhep"
$extension_stdhep_up = ".up.stdhep"
$analysis_filename = [unknown string]
$label = ""
$physical_unit = ""
$title = ""
$description = ""
$xlabel = ""
$ylabel = ""
tolerance =            0.0000000000000000
$fc = "/opt/gcc-4.5/bin/gfortran"
$fcflags = "-g -O2"
?rebuild_library = true
?rebuild_phase_space = true
?rebuild_grids = true
?rebuild_events = true
?recompile_library = false
GF =    1.1663900000000003E-005
mZ =    91.188199999999995
mW =    80.418999999999997
mH =    200.000000000000000
alphas = 0.11780000000000000
me =    5.1099999999999953E-004
mmu =   0.10570000000000000

```

```

mtau =     1.7769999999999999
ms =      9.5000000000000011E-002
mc =      1.2000000000000000
mb =      4.2000000000000002
mtop =    173.09999999999999
wtop =    1.5229999999999999
wZ =      2.4430000000000001
wW =      2.0489999999999999
wH =      1.4190000000000000
khgaz =   0.0000000000000000
khgaga =  0.0000000000000000
khgg =   0.0000000000000000
xi0 =    0.0000000000000000
xipm =   0.0000000000000000
v* =     246.21845810181634
cw* =    0.88190138636358650
sw* =    0.47143392403388212
ee* =    0.3028600000000002
alpha_em_i* = 132.50494581248503
particle* = PDG(0)
D_QUARK* = PDG(1)
d* = PDG(1)
down* = PDG(1)
dbar* = PDG(-1)
D* = PDG(-1)
d~* = PDG(-1)
U_QUARK* = PDG(2)
u* = PDG(2)
up* = PDG(2)
ubar* = PDG(-2)
U* = PDG(-2)
u~* = PDG(-2)
S_QUARK* = PDG(3)
s* = PDG(3)
strange* = PDG(3)
sbar* = PDG(-3)
S* = PDG(-3)
s~* = PDG(-3)
C_QUARK* = PDG(4)
c* = PDG(4)
charm* = PDG(4)
cbar* = PDG(-4)
C* = PDG(-4)
c~* = PDG(-4)
B_QUARK* = PDG(5)
b* = PDG(5)
bottom* = PDG(5)
bbar* = PDG(-5)
B* = PDG(-5)
b~* = PDG(-5)
T_QUARK* = PDG(6)
t* = PDG(6)
top* = PDG(6)
tbar* = PDG(-6)
T* = PDG(-6)
t~* = PDG(-6)
E_LEPTON* = PDG(11)
e-* = PDG(11)
e1* = PDG(11)
electron* = PDG(11)
e* = PDG(11)
e++* = PDG(-11)
E1* = PDG(-11)
positron* = PDG(-11)
E_NEUTRINO* = PDG(12)

```

```

nue* = PDG(12)
n1* = PDG(12)
nu_e* = PDG(12)
ve* = PDG(12)
e-neutrino* = PDG(12)
nuebar* = PDG(-12)
N1* = PDG(-12)
ve~* = PDG(-12)
MU_LEPTON* = PDG(13)
mu-* = PDG(13)
e2* = PDG(13)
muon* = PDG(13)
mu+* = PDG(-13)
E2* = PDG(-13)
MU_NEUTRINO* = PDG(14)
numu* = PDG(14)
nu_mu* = PDG(14)
n2* = PDG(14)
vm* = PDG(14)
mu-neutrino* = PDG(14)
numubar* = PDG(-14)
N2* = PDG(-14)
vm~* = PDG(-14)
TAU_LEPTON* = PDG(15)
tau-* = PDG(15)
e3* = PDG(15)
tau* = PDG(15)
ta-* = PDG(15)
tauon* = PDG(15)
tau+* = PDG(-15)
E3* = PDG(-15)
ta+* = PDG(-15)
TAU_NEUTRINO* = PDG(16)
nutau* = PDG(16)
nu_tau* = PDG(16)
n3* = PDG(16)
vt* = PDG(16)
tau_neutrino* = PDG(16)
nutaubar* = PDG(-16)
N3* = PDG(-16)
vt~* = PDG(-16)
GLUON* = PDG(21)
gl* = PDG(21)
g* = PDG(21)
G* = PDG(21)
gluon* = PDG(21)
PHOTON* = PDG(22)
A* = PDG(22)
gamma* = PDG(22)
photon* = PDG(22)
Z_BOSON* = PDG(23)
Z* = PDG(23)
W_BOSON* = PDG(24)
W+* = PDG(24)
Wp* = PDG(24)
W-* = PDG(-24)
Wm* = PDG(-24)
HIGGS* = PDG(25)
H* = PDG(25)
h* = PDG(25)
Higgs* = PDG(25)
PROTON* = PDG(2212)
p* = PDG(2212)
p+* = PDG(2212)

```

```

pbar* = PDG(-2212)
p-* = PDG(-2212)
HADRON_REMNANT* = PDG(90)
HADRON_REMNANT_SINGLETON* = PDG(91)
HADRON_REMNANT_TRIPLET* = PDG(92)
HADRON_REMNANT_OCTET* = PDG(93)
charged* = PDG(11, 13, 15)
colored* = PDG()
m-* = PDG(13)
m** = PDG(-13)
t-* = PDG(15)
t** = PDG(-15)
n_calls(qedtest_eemm) = [unknown integer]
integral(qedtest_eemm) = [unknown real]
error(qedtest_eemm) = [unknown real]
accuracy(qedtest_eemm) = [unknown real]
chi2(qedtest_eemm) = [unknown real]
efficiency(qedtest_eemm) = [unknown real]
-----
Cut expression:
Evaluation tree:
[empty]
-----
Weight expression:
Evaluation tree:
[empty]
-----
Scale expression:
Evaluation tree:
[empty]
-----
begin type(vamp_grids) :: g
size (g%grids) = 1
sum_integral = 0.1739012656355327999336E+05
sum_weights = 0.5003828047561890457473E+02
sum_chi2 = 0.6043702921481354162097E+07
begin g%weights
    1 0.10000000000000000000000000000000E+01
end g%weights
begin g%num_calls
    1 2000
end g%num_calls
begin g%grids
begin type(vamp_grid) :: g
size (g%div) = 2
num_calls = 2000
calls_per_cell = 71
stratified = T
all_stratified = F
quadrupole = F
    mu(1) = 0.3475364540559422721344E+03
    mu(2) = 0.1998469952394245005567E-01
sum_integral = 0.1739012656355328363134E+05
sum_weights = 0.5003828047561891168016E+02
sum_chi2 = 0.6043702921481356956065E+07
calls = 0.19880000000000000000000000000000E+04
dv2g = 0.1428571428571428526821E-01
jacobi = 0.5030181086519115111338E-03
f_min = 0.2606941812215391678365E+03
f_max = 0.5207462045037691495963E+03
mu_gi = 0.6379018118552520633102E+02
sum_mu_gi = 0.3191950977751858772535E+04
begin g%num_div
    1 1
    2 20

```

```

end g%num_div
begin g%div
begin type(division) :: d
ubound(d%x,1) = 1
d%ng = 1
d%stratified = F
d%dx = 0.10000000000000000000E+01
d%dxg = 0.10000000000000000000E+01
d%x_min = 0.00000000000000000000E+00
d%x_max = 0.10000000000000000000E+01
d%x_min_true = 0.00000000000000000000E+00
d%x_max_true = 0.10000000000000000000E+01
begin d%x
    0 0.00000000000000000000E+00
    1 0.10000000000000000000E+01
end d%x
end type(division)
begin type(division) :: d
ubound(d%x,1) = 14
d%ng = 28
d%stratified = T
d%dx = 0.10000000000000000000E+01
d%dxg = 0.50000000000000000000E+00
d%x_min = 0.00000000000000000000E+00
d%x_max = 0.10000000000000000000E+01
d%x_min_true = 0.00000000000000000000E+00
d%x_max_true = 0.10000000000000000000E+01
begin d%x
    0 0.00000000000000000000E+00
    1 0.7142857142857142460635E-01
    2 0.1428571428571428492127E+00
    3 0.2142857142857142738190E+00
    4 0.2857142857142856984254E+00
    5 0.3571428571428571507873E+00
    6 0.4285714285714285476381E+00
    7 0.50000000000000000000E+00
    8 0.5714285714285713968508E+00
    9 0.6428571428571429047238E+00
    10 0.7142857142857143015746E+00
    11 0.7857142857142856984254E+00
    12 0.8571428571428570952762E+00
    13 0.9285714285714286031492E+00
    14 0.10000000000000000000E+01
end d%x
end type(division)
end g%div
empty g%map
empty g%mu_x
end type(vamp_grid)
end g%grids
end type(vamp_grids)

-----
| Global history [vamp]:
-----
[vamp] it #calls integral average chi2 eff.
[vamp] 1 1988 0.0000E+00(0.00E+00) 0.347536E+03(0.14E+00) 0.0 0.000
-----

-----
| Channel histories [vamp]:
=====
chan#001 it #calls integral average chi2 eff.
chan#001 1 1988 0.3475E+03(0.14E+00) 0.347536E+03(0.14E+00) 0.0 0.667
=====
```

```

=====
| It      Calls  Integral[GeV] Error[GeV] Err[%]   Acc  Eff[%]   Chi2 N[It] |
=====
| 1       2000  3.4753645E+02 1.41E-01    0.04   0.02* 66.74
=====
| 1       2000  3.4753645E+02 1.41E-01    0.04   0.02  66.74
=====
| Phase-space grove weight history: (numbers in %)
| grove | 1
=====
| 1 | 100
=====

=====
=====
=====
| Decay tree:
tries =           1
acceptance probability = 1.0000000000000000
=====
Mother process = 'qedtest_eemm'
=====
Decay chain:
=====
Decay configuration: [undefined]
=====
Evaluator: Color-summed including all decays
Evaluator:
Interaction:     8
Virtual:
Particle 1
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)1
Particle 2
E = 250.00000000000000
P = 0.0000000000000000 0.0000000000000000 -249.9999999947775
mask [fch] = [FTT]
internal links: X => 3 4
source: (2)2
Outgoing:
Particle 3
E = 250.00000000000000
P = 223.64041528426193 -102.79251826657215 43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (5)3
Particle 4
E = 250.00000000000000
P = -223.64041528426193 102.79251826657215 -43.802416227981048
mask [fch] = [FTT]
internal links: 1 2 => X
source: (5)4
State matrix: norm = 8.67160247171198766E-003
[f(11)]
[f(-11)]
[f(13)]
[f(-13)] => ME(1) = 1.000000000000
Matrix-element multiplication
Input interaction 1: 2
Input interaction 2:      5

```

```

ME(1) =
+ ME1(1) x ME2(1)
=====
Evaluator: Color flow components including all decays
Evaluator:
Interaction:      9
Virtual:
Particle 1
E =    250.00000000000000
P =    0.0000000000000000      0.0000000000000000      249.9999999947775
mask [fch] = [FFgT]
internal links: X => 3 4
source: (7)1
Particle 2
E =    250.00000000000000
P =    0.0000000000000000      0.0000000000000000      -249.9999999947775
mask [fch] = [FFgT]
internal links: X => 3 4
source: (7)2
Outgoing:
Particle 3
E =    250.00000000000000
P =    223.64041528426193      -102.79251826657215      43.802416227981048
mask [fch] = [FFgT]
internal links: 1 2 => X
source: (6)3
Particle 4
E =    250.00000000000000
P =    -223.64041528426193     102.79251826657215      -43.802416227981048
mask [fch] = [FFgT]
internal links: 1 2 => X
source: (6)4
State matrix: norm = 1.0000000000000000
[f(11)]
[f(-11)]
[f(13)]
[f(-13)] => ME(1) = 8.671602471712E-03
Matrix-element multiplication
Input interaction 1: 7
Input interaction 2:      6
ME(1) =
+ ME1(1) x ME2(1)
=====
[Process: qedtest_eemm]

Particle set:
Particle 1 [i] f(11)
Momentum:
E =    250.00000000000000
P =    0.0000000000000000      0.0000000000000000      249.9999999947775
T =    2.61124455391836818E-007
Children: 3 4
Particle 2 [i] f(-11)
Momentum:
E =    250.00000000000000
P =    0.0000000000000000      0.0000000000000000      -249.9999999947775
T =    2.61124455391836818E-007
Children: 3 4
Particle 3 [o] f(13)
Momentum:
E =    250.00000000000000
P =    223.64041528426193      -102.79251826657215      43.802416227981048
T =    1.1172489955660817E-002
Parents: 1 2
Particle 4 [o] f(-13)

```

```
Momentum:  
E = 250.0000000000000  
P = -223.64041528426193 102.79251826657215 -43.802416227981048  
T = 1.11724899955660817E-002  
Parents: 1 2  
  
Event weight = 1.000000000000000  
Excess weight = 0.000000000000000  
=====
```

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