HIP — Symbolic High-Energy Physics Calculations

Alexander Hsieh and Eran Yehudai

Stanford Linear Accelerator Center Stanford University, Stanford, California 94309

ABSTRACT

We present HIP — a set of computer packages for symbolic calculation of Feynman diagrams. The packages were designed as an aid in calculating tree-level multi-particle electroweak production processes, though they can be used for a much wider class of calculations. We have used the packages to calculate such processes as $e\gamma \rightarrow W\nu$ and $\gamma\gamma \rightarrow W^+W^-$ at tree level with arbitrary $W\gamma$ couplings. The packages are written in the computer algebra language *Mathematica* which provides a powerful working environment.

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1. Introduction

Present day pertubative calculations in the Standard Model (SM) and its extensions often require tedious algebra calculations. While tree level calculations of two-body production processes in the SM can certainly be done manually, it is often helpful to have a check in the form of an automatic calculation method. Processes involving the production of more than two particles (e.g. $e^+e^- \rightarrow W^+W^-\gamma$) or complicated Feynman rules (e.g. $\gamma\gamma \rightarrow W^+W^-$ with arbitrary $W\gamma$ couplings) involve much more difficult calculations in which computerized aid is almost indispensable.

Two approaches have been used to automate calculations of this type. Hagiwara *et al.*^[1] have written a set of Fortran subroutines that calculate matrix elements numerically. This approach allows the automatic calculation of complicated tree level processes, but it is restricted to numeric results. An alternative approach introduced by Veltman^[2] with *Schoonship* is to allow symbolic manipulation of expressions. While *Schoonship* was written as a special purpose program, with all the necessary algorithms 'hard-wired' into its assembly code, $Reduce^{[3]}$ followed a more general approach. Much more mathematical knowledge (*e.g.* integration rules) is incorporated into the program as higher-level lisp routines. The user, building on that basis of knowledge, can then expand the program by writing his or her own routines. The price for that flexibility is naturally paid in performance.

We follow this latter approach by writing our packages in *Mathematica*^[4] one of the newer and most promising of the new generation of symbolic manipulation languages which also includes *Maple*^[5] Using *Mathematica*'s high-level programming language greatly simplifies the task of writing programs. Additionally, the physics calculations are embedded within a powerful environment in which results can be simplified, calculated numerically and plotted. With the rapid advance in computer performance, the CPU-time needed for the calculations is usually negligible compared with the time needed to prepare the input and process the output of the programs. This approach is also being used by the Würzburg group in their

programs $FeynArts^{[6]}$ and $FeynCalc.^{[7]}$

Unlike Feyn Arts we do not attempt to automatically generate the necessary Feynman diagrams. Typically, tree-level diagrams can be easily generated manually. HIP takes as input expressions describing the Feynman diagrams. HIP then provides the user with a rich set of operators by which to manipulate the physical objects occurring in these expressions. The user may, for example, substitute explicit four-vectors and particle polarizations, square the matrix elements to give traces that can then be evaluated or convert the expressions to spinor techniques.^[8] As an aid in calculating cross-sections and decay widths, phase-space integrals can be automatically constructed and evaluated symbolically, numerically, or converted to a C or Fortran program.

The traditional method of Feynman diagram calculation involves squaring the matrix element symbolically. The number of terms involved increases like the square of the number of Feynman diagrams involved. In contrast, spinor techniques are methods for calculating Feynman diagrams numerically at the matrix element level. The number of terms involved is linear with the number of Feynman diagrams. Photon and fermion polarizations have to be summed explicitly. Spinor techniques are simplest when the fermions involved can be taken to be massless. They are most useful when a large number of Feynman diagrams is involved.

We have used HIP extensively, typically calculating processes with relatively simple topology ($2 \rightarrow 2$ and $2 \rightarrow 3$ tree-level cross-sections) but with complex Feynman rules.

The paper proceeds as follows. In the next section we give a brief overview of HIP. Some of the major objects and functions are mentioned. In sect. 3 we describe in more detail some of HIP's more important functions, presenting the mathematical relations they use and short examples of their application. In sect. 4 we give two complete examples: the calculation of the width for the muon decay $\mu^- \rightarrow \nu_{\mu} e^- \overline{\nu}_e$ with a finite electron mass, and a calculation of the matrix element for $e_R^+ e_R^- \rightarrow Z \rightarrow t\bar{t} \rightarrow W^+ W^- b\bar{b}$, including the W coupling to light fermions, which preserves all spin and angular correlations. In sect. 5 we summarize and give an outlook. The complete listing of HIP functions, with their usage messages (available as on-line documentation) is given in the appendix.

2. Overview

The packages in HIP contain functions that can manipulate various mathematical objects occurring in high-energy physics such as four-vectors, spinors and gamma matrices. Rather than follow one strict path from input to output, the packages allow the user to specify how a calculation proceeds (either interactively or in batch mode). A typical calculation might be to construct a matrix element, square it and sum over polarizations, construct the phase-space integral and evaluate this integral to give a symbolic expression for the total cross-section.

The most fundamental component of any high energy calculation is the manipulation of four-vectors. Basic objects such as the dot-product $(p \cdot q)$ (DotProduct[p, q]), the metric $g^{\mu\nu}$ (G[mu, nu]) and the completely anti-symmetric tensor $\epsilon^{\mu\nu\sigma\tau}$ (Eps[mu, nu, sig, tau]) are defined, with some of their elementary properties (*e.g.* the dot-product is symmetric in its two arguments). Four-vectors can be specified in terms of their components. They can then be boosted (using the function Boost), represented as sum of other four-vectors (Decay), etc. In addition, four-vectors can also be treated without reference to the explicit components. Dot products can be given explicit values (SetDotProduct, SetMass), Mandelstam variables defined (SetMandelstam), Lorentz indices defined (PrepareIndex) and contracted (Contract).

The second component in HIP is the Dirac algebra. The basic objects involved are the Dirac gamma matrices γ^{μ} (DiracGamma[mu]), γ^{5} (DiracGamma5), the projection operators $P_{\lambda} = (1 + \lambda \gamma^{5})/2$ (HelicityProjection[lambda]) and $\not = p_{\mu}\gamma^{\mu}$ (Slash[p]). The Dirac matrix product is represented by the *Mathematica* builtin function NonCommutativeMultiply (aliased to **). The trace of a product of Dirac gamma matrices is computed using the operator GammaTrace. Some programs handling Dirac algebra, notably Reduce, can only deal with gamma matrices. HIP can also work with spinors. The basic spinor objects u(p) and v(p) (SpinorU[p] and SpinorV[p]) and their conjugates $\overline{u}(p)$ and $\overline{v}(p)$ (SpinorUbar[p] and SpinorVbar[p]) are defined. The function AbsSquared is used to square matrix element expressions which may include these spinors.

Expressions involving spinors do not have to be squared before they are calculated numerically. The HIP function ConvertToST converts a suitable expression involving spinors to an expression involving the elementary spinor products $s(p,k) = \overline{u}_R(p)u_L(k)$ and $t(p,k) = \overline{u}_L(p)u_R(k)$ (SpinorS[p, k] and SpinorT[p, k]) defined in reference 8. The expression produced can then be evaluated numerically by giving explicit values to the components of the four-vectors. Alternatively, it can be squared and converted back to an expression involving traces using the function STToTraces.

Given an expression for the matrix element squared associated with a process, the calculation of physical observables such as cross sections and decay widths involves integration over the phase space of the out-going particles. The functions CrossSection and DecayWidth set up the phase-space integral. The functions return a PhaseSpaceIntegral object that can then be evaluated either symbolically using EvaluatePhaseSpaceIntegral or numerically using NEvaluatePhaseSpaceIntegral. Alternatively, one can write a Mathematica program to convert such an object to a C or Fortran program for numeric evaluation. Such a conversion program would be highly specific, depending on the particular programming language, integration routine etc. We have used one such program in our work, but it is not included with HIP.

HIP includes some of the common Feynman rules of the Standard Model which are implemented using the functions Vertex and Propagator. Constants such as $\sin^2 \theta_W$ (Sin[ThetaW]^2) and particle masses (e.g. Mass[ZBoson]) are usually kept as symbolic constants. However, HIP stores a table of their numerical values; these are substituted for the symbolic expression by the *Mathematica* function N.

3. HIP functions

Strictly speaking, Mathematica does not distinguish between data-structures, functions and procedures. In practice, however, the Mathematica objects defined in HIP can be divided into several broad categories. In all cases we try to follow the Mathematica convention of beginning each name with a capital letter. Further, as far as is practical we use full, descriptive English names rather than cryptic acronyms. Using Mathematica utilities, the user can choose his or her own cryptic abbreviations. The major categories are:

- 1. Objects such as gamma matrices or dot products. These are characterized by the property that they usually remain unevaluated.
- 2. Declarations and definitions. These do not typically return anything, but are rather invoked as part of the initialization process.
- 3. Operations such as contracting indices or taking traces. These typically take their input and convert it to an equivalent expression.

In this section we describe the most important members of each class.

Table 1 lists the major functions representing objects with their equivalent in ordinary physical notation.

The most useful declarative functions are:

- PrepareIndex: PrepareIndex[mu, nu] declares μ and ν and Lorentz indices.
- SetMass: SetMass[p1, p2,..., m] sets p1, p2,... to be four-vectors with invariant mass m.*
- SetMandelstam: SetMandelstam[{p1, p2, p3, p4}, {m1, m2, m3, m4}, s, t, u] sets p1, p2, p3 and p4 to be on-shell with masses m1, m2, m3 and

^{*} Note that the p's are used in a dual mode, both as representing momenta and as representing particles. The mass m associated with p is the mass of the particle carrying the momentum p. For off-shell particles, $p^2 \neq m^2$ (DotProduct[p, p] != Mass[p]^2).

HIP Function	Example	Physical Equivalent
{}	{px, py, pz, e}	The four-vector (p_x, p_y, p_z, E)
DotProduct	DotProduct[p, q]	$p\cdot q$
G	G[mu, nu]	$g^{\mu u}$
Eps	Eps[p, q, mu, nu]	$\epsilon_{\tau\sigma\mu\nu}p^{\tau}q^{\sigma}$
DiracGamma	DiracGamma[mu]	γ^{μ}
Slash	Slash[p]	ý
DiracGamma5	DiracGamma5	γ^5
**	DiracGamma[mu]**Slash[p]	$\gamma_{\mu} p$
SpinorU	SpinorU[p, lambda]	$u_{\lambda}(p)$
SpinorUbar	SpinorUbar[p, lambda]	$\overline{u}_{\lambda}(p)$
HelicityProjection	HelicityProjection[Left]	$P_L = (1 - \gamma^5)/2$
SpinorS	SpinorS[p, k]	$s(p,k) = \overline{u}_R(p)u_L(k)$
SpinorT	<pre>SpinorT[p, k]</pre>	$t(p,k) = \overline{u}_L(p)u_R(k)$

Table 1. HIP functions representing objects

m4 respectively and sets the DotProducts of p1, p2, p3 and p4 in terms of the Mandelstam variables s, t and u and the masses:

$$(p_1 \cdot p_2) \to \frac{1}{2}(s - m_1^2 - m_2^2) \qquad (p_3 \cdot p_4) \to \frac{1}{2}(s - m_3^2 - m_4^2) (p_1 \cdot p_3) \to \frac{1}{2}(-t + m_1^2 + m_3^2) \qquad (p_2 \cdot p_4) \to \frac{1}{2}(-t + m_2^2 + m_4^2) (p_1 \cdot p_4) \to \frac{1}{2}(-u + m_1^2 + m_4^2) \qquad (p_2 \cdot p_3) \to \frac{1}{2}(-u + m_2^2 + m_3^2),$$

where m_i is the mass of the particle p_i .

Most of HIP's functionality is implemented as operator-type functions. The main ones are:

— Boost: Boost[fv, rap, dir] gives a four-vector obtained by boosting the four-vector fv by rapidity rap in the direction specified by dir. Example:

In[1]:= Boost[{0, 0, 0, m}, r, {cth, 0}]

Boost the four-vector (0, 0, 0, m) by rapidity r in the direction $\cos \theta = \operatorname{cth}, \phi = 0.$ Out[1]= {Sqrt[1 - cth²] m Sinh[r], 0, cth m Sinh[r], m Cosh[r]}

 $(m\sqrt{1-\cos^2\theta}\sinh r, 0, m\cos\theta\sinh r, m\cosh r).$

- Decay: Decay[v, dir, {m1, m2}] gives two four-vectors v_1 and v_2 such that $v_1 + v_2 = v$, $v_1^2 = m_1^2$, $v_2^2 = m_2^2$ and the direction of v_1 in the v center-of-mass frame is given by dir. Example:

 $In[1] := Decay[{0, 0, 0, m}, {cth, 0}, {m1, 0}]$

Decompose the four-vector p = (0, 0, 0, m) into two four-vectors p_1 and p_2 such that $p_1^2 = m_1^2$, $p_2^2 = 0$ and the direction of p_1 in the *p* center-of-mass frame is given by $\cos \theta = \operatorname{cth}$ and $\phi = 0$. After some rearrangement one gets:

Sqrt[1 - cth]
$$\binom{2}{m}$$
 $\binom{2}{m}$ $\binom{2}{m}$

$$p_{1} = \left(\frac{\sqrt{1 - \cos^{2}\theta}(m^{2} - m_{1}^{2})}{2m}, 0, \frac{\cos\theta(m^{2} - m_{1}^{2})}{2m}, \frac{m^{2} + m_{1}^{2}}{2m}\right),$$
$$p_{2} = \left(\frac{-\sqrt{1 - \cos^{2}\theta}(m^{2} - m_{1}^{2})}{2m}, 0, \frac{-\cos\theta(m^{2} - m_{1}^{2})}{2m}, \frac{m^{2} - m_{1}^{2}}{2m}\right).$$

- Contract: Contract[expr, index] contracts index in expr. Contract with respect to a particular index μ invokes a large set of rules. The basic rules for handling arbitrary tensors and vector are:

$$g^{\mu}_{\mu} \to D$$

$$p_{\mu}q^{\mu} \to p \cdot q \qquad (3.1)$$

$$g^{\tau}_{\mu}T^{\nu_{1}\dots\mu\dots\nu_{n}} \to T^{\nu_{1}\dots\tau\dots\nu_{n}}$$

where D is the dimension of space-time.^{*} For handling the completely antisymmetric ϵ symbol we use

$$\epsilon_{\mu\nu_{1}\cdots\nu_{n}}\epsilon^{\mu\tau_{1}\cdots\tau_{n}} \to (D-n) \begin{vmatrix} g_{\nu_{1}}^{\tau_{1}} & \cdots & g_{\nu_{1}}^{\tau_{n}} \\ \vdots & \ddots & \vdots \\ g_{\nu_{n}}^{\tau_{1}} & \cdots & g_{\nu_{n}}^{\tau_{n}} \end{vmatrix}.$$
(3.2)

The rules associated with γ -matrices are:^[9]

$$p_{\mu}\gamma^{\mu} \rightarrow \not p$$

$$\gamma_{\mu}\gamma^{\mu} \rightarrow D$$

$$\gamma_{\mu}\gamma_{\nu}\gamma^{\mu} \rightarrow (2-D)\gamma_{\nu}$$

$$\gamma_{\mu}\gamma_{\nu_{1}}\gamma_{\nu_{2}}\gamma^{\mu} \rightarrow (D-4)\gamma_{\nu_{1}}\gamma_{\nu_{2}} + 4g_{\nu_{1}\nu_{2}}$$

$$\gamma_{\mu}\gamma_{\nu_{1}}\gamma_{\nu_{2}}\gamma_{\nu_{3}}\gamma^{\mu} \rightarrow -2\gamma_{\nu_{3}}\gamma_{\nu_{2}}\gamma_{\nu_{1}} - (D-4)\gamma_{\nu_{1}}\gamma_{\nu_{2}}\gamma_{\nu_{3}}.$$
(3.3)

^{*} Most of HIP's functions operate well in arbitrary D dimensions. The exceptions are the functions dealing with vectors given in terms of their explicit components (e.g. Boost), functions associated with phase-space integrals and functions treating γ^5 .

For more complicated cases we use

$$\gamma_{\mu}\Gamma^{(n)}\gamma^{\mu} \rightarrow \begin{cases} 2\Gamma^{(n)} + 2\Gamma_{R}^{(n)} & (n \text{ even}) \\ -2\Gamma_{R}^{(n)} & (n \text{ odd}) \\ (-1)^{n} \left((D-4)\Gamma^{(n)} + 2\Gamma_{R}^{(n)} - 2\sum_{i=4}^{n-3}(-1)^{i}\gamma_{\nu_{i}}\Gamma_{i}^{(n)} \right) & (D \neq 4), \end{cases}$$

$$(3.4)$$

where

:

$$\Gamma_{R}^{(n)} = \gamma_{\nu_{1}} \cdots \gamma_{\nu_{n}}$$

$$\Gamma_{R}^{(n)} = \gamma_{\nu_{n}} \cdots \gamma_{\nu_{1}}$$

$$\Gamma_{i}^{(n)} = \gamma_{\nu_{1}} \cdots \gamma_{\nu_{i-1}} \gamma_{\nu_{i+1}} \cdots \gamma_{\nu_{n}}.$$
(3.5)

Example:

In[1]:= Contract[G[mu, nu] p[mu], mu]

Contract the index μ in $g_{\mu\nu}p^{\mu}$.

Out[1]= p[nu]

 p_{ν}

Out[2]= (-4 + SpaceTimeDimension) Slash[p] ** DiracGamma[nu] + > 4 p[nu] $(D-4) \not \! p \gamma_{\nu} + 4p_{\nu}$ where D is the space-time dimension. — GammaTrace: GammaTrace[expr] is the trace (in spinor space) of expr. tr {1} can be left unevaluated as the constant DiracGammaSize, but is usually set to 4. Whenever possible, GammaTrace uses the following simple rules:

$$\operatorname{tr} \{\gamma_{\mu_{1}} \dots \gamma_{\mu_{2n+1}}\} \to 0 \qquad \operatorname{tr} \{\gamma^{5}\} \to 0$$
$$\operatorname{tr} \{\gamma^{5}\gamma_{\mu}\gamma_{\nu}\} \to 0 \qquad \operatorname{tr} \{\gamma^{5}\gamma_{\mu}\gamma_{\nu}\gamma_{\tau}\gamma_{\sigma}\} \to \operatorname{tr} \{1\} g_{\mu\nu} \qquad \operatorname{tr} \{\gamma^{5}\gamma_{\mu}\gamma_{\nu}\gamma_{\tau}\gamma_{\sigma}\} \to \operatorname{tr} \{1\} i\epsilon_{\mu\nu\tau\sigma}.$$
$$\operatorname{tr} \{\gamma_{\mu}\gamma_{\nu}\gamma_{\tau}\gamma_{\sigma}\} \to \operatorname{tr} \{1\} (g_{\mu\nu}g_{\tau\sigma} + g_{\mu\sigma}g_{\nu\tau} - g_{\mu\tau}g_{\nu\sigma}) \qquad (3.6)$$

Traces of longer expressions invoke the following recursive rules:

$$\operatorname{tr}\left\{\gamma_{\mu}\Gamma^{(n)}\right\} \to \sum_{i=1}^{n} (-1)^{(i+1)} g_{\mu\nu_{i}} \operatorname{tr}\left\{\Gamma_{i}^{(n)}\right\}$$
$$\operatorname{tr}\left\{\gamma^{5}\Gamma^{(n)}\right\} \to \sum_{1 \leq i < j < k < l \leq n} (-1)^{(i+j+k+l)} i \epsilon_{\nu_{i}\nu_{j}\nu_{k}\nu_{l}} \operatorname{tr}\left\{\Gamma_{ijkl}^{(n)}\right\}.$$
$$(3.7)$$

Example:

In[1]:= GammaTrace[DiracGamma[mu]**DiracGamma[nu]

$$angle {
m tr}\left\{ \gamma_{\mu}\gamma_{
u}
ight\}$$

Out[1] = 4 G[mu, nu]

 $4g_{\mu\nu}$

In[2]:= GammaTrace[DiracGamma5**DiracGamma[mu]**DiracGamma[nu]**
DiracGamma[sig]**DiracGamma[tau]]

 $\operatorname{tr}\left\{\gamma^5\gamma_{\mu}\gamma_{\nu}\gamma_{\sigma}\gamma_{\tau}\right\}$

Out[2] = 4 I Eps[mu, nu, sig, tau]

 $4i\epsilon_{\mu\nu\sigma\tau}$

In[3]:= GammaTrace[DiracGamma[mu]**Slash[p1]**DiracGamma[nu]**
 Slash[p2]**DiracGamma[mu]**Slash[p3]**DiracGamma[nu]**
 Slash[p4]]

$$-32(p_1 \cdot p_3)(p_2 \cdot p_4)$$

 AbsSquared: AbsSquared[expr] is the absolute value of expr squared. AbsSquared sums over polarization of both external spinors and vectors unless their polarizations are explicitly specified:

$$\begin{aligned} |Xu(p)|^{2} &\to \sum_{\lambda} Xu_{\lambda}(p)\overline{u}_{\lambda}(p)X^{*} \to X(\not p+m)X^{*} \\ |Xv(p)|^{2} &\to \sum_{\lambda} Xv_{\lambda}(p)\overline{v}_{\lambda}(p)X^{*} \to X(\not p-m)X^{*} \\ |\epsilon_{\mu}(p)|^{2} \to \begin{cases} -g_{\mu\mu'} & (m=0) \\ -g_{\mu\mu'} + \frac{p_{\mu}p_{\mu'}}{m^{2}} & (m\neq 0), \end{cases} \end{aligned}$$
(3.8)

where m is the mass associated with p. Example:

In[1]:= AbsSquared[SpinorUbar[p]**SpinorV[q]]

 $\left|\overline{u}(p)v(q)\right|^2$

Out[1]= 4 DotProduct[p,q] - 4 Mass[p] Mass[q]

$$4p \cdot q - 4m_p m_q$$

In[2]:= AbsSquared[SpinorUbar[p, Right]**SpinorV[q, Left]]

 $\left|\overline{u}_{R}(p)v_{L}(q)\right|^{2}$

Out[2]= 2 DotProduct[p,q]

 $2p \cdot q$

In[3]:= AbsSquared[SpinorUbar[p]**DiracGamma[mu]**SpinorU[q]]

 $\left|\overline{u}(p)\gamma_{\mu}u(q)\right|^{2}$

Out[3]= -4 DotProduct[p, q] G[mu, Conjugate[mu]] +

- > 4 G[mu, Conjugate[mu]] Mass[p] Mass[q] +
- > 4 p[Conjugate[mu]] q[mu] + 4 p[mu] q[Conjugate[mu]]

 $-4(p \cdot q)g_{\mu\mu'} + 4g_{\mu\mu'}m_pm_q + 4p_{\mu'}q_{\mu} + 4p_{\mu}q_{\mu'}$

Conjugate[mu] (μ' is the new index needed on squaring the expression).

— ConvertToST: ConvertToST[expr] attempts to convert expr to SpinorS and SpinorT objects. Unless instructed otherwise, the program assumes every massive vector q is the sum of two massless vectors LightlikeVectorDecayedFrom[q, 1] and LightlikeVectorDecayedFrom[q, 2]. Example:

```
In[1]:= ConvertToST[
    SpinorUbar[p, Right]**Slash[q]**SpinorU[k, Right]]
```

Convert $\overline{u}_R(p) \not q u_R(k)$ to spinor techniques. p and k have previously been set massless.

Out[1]= -(SpinorS[p, LightlikeVectorDecayedFrom[q, 1]] *
> SpinorT[k, LightlikeVectorDecayedFrom[q, 1]]) -

```
> SpinorS[p, LightlikeVectorDecayedFrom[q, 2]] *
```

> SpinorT[k, LightlikeVectorDecayedFrom[q, 2]]

The result is $s(p, q_1)t(k, q_1) - s(p, q_2)t(k, q_2)$ where $q_1^2 = q_2^2 = 0$ and $q_1 + q_2 = q$ are assumed.

- Vertex: Vertex[line1, line2, ...] is the Feynman rule for a vertex. Each line is of the form particle or {particle, p} or {particle, p, indices} where particle is the particle type, p is the particle's momentum, and indexes are the appropriate Lorentz or gauge group indexes. Example:

In[1]:= Vertex[Electron, Electron, {ZBoson, p, mu}]

The eeZ vertex for Z boson carrying momentum p_{μ}

```
Out[1]= (-2 I Sqrt[Alpha] Sqrt[Pi] DiracGamma[mu] **
        -HelicityProjection[Left] 2
(-----+ Sin[ThetaW] )) /
```

(Cos[ThetaW] Sin[ThetaW]) >

>

 $\frac{-2i\sqrt{\alpha}\sqrt{\pi}}{\cos\theta_W\sin\theta_W}\gamma_\mu\left(-\frac{1}{2}P_L+\sin^2\theta_W\right)$

where α is the electromagnetic fine structure constant and θ_W is the Weinberg angle.

— CrossSection and DecayWidth: CrossSection[me2, q1->{q1x, q1y, q1z, e1}, q2->{q2x, q2y, q2z, e2}, outGoing] returns an expression for the phase space integral to be evaluated by EvaluatePhaseSpaceIntegral. me2 is the expression for the matrix element squared, {p1x, p1y, p1z, e1} and {p2x, p2y, p2z, e2} are the explicit four-vectors of the incoming particles, and outGoing specifies the order of phase-space evaluation as explained below. DecayWidth[me2, p -> {px, py, pz, e}, out-Going] similarly returns an expression for the phase space integral resulting in the decay width given matrix element squared me2 and initial momentum {px, py, pz, e}.

The formula used for cross-section calculations is

$$\sigma = \frac{(2\pi)^4}{2s} \frac{2|q_1|}{\sqrt{s}} \int |\mathcal{M}|^2 d\Phi_n(q_1 + q_2; p_1, \dots, p_n), \tag{3.9}$$

and for the decay width it is

$$\Gamma = \frac{(2\pi)^4}{2M} \int |\mathcal{M}|^2 d\Phi_n(P; p_1, \dots p_n).$$
(3.10)

Phase-space integration is performed by a recursive use of the relations

$$d\Phi_n(P; p_1, \dots, p_n) = d\Phi_{n-1}(P; p_{12}, p_3, \dots, p_n) d\Phi_2(p_{12}; p_1, p_2) (2\pi)^3 dm_{12}^2$$

= $d\Phi_{n-1}(P; p_{12}, p_3, \dots, p_n) \frac{1}{8(2\pi)^3} \frac{2|p_1|}{m_{12}} d\Omega_{12} dm_{12}^2,$
(3.11)

where $p_{12}^2 = m_{12}^2$ and Ω_{12} represents the direction of the 'decay' of the vector p_{12} to p_1 and p_2 in its center-of-mass frame. The factor $2|p_1|/m_{12}$ is given by

$$\frac{2|p_1|}{m_{12}} = \begin{cases} \frac{\left[\left(m_{12}^2 - (m_1 + m_2)^2\right)\left(m_{12}^2 - (m_1 - m_2)^2\right)\right]^{1/2}}{m_{12}^2} \\ \sqrt{1 - \frac{4m_1}{m_{12}}} \\ \sqrt{1 - \frac{4m_1}{m_{12}}} \\ 1 - \frac{m_1^2}{m_{12}^2} \\ 1 \end{cases} \qquad (m_1 = m_2) \\ (m_1 = m_2 = 0) \\ (m_1 = m_2 = 0). \end{cases}$$
(3.12)

The argument outGoing tells HIP how to build the phase space element $d\Phi_n$. It specifies both the order that the momenta $p_1 \cdots p_n$ are paired (eq. (3.11)) and, optionally, the symmetry of the individual two-body phase-space elements. By default, the complete angular integral over Ω_{12} is constructed. Often, due to the symmetry of the process, one can reduce the dimension of this integral (in the case of cylindrical symmetry), or eliminate it completely (in the case of spherical symmetry). This is done using the keywords Cylindrical and Spherical.

For example, let us consider the decay process $\mu^{-}(p) \rightarrow e^{-}(p_1)\overline{\nu}_e(p_2)\nu_{\mu}(p_3)$. If the μ is unpolarized, the decay process is spherically symmetric. The direction of ν_{μ} may be chosen arbitrarily. Once that is done, the direction of the electron with respect to the $(e^{-}\overline{\nu}_e)$ system has cylindrical symmetry about the ν_{μ} direction. The outGoing argument is given by Spherical[Cylindrical[p1, p2], p3]. If the μ is polarized, the spherical symmetry of the decay is reduced to a cylindrical symmetry about the polarization axis. outGoing is then given by Cylindrical[{p1, p2}, p3].

4. Examples

In this section we give two examples showing step by step how a HIP calculation is carried out. In the first example we compute the decay width of a muon in the process $\mu^- \to e^- \nu_{\mu} \overline{\nu}_e$ and with a non-zero electron mass. In the second example we express the matrix element for the process $e_R^+ e_R^- \to Z \to t\bar{t} \to W^+ W^- b\bar{b}$ as a spinor-technique expression.

4.1 $\Gamma(\mu^- \to e^- \nu_\mu \overline{\nu}_e)$

We use a low-energy approximation in which the W-propagator is a constant and is absorbed, along with the coupling constant g into the Fermi constant G_F . fig. 1 shows the single Feynman diagram for the process.

In[1]:= PrepareIndex[sig]

Instruct Mathematica to treat sig as an index.

In[2]:= SetMass[{pnub, 0}, {pe, me}, {pnu, 0}, {pmu, mmu}]



Figure 1. Feynman diagram for muon decay $\mu^- \rightarrow e^- \overline{\nu}_e \nu_\mu$.

Set the masses of the four external particles. The neutrinos (ν_{μ} and $\overline{\nu}_{e}$ carrying momenta pnu and pnub respectively) have zero mass. The electron (pe) is set to have mass me while the muon (pmu) is set to have mass mmu. Later on, these masses can be given numerical values.

In[3]:= matrixelement = 2 Sqrt[2] FermiGF *
 SpinorUbar[pnu] ** DiracGamma[sig] ** SpinorU[pmu, Left] *
 SpinorUbar[pe] ** DiracGamma[sig] ** SpinorV[pnub, Left];

$$\mathcal{M} = 2\sqrt{2}G_F\overline{u}(p_{\nu})\gamma_{\sigma}u_L(p_{\mu})\overline{u}(p_e)\gamma^{\sigma}u_L(p_{\overline{\nu}})$$

In[4]:= me2 = AbsSquared[matrixelement]/2;

Square the matrix-element using the Mathematica function AbsSquared.

We suppress the printing of the long intermediate result.

In[5]:= me2 = Contract[me2, {sig, Conjugate[sig]}] // Factor

Contract over the indices σ and σ' . // Factor instructs Mathematica to the factor the expression.

Out[5] = 128 FermiGF DotProduct[pe, pnu] DotProduct[pmu, pnub]

 $128G_F^2(p_e\cdot p_{\nu})(p_{\mu}\cdot p_{\overline{\nu}})$

Ask Mathematica to construct the phase-space integral to compute the decay-width. The expression Spherical[Cylindrical[pe, pnub], pnu] indicates a cylindrical symmetry in the phase-space integral over the pair $(e^-, \overline{\nu}_e)$ and a spherical symmetry over pair $(\nu_{\mu}, (e^-, \overline{\nu}_e))$ (here $(e^-, \overline{\nu}_e)$) is the combined system of e^- and $\overline{\nu}_e$.)

```
Out[7] = -PhaseSpaceIntegral-
```

In[8]:= width = EvaluatePhaseSpaceIntegral[width];

Evaluate the phase-space integral symbolically. Again we suppress the long intermediate result.

In[9]:= Factor[width /. me->x mmu /. Log[a_ b_] :> Log[a]+Log[b]] Use some Mathematica rules to tidy up the expression. We express the mass of the electron in terms of the mass of the muon $m_e = xm_{\mu}$ and combine logarithms using the rule $\log(ab) \rightarrow \log a + \log b$.

 $\begin{array}{c} -(\text{FermiGF} \ \text{mmu} \ (-1 + 8 \ \text{x} \ -8 \ \text{x} \ + \ \text{x} \ + \ 12 \ \text{x} \ \text{Log}[\text{x} \])) \\ \text{Out[10]} = & ----- \\ & 192 \ \text{Pi} \end{array}$

$$\Gamma(\mu^- \to e^- \nu_\mu \overline{\nu}_e) = \frac{G_F^2 m_\mu^5 (1 - 8x^2 + 8x^6 - x^8 - 12x^4 \log(x^2))}{192\pi^3}$$

4.2 MATRIX ELEMENT FOR $e_R^+ e_R^- \to Z \to t\bar{t} \to W^+ W^- b\bar{b}$ in spinor technique form

We convert the matrix element for the process $e_R^+ e_R^- \to Z \to t\bar{t} \to W^+ W^- b\bar{b}$ into the spinor technique form^[8]. The advantage of this form is that it can easily be used in a numerical calculation. The spinor technique objects s(p,q) and t(p,q)are defined in table 1. s and t have compact expressions which are easily evaluated numerically.

The relevant diagram is shown in fig. 2. The *Mathematica* computation follows. We have suppressed most of the intermediate output.

In[1]:= PrepareIndex[tau, mu, nu]

Instruct Mathematica to treat tau, mu and nu as indices.

In[2]:= SetMass[{p1, p2, pb, pbb, 0}, {pwp, pwm, Mass[WBoson]}]

Set the masses of the external particles. The e^- and e^+ have momenta p1 and p2 respectively, and are massless. The b and \overline{b} have momenta pb and pbb respectively, and are also massless. The W^+ and W^- have momenta pwp and pwm respectively, and mass Mass[WBoson].

In[3]:= SetDotProduct[{p1, p2, s/2}]

We set the value of $(p_1 \cdot p_2)$ to s/2.



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Figure 2. Feynman diagram for the process $e_R^+ e_R^- \to Z \to t\bar{t} \to W^+ W^- b\bar{b}$.

```
In[4]:= matrixElement =
    HeavyVectorPolarization[pwp, nu] *
    HeavyVectorPolarization[pwm, mu] *
    SpinorVbar[p2, Right] **
    Vertex[{Electron}, {Electron}, {ZBoson, p1+p2, tau}] **
    SpinorU[p1, Right] * Propagator[{ZBoson, p1+p2}] *
    SpinorUbar[pb, Left] **
    Vertex[{BottomQuark}, {TopQuark}, {WBoson, pwp, nu}] **
    FermionPropagator[TopQuark, pwp+pb] **
    Vertex[{TopQuark}, {TopQuark}, {ZBoson, p1+p2, tau}] **
    FermionPropagator[TopQuark, -pwm-pbb] **
    Vertex[{BottomQuark}, {TopQuark}, {WBoson, pwm, mu}] **
    SpinorV[pbb, Left]
```

$$\varepsilon^{\nu}(p_{W^{+}})\varepsilon^{\mu}(p_{W^{-}})\left\{\overline{v}_{R}(p_{2})\mathcal{V}^{\tau}(e,e,Z)u_{R}(p_{1})\right\}\Delta(Z,p_{1}+p_{2})$$

$$\times\left\{\overline{u}_{L}(p_{b})\mathcal{V}_{\nu}(b,t,W)\Delta(t,p_{b}+p_{+})\mathcal{V}_{\tau}(t,t,Z)\Delta(t,-p_{\overline{b}}-p_{-})\mathcal{V}_{\mu}(b,t,W)v_{L}(p_{\overline{b}})\right\},$$
where $\Delta(X,p)$ is the propagator for particle X with momentum p and

where $\Delta(X,p)$ is the propagator for particle X with momentum p and $\mathcal{V}_{\sigma}(A,B,C)$ is the vertex of the particles A, B and C with Lorentz index σ .

In the computation the massive four-vector pwp is replaced by the sum of two massless four-vectors r1 and r2. pwm is similarly replaced by r3+r4. In the spinor technique, r_1 and r_2 are the momenta of the massless fermion and antifermion into which the W decays. The final expression, thus, provides us with the angular distribution of the final state fermions.

In[6]:= convertedMatrixElement = ConvertToST[matrixElement] /. lightlikeVectorRules;

Convert the matrix element to spinor technique form and apply the simplifying rules. After some rearrangement, we get the following expression:

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$$-4i\alpha^{2}\pi V_{tb}^{2}s(p_{\overline{b}}, r_{4})t(p_{b}, r_{1}) \Big[4m_{t}^{2}s(p_{2}, r_{2})t(p_{1}, r_{3}) \\ + (4 - 3/\sin^{2}\theta_{W}) \big(s(p_{b}, r_{2})t(p_{1}, p_{b}) + s(r_{1}, r_{2})t(p_{1}, r_{1})\big) \\ \times \big(s(p_{2}, p_{\overline{b}})t(p_{\overline{b}}, r_{3}) - s(p_{2}, r_{4})t(r_{3}, r_{4})\big)\Big] \Big/ \\ \Big[\cos^{2}\theta_{W}m_{W}^{2} \big(2(p_{b} \cdot p_{+}) - m_{t}^{2} + m_{W}^{2}\big) \big(2(p_{\overline{b}} \cdot p_{-}) - m_{t}^{2} + m_{W}^{2}\big) \big(s - m_{Z}^{2}\big)\Big]$$

5. Conclusion and Outlook

We developed HIP as an aid in the calculation tree-level processes in high energy physics which would otherwise be much more difficult. HIP's main feature is in providing an environment within *Mathematica* in which one can refer to objects and perform operations that occur frequently in this field. One can use HIP interactively to assist with small calculations, or set it up to automatically perform massive 'symbol crunching'.

We have checked HIP against hand calculations of $e^+e^- \to W^+W^-$, $e\gamma \to W^+\nu$, both with arbitrary (C and P conserving) $W\gamma$ couplings, and of numerous simple electroweak processes. We also checked them against published results for $e^+e^- \to W^+W^-\gamma$, $e^+e^- \to W^+W^-Z$, $e^+e^- \to \gamma\gamma\gamma$ and $e^+e^- \to ZZZ$.

In the future, we hope to extend HIP's capabilities into performing loop integrals, calculating color factors and incorporating other techniques for symbolically calculating Feynman diagrams at the matrix element level. We also hope to translate HIP to other symbolic languages such as *Maple*, so as to maximize the group of its potential users.

HIP is available for distribution. The distribution includes the various component Mathematica packages, the online documentation as listed in the appendix, and several files containing sample calculations done by HIP.

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APPENDIX

In this appendix we include the complete listing of the objects defined in HIP, together with their usage messages. These usage messages are also available as on-line documentation. The functions are listed in alphabetical order.

AbsSquared::usage = "AbsSquared[expr] is the absolute value squared of expr."

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Boost::usage = "Boost[fv, rap, dir] gives a four-vector obtained by boosting fv by rapidity rap in the direction specified by dir. If dir is of the form {cth, phi}, cth is the Cosine of the angle between the direction and the z axis and phi is the angle between the direction's projection on the x-y plane and the x axis. If dir is of the form {x, y, z}, the direction is taken to be the direction of the vector {x, y, z}. Boost[fv, rap] boosts fv by amount rap in the z direction."

BoostAmount::usage = "BoostAmount[fv] gives a vector {rap, {x, y,
z}} such that Boost[{0, 0, 0, Mass[fv]}, rap, {x, y, z}] is fv."

CommutativeAllQ::usage = "CommutativeAllQ[expr] is True if expr does not have any non-commuting sub-expressions, and False otherwise."

CommutativeQ::usage = "CommutativeQ[x] is True if x is commutative (the default), and False if x is non-commutative."

Commutator::usage = "Commutator[a, b] is an equivalent expression to a**b, with a and b interchanged (e.g. Commutator[a, b] = b**a + [a, b])."

Commute::usage = "Commute[expr, {x, y}] commutes y from the right of x to the left of x everywhere in expr."

Contract::usage = "Contract[expr, index] contracts index in expr. All subexpressions of form G[index, _] or p_[index, _] can potentially be effected. Contract[expr, {index1, index2, ...}] does the contraction sequentially."

ConvertToChiralFermions::usage = "ConvertToChiralFermions[expr] attempts to convert all occurrences of DiracGamma5 and HelicityProjection in expr into products of chiral spinors."

ConvertToGamma5::usage = "ConvertToGamma5[expr] attempts to convert all occurrences of HelicityProjection and spinors of specific chirality into products involving DiracGamma5."

ConvertToMassless::usage = "ConvertToMassless is an option for ConvertToST."

ConvertToST::usage = "ConvertToST[expr] attempts to convert DiracGamma5 and HelicityProjection occurrences in expr to SpinorS and SpinorT objects." CrossSection::usage = "CrossSection[me2, p1 -> {p1x, p1y, p1z, e1}, p2 -> {p2x, p2y, p2z, e2}, outGoing] returns a PhaseSpaceIntegral. me2 is the matrix element squared, {p1x, p1y, p1z, e1} and {p2x, p2y, p2z, e2} are the explicit four-vectors of the incoming particles, and outGoing specifies the order of phase-space evaluation."

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Cylindrical::usage = "Cylindrical[p1, p2] indicates a cylindrically symmetric two body decay into p1 and p2 about the direction of motion of the decaying particle."

Decay::usage = "Decay[fv, dir, {m1, m2}] gives two four-vectors {fv1, fv2} such that fv1+fv2 is fv, Mass[fv1] is m1, Mass[fv2] is m2 and the direction of fv1 in the fv center-of-mass frame is given by dir. If dir is of the form {cth, phi}, cth is the Cosine of the angle between the direction and the z axis and phi is the angle between the direction's projection on the x-y plane and the x axis. If dir is of the form {x, y, z}, the direction is taken to be the direction of the vector {x, y, z}. Decay[fv, dir] is the same as Decay[fv, dir, {0, 0}]."

DecayWidth::usage = "DecayWidth[me2, p -> {px, py, pz, e}, outGoing] returns a PhaseSpaceIntegral. me2 is the matrix element squared, {px, py, pz, e} is explicit four-vector of the decaying particle, and outGoing specifies the order of phase-space evaluation."

DiracGamma5::usage = "DiracGamma5 is mainly meaningful for SpaceTimeDimension of 4. Some of its features will work in any even spacetime dimension."

DiracGamma::usage = "DiracGamma[mu] is the Dirac Gamma matrix. It is set to be non-commutative. Note that DotProduct[DiracGamma, p] is simplified to Slash[p]."

DiracGammaSize::usage = "DiracGammaSize is the dimension of the Dirac spinors. Its default value is 4."

DotProduct::usage = "DotProduct[p, q] is the Lorentz invariant dot product. It is evaluated explicitly (using the metric) if both p and q are lists of length SpaceTimeDimension."

DotProductRules::usage = "DotProductRules[{p, q, u}] returns {toRule, fromRule}. toRule takes DotProduct[p, q] to u and fromRule takes u to DotProduct[p, q]. DotProductRules[{p1, q1, u1}, {p2, q2, u2},...] returns {toRules, fromRules} where toRules are the rules for converting DotProducts to u's and fromRules are the rules for converting u's to DotProducts. DotProductRules[{p, q, u}, Mandelstam -> {s}] will define fromRules in terms of s."

Energy::usage = "Energy[fv] is the energy of fv."

Eps::usage = "Eps[a, b, c, ...] is the completely anti-symmetric object. Its arguments are automatically sorted alphabetically. Eps[..., p, ...] is the same as Eps[..., mu, ...] p[mu]."

EvaluatePhaseSpaceIntegral::usage =
"EvaluatePhaseSpaceIntegral[PhaseSpaceIntegral[...]] will
evaluate the Phase-space-integral."

ExpandSlash::usage = "ExpandSlash[expr] expands slash of sums to sums of slashes in expr."

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G::usage = "G[mu, nu] is the ordinary spacetime metric. mu and nu range from 1 to SpaceTimeDimension."

GammaTrace::usage = "GammaTrace[expr] is the trace (in spinor space) of expression. Note that no division by DiracGammaSize is carried out, ie GammaTrace[1] gives DiracGammaSize."

HeavyVectorPolarization::usage = "HeavyVectorPolarization[p, mu, pol] represents the polarization vector of a massive vector particle with momentum p, polarization pol and index mu."

HeavyVectorPolarization::usage = "HeavyVectorPolarization[p, mu, pol] represents the polarization vector of a massive vector particle with momentum p, polarization pol and index mu."

HelicityProjection::usage = "HelicityProjection[pol] is the helicity projection operator (1 +/- Gamma5)/2."

KobayashiMaskawa::usage = "KobayashiMaskawa[p1, p2] is the Kobayashi-Moskawa matrix element connecting particles p1 and p2."

LightlikeVectorDecayedFrom::usage = "LightlikeVectorDecayedFrom[p, n] represents a lightlike vector used for the polarizations of massive vector bosons. It obeys p = LightlikeVectorDecayedFrom[p,1] + LightlikeVectorDecayedFrom[p,2] where p is the momentum of the external massive vector boson."

LightlikeVectorNotCollinearWith::usage = "LightlikeVectorNotCollinearWith[p] represents a lightlike vector that is not collinear with p."

LongitudinalPolarization::usage = "LongitudinalPolarization[p] is a four-vector longitudinal polarization of the four-vector p."

Mandelstam::usage = "Mandelstam is an option for specifying a list of Mandelstam-type variables."

MandelstamRules::usage = "MandelstamRules[{p1, p2, p3, p4}, {m1, m2, m3, m4}, s, t, u] returns {toRules, fromRules}. toRules are rules for converting DotProducts of p1, p2, p3 and p4 to expressions containing s, t and u and the masses. fromRules are the rules for converting s, t and u to expressions containing the DotProducts of p1, p2, p3 and p4 and the masses. It sets p1, p2, p3, p4 to be on-shell with masses m1, m2, m3, m4 respectively."

Mass::usage = "Mass[p] is the mass of a four-vector p. It is
evaluated explicitly if p is a list of length SpaceTimeDimension.

MasslessVectorPolarization::usage = "MasslessVectorPolarization[p, mu, pol] represents the polarization vector of a massless vector particle with momentum p, polarization pol and index mu."

MasslessVectorPolarization::usage = "MasslessVectorPolarization[p, mu, pol] represents the polarization vector of a massless vector particle with momentum p, polarization pol and index mu."

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Momentum::usage = "Momentum[fv] is the three-component momentum of fv."

NEvaluatePhaseSpaceIntegral::usage =
"NEvaluatePhaseSpaceIntegral[PhaseSpaceIntegral[...]] will
evaluate the Phase-space-integral numerically."

NonCommutativeExpand::usage = "NonCommutativeExpand[expr] expands NonCommutativeMultiply of sums."

NonCommutativeFactor::usage = "NonCommutativeFactor[expr] attempts to factor sums of NonCommutativeMultiply's."

Opposite::usage = "Opposite[p] represents a four-momentum which has the same energy but opposite three momentum to the four-momentum p."

PerpendicularMomentum::usage = "PerpendicularMomentum[fv] gives the component of fv perpendicular to the z-axis."

PhaseSpaceIntegral::usage = "PhaseSpaceIntegral[integrand, {p1, (p2)}, decays, integrals, extmom] specifies a phase-space integral. Use EvaluatePhaseSpaceIntegral to evaluate the integral symbolically, and NEvaluatePhaseSpaceIntegral to evaluate it numerically."

PolarizationCombinations::usage = "PolarizationCombinations[me] returns a list of the possible polarization combinations of the initial and final particles for the matrix-element me over the polarizations which are not explicitly written in the matrix-element."

PrepareIndex::usage = "PrepareIndex[a, b, c, ...] makes future contractions involving a, b, c, ... work faster."

Propagator::usage = "Propagator[line] is the factor associated with the particle line, where line can be {type, p} or {type, p, mu, nu} where type is the type of particle, p is the momentum, and mu and nu are Lorentz indexes. Propagator[{type, p__}] is the same as Propagator[type][p]"

STToTraces::usage = "STToTraces[expr] attempts to convert products
of SpinorS's and SpinorT's in expr to traces using the rule
SpinorS[p1,p2]*SpinorT[p2, p3]...SpinorS[pn, p1] =
GammaTrace[Slash[p1]**Slash[p2]**...
Slash[pn]HelicityProjection[Right]]"

SetDotProduct::usage = "SetDotProduct[{p, q, u}] sets DotProduct[p, q] to be u. SetDotProduct[{p1, q1, u1}, {p2, q2, u2},...] sets DotProduct[p1, q1] to be u1, DotProduct[p2, q2] to be u2, etc."

SetMandelstam::usage = "SetMandelstam[{p1, p2, p3, p4}, {m1, m2, m3, m4}, s, t, u] sets p1, p2, p3, p4 to be on-shell with masses m1, m2, m3, m4 respectively and sets the DotProducts of p1, p2, p3 and p4 in terms of the Mandelstam variables s, t and u and the masses."

SetMass::usage = "SetMass[{p, m}] sets p to be a four-vector with invariant mass m. SetMass[{p1, p2,..., m}] sets p1, p2,.. to be four-vector with invariant mass m. SetMass[{p1, m1}, {p2, m2},...] applies SetMass to {p1, m1}, {p2, m2},... in turn."

SetNonCommutative::usage = "SetNonCommutative[a, b, c, ...] sets the symbols a, b, c, ... to be non-commutative."

SetReal::usage = "SetReal[a] sets Conjugate[a] = a. SetReal[a, b,...] applies SetReal to a, b,... in turn."

Slash::usage = "Slash[p] is the same as Contract[p[mu]DiracGamma[mu], mu]. Slash is set to be non-commutative."

•

SpaceDirection::usage = "SpaceDirection[fv] gives the pair {cth, phi} describing the direction of the momentum of fv."

SpaceTimeDimension::usage = "SpaceTimeDimension is what you'd
expect."

Spherical::usage = "Spherical[p1, p2] indicates a spherically symmetric two body decay into p1 and p2."

SpinorS::usage = "SpinorS[p, q] is the same as SpinorUbar[p, Right]**SpinorU[q, Left]. Both p and q have to be massless."

SpinorT::usage = "SpinorT[p, q] is the same as SpinorUbar[p, Left]**SpinorU[q, Right]. Both p and q have to be massless."

SpinorU::usage = "SpinorU[p] is a spinor object of a four-vector p.
SpinorU[p, pol] is a spinor object of particular polarization."

SpinorUbar::usage = "SpinorUbar[p] is a spinor object of a
four-vector p. SpinorUbar[p, pol] is a spinor object of particular
polarization."

SpinorV::usage = "SpinorV[p] is a spinor object of an anti-particle
of four-vector p. SpinorV[p, pol] is a spinor object of particular
polarization."

SpinorVbar::usage = "SpinorVbar[p] is a spinor object of an anti-particle of four-vector p. SpinorVbar[p, pol] is a spinor object of particular polarization."

Vertex::usage = "Vertex[line1, line2, ...] is the Feynman rule for the vertex. Each line is of the form {particle} or {particle, p} or {particle, p, indexes} where particle is the particle type, p is the particle's momentum, and indexes are the appropriate Lorentz or gauge group indexes."

ZAxis::usage = "ZAxis is an option used with functions receiving a direction as an argument. It specifies the axis with respect to which the direction is given."

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Fig. 1



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Fig. 2