# GoSam Manual

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## 1 Introduction

#### 1.1 Synopsis

GoSam is a general one-loop evaluator for matrix elements. The program produces Fortran 95 code from a given process description by evaluating Feynman diagrams and translating the associated oneloop diagrams into a numerical representation of the numerator such that it can be evaluated and reduced numerically with either the golem95 library [BGH<sup>+</sup>09, CGH<sup>+</sup>11] or SAMURAI [MORT10] or PJFry [Yun11, FR11].

#### 1.2 Conventions

In this manual, shell commands are indicated by lines starting with a dollar sign (\$) and are given for the **bash** shell only. Lines that are broken for type setting reasons and should continue the previous line(s) start with a  $\hookrightarrow$ .

**Python** program fragments are denoted by the '>>>' and '...' (for continuation lines) prompts.

## 2 Setup

#### 2.1 Prerequisites

The generation of matrix element code using GoSam can be understood as a three step process, although the three steps are not necessarily obvious to the user. In principle, each step could be run on a different machine and the programs listed below only need to be available during the respective step.

- 1. During the setup of the process directory Python and QGraf need to be installed. This phase is initiated by running gosam.py or any user written Python script of similar functionality.
- 2. During the **code generation** only Form and haggies are run. This phase is initiated by running make source.
- 3. During the compilation and running of the matrix element a Fortran 95 compiler and the chosen reduction libraries need to be installed. At the level of the matrix element, this phase is initiated by running make compile. Please note that running make compile will invoke make source if the latter has not been run successfully before that.

Before running the GoSam package, Please ensure that the following programs are available on your system. The numbers indicate during which phase of the code generation the tools will be required.

- QGraf (1) QGraf [Nog93] is required in version 3.1 or higher and can be downloaded from http://cfif.ist.utl.pt/~paulo/qgraf.html.
- Python (1) This program has been tested with Python versions 2.6, 2.7 and 3.1. Please see also http://python.org.
  - Form (2) You will need Form [Ver00] version 3.3 (build 11-aug-2010 or later).
    The most recent version is available from http://www.nikhef.nl/
    ~form/.
- haggies (2) The code generator haggies [Rei10] is included in the GoSam distribution already. Alternatively, it can also be obtained separately from the URL http://sourceforge.net/projects/haggies/. haggies requires Java in version 1.5 or higher. The current version of GoSam requires haggies in version 1.1 or higher.
- golem95/SAMURAI/PJFry For one-loop calculations, at least one of these three libraries is (3) required. If the program is used for the extraction of the  $R_2$  term only, the libraries are not required.

• golem95 can be downloaded from http://projects.hepforge. org/golem/. • SAMURAI can be downloaded from http://projects.hepforge. org/samurai/. • PJFry can be obtained via git from https://github.com/ Vayu/PJFry/. refrep.cls (3) The documentation is based on the LATEX-class refrep, which appears not to be in the default installation of all LATEX distributions. It can be downloaded from http://www.ctan.org/ as part of the refman package. This file is only needed if one intends to run make doc, which generates some documentation for the matrix element.  $! \rightarrow$  Please note that these programs might have license policies which are different from the license applying to GoSam. The authors of GoSam do not take any responsibility for any problems related to the above mentioned software packages. Download The GoSam source code can be downloaded either via subversion or via HTTP download. Subversion

You can check-out a working copy of the repository with the command

\$ svn co http://svn.hepforge.org/gosam/trunk/ gosam-1.0

This will create a folder gosam-1.0 in your current directory. Authenticated users can use the URL

\$ svn co svn+ssh://svn.hepforge.org/hepforge/svn/ →gosam/trunk/ gosam-1.0

to gain read and write access to the project files.

#### 2.2.2 HTTP Download

2.2

2.2.1

Under http://www.hepforge.org/downloads/gosam/ you can download the sources of GoSam using a web browser or a HTTP client like wget or curl. If you received GoSam as a tar-ball you can unpack it using the command

\$ tar xzvf gosam-1.0.tar.gz

#### 2.3 Installation

GoSam is distributed as a Python package. The installation of the source package is done by running the setup script. One of the following scenarios will be encountered most probably:

• If the Python installation resides in /usr or /usr/local and the user has super-user privileges:

\$ sudo python setup.py install

• If the user wants (or has to) maintain an alternative installation path for Python modules. < XXXX > here denotes the name of the alternative installation tree:

\$ python setup.py install --prefix=<XXXX>

The **prefix** option can also be permanently set in the user's **pydistutils** config file<sup>1</sup> by adding the following lines.

[install] prefix=<XXXX>

After successful installation the user should also update the environment variables

PATH=\$PATH:<XXXX>/bin PYTHONPATH=\$PYTHONPATH:<XXXX>/lib/python<version> ⇔/site-packages

#### 2.4 Directory Structure

The GoSam source directory has the structure as described below:

- doc/ This directory contains the documentation and example setup files.You can run make in this directory to generate the document refman.pdf; this is the document you are currently reading.
- models/ For each implemented model this directory contains the QGraf model file (no extension), a Form interface (\*.hh) and a Python module (\*.py). Currently, only the Standard Model (sm) is distributed with GoSam. A second version of the Standard Model (smdiag) implements diagonal flavour structure ( $V_{CKM} = diag\{1, 1, 1\}$ ) The structure of the model files is discussed in more detail in Chapter C.1. Model files for the MSSM based on LanHEP [Sem10] and FeynRules/UFO [DDF<sup>+</sup>11] can be found in the directory examples/model/.
- templates/ Contains templates for the creation of the files in the process directory. The contents are transformed by the class golem.util.parser.Template and its subclasses in golem.templates.\*. The translation of the templates is controled by the file templates.xml of the same directory.
- src/python/ All model independent Python modules can be found in this directory tree.
  - src/form/ Here one finds all Form files which are not part of the template.
    - build This directory is created during building and installation of this

<sup>&</sup>lt;sup>1</sup> On Unix systems and MacOS this file is called **\$HOME/.pydistutils.cfg**, on Windows it is **%HOME%**\pydistutils.cfg.

package by running setup.py. The files in this directory are of temporary nature and can be safely removed.

dist This directory is created by running setup.py with the sdist or bdist command and contains the distributable package files. To create a tar-ball from the working copy, Please run

\$ python setup.py sdist --formats=gztar

For more information please run

\$ python setup.py --help-commands

- examples This directory contains some simple example processes for which GoSam has been compared to the literature.
  - olp Files in this directory are used by gosam.py --olp, which is GoSam's implementation of the Les Houches interface for one-loop programs [BBD<sup>+</sup>10].

## 3 Setup of a Process

#### 3.1 Introduction

This chapter provides a step by step guide how to set up a new process.

In order to generate the matrix element for a given process one has to create a process specific setup file, which we call *process card*.

The syntax of this file is closely related to that of Java .properties files. The detailed syntax and a full list of options are given in Appendix D. Here we first give a commented example, which should be sufficient to explain the most important features of a *process card*.

## 3.2 Example: $e^+e^- \rightarrow t\bar{t}$ at NLO in QCD

It is recommended to generate and modify a template file for the process card instead of starting from scratch. This can be done by invoking the shell command

#### \$ gosam.py --template eett.in

This would generate the file eett.in with some documentation for all accepted options. The options are filled with some default values, which can be set in a global configuration file. The script will search<sup>1</sup> in the GoSam directory, in the user's home directory and in the current working directory for a file named '.golem' or 'golem.in'. Such a file can be generated with the following command:

\$ golem-config.py > golem.in

In the following brief tutorial it is assumed that the process  $e^+e^- \rightarrow t\bar{t}$  should be calculated to order  $\mathcal{O}(\alpha)\mathcal{O}(\alpha_s)$  (virtual corrections); the tree-level process is of order  $\mathcal{O}(\alpha)$ . We neglect the exchange of a Z or a Higgs boson and treat the electron massless. The output directory is assumed to be in the relative path **eett**.

Listing 3.1: eett.in

```
1 process_path=eett
2 process_name=eett
3 in= e+, e-
4 out= t, t~
5 model= sm
```

<sup>&</sup>lt;sup>1</sup> in this order

```
6
   order= gs, 0, 2
7
8
   qgraf.options=nosnails, notadpoles, onshell
9
   qgraf.verbatim=\
10
      true=iprop[Z, 0, 0];\n
11
      true=iprop[H, 0, 0];
12
  zero=me
13
   one=gs,e
14
15
   extensions=samurai
   samurai.fcflags='pkg-config --cflags samurai'
16
17
   samurai.ldflags='pkg-config --libs samurai'
```

The above lines are discussed one by one. The line numbers on the left are only included for better readability and *must not* be included in the setup file.

- 1 The option process\_path specifies the directory to which all generated files and directories are written. The directory which is specified here must already exist. Specification of a process path is mandatory.
- 2 Setting a process name is optional but recommended. All module names will be prefixed with the process name (e.g. precision  $\rightarrow$  eett\_precision). This will avoid name conflicts if at a later stage more than one matrix elements are linked into one executable.
- 3–4 The options in and out specify the particles of the initial and final state. The particle names must be defined in the selected model file. As the model files usually define mnemonics for the particle names there might be several ways of specifying the same process. Instead of 'e+' one could have written 'ep' or 'positron'. For a complete list of alternative particle names please refer to the documentation of the according model file.

Specifying in and out particles is mandatory.

- 5 The option model specifies which model files should be used in order to generate and evaluate the diagrams. This option is mandatory.
- 6 The option order is a comma separated list with three entries. The first entry specifies a symbol that denotes a coupling constant. In the Standard Model file sm the only two possibilities are 'gs' for the strong coupling constant  $g_s$  and 'e' for the electro-weak coupling. The second number is the power of the chosen coupling constant for the tree-level diagrams and the third parameter specifies the power of that coupling constant for the one-loop diagrams. Note that the numbers refer to the powers in the diagrams of the amplitude

 $\rightarrow$  Appendix C

 $! \rightarrow$ 

rather than the squared amplitude. In the above example the string 'gs, 0, 2' specifies that the tree-level diagrams should be of order  $g_s^0$  and the one-loop diagrams should be of order  $g_s^2$  and an unspecified power of e in both cases. If there is no tree level, i.e. the process is loop induced, the keyword NONE should be put as second item in the list, instead of the tree level power of the coupling.

The value of this option is translated into a vsum constraint in the file qgraf.dat.

This option is mandatory.

- 8-11 The option qgraf.options creates the line 'options=...;' in the file qgraf.dat. The value of the option qgraf.verbatim is passed verbatim to the file qgraf.dat. In our example we specify that loops of size one and self-energy insertions at external lines should be omitted in the graph generation. Lines 9-11 suppress the generation of diagrams containing Higgs and Z bosons. As these commands are passed verbatim to QGraf no mnemonic names are allowed here, e.g. the Higgs particle has to be denoted by 'H' and cannot be replaced by 'h'. For a complete list of available options, Please consult the QGraf manual. For a complete list of particle names see Appendix C.2 resp. the documentation of the model file. These options can be omitted.
- 12–13 The keywords zero and one specify a set of symbols that should be treated as zero (resp. one). These simplifications are applied at the symbolical level. Only symbols that appear in the Form interface of the model file should be specified here (masses, couplings, CKM-matrix elements, etc). In the example we specify the electron mass 'me' to be zero and we do not keep the coupling constants in the calculation explicitly ( $g_s = e = 1$ ).

These options can be omitted.

- 15 The option extensions contains a list of extensions to the core of the program.
- 16–17 For each extension one can add options of the form *extension.name*. Currently the program is scanning for options of the form *extension.ldflags* and *extension.fcflags*. These options are copied to the contens of the according variables (FCFLAGS and LDFLAGS) in the makefiles.

In order to populate the specified process directory with files one invokes

\$ gosam.py eett.in

 $! \rightarrow$ 

#### 3.3 Process Directory Structure

After running golem with an appropriate setup file the process directory contains a number of files which are described below.

- codegen/ This directory contains files which are only relevant for code generation. These files will therefore not be included in a tar-ball created with make dist.
- common/ Fortran files which are common to all helicity amplitudes and to the constructed matrix element code. This directory is always compiled first.
  - doc/ Contains all files (apart from pyxotree.tex and pyxovirt.tex)
     which are necessary for creating doc/process.ps, which lists all
     Feynman diagrams of this process, together with colour and helicity
     info.
- helicity\* This directory contains all files for a specific helicity amplitude. The labeling of the helicities can be found in doc/process.ps. Before invoking make source, this directory only contains the makefiles.
  - matrix This folder contains the code to combine the helicity amplitudes into a matrix element. Here one also finds the test program test.f90. This folder is always compiled last.
- - diagrams-[01].hh The diagram files generated by QGraf.
    - config.sh This script facilitates linking with external programs. For details, run

#### \$ sh ./config.sh -help

- process.hh contains the process dependent definitions for Form. This file is used by golem.frm to generate the expressions for each diagram in every helicity configuration.
- process.dat contains the on-shell conditions, the number of incoming particles and an expression for momentum conservation. This file is needed by the program golem-analyzer.py.
  - func.txt Defines dependencies between parameters of the model files.
- Makefile.conf This files contains the settings which might need to be modified by the user. Please check the contents of this file if you have trouble running the makefiles.
- MakefileThese two files are part of each directory. Makefile.source isMakefile.sourceused when calling make source. Running make from the process<br/>directory will pass through all subdirectories. The following targets<br/>of make are recommended for direct use:

- help: lists all major targets.
- source : generate source files, mainly Fortran 95 files.
- compile : compile the Fortran 95 sources.
  - dist : create a tar-ball of the source files.
  - clean : remove object files and intermediate files.
- very-clean : remove files including targets of make source.
  - doc : create various documents related to the process. To obtain a
     description of the topologies, you need to run source before
     make doc.

#### 3.4 Code Generation and Compilation

The Fortran 95 code is generated by the command

\$ make source

and can be compiled using

\$ make compile

Please note that the compile target invokes the source target if necessary.

A simple test program, which gives the value of the amplitude at a randomly generated phase space point, can be found in the directory matrix/, in order to compile and run it, type

\$ cd matrix \$ make test.exe \$ ./test.exe

The program will generate a file \_debug.xml, which, depending on the settings contains the values of helicity amplitudes and diagrams for a set of phase space points.

#### 3.4.1 Customization

Runtime Parameters. Many settings can be changed without recompiling the code, by creating and modifying the file matrix/param.dat. This file has a very simple format:

- Lines starting with a comment character ('!', '#', ';') in the first column and blank lines are ignored.
- All other lines have the format

```
name = float
# or
name = float, float
```

where the first line defines a real number and the second line defines a complex number, and *name* is a parameter de.

• Whitespace is ignored but must not appear inside names or literals. Physical lines can not be continued nor can multiple entries appear on one line.

The list of recognized names can be found in the file common/model.f90. In addition there are some model independent parameters:

samurai_scalar	selects a library of scalar integrals (see SAMU- RAI documentation).
$\mathtt{samurai}_\mathtt{test}$	sets a method to detect unstable points (see
	SAMURAI documentation).
<pre>samurai_verbosity</pre>	sets the verbosity level of SAMURAI; it should
	be set to zero in a production environment (see
	SAMURAI documentation).
renormalisation	An integer number indicating if no renormal-
	isation (0) or $\beta$ -function renormalisation (1,
	QCD only) should be applied. Other values
	are reserved for future extensions.
gauge <i>i</i> o	for the external vector particle with index $i$
	(e.g. gauge10, gauge20), if not defined as
	a constant.
gauge <i>i</i> z	as gaugeio. The polarisation vector is trans-
	formed into

 $\varepsilon^{\mu}(k_i) \rightarrow \texttt{gauge}i \circ \varepsilon^{\mu}(k_i) + \texttt{gauge}i z \cdot k_i^{\mu}$ 

This allows for a quick check of gauge invariance.

Furthermore, all model constants that have not been specified as zero or one can be set in this way. One can can re-set, for example, the value for the Higgs mass using the entry

mH = 124.5

Please note that upper and lower case letters have to be distinguished and that the names need to be spelled exactly as defined in model.py.

**Compile Time Parameters.** Other configuration options can be found in the file common/config.f90 but require the recompilation of the source code (make clean; make compile). Examples of options contained in config.f90 are

ki debug_lo_diagrams	the floating point kind used throughout the calculation. controls if information about the tree level diagrams is written to the output file.
debug_nlo_diagrams	controls if information about the loop-diagrams is written to the output file.
include_eps_terms	controls if terms of order $\varepsilon$ multiplying poles are taken into account.
include_eps2_terms	controls if terms of order $\varepsilon^2$ multiplying double poles are taken into account.
include_color_avg_factor	controls if the color averaging factor for initial state par- tons is multiplied to the final result.
include_helicity_avg_factor	controls if the helicity averaging factor for initial state particles is multiplied to the final result.
$\texttt{include}_\texttt{symmetry}_\texttt{factor}$	controls if the symmetry factor for identical final state particles is multiplied to the final result.
use_sorted_sum	controls if the diagrams are summed using the algorithm Malcolm [Mal70], which reduces the error accumulated in presence of large cancellations.

#### 3.5 Drawing the Feynman Diagrams

In order to print out the diagrams the makefile contains the target doc which produces the file process.ps. We use  $IAT_EX$  plus the package axodraw [Ver94] to create the graphical representation.

The layout of the diagrams is determined by the algorithm used in feynMF [Ohl95], modelling the propagators by springs. The implemented algorithm works in two steps: first, the topology is disentangled by ordering the external legs such that the diagram can be drawn as a planar graph. The coordinates  $e_k$  of the external legs are fixed along a contour around the drawing area.<sup>2</sup> In a second step the remaining degrees of freedom, the coordinates of the vertices  $v_i = (x_i, y_i)$ , are fixed by minimizing the Lagrangian

$$L(v_1, \dots, v_n; e_1, \dots, e_N) = \frac{1}{4} \sum_{i,j=1}^n t_{ij} (v_i - v_j)^2 + \frac{1}{2} \sum_{i=1}^n \sum_{k=1}^N \lambda_{ik} (v_i - e_k)^2 \quad (3.1)$$

Here, n is the number of vertices and N is the number of external legs. Minimization of the Lagrangian leads to a system of linear

 $<sup>^2</sup>$  Currently, this contour is chosen as an ellipse but in principle any convex shape could be used.

equations, which can easily be solved.

$$\frac{\partial L}{\partial v_r} = 0$$
  
$$\Leftrightarrow \frac{1}{2} \sum_{i,j=1}^n t_{ij} \left( v_i - v_j \right) \cdot \left( \delta_{ir} - \delta_{jr} \right) + \sum_{i=1}^n \sum_{k=1}^N \lambda_{ik} \left( v_i - e_k \right) \cdot \delta_{ir} = 0$$
  
$$\Leftrightarrow M_{rj} v_j \equiv \sum_{j=1}^n t_{rj} \left( v_r - v_j \right) + \left( \sum_{k=1}^N \lambda_{rk} \right) v_r = \sum_{k=1}^N \lambda_{rk} e_k$$

In the last step we used the symmetry of  $t_{ij}$ . The matrix M can be written as

$$M_{rc} = \begin{cases} \left(\sum_{i \neq r} t_{ri}\right) + \left(\sum_{k=1}^{N} \lambda_{rk}\right), & r = c \\ -t_{rc}, & \text{otherwise} \end{cases}$$
(3.2)

The symbol  $t_{ij}$  is the sum of the spring constants of all propagators connecting vertices i and j; similarly,  $\lambda_{ik}$  is the spring constant of the leg k if it is connected to vertex i and zero otherwise.

#### 3.6 Import of Model Files

Examples about how to import model files can be found in the subdirectory examples.

#### 3.6.1 Import from FeynRules

A model description in the UFO [?] format consists of a Python package stored in a directory. In order to import the model into GoSam one needs to set the model variable specifying the keyword FeynRules in front of the directory name, where we assume that the model description is in the directory \$HOME/models/MSSM\_UFO.

#### model= FeynRules,\$HOME/models/MSSM\_UF0

#### 3.6.2 Import from LanHEP

In order to use model files generated by LanHEP the following steps have to be taken:

1. When generating the tables using LanHEP, one should include the following option to ensure that the generated tables have the correct headings<sup>3</sup>. The number of spaces in the column headers are irrelevant as long as the columns are wide enough to contain the respective values.

```
prtcformat
fullname: ' fullname ',
name: ' name ',
```

 $<sup>^{3}</sup>$  GoSam relies on the column names rather than some specific order.

```
aname: ' aname ',
spin2: ' spin2 ',
mass: ' mass ',
width: ' width ',
color: ' color ',
aux: ' aux ',
texname: ' texname ',
atexname: ' atexname ',
pdg: ' pdg '.
```

- 2. If the model file is not already equipped with pdg codes the user might want to use the prtcprop command in LanHEP to add the relevant codes.
- 3. In the setup file, one needs to specify the model as a pair of path and integer number. If the table files are under the directory lanhep/ued/ in the tables func7.mdl, lgrng7.mdl, prtcls7.mdl and vars7.mdl, the correct statement in the setup file would be

model=lanhep/ued, 7

4. The use of user defined functions (external\_func in Lan-HEP) requires an adaption of the file codegen/haggies-l0.in. If one wants to use the function double foo(double,double) the following line sould be added.

@define mdlfoo : real, real -> real =
"foo(%2\$s, %3\$s)";

The function also needs to be declared in codegen/functions.out in the subroutine init\_functions

#### 3.7 Handling Big Processes

Although the default settings should work for most cases, very big processes in terms of the number of diagrams and the size of the expressions can cause the compiler to become very slow or even to crash. In this section we discuss solutions which can help to reduce the load for the compiler and to speed-up the code generation. It should be mentioned that some of these measures can have a negative impact on the runtime efficiency of the generated code.

#### 3.7.1 Grouping of Tree Level Diagrams

By default the expressions of all tree-level diagrams are grouped into one file. This has the advantage that subexpressions which appear in several tree-level diagrams can be reused across the amplitude. In some cases it can happen that the sum of all terms of the tree-level diagrams is too big to be compiled in one subroutine. In this case it is recommended to set the option group to false.

#### 3.7.2 Computation of Abbreviations

The constant, i.e. q- and  $\mu^2$  independent parts of the numerators of the one-loop diagrams are factored out from the numerators and computed as abbreviations. In some cases the list of abbreviations is too big to be compiled into one subroutine. One can restrict the number of instructions that go into a single subroutine by setting **abbrev.limit** to a positive number in the setup file. The variable **abbrev.level**, which by default is set to **helicities**, can be set to **groups** or **diagrams** if the list of abbreviations common to a helicity configuration is too large.

If the list of abbreviations causes haggies to crash, one needs to increase the amount of memory reserved for Java. This can be done by adding the -Xmx option to the call of Java. A typical setting of the variable haggies.bin would be

haggies.bin=java -Xmx3g -jar ↔ \${GOLEMPATH}/haggies/haggies.jar

which assigns 3 GB of memory to Java.

#### 3.7.3 Splitting the Process

If a process becomes too big in order to be linked<sup>4</sup> there are some possibilities to split the process into independent programs:

- the generation of a subset of the helicity configurations, e.g. one helicity configuration per process directory.
- the generation of a subset of diagrams. If the diagrams are not split according to gauge invariant subsets the user should ensure that all subsets are called with the same set of phase space points. An easy way of splitting the diagrams into subsets is by using the option select.nlo=(first):(last), where first and last refer to the diagram numbers in process.ps.

#### 3.8 Advanced Usage

The call to the executable gosam.py can be simulated inside more complex Python programs. It is an easy exercise to run the file generation in user defined Python scripts as long as one includes the module files in the environment variable PYTHON\_PATH. The following script emulates the program gosam.py:

>>> from golem.util.config import Properties
>>> from golem.util.main\_misc import \*
>>> props = Properties()

<sup>&</sup>lt;sup>4</sup> Currently, most systems support programs to a size up to 4 GB. Although 64 bit systems can handle a much bigger address space, the current limitation comes from some legacy code in the GNU linker.

```
>>> props.setProperty("in", ["e+", "e-"])
>>> props.setProperty("out", ["t", "t~"])
>>> # ... populate props with further values ...
>>> workflow(props)
>>> generate_process_files(props)
```

#### 3.9 Advanced Diagram Selection

GoSam implements several ways of selecting subsets of diagrams:

- by restricting QGraf,
- by selecting specific diagrams by their number,
- by defining filters using Python.

#### 3.9.1 Restricting the Generation with QGraf

The options for restricting the set of diagrams at the level of the diagram generation is the most efficient way since this happens already at the earliest possible stage. However, QGraf's built-in filters are sometimes too limited in order to express more advanced criteria.

GoSam allows one to pass information to QGraf through the option qgraf.options and through qgraf.verbatim, qgraf.verbatim.lo and qgraf.verbatim.nlo. For the exact syntax the user is referred to the QGraf documentation.

#### 3.9.2 Selecting Diagrams by their Number

An a posteriori selection 'by eye' can be achieved after all (also unwanted) diagrams of a process have been generated and inspected in doc/process.ps. The user can then modify the options select.lo and select.nlo and rerun gosam.py.

#### 3.9.3 Filtering Diagrams in Python

The user can write short Python functions in order to decide whether a specific diagram is to be taken or not. This function should return True for all diagrams which are kept, and False for all diagrams which should be discarded. These functions are passed by the options filter.lo and filter.nlo.

Longer functions should be defined in an external file, which can be passed using filter.module.

When writing a filter the one can use the predefined particle lists QUARKS, LEPTONS, FERMIONS and BOSONS. The underscore (\_) matches any field.

to be	agram object <b>d</b> has the following methods which are inteded e used in filters. Alternative predefined functions and functors also given.
d.rank():	returns the tensor rank of a diagram. $RANK \equiv \lambda d.(d.rank())$
d.loopsize():	returns the number of propagators in the loop of a diagram. $\texttt{LOOPSIZE} \equiv \lambda \texttt{d.}(\texttt{d.loopsize}())$
d.sign():	computes the sign coming from closed fermion loops. $SIGN \equiv \lambda d.(d.sign())$
d.isNf() :	reports if a diagram contains a closed quark loop of size two where all loop propagators are massless. $NF \equiv \lambda d.(d.isNf())$
d.isMassiveQuarkSE() :	returns True if the diagram contains a QCD self energy in- sertion at a massive quark line. $MQSE \equiv \lambda d.(d.isMassiveQuarkSE())$
d.isScaleless() :	returns True if the loop integral associate with this diagram carries no scale. $SCALELESS \equiv \lambda d.(d.isScaleless())$
<pre>d.vertices(f1,f2,) :</pre>	returns the number of vertices in the diagram with the spec- ified fields. The arguments f1, f2, are lists of field names. VERTICES(f1, f2,) $\equiv \lambda d.(d.vertices(f1, f2,))$
<pre>d.loopvertices(f1,f2,) :</pre>	same as vertices, but only counts vertices which have loop propagators attached. LOOPVERTICES(f1, f2,) $\equiv \lambda d.(d.loopvertices(f1, f2,))$
d.iprop(f,**opts) :	returns the number of propagators of the given fields. Optional arguments are momentum to specify the momentum of the propagator, twospin to filter by the $2\times$ the spin, massive to specify whether massive or massless propagators should be considered and color to filter for certain color representations. IPROP() $\equiv \lambda d.(d.iprop())$
d.chord(f,**opts):	same as iprop but only counts loop propagators. $CHORD() \equiv \lambda d.(d.chord())$
d.bridge(f,**opts):	same as iprop but only counts propagators which are not in a loop. BRIDGE() $\equiv \lambda d.(d.bridge())$
d.QuarkBubbleMasses() :	returns a list of all different masses in a closed quark loop of size two or an empty list if the diagram is not a quark bubble. QBMASSES $\equiv \lambda d.(d.QuarkBubbleMasses())$
Furt	hermore, the following predefined filters exist:
NFGEN(f1,f2,) :	for closed quark loops of size two this filter returns true only if all loop propagators belong to one of the fields in the ar-

gument list. For all diagrams which are not quark bubbles it returns True.

- AND(filter1,filter2,...) : returns True if all filters in the argument list return True.
- **OR(filter1,filter2,...)** : returns True if at least one filter in the argument list returns True.
  - NOT(filter) : returns True if the argument evaluates to False.
    - **TRUE** : always returns True.
    - FALSE : always returns False.

## 4 The Binoth Les Houches Accord Interface

#### 4.1 Initialisation Phase

The script gosam.py --olp which comes with GoSam can be used to generate matrix elements compatible with the specifications of the Binoth Les Houches Accord [BBD<sup>+</sup>10]. This script expects at least the name of an order file. This order file is usually but not necessarily created by a Monte Carlo program. An example file for the partonic  $2 \rightarrow 3$  processes of  $pp \rightarrow t\bar{t} + jets$  is given below:

1	MatrixElementSquareType	CHsummed
2	IRregularisation	tHV
3	OperationMode	CouplingsStrippedOff
4	${f SubdivideSubprocess}$	yes
5	AlphasPower	3
6	CorrectionType	QCD
7		
8	# Here comes the list o	f $subprocesses$
9	# specified through PDG	codes
10	$\# g \qquad g \rightarrow t \ t - b  a  r \ g$	
11	$21 \qquad 21 \rightarrow 6  -6 \qquad 21$	
12	$\# \ u \ u - b \ a \ r \ \rightarrow \ t \ t - b \ a \ r \ g$	
13	$2 \qquad -2 \rightarrow 6  -6 \qquad 21$	
14	$\# \ u \qquad g \rightarrow t \ t - b  a  r \ u$	
15	$2 \qquad 21 \rightarrow 6  -6 \qquad 2$	

The line numbers are not part of the file. The arrow ' $\rightarrow$ ' is generated by the two characters '->'. The following options are part of the Standard and accepted by GoSam:

# MatrixElementSquareType : accepts the values Hsummed, Csummed, Haveraged, Caveraged, CHsummed, CHaveraged.

The value **NOTsummed** is not supported. Sensible combinations are also allowed, as in

#### MatrixElementSquareType Hsummed Caveraged

In GoSam this statement is optional. Any quantity which is not explicitly averaged is assumed to be summed

CorrectionType : accepts the values QCD, QED and EW, whereas GoSam does not distinguish between the latter two (this behaviour might change in the future when appropriate model files are available).

This statement is mandatory and must not be omitted.

IRregularisation :	accepts the values <b>tHV</b> ('t Hooft-Veltman scheme) and <b>DRED</b> (dimensional reduction). The value CDR (conventional dimensional regularisation) is not supported and therefore rejected.
	This statement is mandatory and must not be omitted.
MassiveParticleScheme :	accepts the value <b>OnShell</b> only. At the moment this option has no effect on the generation of the matrix element. This statement is optional; if it appears in the order file a warning is issued, reminding the user that no UV-counterterms for massive particles are implemented yet.
IRsubtractionMethod :	accepts the value <b>None</b> only. <b>GoSam</b> does not provide any subtracted output.
	This statement is optional.
ModelFile:	accepts the name of parameter file in the Les Houches Accord format. The script reads the parameter file setting all masses to zero which are not specified explicitly to be non-zero.
	This statement is mandatory.
	It is recommended to use absolute paths here as the file will later be read in the function <b>OLP_Start</b> in the matrix element code, which might be located elsewhere.
OperationMode:	accepts the value <b>CouplingsStrippedOff</b> only.
	This statement is optional. If it is given, the coupling con- stants are stripped off from the amplitude.
SubDivideSubProcess :	accepts logical values ( <b>yes</b> or <b>no</b> ).
	If the value is <b>yes</b> a separate channel for each helicity is assigned. Otherwise there will be one channel per subprocess.
	This statement is optional. Its default value is <b>no</b> .
AlphasPower :	the power of $\alpha_s$ of the Born cross-section. At least one of the options <b>AlphaPower</b> and <b>AlphasPower</b> has to be specified.
AlphaPower :	the power of $\alpha$ of the Born cross-section. At least one of the options <b>AlphaPower</b> and <b>AlphasPower</b> has to be specified.
	options which have been proposed for electro-weak corrections currently not supported.

## 4.1.1 Command Line Arguments of gosam.py --olp

The syntax for the invocation of  $\verb"gosam.py"$  is as follows:

 $gosam.py --olp \{ \langle option \rangle \}$  $\hookrightarrow \langle order \ file \rangle \{ \langle order \ file \rangle \}$   $\hookrightarrow \{\langle key \rangle = \langle value \rangle \}$ 

The allowed options are given below. The list of  $\langle key \rangle = \langle value \rangle$ -pairs supplements the options given in the configuration files.

- -h, --help : Prints a help screen with all available command line options and exits.
- -d, --debug : With this options the script will print lots of extra information to the screen, which is usually not useful for non-experts.
- -v, --verbose : The script will print information e.g. about creating directories and reading files.
  - -w, --warn : Warnings and errors are printed. This is the default setting.
  - -q, --quiet : Only errors are printed, no warnings are issued.
- -lfile, --log-file=file : All messages are written to a log file. When one or more log files are specified the information is still written to the screen with the latest specified level of detail. The following example will read the order file test.olo; messages at the debug level will be written to detailed.log, warnings and errors are written to short.log and only errors are printed to the screen.

\$ gosam.py --olp -d -ldetailed.log -w ↔ -lshort.log -q test.olo

- -cfile, --config=file : Overlay default config files by the specified file. Usually, the script first searches in the default locations for configuration files. Afterwards, all files specified by -c options are read in the order in which they are encountered. Values which are already set by earlier files will be overwritten. See also option '-C'.
- -C, --no-defaults : The script will not search for configuration files (.golem and golem.in) in the standard locations (GoSam installation directory, user's home directory and current working directory).
  - -f, --force : Overwrite contract files without asking. The default behaviour is that contract files are not overwritten. If a contract file already exists the program gives an error message.
- -e, --use-single-quotes : Activates syntax extensions that allow the use of single quotes in order and contract files (See Section 4.1.2).
- -E, --use-double-quotes : Activates syntax extensions that allow the use of double quotes in order and contract files (See Section 4.1.2).
  - -b, --use-backslash : Activates syntax extensions that allow the use of backslash escape sequences in order and contract files (See Section 4.1.2).
    - -i, --ignore-case : Activates syntax estensions which make the parsing of order and contract files case-insensitive (See Section 4.1.2).

-x,ignore-unknown :	Unknown statements or values in order and contract files will be ignored. The default behaviour is that unknown state- ments and/or values will lead to an error message.
-ofile,output-file=file :	Specifies the name of the contract file(s). The following set of wildcard sequences can be used to derive the name of the contract file from the name of the order file. A value of '-' writes to the standard output.
	%f : The full file name (e.g. 'dir/process.olo')
	$\ensuremath{\% F}$ : The file name without any leading path ('process.olo')
	%p: Path name only ('dir/')
	%s: The stem of the file name ('process')
	%e : The extension of the file name ('.olo')
	By default this option is set to '%p%s.olc'.
-D <i>dir</i> ,destination= <i>dir</i> :	Chooses the output directory, to which each process is writ- ten. The same wildcards as above can be used. By default, all output is written to the current working directory. It is therefore not recommended to set this option using wildcards when more than one order file is specified.
$\verb-tpath, \verbtemplates=path:$	Sets an alternative templates directory or template XML-file.
-z, $scratch$ :	Overwrites all process files, including those which otherwise would be preserved (Makefile.conf, config.f90 etc).

#### 4.1.2 GoSam Extensions to the Original Standard

Modern file systems allow for path names which cannot be expressed in the original formulation of the Les Houches accord. Therefore GoSam implements syntax extensions for order and contract files for including special characters in statements, especially in file names (as in ModelFile).

- double quotes : This syntax extension proposes that inside a pair of double quotes (ASCII character #34) special characters lose their special meaning. The backslash acts as escape character, with the following set of escape sequences being allowed:
  - \t expands to a horizontal tabulator character (ASCII character #9),
  - n expands to a new line character (ASCII character #10),
  - f expands to a form feed character (ASCII character #12),
  - r expands to a carriage return character (ASCII character #13),

- - any other character following a backslash expands to itself, in particular \" and \\.
- single quotes : This syntax extension proposes that inside a pair of single quotes (ASCII character #39) all characters lose their special meaning. There is no escape character. A literal single quote is generated by a sequence of two single quotes (Pascal like).
- backslash escapes : This syntax extension proposes that any character following a backslash loses its special meaning.

Different extensions might prove useful on different operating systems. On a Windows system, the file name F:\Golem Files\mssm.slha can only be expressed with the proposed syntax extensions and would have the following three equivalent representations:

- F:\\Golem\ Files\\mssm.slha
- 'F:\Golem Files\mssm.slha'
- "F:\\Golem Files\\mssm.slha"

The three extensions can be switched on by the command line options of gosam.py --olp, '-E', '-e' and '-b' respectively.

#### 4.1.3 Advanced Usage

The core functionality of the script gosam.py --olp is implemented by the function golem.util.olp.process\_order\_file, which has the the following signature:

process\_order\_file(order\_file\_name, out\_file, out\_dir, conf, templates=None, ignore\_case=False, ignore\_unknown=False, single\_quotes=False, double\_quotes=False, backslash\_escape=False)

- order\_file\_name : (character string) name of the order file.
   out\_file : (file object, open for writing) contract file.

  - templates : (character string) template directory or name of an XML-file.
    - ... : all other arguments activate the corresponding syntax extensions.

The return value is zero in case of a success and one if an error occurred.

A list of options read from default config files can be obtained by the function golem.util.main\_misc.find\_config\_files(). The following example suggests the usage of the interface from a Pythonbased Monte Carlo program

import os
import golem

# Monte Carlo program prepars the process # and writes order file proc.olo ... # (not shown in example) conf = golem.util.main\_misc.find\_config\_files() f = open("proc.olc", 'w') os.mkdir("proc/") # Add own options conf[golem.properties.model] = \ "FeynRules,\_\${HOME}/models/mssm\_ufo" conf[golem.properties.fc\_bin] = "gfortran" err\_flag = golem.util.olp.process\_order\_file(\ "proc.olo", f, "proc/", conf) if err\_flag > 0:

print "Problems\_generating\_OLP" print "Please\_consult\_the\_file\_proc.olc"

#### 4.2 Runtime Phase

After the script gosam.py --olp or any equivalent program has been run successfully, the files in the newly created process directories are compiled by invoking make in the respective top-level directory. This generates the object file olp\_module.o which contains all API functions. The library for a given process can be linked using the script config.sh in the same directory. The makefile of a client program would typically contain code similar to the following:

PROCESS\_PATH=path/to/your/process-files LDFLAGS+=\$(shell sh \$(PROCESS\_PATH)/config.sh -libs)

! → The module olp\_module.f90 uses Fortran 2003 extensions (ISO\_C\_BINDING) for establishing a well defined interface for the linker. Older Fortran 95 compilers might therefore not be able to compile this module. Please refer to the compiler documentation for details.

The file olp.h contains the following prototypes.

The first two functions are defined exactly as proposed in [BBD<sup>+</sup>10]. The other two functions extend the original standard. It should, however, be noted that the generated matrix element code can be run without any calls to either OLP\_Finalize or OLP\_Option.

OLP\_Start

void OLP\_Start(char\* contract\_file, int\* success);

This function should be called before the first evaluation of the matrix element. It ensures that all global variables in the matrix element code are initialized properly. The argument contract\_file should receive the (full) name of the contract file which was generated together with the matrix element. The integer success is initialized by OLP\_Start to either the value one, indicating success, or zero, indicating that an error occurred during initialization.

Matrix elements generated with GoSam will try to read the SLHA model file specified by the option **ModelFile** in the contract file. It is not required that the contract file used in the runtime phase points to the same model file as used during the initialisation phase. However, values which were set to zero during initialisation will remain zero during the runtime phase.

OLP\_EvalSubProcess

#### 

This function retrieves the values for a channel of the OLP for a given phase space point. A channel might be a subprocess or a gauge invariant partial amplitude, depending on the settings in the contract file. The channel is labeled by the argument label. The second argument is a one-dimensional array holding the  $5 \times N$  components of the momenta for an N-particle process. They are in the order

$$(E^{(1)}, p_x^{(1)}, p_y^{(1)}, p_z^{(1)}, m^{(1)}, E^{(2)}, p_x^{(2)}, p_y^{(2)}, p_z^{(2)}, m^{(2)}, \dots, m^{(N)})$$

The third argument is the renormalization scale (not its square). A list of scale dependent parameters is passed in the fourth argument. Its first entry is expected to be  $\alpha_s(\mu)$ . Any further entries are user-defined; the user is expected to adapt the subroutine

	<pre>init_event_parameters in olp_module.f90 if he wishes to make use of any additional parameters.</pre>
	The last argument is an array of length four. Its entries are, in this order,
	1. the coefficient of the $1/\varepsilon^2$ pole in the Laurent series of the interference term between virtual and Born amplitude,
	2. the coefficient of the $1/\varepsilon$ pole in the Laurent series of the interference term between virtual and Born amplitude,
	3. the $\mathcal{O}(1)$ term in the Laurent series of the interference term between virtual and Born amplitude,
	4. the square of the Born amplitude.
	Matrix elements generated by GoSam use the convention that in case of an error during the evaluation of the matrix element, the fourth entry is set to $(-1)$ . It is therefore recommended that client programs check for the positiveness of the Born matrix element.
OLP_Finalize	
	<pre>void OLP_Finalize();</pre>
	This function should be called after the last evaluation of the ma- trix element. It allows the OLP to close any open file handles, to release allocated memory and to exit gracefully. Although on most modern operating systems this is done automatically, it is good practice and therefore recommended to always call this function before exiting the program.
OLP_Option	
	<pre>void OLP_Option(char* assignment, int* success);</pre>
	This function can be used to update internal parameters of the OLP which are not part of the standard. The first argument is a character string containing a textual representation of the requested assignment. The second argument will be set by the function according to the success of the request.
	Matrix elements generated with GoSam accept any string which would also be valid as a (non-comment) line in a parameter file (see model.f90). Typical calls would be
	<pre>OLP_Option("samurai_test=3", &amp;flag); /* The previous call requires * reinitialization of the OLP */ OLP_Start(contract_file, &amp;flag); OLP_Option("Nf=5", &amp;flag); /* Setting the Higgs mass: */ OLP_Option("mH=124.5", &amp;flag);</pre>

The necessity to link the client program each time another OLP is used might become cumbersome, especially when one likes to work with more than one OLP at the same time. We have therefore developed a socket protocol which enables any client program to access the same functionality as defined in the Les Houches accord [BBD<sup>+</sup>10] through a TCP/IP connection with a server hosting the OLP. In this way it is possible to access multiple OLPs simultaneously and to load OLPs at runtime.

#### The OLP Socket Server

OLPs generated with GoSam contain additional files in their toplevel directory implementing a server for the OLP Socket protocol. These files are

- olp\_daemon.c : ANSI-C file with service routines especially network related routines,
- olp\_daemon.h : ANSI-C file, header for olp\_daemon.c,
- olp\_protocol.l : Lex/Flex file, part of the grammar definition of the protocol and
- olp\_protocol.y : Yacc/Bison file, part of the grammar definition of the protocol, contains the main program.

These files are compiled with the command

\$ make olp\_daemon EXTRA\_LDFLAGS=...

It is often necessary to specify the variable EXTRA\_LDFLAGS to provide the necessary run-time libraries of the Fortran 95 compiler.

The compiled program can be run with the following options

\$ olp\_daemon [-p port] [-s|-S] [-f] file-name

- -ffile-name : name of a contract file (required).
  - -pport : port at which the program accepts connections, default: 7711.
  - -s/-s: forbid resp. allow the SHUTDOWN command, default: allow.
  - -r/-R: forbid resp. allow the RESTART command, default: allow.
    - -d : detach from terminal (run as daemon).

#### **OLP** Socket Clients

Sample client implementations for C++, Java and Python are provided in the directory olp/contrib/. Below, a brief example for the C++ case is given:

olp::OLPClient OLP\_EvalSubProcess("localhost", 7711); OLP\_EvalSubProcess(0, num\_legs, mom, scale,

```
num_param, param, amp);
OLP_EvalSubprocess.close();
```

The class OLPClient overwrites the operator () emulating the original protocol as closely as possible. For technical reasons, two additional arguments (num\_legs and num\_param) are required, specifying the number of external legs and the length of the array param respectively.

Definition of the Protocol

The protocol consists of statements sent by the client to the server. Each statement is terminated by a newline character. The server responds with one line starting with a three digit number followed by a space and an optional message. The three digit number contains the response code. A response code of 200 signals success, all other values denote an error.

## Appendix A Conventions of the Amplitude

#### A.1 Convention of golem95

The integral library golem95 computes integrals of the form

$$\int \frac{\mu^{2\varepsilon} d^{n}k}{i\pi^{n/2}} \frac{k^{\mu_{1}} \cdots k^{\mu_{r}}}{((k+r_{1})^{2} - m_{1}^{2}) \cdots (k+r_{N})^{2} - m_{N}^{2})} = r_{\Gamma} \cdot \left[\frac{c_{-2}}{\varepsilon^{2}} + \frac{c_{-1}}{\varepsilon} + c_{0} + \mathcal{O}(\varepsilon)\right] \quad (A.1)$$

where  $n = (4 - 2\varepsilon)$  and

$$r_{\Gamma} = \frac{\Gamma(1+\varepsilon)\Gamma^2(1-\varepsilon)}{\Gamma(1-2\varepsilon)}.$$
 (A.2)

The integration measure for the internal momentum k is

$$\frac{\mu^{2\varepsilon} \mathrm{d}^n k}{(2\pi)^n} = \mu^{2\varepsilon} \frac{i}{2^n \pi^{n/2}} \cdot \frac{\mathrm{d}^n k}{i\pi^{n/2}} = \frac{(4\pi)^{\varepsilon} \cdot i}{(4\pi)^2} \cdot \frac{\mu^{2\varepsilon} \mathrm{d}^n k}{i\pi^{n/2}}.$$
 (A.3)

#### A.2 Convention of GoSam

The factor from above which does not go into the integral definition of golem95 can be written as

$$\frac{(4\pi)^{\varepsilon} \cdot i}{(4\pi)^2} = \frac{(4\pi)^{\varepsilon}}{(2\pi)(4\pi)} \frac{i}{2}$$
(A.4)

The factor of i/2 is included in the amplitude definition of GoSam. The factors  $(2\pi)$  and  $(4\pi)$  are later used to build up a factor of  $\alpha_x/2\pi$ , where  $\alpha_x$  is either  $\alpha$  or  $\alpha_s$ .

In the following we assume that the coupling constants<sup>1</sup> have been set to one in the setup of GoSam. This ensures that the one-loop matrix element in QCD is calculated in the  $\overline{\rm MS}$  scheme as

$$|\mathcal{M}|_{1-\text{loop}}^2 = \frac{\alpha_s}{2\pi} \frac{(4\pi)^{\varepsilon}}{\Gamma(1-\varepsilon)} \cdot \left[\frac{c_{-2}}{\varepsilon^2} + \frac{c_{-1}}{\varepsilon} + c_0 + \mathcal{O}(\varepsilon)\right] (g_1^{n_1} \cdots g_q^{n_q})$$
(A.5)

The factor  $(g_1^{n_1} \cdots g_q^{n_q})$  are the coupling constants appearing in the squared tree-level matrix element. GoSam will return the coefficients  $c_{-2}$ ,  $c_{-1}$  and  $c_0$ .

The conversion between different conventions for the  $\Gamma$ -functions is straightforward:

$$\frac{1}{\Gamma(1-\varepsilon)} = r_{\Gamma} + \mathcal{O}(\varepsilon^3) = \left(1 - \frac{\pi^2}{6}\varepsilon^2\right)\Gamma(1+\varepsilon) + \mathcal{O}(\varepsilon^3) \quad (A.6)$$

 $<sup>^{1}</sup>$  e and  $g_s$  in the standard model

The relevant terms in the expansion of  $r_{\Gamma}$  are

$$r_{\Gamma} = e^{-\gamma_E \varepsilon} \left( 1 - \frac{\pi^2}{12} \varepsilon^2 \right) + \mathcal{O}(\varepsilon^3)$$
 (A.7)

If one prefers to pull out a factor of  $e^{-\gamma_E \varepsilon} (4\pi)^{\varepsilon}$  the appropriate definition of the matrix element up to terms of  $\mathcal{O}(\epsilon)$  is

$$\frac{|\mathcal{M}|_{1-\text{loop}}^2}{e^{-\gamma_E \varepsilon} (4\pi)^{\epsilon}} = \frac{\alpha_s}{2\pi} \cdot \left[\frac{c_{-2}}{\varepsilon^2} + \frac{c_{-1}}{\varepsilon} + \left(c_0 - \frac{\pi^2}{12} c_{-2}\right)\right] (g_1^{n_1} \cdots g_q^{n_q})$$
(A.8)

## Appendix B Explicit Reduction of the R<sub>2</sub> Terms

The  $R_2$  term [OPP08] consists of all terms of the numerator containing an explicit  $\varepsilon$  or  $\mu^2$  coming from the Lorentz algebra. For an explicit reduction of these terms, a list of all integrals of the form

$$\int \frac{\mu^{2\varepsilon} \mathrm{d}^n k}{i\pi^{n/2}} \frac{N(\hat{q}) \cdot \mu^{2\alpha} \cdot \varepsilon^{\beta}}{D_0 \cdots D_N} \tag{B.1}$$

where either  $\alpha$  or  $\beta$  is a positive integer number and the denominators are  $D_i = (q + r_i)^2 - m_i^2 + i\delta$ . Note that integrals where both  $\alpha$  and  $\beta$  are non-zero, will not contribute to the final result. We expand the above tensor integral and only consider the term of rank r, similarly to Eq. (208) in Ref. [Rei09]:

$$I_N^{n,\alpha,\beta;\mu_1\dots\mu_r} = (-1)^r \frac{\Gamma(\alpha-\varepsilon)}{\Gamma(-\varepsilon)} \varepsilon^\beta \sum_{l=0}^{\lfloor r/2 \rfloor} \left(-\frac{1}{2}\right)^l \sum_{j_1,\dots,j_{r-2l}=1}^N \times \left[\hat{g}^{\bullet\bullet}\dots\hat{g}^{\bullet\bullet}r_{j_1}^{\bullet}\dots r_{j_{r-2l}}^{\bullet}\right]^{\mu_1\dots\mu_r} I_N^{n+2\alpha+2l}(j_1,\dots,j_{r-2l}). \quad (B.2)$$

Here, the integral  $I_N^d(j_1, j_2, ...)$  denotes a Feynman parameter integral with the parameters  $z_{j_1}, z_{j_2}, ...$  in the numerator,

$$I_{N}^{d}(j_{1},...,j_{p}) = (-1)^{N} \Gamma\left(N - \frac{d}{2}\right) \int d_{\Box}^{N} z \, \delta_{z} \frac{\prod_{\nu=1}^{p} z_{j_{\nu}}}{\left[-\frac{1}{2}z^{\mathsf{T}}Sz - i\delta\right]^{N-d/2}}, \quad (B.3)$$

where  $d_{\Box}^N z = \prod_{j=1}^N dz_j \Theta(z_j) \Theta(1-z_j)$  and  $\delta_z = \delta(1-\sum_i z_i)$ . The square brackets  $[\ldots]^{\mu_1 \ldots \mu_p}$  expand to the sum of all possible assignments of indices to the  $\hat{g}^{\bullet\bullet}$ -tensors where a (one) arbitrary assignment of indices to the momenta  $r_i^{\bullet}$  is chosen.

We only need to consider integrals containing an UV pole, which leads to a rational term when multiplied with  $\varepsilon$  stemming either from  $\varepsilon^{\beta}$  or from

$$\frac{\Gamma(\alpha - \varepsilon)}{\Gamma(-\varepsilon)} = (\alpha - 1)! \left[ -\varepsilon + \mathcal{O}(\varepsilon^2) \right], \quad \text{for } \alpha > 0.$$
 (B.4)

The UV divergence comes from the Gamma function

$$\Gamma\left(N - \frac{n+2\alpha+2l}{2}\right) = \Gamma(\varepsilon - (2+\alpha+l-N)) \equiv \Gamma(\varepsilon - \eta)$$
(B.5)

in the Feynman parameter integral  $I_N^{n+2\alpha+2l}.$  Hence, we examine further the expression

$$\varepsilon \cdot I_N^{n+2l+2\alpha}(l_1, \dots, l_{r-2l}) = \begin{cases} \mathcal{O}(\varepsilon), & \eta < 0\\ (-1)^N \frac{1}{2^\eta \eta!} \int d_{\Box}^N z \delta_z \left[ z^\mathsf{T} S z \right]^\eta \prod_{i=1}^{r-2l} z_{l_i}, & \eta \ge 0 \end{cases}$$
(B.6)

The remaining integration can be understood as a special case of the Feynman parameter identity

$$\frac{1}{\prod_{j=1}^{N} A_j^{\alpha_j}} = \frac{\Gamma(\alpha)}{\prod_{j=1}^{N} \Gamma(\alpha_j)} \int d_{\Box}^N z \, \delta_z \frac{\prod_{j=1}^{N} z_j^{\alpha_j - 1}}{\left(\sum_{j=1}^{N} z_j A_j\right)^{\alpha}} \tag{B.7}$$

for  $A_j = 1$ , in which case one finds

$$\int d_{\Box}^{N} z \, \delta_{z} \prod_{j=1}^{N} z_{j}^{\alpha_{j}-1} = \frac{\prod_{j=1}^{N} \Gamma(\alpha_{j})}{\Gamma(\alpha)} \tag{B.8}$$

All phenomenologically relevant, non-zero cases for renormalizable gauge theories (working in Feynman gauge) are listed below:

$$I_1^{n,0,1} = -\frac{1}{2}S_{11} \tag{B.9}$$

$$I_1^{n,0,1;\mu_1} = \frac{1}{2} S_{11} \cdot r_1^{\mu_1} \tag{B.10}$$

$$I_2^{n,1,0} = -\frac{1}{6} \left( S_{11} + S_{12} + S_{22} \right) \tag{B.11}$$

$$I_2^{n,0,1} = 1 (B.12)$$

$$I_2^{n,0,1;\mu_1} = -\frac{1}{2} \left( r_1^{\mu_1} + r_2^{\mu_1} \right)$$
(B.13)

$$I_{2}^{n,0,1;\mu_{1}\mu_{2}} = \frac{1}{6} \left( 2r_{1}^{\mu_{1}}r_{1}^{\mu_{2}} + r_{1}^{\mu_{1}}r_{2}^{\mu_{2}} + r_{2}^{\mu_{1}}r_{1}^{\mu_{2}} + 2r_{2}^{\mu_{1}}r_{2}^{\mu_{2}} \right) - \frac{1}{12} \hat{g}^{\mu_{1}\mu_{2}} \left( S_{11} + S_{12} + S_{22} \right)$$
(B.14)

$$I_3^{n,1,0} = \frac{1}{2} \tag{B.15}$$

$$I_3^{n,1,0;\mu_1} = -\frac{1}{6} \left( r_1^{\mu_1} + r_2^{\mu_1} + r_3^{\mu_1} \right)$$
(B.16)

$$I_3^{n,0,1;\mu_1\mu_2} = \frac{1}{4}\hat{g}^{\mu_1\mu_2} \tag{B.17}$$

$$I_3^{n,0,1;\mu_1\mu_2\mu_3} = -\frac{1}{12} \sum_{l=1}^3 \left[ \hat{g}^{\bullet\bullet} r^{\bullet} \right]^{\mu_1\mu_2\mu_3} \tag{B.18}$$

$$I_4^{n,1,0;\mu_1\mu_2} = \frac{1}{12}\hat{g}^{\mu_1\mu_2} \tag{B.19}$$

$$I_4^{n,2,0} = -\frac{1}{6} \tag{B.20}$$

$$I_4^{n,0,1;\mu_1\mu_2\mu_3\mu_4} = \frac{1}{4!} \left[ \hat{g}^{\bullet\bullet} \hat{g}^{\bullet\bullet} \right]^{\mu_1\mu_2\mu_3\mu_4} \tag{B.21}$$

All other integrals of that type are identically zero.

# Appendix C The included Model Files

## C.1 Format of the Model Files

GoSa	<b>m</b> expects three files for a proper model definition:
$\langle model  angle$ .hh :	Form file containing the Feynman rules
$\langle model \rangle$ .py :	Python file
$\langle model \rangle$ :	(no extension) QGraf model file

## C.1.1 The Python File

Thy Python file contains the following definitions

model_name :	a variable of string type containing a human-readable name for this model, such as "Standard Model (Feyn. Gauge) w/o Higgs" etc.
particles :	a Python dict that contains all particles <i>and</i> anti-particles of the model. The keys are the QGraf names of the fields; the values are objects of the class Particle. The constructor has the arguments
	Particle(name, two_spin, mass, color_rep, partner, width='0')
mnemonics :	a Python dict of human-readable particle names. The values are objects of the class Particle. It is save to refer to the dictionary particles.
parameters :	a Python dict of model parameters with their default values. Both key and value are strings.
functions :	a Python dict of variable names and initialization expres- sions. Both key and value are strings.
types :	the types of all parameters and functions indicated by ' $R$ ' for real numbers and 'C' for complex numbers.
latex_names :	a Python dict assigning
line_styles :	a Python dict assigning line styles to field names. The line style used when drawing Feynman diagrams. Allowed values are photon, ghost, scalar, gluon, fermion.

The propagators in the QGraf file must contain the following functions:

- TWOSPIN : twice the spin of the particle.
  - COLOR: the color representation of the particle  $\in \{1, 3, 8\}$ .
  - MASS : the mass of the particle.
  - WIDTH: the width of the particle (currently not used).
    - AUX : must be zero for most fields. Tensor Ghosts, as introduced by CalcHep have the value 1 here.
  - CONJ: for self-conjugate particles the value is ('+'), otherwise it is ('+', '-').

The vertices must provide all fields that should be accessible in VSUM statements and therefore also the ones that GoSam uses in the order option.

#### C.1.3 The Form File

There are two possible ways of specifying the Feynman rules in the Form file. If a model contains only Standard Model like interactions one can make use of the file src/form/vertices.hh in the GoSam directory and just define the coefficients CL and CR in front of the vertices. This strategy is implemented by the modelfiles models/sm. The file Form contains a procedure VertexConstants which replaces the the vertex constants by their symbols. A QED example would be

#### **#Procedure** VertexConstants

```
Id CL([field.em], [field.ep], [field.ph]) = e;
Id CR([field.em], [field.ep], [field.ph]) = e;
#EndProcedure
```

In the header of the Form file all model specific symbols and functions need to be defined. For this simple model we have the fields and the coupling constant as only new symbols.

Symbols [field.em], [field.ep], [field.ph], e;

Instead of using the file vertices.hh one can also use his own vertex definitions. In this case the Form file must contain the definition

**#Define** USEVERTEXPROC "1"

and it must define the procedure ReplaceVertices. An example for QED is given below.

```
#Procedure ReplaceVertices
Identify Once vertex(iv?,
        [field.ep], idx1?, -1, k1?, idx1L1?, -1, idx1C1?,
        [field.em], idx2?, 1, k2?, idx2L1?, 1, idx2C1?,
        [field.ph], idx3?, 2, k3?, idx3L2?, 1, idx3C1?) =
PREFACTOR(i_ * e) *
        NCContainer(Sm(idx3L2), idx1L1, idx2L1) *
        node(idx1, idx2, idx3);
#EndProcedure
```

It should be noted that GoSam expects the procedure VertexConstants to exist in both cases. If all the constants are already substituted inside ReplaceVertices the file must still provide a possibly empty empty implementation of VertexConstants. GoSam ensures that VertexConstants is always called after ReplaceVertices.

It is recommended to wrap any factors that are global prefactors to the diagram into the argument of the function PREFACTOR as GoSam scans for these functions and brackets them out. Each vertex definition must contain a factor node which contains the indices<sup>1</sup> of the fields at this vertex.

The QGraf style file generates vertex functions as follows:

#### vertex(vertex index,

field<sub>1</sub>, index<sub>1</sub>,  $\pm 2$ spin<sub>1</sub>, momentum<sub>1</sub>,  $\mu_1$ ,  $\pm$ color rep<sub>1</sub>, color index<sub>1</sub>, field<sub>2</sub>, index<sub>2</sub>,  $\pm 2$ spin<sub>2</sub>, momentum<sub>2</sub>,  $\mu_2$ ,  $\pm$ color rep<sub>2</sub>, color index<sub>2</sub>,

field<sub>n</sub>, index<sub>n</sub>,  $\pm 2$ spin<sub>n</sub>, momentum<sub>n</sub>,  $\mu_n$ ,  $\pm$ color rep<sub>n</sub>, color index<sub>n</sub>)

The entries are:

vertex index :	The unique index of this vertex. $(iv1, iv2,)$	
fold .	The field name of the <i>i</i> th partials. These names	~

- field<sub>i</sub>: The field name of the *i*-th particle. These names are constructed from the QGraf field name as [field. $\langle name \rangle$ ].
- index<sub>i</sub>: A unique name for this "ray" (at index 1 they are idx1r1, idx1r2,...)
- $\pm 2 \operatorname{spin}_i$ : twice the spin of the *i*-th particle. The sign distinguishes particles (+) from antiparticles (-).

momentum $_i$ : the incoming momentum of the *i*-th particle.

 $<sup>^1</sup>$  In  ${\tt QGraf}\xspace's terminology these indices are a combination of vertex and ray index of the field.$ 

- $\mu_i$ : the Lorentz index of the *i*-th particle. Depending on the spin of the particle this is a spinor index (spin 1/2), a Lorentz index (spin 1) or a dummy index (spin 0). For higher spins this index must be split into its components using the function SplitLorentzIndex. For its proper definition the reader is referred to the document src/form/lorentz.pdf.
- $\pm$ color rep<sub>i</sub>: the color representation of the *i*-th particle. Allowed values currently are  $\pm 1, \pm 3, \pm 8$ , although the sign only really makes sense for the fundamental representation 3 and its conjugate  $\bar{3} \equiv -3$ .
- $\operatorname{colorindex}_i$ : The color index of the *i*-th particle. Depending on the color representation this is an index in the fundamental, the adjoint or the trivial representation.

All symbols defined in src/form/symbols.hh are also accessible
 ! → in this Form file. Note: until recently the definition of Sqrt2 and sqrt2 were part of the model file. Now these symbols are part of src/form/symbols.hh and must not be redefined.

!  $\rightarrow$  All Dirac matrices and metric tensors must use the notation introduced by spinney. The metric tensor is  $g^{\mu\nu} = d(\mu, \nu)$  and  $\gamma^{\mu} =$  $Sm(\mu), \gamma_5 = Gamma5, \Pi_+ = ProjPlus, \Pi_- = ProjMinus.$  All noncommuting objects must reside inside the function NCContainter (see example).

The color structure must use the objects  $t_{ij}^A = T(A, i, j)$  (where the color flow is such that j is the index of an anti-quark),  $f^{ABC} = f(A, B, C)$  and  $f^{ABE}f^{CDE} = f4(A, B, C, D)$ . At vertices coupling colored with colorless particles it might be necessary to use the d<sub>-</sub> tensor to file the color flow through the vertex.

! → Note that all propagators and wave functions are defined in a model independent way in the files src/form/propagators.hh and src/form/legs.hh. Please, refrain from modifying these files directly but make all changes to src/form/lorentz.nw.

In theories with Maiorana fermions the model file should include the following line:

**#Define** DISPOSEQGRAFSIGN "1"

### C.2 Standard Model (sm)

### C.2.1 Synopsis

The model 'sm' contains the Feynman rules for the Standard Model in Feynman gauge as described in [BDJ01, Appendix A].

# C.2.2 Particle Content

	Name		Alternative Name	es Mas	$\mathbf{SS}$	Comment
	ер		positron e+	me		$e^+$
	em		electron e-	me		$e^-$
	ne			0		$ u_e $
	nebar		ne~	0		$\bar{ u}_e$
	mup		mu+	mmu		$\mu^+$
Leptons	mum		mu-	mmu		$\mu^{-}$
	nmu			0		$ u_{\mu}$
	nmubar		nmu~	0		$ar{ u}_{\mu}$
	taup		tau+	mta	u	$e^+$
	taum		tau-	mta	u	$e^{-}$
	ntau			0		$ u_{ au}$
	ntaubar		ntau~	0		$ar u_ au$
			1	2.6		·
	Name	A	lternative Names	Mass		Comment
	U	u		mU	u	
	Ubar	u		mU	$ \bar{u}$	
	D	d		mD	$\begin{vmatrix} d \\ \overline{d} \end{vmatrix}$	
	Dbar	d~		mD	$\bar{d}$	
	S	S		mS	u	
Quarks	Sbar	s~		mS	$ \bar{u} $	
	C	С		mC	$d_{\overline{a}}$	
	Cbar	с		mC	$ \bar{d}$	
	Т	t		mΤ	$\frac{t}{-}$	
	Tbar	t		mΤ	$\overline{t}$	
	В	b		mB	$  \frac{b}{-}  $	
	Bbar	b	~	mB	$ \bar{b}$	
	Name	А	lternative Names	Mass	C	Comment
	g	g	luon	0	g	
Course Doorse	A	p	hoton gamma	0	$ \gamma$	
Gauge Bosons	Z			mZ Z		
	Wp	W	+	mW	V	V+
	Wm	W	-	mW	V	V-
Scalar Bosons	Name		lternative Names	Mass	L	Comment
	Н		higgs	mH	H	
	phim	_	hi-	mW	$\phi$	
	phip	p	hi+	mW	$  \phi$	+
	chi			mΖ	$\chi$	

Ghost Fields	Name	Alternative Names	Mass	Comment
	gh		0	$u^g$
	ghbar		0	$\bar{u}^g$
	ghA		0	$u^A$
	ghAbar		0	$\bar{u}^A$
	ghZ		mΖ	$u^Z$
	ghZbar		mΖ	$\bar{u}^Z$
	ghWp		mW	$u^+$
	ghWpbar		mW	$\bar{u}^+$
	ghWm		mW	$u^-$
	ghWmbar		mW	$\bar{u}^-$

## C.2.3 Parameters

This section lists all model parameters which are not already listed as particle masses.

Name	Symbol	Description
NC	$N_C$	Number of colors in QCD
е	e	electro-weak coupling constant: $\alpha = e^2/(4\pi)$
gs	$g_s$	strong coupling constant: $\alpha_s = g_s^2/(4\pi)$
SW	$s_w = \sin \theta_w$	sine of weak mixing angle
CW	$c_w = \cos \theta_w$	cosine of weak mixing angle
VUD	$V_{ud}$	CKM mixing matrix element
CVDU	$V_{du}^{\dagger}$	"
VUS	$V_{us}$	"
CVSU	$V_{su}^{\dagger}$	"
VUB	$V_{ub}$	"
CVBU	$V_{bu}^{\dagger}$	"
VCD	$V_{cd}$	"
CVDC	$V_{dc}^{\dagger}$	"
VCS	$V_{cs}^{ac}$	"
CVSC	$V_{sc}^{\dagger}$	"
VCB	$V_{cb}$	"
CVBC	$V_{bc}^{\dagger}$	"
VTD	$V_{td}^{oc}$	"
CVTD	$V_{dt}^{\dagger}$	"
VTS	$V_{to}$	"
CVST	$V_{st}^{\dagger}$	"
VTB	$V_{tb}$	"
CVTB	$V_{bt}^{\dagger}$	"

# Appendix D Template for a Process Setup File

 $\rightarrow$  Chapter 3.1

In order to create a new process setup file one can invoke

\$ gosam.py --template your\_new\_file.in

This is the recommended way of obtaining the most recent documentation of the available options.

The syntax of a general process setup file should obey the following rules:

- A setup file (*process card*) consists of a sequence of lines representing key-value pairs. A key-value pair can span across several lines if each of the lines except the last line is terminated by a backslash.
- A setup file is allowed to contain any number of blank lines or comment lines, indicated by a '!' or a '#' as its first non-blank character.
- The key and the value are separated by a blank, a colon ':' or an equals sign '='. Notice that the line 'key\_=value' will be interpreted as the key 'key' followed by the value '=\_value' as the terminator of the key is the blank and not the equals sign. In order to produce one of the terminators literally as a part of the key one has to escape it with a backslash, e.g. 'very\\_long\\_name:value' would translate to the key 'very\_long\_name' and the value 'value'.
- The escape characters '\\', '\n', '\r', '\f' and '\t' work as usual. Backslashes in front of any other character are just dropped.
- Leading and trailing blanks are removed from the key and the value by default and must be escaped to preserve them. Whitespace is also removed in front and after commas if the value is interpreted as a comma separated list.
- If an option expects a logical value, the literals '1', 'true', '.true.', 't', '.t.', 'yes' and '.y.' are recognized as the value *true*. These values are interpreted case-insensitively. If a value is not recognized as *true* it corresponds to *false*.
- $! \rightarrow$  Note, that deviating from the Java standard, unicode escapes, such as '\u10EF', have not been implemented; neither are octal and hexadecimal escape sequences recognized.

```
process_name : (text)
     A symbolic name for this process. This name will be used
     as a prefix for the Fortran modules.
     Golem will insert an underscore after this prefix.
     If the process name is left blank no prefix will be used
     and no extra underscore will be generated.
process_path: (text)
     The path to which all Form output is written.
     If no absolute path is given, the path is interpreted relative
     to the working directory from which golem-main.py is run.
     Example:
    process_path=/scratch/golem_processes/process1
in: (comma separated list)
     A comma-separated list of initial state particles.
     Which particle names are valid depends on the
    model file in use.
    Examples (Standard Model):
     1) in=u,u~
     2) in=e+,e-
    3) in=g,g
out : (comma separated list)
     A comma-separated list of final state particles.
     Which particle names are valid depends on the
    model file in use.
    Examples (Standard Model):
     1) out=H,u,u~
     2) out=e+,e-,gamma
    3) out=b,b<sup>~</sup>,t,t<sup>~</sup>
model : (comma separated list)
     This option allows the selection of a model for the
     Feynman rules. It has to conform with one of four possible
     formats:
     1) model=<name>
     2) model=<path>, <name>
    3) model=<path>, <number>
     4) model=FeynRules, <path>
    Format 1) searches for the model files <name>, <name>.hh
```

and <name>.py in the models/ directory under the installation

```
path of Golem.
    Format 2) is similar to format 1) but <path> is used instead
     of the models/ directory of the Golem installation
    Format 3) expects the files func<number>.mdl, lgrng<number>.mdl,
    prtcls<number>.mdl and vars<number>.mdl in the directory <path>.
     These files need to be in CalcHEP/CompHEP format.
     Format 4) expects files according to the new FeynRules Python
     interface in the directory specified by <path>.
     (Not fully implemented yet)
model.options : (comma separated list)
     If the model in use supports options they can be passed via this
    property.
order : (comma separated list)
     A 3-tuple <coupling>, <born>, <virt> where <coupling> denotes
     a function of the qgraf style file which can be used as
     an argument in a 'vsum' statement. For the standard model
     file 'sm' there are two such functions, 'gs' which counts
    powers of the strong coupling and 'gw' which counts powers
     of the weak coupling. <born> is the sum of powers for the
     tree level amplitude and <virt> for the virtual amplitude.
     The line
        order = gs, 4, 6
     would select all diagrams which have (gs)<sup>4</sup> at tree level
     and all loop graphs with (gs)<sup>6</sup>.
     Note: The line
        order = gw, 2, 2
     does not imply that no virtual corrections are calculated.
     Instead, for the virtual corrections diagrams are chosen
     with the same order in gw but higher order in gs.
     In other models with more than two different coupling
     constants additional 'vsum' statements, which can be passed
     via the qgraph.verbatim option, might be needed
     to select the correct set of diagrams.
     If the last number is omitted no virtual corrections are
     calculated.
     See also: qgraf.options, qgraf.verbatim
zero: (comma separated list)
     A list of symbols that should be treated as identically
     zero throughout the whole calculation. All of these
```

```
symbols must be defined by the model file.
    Examples:
    1) # Light masses are set to zero here:
       zero=me,mU,mD,mS
    2) # Diagonal CKM matrix:
       zero=VUS, VUB, CVDC, CVDT, \backslash
            VCD, VCB, CVSU, CVST, \
            VTD, VTS, CVBU, CVBC
       one= VUD, VCS, VTB, \
            CVDU, CVSC, CVBT
    See also: model, one
one : (comma separated list)
    A list of symbols that should be treated as identically
    one throughout the whole calculation. All of these
    symbols must be defined by the model file.
    Example:
    one=gs, e
    See also: model, zero
helicities : (comma separated list)
    A list of helicities to be calculated. An empty list
    means that all possible helicities should be generated.
    The helicities are specified as a string of characters
    according to the following table:
       spin massive | 'm' '-' '0'
                                     ·+ ·
                                             'k'
             YES/NO | ---- 0 ----
         0
             YES/NO | ---- -1/2 ---- +1/2 ----
       1/2
                    | ---- -1 ----
         1
               NO
                                      +1 ----
                    | ---- -1 0 +1 ----
         1
              YES
                    | -3/2 ---- +3/2
       3/2
               NO
                    | -3/2 -1/2 ---- +1/2 +3/2
       3/2
              YES
                    | -2 ---- ----
         2
              NO
                                            +2
              YES
                    | -2 -1
         2
                                0
                                     +1
                                             +2
```

Please, note that 'k' and 'm' are not in use yet but reserved for future extensions to higher spins.

The characters correspond to particle 1, 2, ... from left to right.

Examples: # e+, e- --> gamma, gamma:

```
# Only three helicities required; the other ones are
       # either zero or can be obtained by symmetry
       # transformations.
       helicities=+-++,+-+-,+---;
    Multiple helicities can be encoded in patterns, which are expanded
    at the time of code generation. Patterns can have one of the following
    forms:
        [+-], [+-0], [+0] etc. : the bracket expands to one of the symbols
             in the bracket at a time.
       EXAMPLE
             helicities=[+-]+[+-0]
             # expands to 6 different helicities:
             # helicities=+++, ++-, ++0, -++, -+-, -+0
        [a=+-], etc. : as above, but the helicity is also assigned to the
             symbol and can be reused.
       EXAMPLE
             helicities=[i=+-]+i+
             # expands to two helicities
             # helicities=++++, -+-+
        [ab=+-0], etc. : as above, the first symbol is assigned the helicity,
             the second is minus the helicity
       EXAMPLE
             helicities=[qQ=+-][pP=+-]PQ[+-0]
             # expands to 12 helicities
             # helicities=++--+,++---0,+-++,+-+--,+-+-0,\
                           -+-++,-+-+-,-+++,--+++,--++0
             #
ggraf.options : (comma separated list)
    A list of options which is passed to qgraf via the 'options' line.
    Possible values (as of qgraf.3.1.1) are zero, one or more of:
       onepi, onshell, nosigma, nosnail, notadpole, floop
       topol
    Please, refer to the QGraf documentation for details.
qgraf.verbatim : (text)
    This option allows to send verbatim lines to
    the file qgraf.dat. This can be useful if the user
    wishes to put additional restricitons to the selected diagrams.
    This option is mainly inteded for the use of the operators
       rprop, iprop, chord, bridge, psum
    Note, that the use of 'vsum' might interfer with the
    option qgraf.power.
    Example:
    qgraf.verbatim=\
       # no top quarks: \n\
       true=iprop[T, 0, 0];\n
```

```
# at least one Higgs:\n\
       false=iprop[H, 0, 0];\n
    Please, refer to the QGraf documentation for details.
     See also: qgraf.options, order
qgraf.verbatim.lo : (text)
     Same as ggraf.verbatim but only applied to LO diagrams.
     See also: qgraf.verbatim, qgraf.verbatim.nlo
qgraf.verbatim.nlo : (text)
    Same as qgraf.verbatim but only applied to LO diagrams.
     See also: qgraf.verbatim, qgraf.verbatim.nlo
qgraf.bin : (text)
    Points to the QGraf executable.
     Example:
     qgraf.bin=/home/my_user_name/bin/qgraf
     Default: ggraf
form.bin : (text)
    Points to the Form executable.
    Examples:
     1) # Use TForm:
       form.bin=tform
     2) # Use non-standard location:
       form.bin=/home/my_user_name/bin/form
     Default: form
form.tempdir : (text)
     Temporary directory for Form. Should point to a directory
     on a local disk.
     Examples:
     form.tempdir=/tmp
     form.tempdir=/scratch
    Default: /tmp
haggies.bin: (text)
    Points to the Haggies executable.
    Haggies is used to transform the expressions of the diagrams
     into optimized Fortran90 programs. It can be obtained from
```

http://www.nikhef.nl/~thomasr/download.php

#### Examples:

1) haggies.bin=/home/my\_user\_name/bin/haggies

2) haggies.bin=/usr/bin/java -Xmx50m -jar ./haggies.jar

Default: java -jar /home/gionata/share/golem/haggies/haggies.jar

fc.bin : (text)

Denotes the executable file of the Fortran90 compiler.

Default: gfortran

group : (*true/false*)

Flag whether or not the tree-level diagrams should be grouped into a single file.

Default: True

extensions : (comma separated list)

A list of extension names which should be activated for the code generation. These names are not standardised at the moment.

One option which is affected by this is LDFLAGS. In the following example only ldflags.looptools is added to the LDFLAGS variable in the makefiles whereas the variable ldflags.qcdloop is ignored.

extensions=golem95,samurai

ldflags.qcdloops=-L/usr/local/lib -lqcdloop

NOTE: Make sure you activate at least one of 'samurai' and 'golem95'.

Currently active extensions:

samurai	use Samurai for the reduction
golem95	use Golem95 for the reduction
pjfry	use PJFry for the reduction (experimental)
dred	use four dimensional algebra (dim. reduction)
fr5	finite renormalisation for gamma_5
powhegbox	generate additional code specific to PowHEGBox
autotools	use Makefiles generated by autotools
qshift	apply the shift of Q already at the FORM level
topolynomial	(with FORM >= $4.0$ ) use the ToPolynomial command
gaugecheck	modify gauge boson wave functions to allow for
	a limited gauge check (introduces gauge*z variables)
olp_daemon	(OLP interface only): generates a C-program providing
	network access to the amplitude

templates : (text)

```
Path pointing to the directory containing the template
     files for the process. If not set golem uses the directory
     <golem_path>/templates.
     The directory must contain a file called 'template.xml'
debug : (comma separated list)
     A list of debug flags.
     Currently, the words 'lo', 'nlo' and 'all' are supported.
golem95.fcflags : (text)
     FCFLAGS required to compile with golem95.
     Example:
     golem95.fcflags=-I/usr/local/include/golem95
     Default: 'pkg-config --cflags golem'
golem95.ldflags : (text)
    LDFLAGS required to link golem95.
    Example:
     golem95.ldflags=-L/usr/local/lib/ -lgolem-gfortran-double
     Default: 'pkg-config --libs golem'
samurai.fcflags : (text)
     FCFLAGS required to compile with samurai.
    Example:
     samurai.fcflags=-I/usr/local/include/samurai
samurai.ldflags : (text)
    LDFLAGS required to link samurai.
    Example:
    samurai.ldflags=-L/usr/local/lib/ -lsamurai-gfortran-double
samurai.version : (text)
     The version of the samurai library in use.
     Example:
     samurai.version=2.1.0
    Default: 2.0
select.lo : (comma separated list)
    A list of integer numbers, indicating leading order diagrams to be
     selected. If no list is given, all diagrams are selected.
     Otherwise, all diagrams not in the list are discarded.
```

```
The list may contain ranges:
    select.lo=1,2,5:10:3, 50:53
    which is equivalent to
    select.lo=1,2,5,8,50,51,52,53
    See also: select.nlo, filter.lo, filter.nlo
    Default: ,
select.nlo : (comma separated list)
    A list of integer numbers, indicating one-loop diagrams to be selected.
    If no list is given, all diagrams are selected.
    Otherwise, all diagrams
                             not in the list are discarded.
    The list may contain ranges:
    select.nlo=1,2,5:10:3, 50:53
    which is equivalent to
    select.nlo=1,2,5,8,50,51,52,53
    See also: select.lo, filter.lo, filter.nlo
    Default: ,
filter.lo: (text)
    A python function which provides a filter for tree diagrams.
    filter.lo=lambda d: d.iprop(Z) == 1 \
        and d.vertices(Z, U, Ubar) == 0
    The following methods of the diagram class can be used:
    * d.rank() = the maximum rank in Q possible for this diagram
    * d.loopsize() = the number of propagators in the loop
    * d.vertices(field1, field2, ...) = number of vertices
        with the given fields
    * d.loopvertices(field1, field2, ...) = number of vertices
        with the given fields; only those vertices which have
         at least one loop propagator attached to them
    * d.iprop(field, momentum="...", twospin=..., massive=True/False,
                                                               color=...) =
        the number of propagators with the given properties:
         - field: a field or list of fields
          - momentum: a string denoting the momentum through this propagator,
```

```
such as "k1+k2"
         - twospin: two times the spin (integer number)
         - massive: select only propagators with/without a non-zero mass
         - color: one of the numbers 1, 3, -3 or 8, or a list of
                  these numbers
    * d.chord(...) = number of loop propagators with the given properties;
        the arguments are the same as in iprop
    * d.bridge(...) = number of non-loop propagators with the given
        properties; the arguments are the same as in iprop
    See also: filter.nlo, select.lo, select.nlo
filter.nlo: (text)
    A python function which provides a filter for loop diagrams.
    See filter.lo for more explanation.
filter.module : (text)
    A python file of predefined functions which should be available
    in filters.
    Example:
    filter.module=filter.py
    filter.nlo=my_nlo_filter("vertices.txt")
    filter.lo=my_nlo_filter("vertices.txt")
    ----- filter.py -----
    class my_nlo_filter_class:
       def __init__(self, fname):
          self.fields = []
          f = open(fname, 'r')
          for line in f.readlines():
             fields = map(lambda s: s.strip(),
                   line.split(","))
             self.fields.append(fields)
          f.close()
       def __call__(self, diag):
          for lst in self.fields:
             if diag.vertices(*lst) > 0:
                return False
          return True
      ------
```

See filter.lo, filter.nlo

renorm\_beta : (true/false) Set the name of the same variable in config.f90 Activates or disables beta function renormalisation QCD only Default: True renorm\_mqwf : (true/false) Set the name of the same variable in config.f90 Activates or disables UV countertems coming from external massive quarks QCD only Default: True renorm\_decoupling : (*true/false*) Set the name of the same variable in config.f90 Activates or disables UV counterterms coming from massive quark loops QCD only Default: True renorm\_mqse : (true/false) Set the name of the same variable in config.f90 Activates or disables the UV counterterm coming from the massive quark propagators QCD only Default: True renorm\_logs : (true/false) Set the name of the same variable in config.f90 Activates or disables the logarithmic finite terms of all UV counterterms QCD only Default: True renorm\_gamma5 : (true/false)

Set the same variable in config.f90

Activates finite renormalisation for axial couplings in the 't Hooft-Veltman scheme

QCD only, works only with built-in model files.

Default: True

reduction\_interoperation: (integer number)

Set the same variable in config.f90. A value of '-1' lets gosam decide depending on the specified extensions.

See common/config.f90 for details.

Default: -1

reference-vectors: (comma separated list)

A list of reference vectors for massive and higher spin particles. For vectors which are not assigned here, the program picks a reference vector automatically.

Each entry of the list has to be of the form <index>:<index>

EXAMPLE

```
in=g,u
out=t,W+
reference-vectors=1:2,3:4,4:3
```

In this example, the gluon (particle 1) takes the momentum k2 as reference momentum for the polarisation vector. The massive top quark (particle 3) uses the light-cone projection 14 of the W-boson as reference direction for its own momentum splitting. Similarly, the momentum of the W-boson is split into a direction 14 and one along 13.

If cycles are generated in the list (13 has to be known in order to determine 14 and vice versa in the above example) they must be at most of length two. For the reference momenta of lightlike gauge bosons the length of cycles does not matter, e.g.

```
in=g,g
out=g,g
reference-vectors=1:2,2:3,3:4,4:1
```

abbrev.limit: (*text*)

Maximum number of instructions per subroutine when calculating abbreviations, if this number is positive.

```
Default: 0
```

```
abbrev.level: (text)
     The level at which abbreviations are generated. The value should be
     one of:
                       generates files helicity<X>/abbrevh<X>.f90
        helicity
                       generates files helicity<X>/abbrevg<G>h<X>.f90
        group
                       generates files helicity<X>/abbrevd<D>h<X>.f90
        diagram
     Default: helicity
r2: (text)
     The algorithm how to treat the R2 term:
     implicit
                 -- mu^2 terms are kept in the numerator and reduced
                    at runtime
     explicit
                 -- mu<sup>2</sup> terms are reduced analytically
                 -- same as 'explicit' but only the R2 term is kept in
     only
                    the result
     off
                 -- all mu<sup>2</sup> terms are set to zero
     Default: implicit
crossings: (comma separated list)
     A list of crossed processes derived from this process.
     For each process in the list a module similar to matrix.f90 is
     generated.
     Example:
    process_name=ddx_uux
     in=1,-1
     out=2,-2
     crossings=dxd_uux: -1 1 > 2 -2, ud_ud: 2 1 > 2 1
pyxodraw : (true/false)
     Specifies whether to draw any diagrams or not.
     Default: True
```

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