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# Guide to *FeynCalc* 1.0

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# Contents



# 0 Introduction to FeynCalc

*FeynCalc* 1.0 is a *Mathematica* 2.0 package for algebraic calculations in high energy physics. The basic idea of *FeynCalc* is to provide convenient tools for radiative corrections in the standard model. The input for *FeynCalc*, the analytical expressions for the diagrams, can be entered by hand or can be taken directly from the output of another package, *FeynArts* [1], which produces all diagrams for a given process. The user can provide certain additional information about the process under consideration, *i.e.*, the kinematics and the choice of the standard matrix elements may be defined. Once this is done, *FeynCalc* performs the algebraic calculations like tensor algebra, tensor integral decomposition and reduction, yielding a polynomial in standard matrix elements, special functions, kinematical variables and scalars. With an appropriate option setting this polynomial is directly converted to an optimized Fortran file.

The computational methods and algorithms implemented in *FeynCalc* can be found in [2] and [3], where further references are given.

*FeynCalc* provides also calculator-type like features. You enter a Dirac trace in a very similar notation you use by hand and get back an answer, suitable for further manipulation with *Mathematica* or ready to translate into Fortran. Apart from these basic operations like contraction of tensors, simplification of products of Dirac matrices and trace calculation no attempt has been made to provide tools for tree-level calculations in *FeynCalc* 1.0. For this purpose a lot of programs exist, among them another *Mathematica* package, HIP [4], developed at SLAC.

A free copy of *FeynCalc* may be obtained from the author. You may redistribute *FeynCalc* with no charge, but please do not change the program. *FeynCalc* 1.0 is the first public release of a set of programs which were developed and used at the University of Würzburg since 1988. Though a lot of testing of the code has been done, there is absolutely no claim that *FeynCalc* is bug-free. You should be sceptical about the results, and when you are sure the program returned a wrong answer, you are encouraged to send email to rolfm@wri.com or mertig@vax.rz.uni-wuerzburg.dbp.de . Suggestions and improvements are also welcome.

The explanations in this preliminary manual are rather short. The user is assumed to be skilled in high energy physics and *Mathematica*. It is strongly recommended to study Sections 1.2, 1.4, 1.7, 1.8, 2.1 - 2.4 and 3.3 of the book *Mathematica*, Second Edition, by Stephen Wolfram.

In *FeynCalc* 1.0 no facilities for the numeric evaluation of Feynman diagrams are provided. Planned extensions of *FeynCalc* are tree level and two-loop facilities, the use of the program *MathLink* for the symbolic-numeric interface, and links to other programs for the evaluation of Feynman diagrams.

## ■ 0.1 Acknowledgments

I would like to thank M. Böhm and A. Denner for their support and helpful discussions, which contributed considerably to the algorithms implemented in *FeynCalc*. I am grateful to J. Küblbeck and H. Eck for providing *FeynArts*, which was used for the graphics. Testing of the results of *FeynCalc* has been done by A. Denner, S. Dittmaier, J. Küblbeck, G. Weiglein and T. Sack.

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## ■ 0.2 Installation

The file `FeynCalc.m` should be put in a directory called `/usr/users/mydir/FeynCalc`. In this directory you have to open a subdirectory `Code` and put the subpackages `GellMann.m`, `General.m`, `Main.m`, `OneLoop.m` and `PaVe.m` into this directory (`/usr/users/mydir/FeynCalc/Code`).

From the directory `/usr/users/mydir/FeynCalc` you can start a *Mathematica* session and load *FeynCalc* by typing `<<FeynCalc.m`.

If you want to work from another directory, you should add the following command in the `init.m` (or `FeynCalc.m`) file:

```
$Path = Append[$Path, "/usr/users/mydir"]
```

If you add the command `<</usr/users/mydir/FeynCalc/FeynCalc.m` to your initialization file `init.m`, *FeynCalc* will be loaded automatically upon invoking *Mathematica*. Throughout this manual *FeynCalc* is always automatically loaded in this way.

# 1 Input Functions

The *FeynCalc* syntax for metric tensors, four-vectors, Dirac matrices, etc., is such that the positioning of the arguments automatically defines the nature of the variables. You can only enter algebraic objects; there is no possibility of working with explicit components of vectors or with specific Dirac matrices like  $\gamma_0$  or  $\gamma_2$ . Note that no declaration of Lorentz indices or four-vectors has to be made. Covariant and contravariant indices are treated on an equal basis, assuming summation over repeated indices. Since explicit components of tensors will not be used at all, this convention causes no problems. The ambiguity in the sign of the Levi-Civita tensor can be fixed by an option of the function `DiracTrace` and `EpsChisholm`, see page 17.

The input functions are macros that translate into the internal representation of *FeynCalc*. But these sometimes lengthy internal representations are difficult to recognize when working interactively. Therefore as a default in *FeynCalc* the global *Mathematica* variable `$PrePrint` is set to `FeynCalcForm`, forcing the display to be in a more readable manner. It is important to realize that the display given by `FeynCalcForm` is always different from the input syntax. Thus you may not substitute, for example, a scalar product in the result of a trace calculation by copying the form you see on the screen.

The internal structure of *FeynCalc*, which you may explore by clearing `$PrePrint`, is explained briefly in Section 4 and in the Reference Guide. Usually it is not necessary to know it. Thus there are three layers of *FeynCalc*: the input set of functions, a `FeynCalcForm` representation of those and the internal structure.

In order to do calculations in the framework of  $D$ -dimensional regularization, *FeynCalc* can be used for calculations in 4,  $D$  or  $D - 4$  dimensions. The dimension of metric tensors, Dirac matrices and four-vectors is specified by setting the option `Dimension` of the corresponding input functions. The default dimension is four. If *FeynCalc* is used for interactive  $D$ -dimensional calculations you should change the option of the input functions. Note that Lorentz indices, momenta and Dirac matrices carry their dimension locally, *i.e.*, for calculating a Dirac trace in  $D$  dimensions the dimension of the Dirac matrices has to be set accordingly, thus there is no need of a dimension option for the function `DiracTrace`.

For the automatic calculation of one-loop amplitudes the input for the function `OneLoop` may always be given in four dimensions. `OneLoop` extends the amplitude to  $D$  dimensions automatically.

## 1.1 Entering Tensors and Scalar Products

<code>FourVector[p, mu]</code>	four-vector $p^\mu$
<code>LeviCivita[mu, nu, ro, si]</code>	Levi-Civita tensor $\epsilon^{\mu\nu\rho\sigma}$
<code>MetricTensor[mu, nu]</code>	metric tensor $g^{\mu\nu}$
<code>ScalarProduct[p, q]</code>	$p \cdot q$

Basic input functions.

Levi-Civita tensors and polarization vectors are defined in *FeynCalc* in four dimensions only. For `MetricTensor`, `FourVector` and `ScalarProduct` other dimensions may be specified with the option `Dimension`.

<i>option name</i>	<i>default value</i>	
<code>Dimension</code>	4	space-time dimension

Option for `FourVector`, `MetricTensor` and `ScalarProduct`.

The following examples show the input syntax and the result, which is displayed by *FeynCalc* in an abbreviated form with the default setting of `$PrePrint` to `FeynCalcForm`. Thus `FeynCalcForm` is similar to the *Mathematica* `OutputForm`, but specialized to the new objects defined in *FeynCalc*.

Enter a scalar product $p \cdot q$ .	<code>In[1]:= ScalarProduct[p, q]</code>
This is the <code>FeynCalcForm</code> .	<code>Out[1]= p . q</code>
This is the input for $g^{\mu\nu}$ .	<code>In[2]:= MetricTensor[mu, nu]</code> <code>Out[2]= g[mu, nu]</code>
A metric tensor $g^{\alpha\beta}$ in $D$ dimensions is entered in this way. The dimension is displayed as an index.	<code>In[3]:= MetricTensor[al, be, Dimension -&gt; D]</code> <code>Out[3]= g [al, be]</code> $D$
Changing $D$ to 4 recovers the default <code>FeynCalcForm</code> for a four-dimensional $g^{\alpha\beta}$ .	<code>In[4]:= % /. D -&gt; 4</code> <code>Out[4]= g[al, be]</code>
In four dimensions $g_{\alpha}^{\alpha} = 4$ .	<code>In[5]:= MetricTensor[al, al]</code> <code>Out[5]= 4</code>
While in $D$ dimensions $g_{\alpha}^{\alpha} = D$ .	<code>In[6]:= MetricTensor[al, al, Dimension -&gt; D]</code> <code>Out[6]= D</code>
Enter a four-dimensional $p^{\mu}$ .	<code>In[7]:= FourVector[p, mu]</code> <code>Out[7]= p[mu]</code>
You may also enter a sum of four-vectors: $(p - 2q)^{\mu}$ .	<code>In[8]:= FourVector[p - 2 q, mu]</code> <code>Out[8]= (p - 2 q)[mu]</code>
Providing $(2q - p)^{\mu}$ yields $-(p - 2q)^{\mu}$ .	<code>In[9]:= FourVector[2 q - p, mu]</code> <code>Out[9]= -(p - 2 q)[mu]</code>
In general numerical factors are pulled out in a standard way from <code>FourVector</code> .	<code>In[10]:= FourVector[2 p, mu]</code> <code>Out[10]= 2 p[mu]</code>
This is a $D$ -dimensional $q^{\alpha}$ .	<code>In[11]:= FourVector[q, al, Dimension -&gt; D]</code> <code>Out[11]= q [al]</code> $D$
A polarization vector $\varepsilon_{\mu}(k)$ is a special four-vector.	<code>In[12]:= PolarizationVector[k, mu]</code> <code>Out[12]= ep[k, mu]</code>
This is how to enter a $\varepsilon_{\mu}^*(k)$ .	<code>In[13]:= Conjugate[PolarizationVector[k, mu]]</code> <code>Out[13]= ep[*][k, mu]</code>

Polarization vectors are entered in a special way. The transversality condition  $\varepsilon(k) \cdot k = 0$  is automatically fulfilled, see page 13.

<code>PolarizationVector[k, mu]</code>	$\varepsilon_{\mu}(k)$
<code>Conjugate[PolarizationVector[k, mu]]</code>	$\varepsilon_{\mu}^*(k)$
<code>Polarization[k]</code>	$\varepsilon(k)$

The input for a polarization vector and polarization.



## ■ 1.2 Entering Dirac Matrices

<code>DiracMatrix[mu, nu, ...]</code>	Dirac matrices $\gamma^\mu \gamma^\nu \dots$
<code>DiracSlash[p, q, ...]</code>	Feynman slashes $\not{p} \not{q} \dots$
<code>DiracMatrix[5]</code>	$\gamma^5$
<code>ChiralityProjector[+1]</code>	$\omega_+ = (1 + \gamma^5)/2$
<code>DiracMatrix[6]</code>	$\gamma^6 = (1 + \gamma^5)/2$
<code>ChiralityProjector[-1]</code>	$\omega_- = (1 - \gamma^5)/2$
<code>DiracMatrix[7]</code>	$\gamma^7 = (1 - \gamma^5)/2$

Various input functions for Dirac matrices.

A Dirac matrix  $\gamma^\mu$  is represented by `DiracMatrix[mu]`. For  $\not{p} = p_\mu \gamma^\mu$  you may use `DiracSlash[p]`. Products of Dirac matrices or slashes can either be entered by adding subsequent arguments, *i.e.*, `DiracMatrix[mu, nu, ...]` and `DiracSlash[p, q, ...]`, or by multiplying the `DiracMatrix` and `DiracSlash` with the *Mathematica* Dot `“.”`.

The *Mathematica* Dot `“.”` is always used in the input as noncommutative multiplication operator for the objects `DiracMatrix`, `DiracSlash`, `Spinor`, `LeptonSpinor`, `QuarkSpinor` and `GellMannMatrix`. The `“.”` may also be used as a delimiter instead of the `“,”` within the functions `DiracMatrix`, `DiracSlash` and `GellMannMatrix`.

In the output with `FeynCalcForm` the `“.”` as noncommutative multiplication operator is suppressed.

This is how you enter  $(\not{p} + \not{q} + m) \gamma^\mu$ .

```
In[1]:= (DiracSlash[p + q] + m) . DiracMatrix[mu]
Out[1]= (gs[p + q] + m) ga[mu]
```

Here is  $\gamma^\rho \gamma^5 \gamma^\mu \gamma^6 \gamma^\rho \gamma^7$ .

```
In[2]:= DiracMatrix[ro, 5, mu, 6, al, 7]
Out[2]= ga[ro] ga[5] ga[mu] ga[6] ga[al] ga[7]
```

This is an alternative input variety. Note that the spaces around the `“.”` are essential when  $\gamma^5$ ,  $\gamma^6$  and  $\gamma^7$  are involved.

```
In[3]:= DiracMatrix[mu . 5 . nu . 6 . ro . 7]
Out[3]= ga[mu] ga[5] ga[nu] ga[6] ga[ro] ga[7]
```

This is a four-dimensional product:  
 $2\not{b} \not{a} (\not{d} - \not{c}) (6\not{q} - 3\not{p})$ . `DiracSlash` pulls common numerical factors out.

```
In[4]:= DiracSlash[2 b, a, 2 (d - c), 6 q - 3 p]
Out[4]= 12 gs[b] gs[a] gs[c - d] gs[p - 2 q]
```

Here is an alternative input with `Dot`. Commutative products have to be grouped with parentheses here.

```
In[5]:= DiracSlash[(2 b) . a . (2 (d - c)) . (6 q - 3 p)]
Out[5]= 12 gs[b] gs[a] gs[c - d] gs[p - 2 q]
```

Slashed polarization vectors  $\not{\epsilon}(k)$  are entered in this way.

```
In[6]:= DiracSlash[Polarization[k]]
Out[6]= gs[ep[k]]
```

<i>option name</i>	<i>default value</i>
Dimension	4 space-time dimension

Option for `DiracMatrix` and `DiracSlash`.

As settings for Dimension 4, a *Mathematica* Symbol `dim`, or `dim - 4` are possible.

Entering of a D-dimensional  $\gamma^\mu$ .

```
In[7]:= DiracMatrix[mu, Dimension -> D]
```

The dimension is displayed as an index in the FeynCalcForm.

```
Out[7]= ga [mu]
         D
```

This is  $\gamma^\mu \gamma^\mu$  in  $D - 4$  dimensions.

```
In[8]:= DiracMatrix[mu, mu, Dimension -> D - 4]
```

```
Out[8]= ga [mu] ga [mu]
         D-4   D-4
```

### ■ 1.3 Entering Gell-Mann Matrices and SU(3) Structure Constants

*FeynCalc* is not a package especially designed for QCD calculations. But some basic features for Gell-Mann matrices are included.

<code>GellMannMatrix[a, b, ...]</code>	Gell-Mann matrices $\lambda_a \lambda_b \dots$
<code>SU3Delta[a, b]</code>	Kronecker $\delta_{ab}$ with color indices $a, b$
<code>SU3F[a, b, c]</code>	structure constants $f_{abc}$ of SU(3)

Input of Gell-Mann matrices,  $\delta_{ab}$  and SU(3) structure constants.

For noncommutative multiplication use the *Mathematica* Dot “.”.

Products of `GellMannMatrix` have to be separated by a “.”.

```
In[9]:= GellMannMatrix[b] . GellMannMatrix[a]
```

```
Out[9]= la[b] la[a]
```

This is how you enter  $\lambda_c \lambda_b \lambda_a \delta_{de}$ .

```
In[10]:= GellMannMatrix[c, b, a] SU3Delta[d, e]
```

```
Out[10]= d[d, e] la[c] la[b] la[a]
```

The default for the  $f_{abc}$  is:  
 $i/4 [tr(\lambda_a \lambda_c \lambda_b) - tr(\lambda_a \lambda_b \lambda_c)]$ .

```
In[11]:= SU3F[a, b, c]
```

```
Out[11]= I
         4 (-tr[la[a] la[b] la[c]] + tr[la[a] la[c] la[b]])
```

The trace in the output is the FeynCalcForm for `GellMannTrace`.

<i>option name</i>	<i>default value</i>	
<code>SU3FToTraces</code>	True	whether to replace SU3F by traces

An option for the structure constants SU3F.

The structure constants are totally antisymmetric:  $f_{acb} = -f_{abc}$ .

```
In[12]:= SU3F[a, c, b, SU3FToTraces -> False]
```

```
Out[12]= -f[a, b, c]
```

$f_{aab} = 0$ .

```
In[13]:= SU3F[a, a, b]
```

```
Out[13]= 0
```

### ■ 1.4 Entering Spinors

In *FeynCalc* 1.0 two different types of spinors can be entered. For leptonic spinors you can use `LeptonSpinor` or `Spinor`. The convention for a quark spinor is to suppress the color index. The only difference between `QuarkSpinor` and `LeptonSpinor` is that Gell-Mann matrices do not commute with quark spinors.

<code>Spinor[p, m]</code>	$u(p, m)$ or $\bar{u}(p, m)$ for leptons
<code>Spinor[-p, m]</code>	$v(p, m)$ or $\bar{v}(p, m)$ for leptons
<code>LeptonSpinor[p, m]</code>	$u(p, m)$ or $\bar{u}(p, m)$ for leptons
<code>LeptonSpinor[-p, m]</code>	$v(p, m)$ or $\bar{v}(p, m)$ for leptons
<code>QuarkSpinor[p, m]</code>	$u(p, m)$ or $\bar{u}(p, m)$ for quarks
<code>QuarkSpinor[-p, m]</code>	$v(p, m)$ or $\bar{v}(p, m)$ for quarks

The functions for leptonic and hadronic spinors.

Which of the Dirac spinors  $u, v$  and the conjugate spinors  $\bar{u}, \bar{v}$  are understood, depends on the position of the spinors in the Dirac chain and the sign of the momentum argument.

This is a leptonic spinor  $u(p, m)$  or  $\bar{u}(p, m)$ .

```
In[14]:= Spinor[p, m]
Out[14]= u[p, m]
```

The minus sign of the first argument indicates that  $v(p, m)$  or  $\bar{v}(p, m)$  is understood.

```
In[15]:= Spinor[-p, m]
Out[15]= v[p, m]
```

This is  $\bar{u}(p, m) \not{p}$ .

```
In[16]:= Spinor[p, m] . DiracSlash[p]
Out[16]= u[p, m] gs[p]
```

Since  $\not{p}$  is now left of the spinor,  $\not{p} u(p, m)$  is understood.

```
In[17]:= DiracSlash[p] . QuarkSpinor[p, m]
Out[17]= gs[p] u[p, m]
```

This is  $\bar{v}(p, m) \not{q} \not{p}$ .

```
In[18]:= Spinor[-p, m] . DiracSlash[q, p]
Out[18]= v[p, m] gs[q] gs[p]
```

Here we have  $\not{p} v(p, m)$ .

```
In[19]:= DiracSlash[p] . QuarkSpinor[-p, m]
Out[19]= gs[p] v[p, m]
```

Enter a spinor obeying the massless Dirac equation. The `FeynCalcForm` suppresses the 0.

```
In[20]:= Spinor[p, 0]
Out[20]= u[p]
```

You may also omit the second argument for a massless spinor.

```
In[21]:= Spinor[p]
Out[21]= u[p]
```

## ■ 1.5 Entering Denominators of Propagators

The denominators of propagators are entered in a special way. Each one-loop Feynman amplitude has factors of the form

$$df = \frac{1}{[q^2 - m_0^2][(q + p_1)^2 - m_1^2][(q + p_2)^2 - m_2^2] \dots}$$

with  $q$  as loop momentum and the  $p_i$  denoting linear combinations of the external four-vectors.

```
FeynAmpDenominator[PropagatorDenominator[q, m_0],
PropagatorDenominator[q + p, m_1], ...]
1/([q^2 - m_0^2][(q + p)^2 - m_1^2]) ...
```

Entering denominators of propagators.

This is  $1/([q^2 - m_1^2][(q + p)^2 - m_2^2])$ .

The `FeynCalcForm` imitates the usual notation.

```
In[22]:= FeynAmpDenominator[ PropagatorDenominator[q, m1],
                               PropagatorDenominator[q + p, m2]]
```

$$Out[22]= \frac{1}{(q^2 - m1^2) ((p + q)^2 - m2^2)}$$

## ■ 1.6 Entering Small Variables

For radiative corrections in the standard model fermionic masses often can be neglected with respect to the gauge boson masses. Since however some of the Passarino-Veltman integrals may be infrared divergent, the fermionic (and evtl. photonic) small masses must remain as arguments of them and cannot be set to 0. In order to achieve this behavior these small masses have to be entered with head `Small`. The basic idea is that every variable with head `Small` evaluates to 0, unless it is an argument of a scalar integral.

You can avoid to write explicitly the head `Small` around a mass by using the option `SmallVariables` of the function `OneLoop`.

<code>Small[m]</code> head of a small mass $m, m \ll M$
---

A head for small variables.

In spinors a mass with head `Small` is replaced by 0.

```
In[23]:= Spinor[p, Small[m]]
Out[23]= u[p]
```

## 2 Elementary Calculations

You can use *FeynCalc* for basic calculations like Trace evaluations. In the following examples *FeynCalc* was loaded automatically by including `<<FeynCalc.m` in the `init.m` file. The output is given in `FeynCalcForm`, i.e., the default of `$PrePrint` is `FeynCalcForm`.

### ■ 2.1 Contraction of Metric Tensors, Four-Vectors and Levi-Civita Tensors

The `Contract` function contracts equal Lorentz indices, if at least one belongs to a metric tensor, a four-vector or a Levi-Civita tensor.

<code>Contract [expr]</code>	contracts double Lorentz indices
------------------------------	----------------------------------

The function for contraction of tensors.

Contract $g^{\alpha\beta} p^\beta$ .	<code>In[1]:= Contract[MetricTensor[a, b] FourVector[p, b]]</code> <code>Out[1]= p[a]</code>
Contract $q^\alpha (p - q)^\alpha$ .	<code>In[2]:= Contract[FourVector[q, a] FourVector[p - q, a]]</code> <code>Out[2]= (p - q) . q</code>
This contracts $g^{\alpha\beta} \gamma^\alpha$ .	<code>In[3]:= Contract[MetricTensor[a], DiracMatrix[a]]</code> <code>Out[3]= g[a]</code>
Contracting $q^\alpha \gamma^\alpha$ yields a Feynman slash.	<code>In[4]:= Contract[FourVector[q, a] DiracMatrix[a]]</code> <code>Out[4]= gs[q]</code>
Contracting $\varepsilon^{\mu\nu\rho\sigma} p^\sigma$ gives $\varepsilon^{\mu\nu\rho\sigma}$ .	<code>In[5]:= Contract[LeviCivita[mu, nu, ro, si] FourVector[p, si]]</code> <code>Out[5]= eps[mu, nu, ro, p]</code>
The contraction of $\varepsilon^{\alpha\nu\rho\sigma} \varepsilon^{\beta\nu\rho\sigma}$ yields $-6g^{\alpha\beta}$ .	<code>In[6]:= Contract[LeviCivita[a, n, r, s] LeviCivita[b, n, r, s],</code> <code style="padding-left: 40px;">EpsContract -&gt; True ]</code> <code>Out[6]= -6 g[a, b]</code>

<i>option name</i>	<i>default value</i>	
<code>EpsContract</code>	<code>False</code>	whether to contract Levi-Civita Eps
<code>Expanding</code>	<code>True</code>	whether to expand the input
<code>Factoring</code>	<code>False</code>	whether to factor canonically

Options for `Contract`.

Contracting only $g^{\alpha\sigma} p^\alpha p^\sigma$ in $g^{\alpha\sigma} p^\alpha p^\sigma (q^\beta + r^\beta) (p^\beta - s^\beta)$ .	<code>In[7]:= Contract[ MetricTensor[a], si] *</code> <code style="padding-left: 40px;">FourVector[p, a] FourVector[p, si] *</code> <code style="padding-left: 80px;">( FourVector[q, be] + FourVector[r, be] ) *</code> <code style="padding-left: 80px;">( FourVector[p, be] - FourVector[q, be] ),</code> <code style="padding-left: 40px;">Expanding -&gt; False ]</code> <code>Out[7]= p.p (p[be] - q[be]) (q[be] + r[be])</code>
<i>FeynCalc</i> uses the transversality condition $(k^\mu \cdot \varepsilon^\mu(k)) = 0$ for polarization vectors.	<code>In[8]:= Contract[FourVector[k, mu] PolarizationVector[k, mu]]</code> <code>Out[8]= 0</code>

## ■ 2.2 Simplification of Scalar Products and Four-Vectors

<code>ExpandScalarProduct[expr]</code>	expand scalar products and four-vectors in <i>expr</i>
--	--

A function for expansion of scalar products and four-vectors.

As an example expand  $(a + b) \cdot (c - 2d)$ .

```
In[1]:= ExpandScalarProduct[ScalarProduct[a + b, c - 2 d]]
Out[1]= a.c - 2 a.d + b.c - 2 b.d
```

Consider again  $q^\alpha (p - q)^\alpha$ .

```
In[2]:= Contract[FourVector[q, al] FourVector[p - q, al]]
Out[2]= (p - q) . q
```

This expands the scalar products.

```
In[3]:= ExpandScalarProduct[%]
Out[3]= p.q - q.q
```

This is how you can substitute  $q^2 \rightarrow 0$  afterwards.

```
In[4]:= % /. ScalarProduct[q, q] -> 0
Out[4]= p.q
```

Instead of substituting scalar products at the end of the calculation another possibility is to assign special values for scalar products first. These special values are inserted immediately whenever possible during the calculation.

Set  $q^2 = 0$  before a calculation.

```
In[5]:= ScalarProduct[q, q] = 0
Out[5]= 0
```

Expanding the result of the contraction  $q^\alpha (p - q)^\alpha$  now yields  $(p \cdot q)$ .

```
In[6]:= ExpandScalarProduct[Out[2]]
Out[6]= p.q
```

This expands  $(a + b)^\mu$  to  $a^\mu + b^\mu$ .

```
In[7]:= ExpandScalarProduct[FourVector[a + b, mu]]
Out[7]= a[mu] + b[mu]
```

## ■ 2.3 Simplification of Products of Dirac Matrices and Spinors

For the simplification of noncommutative products of Dirac matrices and spinors, two functions are provided: `DiracOrder`, which orders products of Dirac matrices in a canonical way, and `DiracSimplify`, which contracts Dirac matrices with equal indices, moves  $\gamma^5, \gamma^6$  and  $\gamma^7$  to the right end applies the Dirac equation and expands noncommutative products, see section 4.4.11. The Dirac matrices in the result of `DiracSimplify` are only ordered in a canonical way if they are between spinors.

<code>DiracOrder[expr]</code>	alphabetical ordering of Dirac matrices
<code>DiracOrder[expr, {a, b, ...}]</code>	ordering according to <i>a, b, ...</i>
<code>DiracSimplify[expr]</code>	contract all Lorentz indices and simplify

Simplification functions for Dirac matrices and spinors.

Both functions take as *expr* either subsequent `DiracMatrix` or `DiracSlash` separated by "," or any expression with "." as the noncommutative multiplication operator between Dirac matrices or Dirac slashes.

Order the product  $\gamma^\beta \gamma^\alpha \rightarrow 2g^{\alpha\beta} - \gamma^\alpha \gamma^\beta$ .

```
In[1]:= DiracOrder[DiracMatrix[be, al]]
Out[1]= 2 g[al, be] - ga[al] ga[be]
```

Anticommutate back to  $\gamma^\beta \gamma^\alpha$ .

```
In[2]:= DiracOrder[%, {be, al}]
Out[2]= ga[be] ga[al]
```

Simplifications like  $\gamma^\mu \gamma^\mu \not{p} \not{p} = 4p^2$  are built in.

```
In[3]:= DiracOrder[DiracMatrix[mu, mu], DiracSlash[p, p]]
Out[3]= 4 p.p
```

$\gamma^\alpha \gamma^\mu \gamma^\alpha = (2 - D) \gamma^\mu$  in  $D$  dimensions.

```
In[4]:= DiracOrder[DiracMatrix[a, m, a, Dimension -> D]]
Out[4]= (2 - D) ga[m]
```

$-\not{p} \not{q} \not{p} = \not{q} p^2 - 2\not{p} (p \cdot q)$ .

```
In[5]:= DiracOrder[DiracSlash[-p, q, p]]
Out[5]= gs[q] p.p - 2 gs[p] p.q
```

Basically `DiracOrder` is just the implementation of the anticommutator relation  $\{\gamma^\mu, \gamma^\nu\} = 2g^{\mu\nu}$ .

For calculations involving  $\gamma^5$  you can use the function `DiracSimplify`. All  $\gamma^5, \gamma^6$  and  $\gamma^7$  are moved to the right end of the product of Dirac matrices; see section 4.4.11 for the treatment of  $\gamma^5$  in  $D$  dimensions. `DiracSimplify` applies the Dirac equation. The result of `DiracSimplify` is not canonically ordered.

This is  $\gamma^\mu \gamma^\mu = D$ .

```
In[1]:= DiracSimplify[DiracMatrix[mu, mu, Dimension -> D]]
Out[1]= D
```

Here the Kahane algorithm is used.

$\gamma^\mu \gamma^\nu \gamma^\rho \gamma^\sigma \gamma^\mu = -2\gamma^\sigma \gamma^\rho \gamma^\nu$ .

```
In[2]:= DiracSimplify[DiracMatrix[mu, nu, ro, si, mu]]
Out[2]= -2 ga[si] ga[ro] ga[nu]
```

Kahane also gives this identity:

$\frac{1}{2} \gamma^\mu \gamma^\alpha \gamma^\beta \gamma^\gamma \gamma^\delta \gamma^\mu =$   
 $\gamma^\gamma \gamma^\beta \gamma^\alpha \gamma^\delta + \gamma^\delta \gamma^\alpha \gamma^\beta \gamma^\gamma$ .

```
In[3]:= DiracSimplify[1/2 DiracMatrix[mu, a, b, c, d, mu]]
Out[3]= ga[c] ga[b] ga[a] ga[d] + ga[d] ga[a] ga[b] ga[c]
```

This is

$\not{p} (m - \not{p}) \not{p} = \not{p} p^2 + p^2 m - 2\not{p} (p \cdot q)$ .

```
In[4]:= DiracSimplify[DiracSlash[p], DiracSlash[-q] + m,
DiracSlash[p]]
Out[4]= gs[q] p.p + m p.p - 2 gs[p] p.q
```

This is  $\gamma^5 \gamma^\mu = -\gamma^\mu \gamma^5$ .

```
In[5]:= DiracSimplify[DiracMatrix[5], DiracMatrix[mu]]
Out[5]= -ga[mu] ga[5]
```

$\gamma^6 \gamma^\nu \gamma^7 \gamma^\mu = \gamma^\nu \gamma^\mu \gamma^6$ .

```
In[6]:= DiracSimplify[DiracMatrix[6, nu, 7, mu]]
Out[6]= ga[nu] ga[mu] ga[6]
```

This is  $(\not{p} - m) u(p) = 0$ .

```
In[7]:= DiracSimplify[(DiracSlash[p] - m) . Spinor[p, m]]
Out[7]= 0
```

Here is the Dirac equation for  $v(p)$ :

$(\not{p} + m) v(p) = 0$ .

```
In[8]:= DiracSimplify[(DiracSlash[p] + m) . Spinor[-p, m]]
Out[8]= 0
```

For the conjugate spinor:  $\bar{u}(\not{p} - m) = 0$ .

```
In[9]:= DiracSimplify[Spinor[p, m] . (DiracSlash[p] - m)]
Out[9]= 0
```

This is  $\bar{v} \not{q} (\not{p} - m) = 2\bar{v} p \cdot q$ .

```
In[10]:= DiracSimplify[Spinor[-p, m] . DiracSlash[q] .
(DiracSlash[p] - m)]
Out[10]= 2 v[p, m] p.q
```

Also more complicated structures are

simplified; for example,

$\bar{v}(p) \not{q} \not{p} u(q) = \bar{v}(p) u(p) [2(p \cdot q) + m M]$ .

```
In[11]:= DiracSimplify[Spinor[-p, m] . DiracSlash[q, p] .
Spinor[q, M] // Factor
Out[11]= v[p, m] u[q, M] (m M + 2 p.q)
```

`DiracSimplify` orders products of Gell-Mann matrices from the Dirac structure.

```
In[12]:= DiracSimplify[QuarkSpinor[-p1] . GellMannMatrix[b] .
DiracSlash[q] . GellMannMatrix[a] .
QuarkSpinor[p2]]
Out[12]= u[p1] la[b] la[a] gs[q] u[p2]
```

## ■ 2.4 Dirac Traces

<code>DiracTrace[expr]</code>	head of a Dirac trace
<code>Tr[expr]</code>	calculate the trace directly

Two functions for Dirac traces.

The function `DiracTrace` takes as *expr* either subsequent `DiracMatrix` or `DiracSlash` separated by ";" or any expression with "." as noncommutative multiplication operator.

The default of `DiracTrace` is not to evaluate Dirac traces directly. For direct calculation the function `Tr` should be used.

```

tr(γα γβ) = 4 gαβ.
In[1]:= Tr[DiracMatrix[a1, b1]]
Out[1]= 4 g[a1, b1]

tr(⧸p ⧸k ⧸p) =
4{(a·d)(b·c) - (a·c)(b·d) + (a·b)(c·d)}.
In[2]:= Tr[DiracSlash[a, b, c, d]]
Out[2]= 4 a.d b.c - 4 a.c b.d + 4 a.b c.d

tr(γa γb γc γd γ5) = -4 i εabcd.
In[3]:= Tr[DiracMatrix[a, b, c, d, 5]]
Out[3]= -4 I eps[a, b, c, d]

You may include metric tensors or
four-vectors, for example,
tr(¼ gαβ γμ γα pμ) = pβ.
In[4]:= Tr[MetricTensor[a1, b1]/4 DiracMatrix[mu],
           DiracMatrix[a1] FourVector[p, mu]]
Out[4]= p[b1]
```

If you want to do more complicated traces it is convenient to introduce abbreviations for the several objects. The following examples show how to do this. The CPU time needed for each trace calculation is for all examples less than one minute (on a NeXT). Some of the examples here verify the results given in [9].

Consider a trace corresponding to the square of the s-channel diagram for  $\gamma e$  scattering:

$$T_1 = \frac{1}{16} \text{tr}[(\not{p}' + m) \gamma^\alpha (\not{p} + \not{k} + m) \gamma^\beta (\not{p} + m) \gamma^\beta (\not{p} + \not{k} + m) \gamma^\alpha]$$

```

Set the abbreviations for Dirac matrices
and slashes here.
In[5]:= ( PP = DiracSlash[p'];
          P = DiracSlash[p];   K = DiracSlash[k];
          A = DiracMatrix[a1]; B = DiracMatrix[b1]
        )
Out[5]= ga[b1]

This is the input for trace T1.
In[6]:= Tr[PP + m,A,P + K + m,B,P + m,B,P + K + m,A/16]
The CPU time needed for the calculation
is of the order of seconds.
Out[6]= 4 m4 + 4 m2 k.k + 4 m2 k.p - 4 m2 k.p' +
        2 k.p k.p' + 2 k.p' p.p - 3 m2 p.p' - k.k p.p' + p.p p.p'
```

Another nontrivial example is a  $D$ -dimensional trace involving 14 Dirac matrices:

$$T_2 = \text{tr}(\gamma^\beta \gamma^\alpha \not{p}_1 \not{p}_2 \gamma^\nu \gamma^\beta \not{p}_2 \not{p}_3 \gamma^\alpha \not{p}_1 \gamma^\nu \not{p}_3 \not{p}_1 \not{p}_2)$$

```

This defines abbreviations for trace T2.
a, b, n denote γα, γβ, γν in D dimensions.
The last command sets P1 = ⧸p1, P2 = ⧸p2,
P3 = ⧸p3.
In[7]:= ( a = DiracMatrix[a1, Dimension -> D];
          b = DiracMatrix[b1, Dimension -> D];
          n = DiracMatrix[nu, Dimension -> D];
          {P1, P2, P3} = Map[ DiracSlash, {p1, p2, p3} ]
        )
Out[7]= {gs[p1], gs[p2], gs[p3]}
```



Here is the input for trace  $T_2$ .

The result is again collected with respect to scalar products.

```
In[8]:= Tr[b,a,P1,P2,n,b,P2,P3,a,P1,n,P3,P1,P2]
Out[8]= (-1152 + 896 D - 224 D2 + 16 D3) p1.p2 p1.p32
        p2.p2 + (1024 - 512 D + 64 D2) p1.p22 p1.p3 p2.p3 +
        (448 - 416 D + 112 D2 - 8 D3) p1.p1 p1.p3 p2.p2 p2.p3 +
        (-512 + 256 D - 32 D2) p1.p1 p1.p2 p2.p32 +
        (-512 + 256 D - 32 D2) p1.p23 p3.p3 +
        (672 - 416 D + 80 D2 - 4 D3) p1.p1 p1.p2 p2.p2 p3.p3
```

This calculates  $T_2$  in four dimensions. Since the "." is used, the replacement  $D \rightarrow 4$  applies to all Dirac matrices. The time needed would be twice as much without calculating the D-dimensional case before.

```
In[9]:= Tr[b.a.P1.P2.n.b.P2.P3.a.P1.n.P3.P1.P2 /. D -> 4]
Out[9]= -128 p1.p2 p1.p32 p2.p2 +
        64 p1.p1 p1.p3 p2.p2 p2.p3 + 32 p1.p1 p1.p2 p2.p2 p3.p3
```

Sometimes you do not want a trace to be evaluated immediately. Here you get the input  $tr(\gamma^\alpha \gamma^\beta \gamma^\rho \gamma^\sigma)$  back.

```
In[10]:= DiracTrace[DiracMatrix[al, be, ro, si]]
Out[10]= tr[ga[al] ga[be] ga[ro] ga[si]]
```

You may then contract, e.g., with  $g^{\alpha\beta}$ .

```
In[11]:= Contract[% MetricTensor[al, be]]
Out[11]= tr[ga[be] ga[be] ga[ro] ga[si]]
```

This evaluates the Dirac trace.

```
In[12]:= EvaluateDiracTrace[%]
Out[12]= 16 g[ro, si]
```

<code>EvaluateDiracTrace[expr]</code>	evaluate DiracTrace in <i>expr</i>
---------------------------------------	------------------------------------

Evaluation of Dirac traces.

<i>option name</i>	<i>default value</i>	
<code>DiracTraceEvaluate</code>	<code>False</code>	whether to evaluate the trace
<code>LeviCivitaSign</code>	<code>-1</code>	which sign convention to use in the result of $tr(\gamma^a \gamma^b \gamma^c \gamma^d \gamma^5)$ . The default gives $(-1)4i \epsilon^{abcd}$
<code>Factoring</code>	<code>False</code>	whether to factor canonically
<code>Mandelstam</code>	<code>{}</code>	utilize the Mandelstam relation
<code>PairCollect</code>	<code>True</code>	whether to collect Pair

Options for DiracTrace.

`Tr` takes the same options as `DiracTrace`, but the default setting of `DiracTraceEvaluate` is `True`.

The option `PairCollect` determines whether the resulting polynomial is collected with respect to metric tensors, four-vectors and scalar products. In the internal representation these three objects have the same head `Pair`, hence the name `PairCollect`.

For  $2 \rightarrow 2$  processes the traces are often expressed in terms of Mandelstam variables. In order to replace the scalar products you can use `SetMandelstam`.

```
SetMandelstam[s, t, u, p1, p2, p3, p4, m1, m2, m3, m4]
```

define scalar products in terms of Mandelstam variables and put the  $p_i$  on-shell

A function for introducing Mandelstam variables.

Assuming all  $p_i$  incoming, *i.e.*,  $p_1 + p_2 + p_3 + p_4 = 0$ , the Mandelstam variables are defined by:  $s = (p_1 + p_2)^2$ ,  $t = (p_1 + p_3)^2$ ,  $u = (p_1 + p_4)^2$ . Using these three equations and the on-shell conditions  $p_i^2 = m_i^2$ , `SetMandelstam` sets the 10 possible scalar products  $(p_i \cdot p_j)$  in terms of  $s, t, u$  and  $m_i^2$ .

For calculation of traces often the Mandelstam relation  $s + t + u = m_1^2 + m_2^2 + m_3^2 + m_4^2$  can be used to get a compact result. If you set the option `Mandelstam -> {s, t, u, m1^2 + m2^2 + m3^2 + m4^2}`, *FeynCalc* figures out the best choice of  $s, t$  or  $u$  in each factor of the result.

As an example for calculating a trace in terms of Mandelstam variables consider the following squared amplitude from the process  $gg \rightarrow t\bar{t}$ , with  $\Sigma_1^{\alpha\rho}$  and  $\Sigma_2^{\beta\sigma}$  as polarization sums for the gluons.

$$\begin{aligned} T_3 &= \text{tr}(\gamma^\sigma (\not{k}_1 - \not{p}_1 - m_t) \gamma^\rho (\not{p}_1 + m_t) (\not{p}_2 - m_t)) p_1^\alpha p_2^\beta \Sigma_1^{\alpha\rho} \Sigma_2^{\beta\sigma}, \text{ where} \\ \Sigma_1^{\alpha\rho} &= -g^{\alpha\rho} + \frac{4}{(u-t)^2} (4m_t^2 - s) k_1^\alpha k_1^\rho + \frac{2}{u-t} [k_1^\rho (p_1 - p_2)^\alpha + k_1^\alpha (p_1 - p_2)^\rho] \\ \Sigma_2^{\beta\sigma} &= -g^{\beta\sigma} + \frac{4}{(t-u)^2} (4m_t^2 - s) k_2^\beta k_2^\sigma + \frac{2}{t-u} [k_2^\sigma (p_1 - k_2)^\beta + k_2^\beta (p_1 - p_2)^\sigma] \end{aligned}$$

Set up  $s, t, u$  for  $gg \rightarrow t\bar{t}$ , with  $k_1, k_2$  as gluon and  $p_1, p_2$  as fermion momenta. Again abbreviations with capital letters are introduced for the Dirac matrices and slashes. *polsum1* and *polsum2* are the polarization sums for the gluons. As external momentum the choice  $n = p_1 - p_2$  has been made.

```
In[13]:= (SetMandelstam[s,t,u, k1,k2,-p1,-p2, 0,0,mt,mt];
          {K1, P1, P2} = Map[ DiracSlash, {k1, p1, p2} ];
          {SI, R0} = Map[ DiracMatrix, {si, ro} ];
          polsum1 = PolarizationSum[al, ro, k1, p1 - p2];
          polsum2 = PolarizationSum[be, si, k2, p1 - p2];
          p1al = FourVector[p1, al];
          p2be = FourVector[p2, be])
```

```
Out[13]= p2[be]
```

This is a possible input for trace  $T_3$ . *FeynCalc* contracts first all Lorentz indices and then calculates the trace.

```
In[14]:= Tr[ polsum1 polsum2 p1al p2be SI,
            K1 - P1 - mt, R0, P1 + mt, P2 - mt,
            Mandelstam -> {s, t, u, 2 mt^2} ]
```

Since the option `Mandelstam` has been specified, the result is given in a factored form, where in each factor one of  $s, t$  or  $u$  is eliminated via the Mandelstam relation. Note that a factor  $(t - u)$  has been cancelled.

$$\text{Out[14]} = \frac{-2 \text{mt} s (\text{mt}^4 - t u) (8 \text{mt}^4 - t^2 - 6 t u - u^2)}{(t - u)^3}$$

An alternative method would be to first calculate the trace without the polarization sums.

```
In[15]:= temp = Tr[SI, K1 - P1 - mt, R0, P1 + mt, P2 - mt]
Out[15]= (-2 mt t + 2 mt u) g[ro, si] - 4 mt k1[si] p1[ro] -
          4 mt k1[ro] p1[si] + 8 mt p1[ro] p1[si] +
          4 mt k1[si] p2[ro] + 4 mt k1[ro] p2[si] -
          8 mt p1[ro] p2[si]
```

Then contract the result with the polarization sums, expand the scalar products and use `TrickMandelstam` in order to get the tricky Mandelstam variable substitution.

```
In[16]:= TrickMandelstam[ ExpandScalarProduct[ Contract[
          temp polsum1 polsum2 p1al p2be]],
          {s, t, u, 2 mt^2} ]
```

This method is faster; but that is not the case for all trace calculations.

$$\text{Out[16]} = \frac{-2 \text{mt} s (\text{mt}^4 - t u) (8 \text{mt}^4 - t^2 - 6 t u - u^2)}{(t - u)^3}$$

The function `TrickMandelstam` is explained in section 4.4.7.

Since Dirac matrices can be given in any dimensions, *FeynCalc* is also able to calculate traces in  $D - 4$  dimensions.

Defining  $T(n) = \text{Tr}(\gamma_{\mu_1} \gamma_{\mu_2} \dots \gamma_{\mu_n} \gamma_{\mu_1} \gamma_{\mu_2} \dots \gamma_{\mu_n})$  we give a list of timings and results for  $T(8)$  to  $T(11)$ . The trace  $T(10)$  is a verification of the result given in [8].

This is a little program defining  $T$ . The dimension of each particular Dirac matrix is set to  $d - 4$ .

The timings are also displayed in `FeynCalcForm`, which yields more usual time units.

```
In[1]:= T[n_] := T[n] = Block[{gammas, calc},
  gammas = Dot @@ Table[
    DiracMatrix[a[i], Dimension -> (d - 4)],
    {i, 1, n} ];
  calc = Timing[ Tr[ gammas . gammas ] ];
  Print["Time = ", calc[[1]]//FeynCalcForm ];
  calc[[2]]];
```

This calculates a trace of 16 matrices.

```
In[2]:= T[8]
Time = 38. s

Out[2]= 123469824 - 135962624 d + 63224832 d2 -
  16145920 d3 + 2461760 d4 - 227584 d5 + 12320 d6 - 352 d7 +
  4 d8
```

Here we have 18.

```
In[3]:= T[9]
Time = 1.8 min

Out[3]= -1879576576 + 2220901376 d - 1127626752 d2 +
  321806848 d3 - 56625408 d4 + 6331584 d5 - 446208 d6 +
  18912 d7 - 432 d8 + 4 d9
```

The trace of 20 Dirac matrices.

```
In[4]:= T[10]
Time = 7.2 min

Out[4]= -31023169536 + 38971179008 d - 21328977920 d2 +
  6679521280 d3 - 1320732160 d4 + 171464832 d5 -
  14710080 d6 + 816960 d7 - 27840 d8 + 520 d9 - 4 d10
```

With 22 Dirac matrices it gets slow.

```
In[5]:= T[11]
Time = 55. min

Out[5]= 551768735744 - 731506905088 d + 427299186688 d2 -
  144858475520 d3 + 31576821760 d4 - 4629805312 d5 +
  463655808 d6 - 31521600 d7 + 1415040 d8 - 39600 d9 +
  616 d10 - 4 d11
```

## ■ 2.5 Gell-Mann Traces and Contraction of Color Indices

`GellMannTrace[expr]` calculates traces of Gell-Mann matrices

The function for trace calculation of  $\lambda_a$  matrices.

The matrices  $\frac{1}{2}\lambda_a$  are the infinitesimal generators of the group  $SU(3)$ . Similar to the case of Dirac traces, also traces of the Gell-Mann matrices  $\lambda_a$  are calculated algebraically. The Cvitanovic algorithm [6] is implemented similar to [7]. Traces with no free color indices are always evaluated to a number. Otherwise only special cases are simplified.

```

tr( $\lambda_a \lambda_b$ ) = 2  $\delta_{ab}$ .
In[1]:= GellMannTrace[GellMannMatrix[a, b]]
Out[1]= 2 d[a, b]

tr(3  $\lambda_a \lambda_b \lambda_a \lambda_b$ ) = -32.
In[2]:= GellMannTrace[ 3 GellMannMatrix[a, b, a, b] ]
Out[2]= -32

tr( $\lambda_a \lambda_b \lambda_c$ ) stays as it is.
In[3]:= GellMannTrace[GellMannMatrix[a, b, c]]
Out[3]= tr[1a[a] 1a[b] 1a[c]]

tr( $\lambda_a \lambda_b \lambda_c f_{abc}$ ) = 48 i.
In[4]:= GellMannTrace[SU3F[a, b, c] GellMannMatrix[a, b, c]]
Out[4]= 48 I

tr( $f_{ars} f_{brs}$ ) = 3  $f_{ars} f_{brs}$  = 9  $\delta_{ab}$ .
In[5]:= GellMannTrace[SU3F[a, r, s] SU3F[b, r, s]]
Out[5]= 9 d[a, b]

A non trivial case:
tr( $\lambda_a \lambda_c \lambda_e \lambda_d f_{abe} f_{bcd}$ ) = 0.
In[6]:= GellMannTrace[ GellMannMatrix[a, c, e, d] *
                    SU3F[a, b, e] SU3F[b, c, d] ]
Out[6]= 0

This takes a few seconds.

```

Repeated indices of  $\delta_{ab}$  and  $f_{abc}$  can always be contracted with the Cvitanovic algorithm.

```

16  $f_{ars} f_{brs}$  = 48  $\delta_{ab}$ .
In[7]:= 16 SU3F[a, r, s] SU3F[b, r, s]
Out[7]= -((-tr[1a[a] 1a[r] 1a[s]] + tr[1a[a] 1a[s] 1a[r]])
        (-tr[1a[b] 1a[r] 1a[s]] + tr[1a[b] 1a[s] 1a[r]]))

The structure constants are replaced by
traces.
Expanding forces the products of traces to
simplify, if there are indices to contract.
In[8]:= ExpandAll[%]
Out[8]= 48 d[a, b]

Define f as SU3F.
In[9]:= f[x_..] := SU3F[x]

 $f_{abr} f_{rcs} + f_{bcr} f_{ras} + f_{car} f_{rbs} = 0$ .
In[10]:= ExpandAll[ f[a,b,r] f[r,c,s] + f[b,c,r] f[r,a,s] +
                    f[c,a,r] f[r,b,s] ]
Out[10]= 0

 $\lambda_a \lambda_b \lambda_a = -\frac{2}{3} \lambda_b$ .
In[11]:= GellMannMatrix[a, b, a]
Out[11]=  $\frac{-2}{3} 1a[b]$ 

This is  $f_{abc} \lambda_b \lambda_c = 3 i \lambda_a$ .
In[12]:= SU3F[c, a, b, SU3FToTraces -> False] *
        GellMannMatrix[b, c]
Out[12]= 3 I 1a[a]

```

## 3 One-Loop Calculations

The methods and conventions implemented in *FeynCalc* for the evaluation of one-loop diagrams are described in [2] and [3]. The usual Passarino-Veltman scheme for the one-loop integrals is adapted to a large extent [2]. The coefficient functions of the tensor integrals are defined similar to [2], except that the Passarino-Veltman integrals take internal masses squared as arguments. A wrapper Fortran program to link FF — a Fortran program by G.J. van Oldenborgh [5] for the evaluation of the scalar  $n$ -point integrals — and *FeynCalc* is available from the author. Currently *FeynCalc* is limited to four-point integrals.

### 3.1 Passarino-Veltman Integrals and Reduction of Coefficient Functions

The scalar integrals  $A_0, B_0, C_0$  and  $D_0$  are represented in *FeynCalc* as functions with all arguments consisting of scalar products or masses squared. Thus  $A_0$  has one,  $B_0$  three,  $C_0$  six, and  $D_0$  ten arguments. The symmetry properties of the arguments are implemented, *i.e.*, a standard representative of all possible argument permutations of each  $B_0, C_0$  and  $D_0$  is returned. For example,  $B_0 [pp, m_2^2, m_1^2] \rightarrow B_0 [pp, m_1^2, m_2^2]$ , where  $pp$  denotes the scalar product  $(p \cdot p) = p^2$ .

$A_0 [m_0^2]$	$(i\pi^2)^{-1} \int d^D q (q^2 - m_0^2)^{-1}$
$B_0 [p_1^2, m_0^2, m_1^2]$	$(i\pi^2)^{-1} \int d^D q [(q^2 - m_0^2)((q + p_1)^2 - m_1^2)]^{-1}$
$DB_0 [p_1^2, m_0^2, m_1^2]$	$\partial B_0(p_1^2, m_0^2, m_1^2) / \partial p_1^2$
$C_0 [p_1^2, p_2^2, p_3^2, m_0^2, m_1^2, m_2^2]$	$(i\pi^2)^{-1} \int d^D q [(q^2 - m_0^2)((q + p_1)^2 - m_1^2)((q + p_2)^2 - m_2^2)]^{-1}$
$D_0 [p_1^2, p_2^2, p_3^2, p_4^2, p_5^2, p_6^2, m_0^2, m_1^2, m_2^2, m_3^2]$	$(i\pi^2)^{-1} \int d^D q [(q^2 - m_0^2)((q + p_1)^2 - m_1^2)((q + p_2)^2 - m_2^2)((q + p_3)^2 - m_3^2)]^{-1}$

Scalar Passarino-Veltman functions  $A_0, B_0, B_0', C_0$  and  $D_0$ .

In *FeynCalc* and in the mathematical definitions given above the factor  $(2\pi\mu)^{(4-D)}$  with the scaling variable  $\mu$  is suppressed.

The convention for the scalar arguments is  $pi0 = p_i^2$   $pij = (p_i - p_j)^2$ ,  $mi2 = m_i^2$ .

The  $B_0$  function is symmetric in the mass arguments.  $In [1] := B_0 [s, mz2, mw2]$   
 $Out [1] = B_0 [s, mw2, mz2]$

Taking the derivative with respect to the first argument yields  $DB_0$ .  $In [2] := D [B, s]$   
 $Out [2] = DB_0 [s, mw2, mz2]$

The tensor-integral decomposition is automatically done by *FeynCalc* when calculating one-loop amplitudes, but extra functions are provided to reduce the coefficients of the tensor-integral decomposition.

For fixing the conventions of the coefficient functions the definitions of the tensor-integrals and the decomposition are given below. In general the one-loop tensor integral is

$$T_{\mu_1 \dots \mu_p}^N(p_1, \dots, p_{N-1}, m_0, \dots, m_{N-1}) = \frac{(2\pi\mu)^{4-D}}{i\pi^2} \int d^D q \frac{q_{\mu_1} \dots q_{\mu_p}}{\mathcal{D}_0 \mathcal{D}_1 \dots \mathcal{D}_{N-1}}$$

with the denominator factors

$$\mathcal{D}_0 = q^2 - m_0^2 \quad \mathcal{D}_i = (q + p_i)^2 - m_i^2, \quad i = 1, \dots, N-1$$

originating from the propagators in the Feynman diagram. The  $i\epsilon$  part of the denominator factors is suppressed.

The tensor integral decompositions for the integrals that *FeynCalc* can do are listed below. The coefficient functions  $B_i, B_{ij}, C_i, C_{ij}, C_{ijk}, D_i, D_{ij}, D_{ijk}$  and  $D_{ijkl}$  are totally symmetric in their indices.

$$B_\mu = p_{1\mu} B_1$$

$$B_{\mu\nu} = g_{\mu\nu} B_{00} + p_{1\mu} p_{1\nu} B_{11}$$

$$C_\mu = p_{1\mu} C_1 + p_{2\mu} C_2 = \sum_{i=1}^2 p_{i\mu} C_i$$

$$\begin{aligned} C_{\mu\nu} &= g_{\mu\nu} C_{00} + p_{1\mu} p_{1\nu} C_{11} + p_{2\mu} p_{2\nu} C_{22} + (p_{1\mu} p_{2\nu} + p_{2\mu} p_{1\nu}) C_{12} \\ &= g_{\mu\nu} C_{00} + \sum_{i,j=1}^2 p_{i\mu} p_{j\nu} C_{ij} \end{aligned}$$

$$\begin{aligned} C_{\mu\nu\rho} &= (g_{\mu\nu} p_{1\rho} + g_{\nu\rho} p_{1\mu} + g_{\mu\rho} p_{1\nu}) C_{001} + (g_{\mu\nu} p_{2\rho} + g_{\nu\rho} p_{2\mu} + g_{\mu\rho} p_{2\nu}) C_{002} \\ &\quad + p_{1\mu} p_{1\nu} p_{1\rho} C_{111} + p_{2\mu} p_{2\nu} p_{2\rho} C_{222} \\ &\quad + (p_{1\mu} p_{1\nu} p_{2\rho} + p_{1\mu} p_{2\nu} p_{1\rho} + p_{2\mu} p_{1\nu} p_{1\rho}) C_{112} \\ &\quad + (p_{2\mu} p_{2\nu} p_{1\rho} + p_{2\mu} p_{1\nu} p_{2\rho} + p_{1\mu} p_{2\nu} p_{2\rho}) C_{122} \\ &= \sum_{i=1}^2 (g_{\mu\nu} p_{i\rho} + g_{\nu\rho} p_{i\mu} + g_{\mu\rho} p_{i\nu}) C_{00i} + \sum_{i,j,k=1}^2 p_{i\mu} p_{j\nu} p_{k\rho} C_{ijk} \end{aligned}$$

$$D_\mu = \sum_{i=1}^3 p_{i\mu} D_i$$

$$D_{\mu\nu} = g_{\mu\nu} D_{00} + \sum_{i,j=1}^3 p_{i\mu} p_{j\nu} D_{ij}$$

$$D_{\mu\nu\rho} = \sum_{i=1}^3 (g_{\mu\nu} p_{i\rho} + g_{\nu\rho} p_{i\mu} + g_{\mu\rho} p_{i\nu}) D_{00i} + \sum_{i,j,k=1}^3 p_{i\mu} p_{j\nu} p_{k\rho} D_{ijk}$$

$$\begin{aligned} D_{\mu\nu\rho\sigma} &= (g_{\mu\nu} g_{\rho\sigma} + g_{\mu\rho} g_{\nu\sigma} + g_{\mu\sigma} g_{\nu\rho}) D_{0000} \\ &\quad + \sum_{i,j=1}^3 (g_{\mu\nu} p_{i\rho} p_{j\sigma} + g_{\nu\rho} p_{i\mu} p_{j\sigma} + g_{\mu\rho} p_{i\nu} p_{j\sigma} + g_{\mu\sigma} p_{i\nu} p_{j\rho} + g_{\nu\sigma} p_{i\mu} p_{j\rho} + g_{\rho\sigma} p_{i\mu} p_{j\nu}) D_{00ij} \\ &\quad + \sum_{i,j,k,l=1}^3 p_{i\mu} p_{j\nu} p_{k\rho} p_{l\sigma} D_{ijkl} \end{aligned}$$

All coefficient functions and the scalar integrals are summarized in one generic function, `PaVe`.

`PaVe[i, j, ..., {P10, P12, ...}, {m02, m12, ...}]`

Passarino-Veltman coefficient functions

Passarino-Veltman coefficient functions of the tensor integral decomposition.

The first set of arguments  $i, j, \dots$  are exactly those indices of the coefficient functions of the tensor integral decomposition. If only a 0 is given as first argument, the scalar integrals are understood. The last argument, the list of inner masses  $m_0^2, m_1^2, \dots$ , determines whether a one-, two-, three- or four-

point function is meant. PaVe is totally symmetric in the  $i, j, \dots$  arguments. The foremost argument is the list of scalar products of the  $p_i$ . They are the same as defined above for the scalar  $B_0, C_0$  and  $D_0$  functions. For  $A_0$  an empty list has to be given.

A certain set of special PaVe shown in the following examples simplify to the usual notation.

To shorten the input squared masses are abbreviated with a suffix 2, *i.e.*, a mass  $m^2$  is denoted by m2. The scalar quantity  $p^2$  is entered as pp,  $p_i^2$  as pi0 and  $(p_i - p_j)^2$  as pij.

For the following examples some options are set.	<code>In[3]:= (SetOptions[A0, A0ToB0 -&gt; False]; SetOptions[{B1, B00, B11}, BReduce -&gt; False]);</code>
This is the scalar Passarino-Veltman one-point function.	<code>In[4]:= PaVe[0, {}, {m02}] Out[4]= A0[m02]</code>
This is the two-point function $B_0(p^2, m_0^2, m_1^2)$ .	<code>In[5]:= PaVe[0, {pp}, {m02, m12}] Out[5]= B0[pp, m02, m12]</code>
Here is the coefficient function $B_1(p^2, m_0^2, m_1^2)$ .	<code>In[6]:= PaVe[1, {pp}, {m12, m22}] Out[6]= B1[pp, m12, m22]</code>
Coefficient functions of metric tensors have two "0" indices for each $g^{\mu\nu}$ .	<code>In[7]:= PaVe[0,0, {pp}, {m02, m12}] Out[7]= B00[pp, m02, m12]</code>
This is $B_{11}(p_1^2, m_0, m_1)$ , the coefficient function of $p_{1\mu} p_{1\nu}$ of $B_{\mu\nu}$ .	<code>In[8]:= PaVe[1,1, {p10}, {m12, m22}] Out[8]= B11[p10, m12, m22]</code>
PaVe with one 0 as first argument are scalar Passarino-Veltman integrals.	<code>In[9]:= PaVe[0, {p10, p12, p20}, {m12, m22, m32}] Out[9]= C0[p10, p12, p20, m12, m22, m32]</code>
This is $D_0$ with 10 arguments.	<code>In[10]:= PaVe[0, {p10, p12, p23, p30, p20, p13}, {m12, m22, m32, m42}] Out[10]= D0[p10, p12, p23, p30, p20, p13, m12, m22, m32, m42]</code>

<code>B1 [p10, m02, m12]</code>	coefficient function $B_1(p_1^2, m_0^2, m_1^2)$
<code>B00 [p10, m02, m12]</code>	coefficient function $B_{00}(p_1^2, m_0^2, m_1^2)$
<code>B11 [p10, m02, m12]</code>	coefficient function $B_{11}(p_1^2, m_0^2, m_1^2)$

Two-point coefficient functions.

The two-point coefficient functions can be reduced to lower-order ones. For special arguments also  $B_0$  is expressed in terms of  $A_0$ , if the option BReduce is specified. Setting the option B0Unique to True simplifies  $B_0(m^2, 0, m^2) \rightarrow 2 + B_0(0, m^2, m^2)$  and  $B_0(0, 0, m^2) \rightarrow 1 + B_0(0, m^2, m^2)$ .

<i>option name</i>	<i>default value</i>	
A0ToB0	True	whether to express $A_0(m^2)$ by $m^2(1 + B_0(0, m^2, m^2))$
BReduce	True	reduce B0, B1, B00, B11 to A0 and B0
B0Unique	False	whether to simplify $B_0(a, 0, a)$ and $B_0(0, 0, a)$

Reduction options for A0 and the two-point functions.

The default is to reduce  $B_1$  to  $B_0$ .

$$\begin{aligned} \text{In}[1] &:= \text{B1}[\text{pp}, \text{m12}, \text{m22}] \\ \text{Out}[1] &= \frac{-\text{B0}[\text{pp}, \text{m12}, \text{m22}]}{2} + \\ &\quad \frac{(-\text{m12} + \text{m22}) (-\text{B0}[0, \text{m12}, \text{m22}] + \text{B0}[\text{pp}, \text{m12}, \text{m22}])}{2 \text{ pp}} \end{aligned}$$

Arguments of two-point functions with head `Small` are replaced by 0, if the other arguments have no head `Small` and are nonzero.  $A0[0]$  and  $A0[\text{Small}[m]^2]$  simplify to 0.

The small mass  $m$  is set to 0, since the other arguments are non-zero and not `Small`.

$$\begin{aligned} \text{In}[2] &:= \text{B1}[\text{pp}, \text{Small}[\text{me2}], \text{m22}] \\ \text{Out}[2] &= \frac{-\text{B0}[\text{pp}, 0, \text{m22}]}{2} + \\ &\quad \frac{\text{m22} (-\text{B0}[0, 0, \text{m22}] + \text{B0}[\text{pp}, 0, \text{m22}])}{2 \text{ pp}} \end{aligned}$$

But in this case no arguments are replaced by 0.

$$\begin{aligned} \text{In}[3] &:= \text{B1}[\text{Small}[\text{me2}], \text{Small}[\text{me2}], 0] \\ \text{Out}[3] &= -\frac{1}{2} - \frac{\text{B0}[\text{Small}[\text{me2}], 0, \text{Small}[\text{me2}]]}{2} \end{aligned}$$

<code>PaVeReduce[expr]</code>	reduces coefficient functions PaVe to A0, B0, C0, D0
<code>K[i]</code>	abbreviations in HoldForm in the result of PaVeReduce

Reduction function for Passarino-Veltman coefficient functions.

Depending on the option `BReduce`  $B_1$ ,  $B_{00}$  and  $B_{11}$  may also remain in the result of `PaVeReduce`.

<i>option name</i>	<i>default value</i>	
<code>IsolateHead</code>	<code>False</code>	whether to use <code>Isolate</code> with this option
<code>Mandelstam</code>	<code>{}</code>	Mandelstam relation, e.g., $\{s, t, u, 2 \text{ mw}^2\}$

Options for `PaVeReduce`.

The function `Isolate` is explained in section 4.4.2.

Reduce  $C_2(m_c^2, m_w^2, t, m_c^2, 0, m_w^2)$  to scalar integrals.

$$\begin{aligned} \text{In}[4] &:= \text{PaVeReduce}[\text{PaVe}[2, \{\text{Small}[\text{me2}], \text{mw2}, \text{t}\}, \\ &\quad \{\text{Small}[\text{me2}], 0, \text{mw2}\}]] \\ \text{Out}[4] &= \frac{\text{B0}[\text{mw2}, 0, \text{mw2}]}{\text{mw2} - \text{t}} - \frac{\text{B0}[\text{t}, 0, \text{mw2}]}{\text{mw2} - \text{t}} \end{aligned}$$

Break down the coefficient function  $C_{12}(s, m^2, m^2, m^2, m^2, M^2)$ .

$$\text{In}[5] := \text{c12} = \text{PaVeReduce}[\text{PaVe}[1, 2, \{s, \text{m2}, \text{m2}\}, \{\text{m2}, \text{m2}, \text{M2}\}], \text{IsolateHead} \rightarrow \text{K}]$$

This is the result in `HoldForm`.

$$\text{Out}[5] = \text{K}[11]$$

The `FullForm` of the assignment to `c12` is `HoldForm[K[11]]`. If you want to get the value of `K[11]`, you can either type `ReleaseHold[c12]` or `K[11]` as done in the example.

$$\begin{aligned} \text{In}[6] &:= \text{K}[11] \\ \text{Out}[6] &= \frac{\text{K}[6]}{2} + \frac{\text{M2} \text{K}[4] \text{K}[7]}{\text{K}[5]^2} + \frac{\text{K}[1] \text{K}[8]}{2 \text{m2} \text{K}[5]} + \frac{\text{K}[3] \text{K}[9]}{2 \text{K}[5]^2} - \\ &\quad \frac{\text{K}[2] \text{K}[10]}{2 \text{m2} \text{K}[5]^2} \end{aligned}$$



Repeated application of `ReleaseHold` reinserts all `K`.

$$\begin{aligned} \text{In}[7] := & \text{FixedPoint}[\text{ReleaseHold}[\#] \&, c12] \\ \text{Out}[7] = & \frac{1}{2(4m_2 - s)} + \frac{(m_2 - M_2) \text{B0}[0, m_2, M_2]}{2m_2(4m_2 - s)} - \\ & \frac{(8m_2^2 - 10m_2M_2 - 2m_2s + M_2s) \text{B0}[m_2, m_2, M_2]}{2m_2(4m_2 - s)^2} + \\ & \frac{(4m_2 - 6M_2 - s) \text{B0}[s, m_2, m_2]}{2(4m_2 - s)^2} + \\ & \frac{M_2(8m_2 - 3M_2 - 2s) \text{C0}[m_2, m_2, s, m_2, M_2, m_2]}{(4m_2 - s)^2} \end{aligned}$$

Take a coefficient function from  $D_{\mu\nu\rho}$ :  $D_{122}(m_c^2, m_w^2, m_w^2, m_c^2, s, t, 0, m_c^2, 0, m_c^2)$ .

```
In[8] := d122 = PaVeReduce[
  PaVe[1, 2, 2, {Small[ME2], MW2, MW2, Small[ME2], S, T},
    {0, Small[ME2], 0, Small[ME2]}],
  Mandelstam -> {S, T, U, 2 MW2}, IsolateHead -> F ]
Out[8] = F[17]
```

Write the result out into a Fortran file.

```
In[9] := Write2[ "d122.for", d122res = d122,
  FormatType -> FortranForm ];
```

This shows the resulting Fortran file.

The first abbreviations are always the scalar integrals.

The partially recursive definitions of the abbreviations are not fully optimized. If you want to have a more optimized Fortran output you may use the settings `MacsymaForm` or `MapleForm` of the option `FormatType` for creating an output file with the syntax of the computer algebra systems *Macsyma* and *Maple*, respectively. Then you can use the Fortran code generators of *Maple* or *Macsyma*. In some versions of *Macsyma* the excellent package *Gentran* by B. Gates and H. van Hulzen is available.

The function `Write2` is explained in Section 4.

Note that the head `Small` is eliminated in the Fortran output automatically.

```
In[10] := !!d122.for
F(1) = BO(MW2,ODO,ODO)
F(2) = BO(T,ODO,ODO)
F(3) = BO(S,ODO,ODO)
F(4) = CO(MW2,MW2,T,ME2,ODO,ME2)
F(5) = CO(MW2,S,ME2,ME2,ODO,ODO)
F(6) = CO(T,ME2,ME2,ME2,ME2,ODO)
F(7) = DO(ME2,MW2,MW2,ME2,S,T,ODO,ME2,ODO,ME2)
F(8) = MW2 + S
F(9) = 4*MW2 - T
F(10) = MW2**2 - S*U
F(11) = F(8)/(F(9)*F(10))
F(12) = -MW2 + S
F(13) = 4*MW2**5 - 5*MW2**4*S - 16*MW2**3*S**2 +
- 4*MW2**2*S**3 + 4*MW2*S**4 - MW2**4*U -
- 4*MW2**2*S**2*U + 8*MW2*S**3*U +
- 4*MW2**2*S*U**2 + S**3*U**2 + S**2*U**3
F(14) = 4*MW2**3 - 9*MW2**2*S + 2*S**3 - MW2**2*U -
- 4*MW2*S*U + 5*S**2*U + 3*S*U**2
F(15) = MW2**2 - 4*MW2*S + 2*S**2 + S*U
F(16) = 2*MW2**6 - 8*MW2**5*S + 12*MW2**4*S**2 -
- 8*MW2**3*S**3 + 2*MW2**2*S**4 -
- 2*MW2**5*T + 20*MW2**4*S*T -
- 36*MW2**3*S**2*T + 20*MW2**2*S**3*T -
- 2*MW2*S**4*T - 6*MW2**3*S**2*T**2 +
- 6*MW2**2*S**2*T**2 - 6*MW2*S**3*T**2 +
- 4*MW2*S**2*T**3 - S**2*T**4
F(17) = -F(11)/2 + S**2*T**2*F(6)*F(12)/(2*F(10)**3) -
- S**3*T**2*F(7)*F(12)/(2*F(10)**3) +
- S**2*T*F(5)*F(12)**2/F(10)**3 +
- F(1)*F(13)/(2*F(9)**2*F(10)**2*F(12)) +
- F(2)*F(8)*F(14)/(2*F(9)**2*F(10)**2) -
- S*F(3)*F(15)/(2*F(10)**2*F(12)) -
- F(4)*F(8)*F(16)/(2*F(9)**2*F(10)**3)
d122res = F(17)
```

The Fortran code generated by `Write2` should be checked with care. All integer numbers (except 0 as argument of `B0`, `C0`, `D0`) are translated to integers. This causes problems when translating variables with rational powers and must be corrected in the Fortran output by hand.

### ■ 3.2 A One-Loop Self Energy Diagram

The function `OneLoop` performs the algebraic simplifications of a given amplitude. The result is given in a polynomial of standard matrix elements, invariants of the process under consideration, and Passarino-Veltman integrals.

<code>OneLoop [q, amp]</code>	calculates the one-loop amplitude <i>amp</i> with <i>q</i> as loop momentum
<code>OneLoop [name, q, amp]</code>	calculates the one-loop amplitude <i>amp</i> and gives it a name

Calculating one-loop amplitudes.

The first argument to `OneLoop` is optional. It indicates a name for the amplitude for bookkeeping reasons. The second argument *q* is the loop momentum, *i.e.*, the integration variable.

As last argument the analytical expression for the graph is given. It may be given in four dimensions. `OneLoop` performs the necessary extension to *D* dimensions automatically.

This is  
 $A_0 = -i \pi^{-2} (2\pi\mu)^{4-D} \int d^D q (q^2 - m^2)^{-1}$ ,  
 corresponding to a tadpole diagram.

The scaling variable  $\mu$  is suppressed in *FeynCalc*.

This calculates the tadpole diagram.

```
In[1]:= -I/Pi^2 FeynAmpDenominator[
      PropagatorDenominator[q, m]]
Out[1]= 
$$\frac{-I}{\text{Pi}^2 (q^2 - m^2)}$$

```

```
In[2]:= OneLoop[q, %]
Out[2]= 
$$\frac{1}{m^2 + m^2} \text{B0}[0, m^2, m^2]$$

```

For a most compact result the factoring option of `OneLoop` is set. For a description of all options of `OneLoop` see Section 3.3.4.

```
In[3]:= SetOptions[OneLoop, Factoring -> True,
      Dimension -> D];
```

This is the transversal part of a photon self energy diagram with a fermion loop.

$i e^2 / ((2\pi)^4 (1 - D))$

$\int d^4 q [q^2 - m_f^2]^{-1} [(q - k)^2 - m_f^2]^{-1}$

$\text{tr}[(m_f + \not{q} - \not{k}) \gamma^\nu (\not{q} + m_f) \gamma^\nu]$

=

$-[e^2(k^2 + 6m_f^2 B_0(0, m_f^2, m_f^2) -$

$-3(k^2 + 2m_f^2) B_0(k^2, m_f^2, m_f^2)) / (36\pi^2)$

```
In[4]:= OneLoop[ q, (I e1^2)/(16 Pi^4)/(1 - D) *
      FeynAmpDenominator[
      PropagatorDenominator[q, mf],
      PropagatorDenominator[q - k, mf] ] *
      DiracTrace[(mf + DiracSlash[q - k]) . DiracMatrix[mu] .
      (mf + DiracSlash[q]) . DiracMatrix[mu]]
      ] /. ScalarProduct[k, k] -> k2 /. (mf^2) -> mf2
Out[4]= 
$$-\frac{e^2}{3} \frac{(k^2 + 6 m_f^2 B_0[0, m_f^2, m_f^2] - 3(k^2 + 2 m_f^2) B_0[k^2, m_f^2, m_f^2])}{(36 \text{Pi}^2)}$$

```

Note that in this example, where the dimension is entered explicitly as a parameter (*D*), the option `Dimension` of `OneLoop` must also be set to *D*.

### ■ 3.3 Generic Diagrams for $W \rightarrow f_j \bar{f}_j$ with OneLoop

As an example for calculating triangle diagrams the result for two generic one-loop diagrams of the decay  $W \rightarrow f_j \bar{f}_j$  for massless fermions given in [2] is verified with *FeynCalc*.

For the two diagrams different approaches are taken. In the first one *FeynCalc* introduces standard matrix elements, *i.e.*, that part of the diagram containing polarization dependencies. In the other approach the set of standard matrix elements is defined by the user before *FeynCalc* calculates the diagrams. The

last possibility is usually preferable, since the choices of *FeynCalc* for the standard matrix elements may have physical significance only by accident.

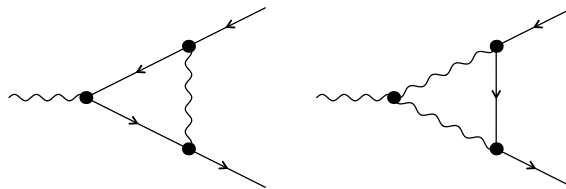


Figure 1: Two generic diagrams for the decay of  $W \rightarrow f_i \bar{f}_j$ , generated by *FeynArts*.

This factors the result.

```
In[1]:= SetOptions[OneLoop, Factoring -> True];
```

This defines a function for abbreviation purposes:  $g_i^- = g_i^- \omega_- + g_i^+ \omega_+$ .

```
In[2]:= (gc[i_] := g[i, "-"] DiracMatrix[7] +
          g[i, "+"] DiracMatrix[6];
```

Set the  $p_i$  on-shell and  $(p_1 \cdot p_2) = k^2/2$ , where  $k$  denotes the momentum of the  $W$ .

```
ScalarProduct[p1, p1] = 0;
ScalarProduct[p2, p2] = 0;
ScalarProduct[p1, p2] = k2/2);
```

The analytical expression for the generic diagram is:

```
In[3]:= Wff1 = OneLoop[ q, I/(2 Pi)^4 FeynAmpDenominator[
          PropagatorDenominator[q, M],
          PropagatorDenominator[q + p1],
          PropagatorDenominator[q - p2]          ] *
          Spinor[p1] . DiracMatrix[nu] . gc[1] .
          DiracSlash[q + p1] .
          DiracSlash[Polarization[k]] .
          gc[3] . DiracSlash[q - p2] .
          DiracMatrix[nu] . gc[2] .
          Spinor[p2] ] /. (M^2) -> M2
```

$$\delta \mathcal{M}_1 = i(2\pi)^{-4} \int (2\pi\mu)^{4-D} d^D q$$

$$[(q^2 - M^2)(q + p_1)^2(q - p_2)^2]^{-1}$$

$$\bar{u}(p_1) \gamma^\nu (g_1^- \omega_- + g_1^+ \omega_+) (\not{q} + \not{p}_1)$$

$$\not{\epsilon} (g_3^- \omega_- + g_3^+ \omega_+) (\not{q} - \not{p}_2) \gamma^\nu (g_2^- \omega_- + g_2^+ \omega_+) v(p_2)$$

Translated into the usual notation the result reads:

$$(1 - 2B_0(k^2, 0, 0) + 2k^2 C_0(0, 0, k^2, 0, M^2, 0) + 2k^2 C_1(k^2, 0, 0, 0, 0, M^2) + 4C_{00}(k^2, 0, 0, 0, 0, M^2)(g_1^+ g_2^+ g_3^+ \bar{u}(p_1) \not{\epsilon} \omega_+ v(p_2) + g_1^- g_2^- g_3^- \bar{u}(p_1) \not{\epsilon} \omega_- v(p_2)))/(16\pi^2)$$

The remaining Dirac structure is wrapped with the head `StandardMatrixElement`.

```
Out[3]= ((1 - 2 B0[0, 0, M2] + 2 k2 C0[0, 0, k2, 0, M2, 0] +
          2 k2 PaVe[1, {k2, 0, 0}, {0, 0, M2}] +
          4 PaVe[0, 0, {k2, 0, 0}, {0, 0, M2}])
          (g[1, +] g[2, +] g[3, +]
          StandardMatrixElement[u[p1] gs[ep[k]] ga[6] u[p2]] +
          g[1, -] g[2, -] g[3, -]
          StandardMatrixElement[u[p1] gs[ep[k]] ga[7] u[p2]])) \
          / (16 Pi^2)
```

This reduces the result to scalar integrals.

```
In[4]:= Wff1 = PaVeReduce[Wff1] // Combine
```

The default result of `PaVeReduce` is not put over a common denominator. This is achieved by using `Combine`.

```
Out[4]= ((2 k2 - 2 (2 k2 + M2) B0[0, 0, M2] +
          (3 k2 + 2 M2) B0[k2, 0, 0] +
          2 (k2 + M2)^2 C0[0, 0, k2, 0, M2, 0])
          (g[1, +] g[2, +] g[3, +]
          StandardMatrixElement[u[p1] gs[ep[k]] ga[6] u[p2]] +
          g[1, -] g[2, -] g[3, -]
          StandardMatrixElement[u[p1] gs[ep[k]] ga[7] u[p2]])) \
          / (16 k2 Pi^2)
```

With this command you can extract the standard matrix elements.

```
In[5]:= var = Select[Variables[Wff1],
  (Head[#]==StandardMatrixElement)&]
Out[5]= {StandardMatrixElement[u[p1] gs[ep[k]] ga[6] u[p2]],
  StandardMatrixElement[u[p1] gs[ep[k]] ga[7] u[p2]]}
```

Here the `StandardMatrixElement` are set to some abbreviations.

```
In[6]:= Set @@ {var, {MA[1], MA[2]} }
Out[6]= {MA[1], MA[2]}
```

In this way you can generate a Fortran file.

```
In[7]:= Write2["Wff1.for", vert =
  Wff1 /. g[i_, "+"] -> gp[i] /. g[i_, "-"] ->
  gm[i],
  FormatType -> FortranForm
]
```

With replacements you can adapt the result to your other Fortran code.

```
Out[7]= Wff1.for
```

Show the content of the file.

```
In[8]:= !!Wff1.for
vert = (2*k2 + (3*k2 + 2*M2)*B0(k2,Null,Null) -
- 2*(2*k2 + M2)*B0(Null,M2,Null) +
- 2*(k2 + M2)**2*
- CO(Null,Null,k2,Null,M2,Null))*
- (gp(1)*gp(2)*gp(3)*MA(1) +
- gm(1)*gm(2)*gm(3)*MA(2))/(16*k2*Pi**2)
```

```
StandardMatrixElement[expr]   head of a standard matrix element
SetStandardMatrixElements[{ {sm1 -> abb1}, {sm2 -> abb2}, ...}]
                               set abbreviations for standard matrix elements
SetStandardMatrixElements[{ {sm1 -> abb1}, {sm2 -> abb2}, ...}, k2 -> p1 + p2 - k1]
                               set abbreviations for standard matrix elements by using
                               energy momentum conservation
```

A head for identifying standard matrix elements;  $sm1, sm2$  are the standard matrixelements,  $abb1, abb2$  the abbreviations.

The function `SetStandardMatrixElements` introduces `StandardMatrixElement[abb1]` for  $sm1$ . The abbreviations  $abb1, abb2, \dots$  may be numbers or strings.

For calculating the generic triangle diagram with a non-abelian gauge coupling the standard matrix elements are set ahead using `SetStandardMatrixElements`.

In addition to the first example the option `ReduceToScalars` is set to `True`, which will produce directly a result in terms of  $B_0$  and  $C_0$ .

The other definitions are convenient abbreviations:  $R$  for the right-handed projection operator  $\gamma_6$ ,  $L$  for the left-handed projection operator  $\gamma_7$ , short mnemonic functions like  $G, FV$  and `feynden` stand for metric tensors, four-vectors and denominators of propagators.

```
In[1]:= (R = DiracMatrix[6]; L = DiracMatrix[7];
  ScalarProduct[p1, p1] = ScalarProduct[p2, p2] = 0;
  ScalarProduct[p1, p2] = k2/2 ;
  SetOptions[ OneLoop,
    Factoring -> True, FormatType -> FortranForm,
    ReduceToScalars -> True, WriteOut->True,
    FinalSubstitutions -> {g[i_, "+"] -> gp[i],
      g[i_, "-"] -> gm[i],
      StandardMatrixElement -> MAT} ];
  G[x_ y_] := MetricTensor[x, y];
  FV[p_, m_] := FourVector[p, m];
  feynden[x:{_, _}..] := FeynAmpDenominator @@
  Map[Apply[PropagatorDenominator, #]&, {x}] );
```

This sets the standard matrix elements:

$$\mathcal{M}_1^+ = \bar{u}(p_1) \not{\epsilon}_1 \omega_+ v(p_2)$$

$$\mathcal{M}_1^- = \bar{u}(p_1) \not{\epsilon}_1 \omega_- v(p_2)$$

```
In[2]:= SetStandardMatrixElements[
  { ( Spinor[p1] . DiracSlash[Polarization[k]] .
    R . Spinor[p2] ) -> {1},
    ( Spinor[p1] . DiracSlash[Polarization[k]] .
    L . Spinor[p2] ) -> {2}
  }
];
```

Here is the second generic diagram.

Note that in the result `StandardMatrixElement` is replaced by `MAT`, as specified in the option `FinalSubstitutions` of `OneLoop` on the previous page.

```
In[3]:= Wff2 = OneLoop[wff2, q, I/(2 Pi)^4 *
  feynden[{q, 0}, {q + p1, M1}, {q - p2, M2}] *
  Spinor[p1] . DiracMatrix[nu] .
  (g[1, "-"] L + g[1, "+"] R) .
  DiracSlash[-q] . DiracMatrix[ro] .
  (g[2, "-"] L + g[2, "+"] R) . Spinor[p2] *
  g3 ( G[ro nu] FV[p1 + 2 p2 - q, nu] -
    G[mu nu] FV[2 p1 + p2 + q, ro] +
    G[nu ro] FV[2 q + p1 - p2, mu] ) *
  PolarizationVector[k, mu]
  ] /. (M1^2) -> M12 /. (M2^2) -> M22
```

```
Out[3]= -(g3 ((2 k2 + M12) B0[0, 0, M12] +
  (2 k2 + M22) B0[0, 0, M22] -
  (k2 + M12 + M22) B0[k2, M12, M22] +
  2 (k2 M12 + k2 M22 + M12 M22)
  CO[0, 0, k2, M12, 0, M22])
  (gp[1] gp[2] MAT[1] + gm[1] gm[2] MAT[2])) / (16 k2 Pi^2)
```

As specified above in the options for `OneLoop`, a Fortran output is written into a file.

```
In[4]:= !!wff2.for
wff2 = -(g3*(-((k2 + M1**2 + M2**2)*B0(k2,M1**2,M2**2)) +
- (2*k2 + M1**2)*BO(Null,M1**2,Null) +
- (2*k2 + M2**2)*BO(Null,M2**2,Null) +
- 2*(k2*M1**2 + k2*M2**2 + M1**2*M2**2)*
- CO(Null,Null,k2,M1**2,Null,M2**2))*
- (gp(1)*gp(2)*MAT(1) +
- gm(1)*gm(2)*MAT(2))/(16*k2*Pi**2)
```

The result is also automatically assigned to `OneLoopResult[name]`.

```
In[4]:= OneLoopResult[wff2]
Out[4]= -(g3 ((2 k2 + M1^2) B0[0, 0, M1^2] +
  (2 k2 + M2^2) B0[0, 0, M2^2] -
  (k2 + M1^2 + M2^2) B0[k2, M1^2, M2^2] +
  2 (k2 M1^2 + k2 M2^2 + M1^2 M2^2)
  CO[0, 0, k2, M1^2, 0, M2^2])
  (gp[1] gp[2] MAT[1] + gm[1] gm[2] MAT[2])) / (16 k2 Pi^2)
```

<code>OneLoopResult[name]</code> is set to the result of <code>OneLoop[name, ...]</code>
--

`OneLoopResult[name]` gets value as a side effect of `OneLoop`.

### ■ 3.4 The Options of OneLoop

Several options of `OneLoop` have already been introduced in the previous section. Here the full list of available options is briefly discussed. The example in Section 3.3.6 shows the use of some options.

<i>option name</i>	<i>default value</i>	
<code>CancelQ2</code>	<code>True</code>	whether to cancel $q^2$
<code>CancelQP</code>	<code>True</code>	whether to cancel $q \cdot p$
<code>DenominatorOrder</code>	<code>True</code>	order the arguments of <code>FeynAmpDenominator</code> canonically
<code>Dimension</code>	<code>True</code>	whether to change the dimension to $D$
<code>Factoring</code>	<code>False</code>	whether to factor the result
<code>FinalSubstitutions</code>	<code>{}</code>	substitutions done at the end of the calculation
<code>FormatType</code>	<code>InputForm</code>	in which language to write out the result file
<code>InitialSubstitutions</code>	<code>{}</code>	substitutions done at the beginning of the calculation, <i>i.e.</i> , energy momentum conservation
<code>IsolateHead</code>	<code>False</code>	whether to use <code>Isolate</code> on the result
<code>Mandelstam</code>	<code>{}</code>	indicate the Mandelstam relation
<code>Prefactor</code>	<code>1</code>	additional prefactor of the amplitude
<code>ReduceGamma</code>	<code>False</code>	insert for $\gamma_6$ and $\gamma_7$ their definitions
<code>ReduceToScalars</code>	<code>False</code>	reduce to $B_0, C_0, D_0$
<code>SmallVariables</code>	<code>{}</code>	a list of masses, which will get wrapped around the head <code>Small</code>
<code>WriteOut</code>	<code>True</code>	write out a result file

Options of `OneLoop`.

In the automatic calculation of one-loop amplitudes it does not matter in which order the arguments of `FeynAmpDenominator` are given. Therefore the default setting of `DenominatorOrder` is `True`. In case you want to verify a result obtained by hand calculation, you can set this option to `False`, which will preserve the order of the propagators as entered. If you want to include the dimension  $D$  explicitly in the input, as in the example in Section 3.3.2, you have to set `Dimension -> D`.

With the default setting of `Dimension` you can enter four-dimensional objects to `OneLoop`, which are automatically extended to  $D$  dimensions inside `OneLoop`. In case you want to calculate a finite amplitude, you can set `Dimension -> 4`.

The option `FinalSubstitutions` indicates substitutions that are done at the very end of the calculation, which may be useful to adapt the output to your Fortran program.

The `Factoring` option should be used only for relatively small problems, since it may be very time consuming to factor the result. Unless the result of `OneLoop` is very short, only the coefficients of `StandardMatrixElement` are factored.

`FormatType` takes `InputForm`, `FortranForm`, `MacsymaForm` or `MapleForm` as settings. If the option `WriteOut` is set to `True`, the result is written out into a file using `Write2` with the setting of `FormatType`.

Replacements are done with `InitialSubstitutions` and `FinalSubstitutions`. Especially energy momentum conservation should be included, e.g., `InitialSubstitutions -> {k2 -> - k1 + p1 + p3}`. Note that the rules listed in `FinalSubstitutions` are not applied as one list of rules, but sequentially in a loop.

If `IsolateHead` is set to `K`, for example, the result will be given as a `K[i]` in `HoldForm`. See `Isolate` for more information. The setting of `Mandelstam` may be, e.g.,

`Mandelstam -> {s, t, u, m1^2 + m2^2 + m3^2 + m4^2}`, where  $s + t + u = m_1^2 + m_2^2 + m_3^2 + m_4^2$ .

The option `CancelQ2` may be altered to `False`, inhibiting then the cancellation of the first propagator in the denominator by replacing  $q^2$  in the numerator by  $q^2 \rightarrow (q^2 - m^2) + m^2$ .

The option `ReduceToScalars` should not be set to `True` when calculating several complicated diagrams involving  $D_{\mu\nu\rho}$  or  $D_{\mu\nu\rho\sigma}$ . Depending on the computer you are using it may nevertheless work, but it is usually better to use `OneLoopSum` with the appropriate options. Note that depending on the setting of the option `BReduce` also two-point coefficient functions may remain in the result.

For processes with light external fermions it is best not to neglect the fermion masses everywhere, but to keep them in the arguments of the scalar Passarino-Veltman functions. This set of masses should be supplied as a list to the option `SmallVariables`, see section 3.3.6.

If `WriteOut` is set to `True`, the result is written out into a file composed of the first argument of `OneLoop`, i.e., the *name*. In which language, i.e., *Mathematica*, *Fortran*, *Macsyma* or *Maple* the result is written, depends on the setting of the option `FormatType`. You may also set `WriteOut` to a string, which denotes the directory to write the result files to.

### ■ 3.5 OneLoopSum and its Options

If you want to sum a list of amplitudes two different possibilities for summing amplitudes are provided with `OneLoopSum`. Either the provided list of amplitudes is calculated and subsequently summed, or the summation occurs partially before calculation. This can be specified with the option `CombineGraphs`.

<pre>OneLoopSum[FeynAmpList[...][FeynAmp[GraphName[... , N1], q, amp1], FeynAmp[GraphName[... , N2], q, amp2], ...]]                                 calculate a list of amplitudes</pre>
<pre>OneLoopSum[expr]                                 sum already calculated amplitudes</pre>

A general function for summing amplitudes. `FeynAmpList`, `FeynAmp` and `GraphName` are heads of *FeynArts*.

The input of `OneLoopSum` is adapted to the output of *FeynArts*. After saving a list of Feynman diagrams created in *FeynArts* with the function `CreateFeynAmp`, i.e. `CreateFeynAmp[ins]>>"eezhhb.amp"`, you can start a new *Mathematica* session<sup>1</sup>, load *FeynCalc* and get the amplitudes by `am = <<"eezhhb.amp"`<sup>2</sup>.

Instead of supplying a list of not yet calculated amplitudes you can also give the sum of already calculated ones as argument (*expr*) to `OneLoopSum`.

<sup>1</sup>It is currently not possible to load *FeynCalc* and *FeynArts* simultaneously into one *Mathematica* session.

<sup>2</sup>Sometimes you will get an error message when loading a file. This is a bug in the *Mathematica* saving routine and may be fixed by editing the file and enclosing the whole expression in two round brackets (...). Do not put a ";" after the last round bracket.

<i>option name</i>	<i>default value</i>	
<code>CombineGraphs</code>	<code>{}</code>	whether to combine amplitudes
<code>IsolateHead</code>	<code>K</code>	whether to <code>Isolate</code> the result
<code>Mandelstam</code>	<code>{}</code>	whether to use the Mandelstam relation
<code>SelectGraphs</code>	<code>All</code>	which graphs to select
<code>ReduceToScalars</code>	<code>True</code>	whether to reduce to scalar integrals
<code>WriteOutPaVe</code>	<code>False</code>	whether to write out the reduced PaVe

Options of `OneLoopSum`

With the default options `OneLoopSum` calculates each amplitude separately by substituting `OneLoop` for `FeynAmp`. Then each single PaVe is reduced to scalar integrals. The hard final part consists in the simplification of the rational coefficients of the scalar integrals. This may involve thousands of factorizations and can therefore take hours of CPU time. But the algebraic simplifications achieved by putting all coefficients of the scalar integrals over a common denominator and to factor them, possibly cancelling factors and reducing the singularity structure, may be very significant. These calculations may need quite a lot of RAM space, therefore the options of `OneLoopSum` allow you to split up the task of summing lots of diagrams.

First you can select a certain subclass of diagrams with the option `SelectGraphs`. You may set, e.g., `SelectGraphs -> {1, 2, 5, 8}`, which selects the amplitudes at positions 1, 2, 5 and 8 of the argument list of `OneLoopSum`. The setting `SelectGraphs -> {1, 2, 5, 8, {10, 40}}` also includes the range of all amplitudes from position 10 to 40.

With the option `CombineGraphs` a possibility is given to sum the graphs before calculation. This is especially useful for combining a graph with its crossed counterpart. In general it makes sense to combine all graphs with the same propagators before calculation, but for very big sums this may reduce the performance considerably. The possible settings for `CombineGraphs` are the same as for `SelectGraphs`. If you use the *FeynArts* syntax for the first argument of `FeynAmp`, i.e. `GraphName[... , N1]`, the last arguments of `GraphName` for combined graphs are concatenated and a new `GraphName` for the summed amplitude is created.

With the setting of the option `WriteOutPaVe -> ""` you can save the result of the reduction of each PaVe in a file for later use. The names of the corresponding files are generated automatically. In case you use `OneLoopSum` several times it recognizes previously saved reductions and loads these results automatically. This may save a considerable amount of time. Instead of setting the option `WriteOutPaVe` to an empty string (which means that the files are written in the current directory), you can specify another directory.

Note that these options together with the possibility of using `OneLoopSum` on already calculated graphs gives you a lot of freedom to split up the calculation, which may be necessary in order to avoid memory overflow.



### ■ 3.6 The Box Graphs of $e^+e^- \rightarrow ZH$

In this section it is shown how to calculate a sum of amplitudes with `OneLoopSum`. The input consists of one page of process-dependent definitions. The unmodified output of *FeynArts* for the amplitudes is directly used as input for *FeynCalc*. The resulting Fortran file is then obtained without any interactive action.

The six standard matrix elements are:

$$\begin{aligned} \mathcal{M}_0^1 &= \bar{v}(p_1) \not{\epsilon}_+ u(p_2) \\ \mathcal{M}_0^2 &= \bar{v}(p_1) \not{\epsilon}_- u(p_2) \\ \mathcal{M}_1^1 &= \bar{v}(p_1) \not{k}_1 \omega_+ u(p_2) \varepsilon p_1 \\ \mathcal{M}_1^2 &= \bar{v}(p_1) \not{k}_1 \omega_- u(p_2) \varepsilon p_1 \\ \mathcal{M}_2^1 &= \bar{v}(p_1) \not{k}_1 \omega_+ u(p_2) \varepsilon p_2 \\ \mathcal{M}_2^2 &= \bar{v}(p_1) \not{k}_1 \omega_- u(p_2) \varepsilon p_2 \end{aligned}$$

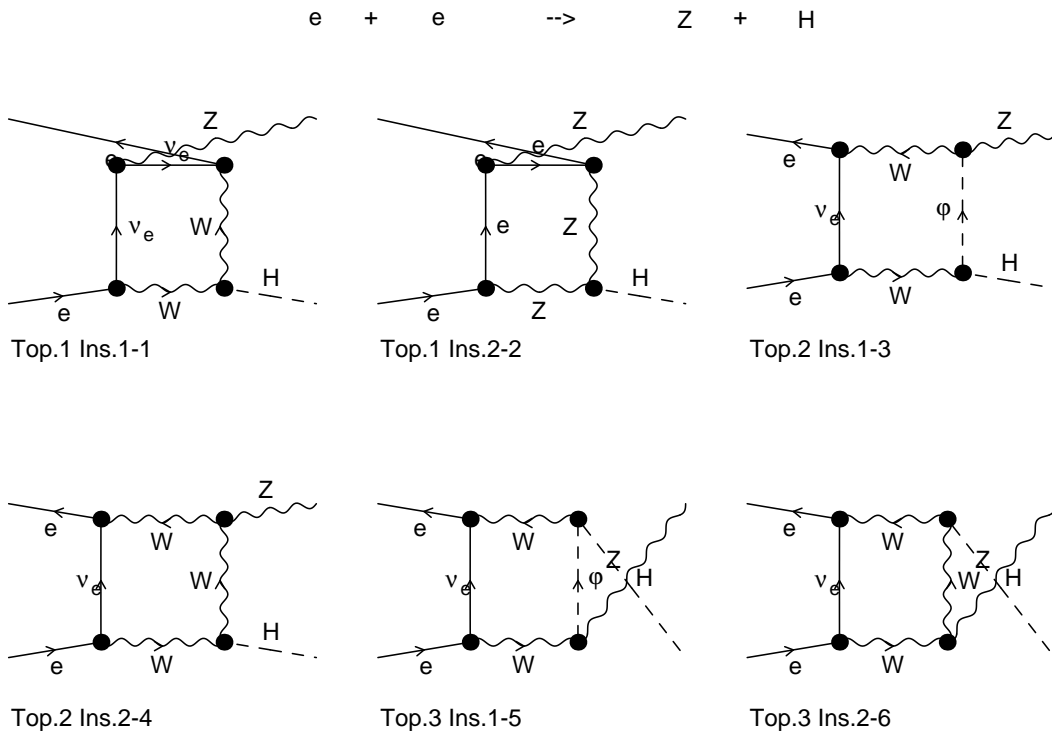


Figure 2: The six box diagrams of  $e^+e^- \rightarrow ZH$ , generated by *FeynArts*.  $\phi$  denotes the unphysical charged Higgs.

The *FeynArts* convention for the momenta is to assign  $p_1$  to the particle north west and count down on the incoming side. The next momentum is then given to the most north eastern particle, here the Z-boson ( $k_1$ ), and the Higgs-boson gets momentum  $k_2$ .

```

(* Define the Mandelstam variables and put momenta on-shell *)
SetMandelstam[S, T, U, p1, p2, -k1, -k2, Small[ME], Small[ME], MZ, MH];

(* Set the option for the ordering of D0's *)
SetOptions[PaVeOrder, PaVeOrderList-> { {S, T}, {S, U}, {T, U} } ];

(* Set the options for OneLoop *)
SetOptions[OneLoop, Mandelstam -> {S, T, U, MH^2 + MZ^2},
  Prefactor -> 1/ALPHA2,
  InitialSubstitutions ->
    { k2 -> p1 + p2 -k1, CW -> MW/MZ,
      EL -> Sqrt[ 4 Pi Sqrt[ ALPHA2] ]
    },
  SmallVariables -> {ME}
];

(* The option for OneLoopSum introduces abbreviations at the end *)
SetOptions[OneLoopSum,
  Prefactor -> 2 ALP4PI FLUFAC,
  Mandelstam -> {S, T, U, MH^2 + MZ^2},
  FinalSubstitutions ->
    { SW -> Sqrt[SW2], ME -> Sqrt[ME2],
      MW -> Sqrt[MW2], MZ -> Sqrt[MZ2], MH -> Sqrt[MH2],
      ME2^n_ -> ME^(2 n)/;Head[n]!=Integer,
      MZ2^n_ -> MZ^(2 n)/;Head[n]!=Integer,
      MW2^n_ -> MW^(2 n)/;Head[n]!=Integer,
      MH2^n_ -> MH^(2 n)/;Head[n]!=Integer,
      SW2^n_ -> SW^(2 n)/;Head[n]!=Integer,
      StandardMatrixElement -> MBM
    }, WriteOutPaVe-> ""
];

(* Define the standard matrixelements *)
SetStandardMatrixElements[
  { Spinor[p1] . DiracSlash[Conjugate[Polarization[k1]]] .
    ChiralityProjector[+1] . Spinor[p2] -> {0, 1},

    Spinor[p1] . DiracSlash[Conjugate[Polarization[k1]]] .
    ChiralityProjector[-1] . Spinor[p2] -> {0, 2},

    ScalarProduct[ Conjugate[Polarization[k1]], p1] *
    Spinor[p1] . DiracSlash[k1] . ChiralityProjector[+1] .
    Spinor[p2] -> {1, 1},

    ScalarProduct[ Conjugate[Polarization[k1]], p1] *
    Spinor[p1] . DiracSlash[k1] . ChiralityProjector[-1] .
    Spinor[p2] -> {1, 2},

    ScalarProduct[ Conjugate[Polarization[k1]], p2] *
    Spinor[p1] . DiracSlash[k1] . ChiralityProjector[+1] .
    Spinor[p2] -> {2, 1},

    ScalarProduct[ Conjugate[Polarization[k1]], p2] *
    Spinor[p1] . DiracSlash[k1] . ChiralityProjector[-1] .
    Spinor[p2] -> {2, 2}
  }, {k2 -> (p1 + p2 - k1)} ];

(* ***** *)

(* get the amplitudes, which have been written to a file by FeynArts *)
eezhamp = <<eezhb.amp;

(* This calculates the amplitudes and sums them up *)
eezhboxes = OneLoopSum[ eezhamp, CombineGraphs -> {1,2,3,4,5,6} ];

(* Here the result is saved and written into a Fortran file *)
Write2["eezhb.s", EEZHBOXES = eezhboxes];
Write2["eezhb.for", EEZHBOXES = eezhboxes, FormatType -> FortranForm];

```

Input program for boxes of  $e^+e^- \rightarrow ZH$ .

```

K(1) = 2*MH2 - MZ2
K(2) = MH2 - 5*MZ2
K(3) = 1 - 2*SW2
K(4) = MH2 + MZ2
K(5) = MH2 + 2*MZ2
K(6) = 4*MH2 + 3*MZ2
K(7) = MH2 - 3*MZ2
K(8) = MH2 - MZ2
K(9) = 2*MH2**2 + 4*MH2*MZ2 + MZ2**2
K(10) = 2*MH2 + MZ2
K(11) = MH2 - 7*MZ2
K(12) = 2*MH2 - 5*MZ2
K(13) = 3*MH2 - 2*MZ2
K(14) = MH2**2 - 2*MH2*MZ2 - MZ2**2
K(15) = 2*MW2 - MZ2
K(16) = 2*MW2**2 - MZ2**2 + 6*MZ2**2*SW2 -
- 12*MZ2**2*SW2**2 + 8*MZ2**2*SW2**3
K(17) = 3*MH2 + 4*MZ2
K(18) = 5*MH2 + 4*MZ2
K(19) = MH2 + 5*MZ2
K(20) = 3*MH2 + 2*MZ2
K(21) = MH2 + 4*MZ2
K(22) = 7*MH2 + 4*MZ2
K(23) = MH - 2*MZ
K(24) = MH + 2*MZ
K(25) = 3*MH2 + 8*MZ2
K(26) = MH2 - 6*MZ2
K(27) = 5*MH2 + 6*MZ2
K(28) = MH2 + 6*MZ2
K(29) = 4*MH2 + MZ2
K(30) = 2*MH2**2 + 13*MH2*MZ2 + 7*MZ2**2
K(31) = 8*MH2 - MZ2
K(32) = 6*MH2**2 + 14*MH2*MZ2 + 3*MZ2**2
K(33) = 6*MH2 + 5*MZ2
K(34) = B0(MH2,MZ,MZ)
K(35) = B0(MZ2,ODO,ODO)
K(36) = B0(MZ2,MW,MW)
K(37) = B0(MH2,MW,MW)
K(38) = D0(MH2,ME2,MZ2,ME2,T,U,MZ2,MZ2,ME2,ME2)
K(39) = D0(MH2,MZ2,ME2,ME2,S,T,MW2,MW2,MW2,ODO)
K(40) = D0(MH2,MZ2,ME2,ME2,S,U,MW2,MW2,MW2,ODO)
K(41) = D0(MH2,ME2,MZ2,ME2,T,U,MW2,MW2,ODO,ODO)
K(42) = C0(MH2,T,ME2,MZ,MZ,ME)
K(43) = C0(MH2,U,ME2,MZ,MZ,ME)
K(44) = C0(MZ2,T,ME2,ME,ME,MZ)
K(45) = C0(MZ2,U,ME2,ME,ME,MZ)
K(46) = C0(MH2,MZ2,S,MW,MW,MW)
K(47) = C0(MH2,T,ME2,MW,MW,ODO)
K(48) = C0(MH2,U,ME2,MW,MW,ODO)
K(49) = C0(MZ2,T,ME2,ODO,ODO,MW)
K(50) = C0(MZ2,T,ME2,MW,MW,ODO)
K(51) = C0(MZ2,U,ME2,ODO,ODO,MW)
K(52) = C0(MZ2,U,ME2,MW,MW,ODO)
K(53) = C0(S,ME2,ME2,MW,MW,ODO)
K(54) = B0(T,ODO,MZ)
K(55) = B0(U,ODO,MZ)
K(56) = B0(T,ODO,MW)
K(57) = B0(U,ODO,MW)
K(58) = MH2*MZ2 - T*U
K(59) = 4*MH2*MZ2**2 - 3*MH2*MZ2*T + MH2*MZ2*U -
- 3*MZ2*T*U + 2*T**2*U - MZ2*U**2
C ---> ... 356 lines omitted ...
K(241) = K(235)*MBM(0,1) + K(236)*MBM(0,2) +
- K(237)*MBM(1,1) + K(238)*MBM(1,2) +
- K(239)*MBM(2,1) + K(240)*MBM(2,2)
K(242) = 2*ALP4PI*FLUFAC*K(241)
EEZHBOXES = K(242)

```

Fortran output file "eezhb.for" generated by OneLoopSum for  $e^+e^- \rightarrow ZH$ .

```

K[1] = 2*MH2 - MZ2
K[2] = MH2 - 5*MZ2
K[3] = 1 - 2*SW2
K[4] = MH2 + MZ2
K[5] = MH2 + 2*MZ2
K[6] = 4*MH2 + 3*MZ2
K[7] = MH2 - 3*MZ2
K[8] = MH2 - MZ2
K[9] = 2*MH2^2 + 4*MH2*MZ2 + MZ2^2
K[10] = 2*MH2 + MZ2
K[11] = MH2 - 7*MZ2
K[12] = 2*MH2 - 5*MZ2
K[13] = 3*MH2 - 2*MZ2
K[14] = MH2^2 - 2*MH2*MZ2 - MZ2^2
K[15] = 2*MW2 - MZ2
K[16] = 2*MW2^2 - MZ2^2 + 6*MZ2^2*SW2 - 12*MZ2^2*SW2^2 + 8*MZ2^2*SW2^3
K[17] = 3*MH2 + 4*MZ2
K[18] = 5*MH2 + 4*MZ2
K[19] = MH2 + 5*MZ2
K[20] = 3*MH2 + 2*MZ2
K[21] = MH2 + 4*MZ2
K[22] = 7*MH2 + 4*MZ2
K[23] = MH - 2*MZ
K[24] = MH + 2*MZ
K[25] = 3*MH2 + 8*MZ2
K[26] = MH2 - 6*MZ2
K[27] = 5*MH2 + 6*MZ2
K[28] = MH2 + 6*MZ2
K[29] = 4*MH2 + MZ2
K[30] = 2*MH2^2 + 13*MH2*MZ2 + 7*MZ2^2
K[31] = 8*MH2 - MZ2
K[32] = 6*MH2^2 + 14*MH2*MZ2 + 3*MZ2^2
K[33] = 6*MH2 + 5*MZ2
K[34] = B0[MH2, MZ2, MZ2]
K[35] = B0[MZ2, 0, 0]
K[36] = B0[MZ2, MW2, MW2]
K[37] = B0[MH2, MW2, MW2]
K[38] = D0[MH2, Small[ME2], MZ2, Small[ME2], T, U, MZ2, MZ2, Small[ME2], Small[ME2]]
K[39] = D0[MH2, MZ2, Small[ME2], Small[ME2], S, T, MW2, MW2, MW2, 0]
K[40] = D0[MH2, MZ2, Small[ME2], Small[ME2], S, U, MW2, MW2, MW2, 0]
K[41] = D0[MH2, Small[ME2], MZ2, Small[ME2], T, U, MW2, MW2, 0, 0]
K[42] = C0[MH2, T, Small[ME2], MZ2, MZ2, Small[ME2]]
K[43] = C0[MH2, U, Small[ME2], MZ2, MZ2, Small[ME2]]
K[44] = C0[MZ2, T, Small[ME2], Small[ME2], Small[ME2], MZ2]
K[45] = C0[MZ2, U, Small[ME2], Small[ME2], Small[ME2], MZ2]
K[46] = C0[MH2, MZ2, S, MW2, MW2, MW2]
K[47] = C0[MH2, T, Small[ME2], MW2, MW2, 0]
K[48] = C0[MH2, U, Small[ME2], MW2, MW2, 0]
K[49] = C0[MZ2, T, Small[ME2], 0, 0, MW2]
K[50] = C0[MZ2, T, Small[ME2], MW2, MW2, 0]
K[51] = C0[MZ2, U, Small[ME2], 0, 0, MW2]
K[52] = C0[MZ2, U, Small[ME2], MW2, MW2, 0]
K[53] = C0[S, Small[ME2], Small[ME2], MW2, MW2, 0]
K[54] = B0[T, 0, MZ2]
K[55] = B0[U, 0, MZ2]
K[56] = B0[T, 0, MW2]
K[57] = B0[U, 0, MW2]
K[58] = MH2*MZ2 - T*U
K[59] = 4*MH2*MZ2^2 - 3*MH2*MZ2*T + MH2*MZ2*U - 3*MZ2*T*U + 2*T^2*U - MZ2*U^2
K[60] = (MZ^5*SW2*HoldForm[K[59]])/(MW2^2*S*HoldForm[K[58]])
(* ..... 338 lines omitted ..... *)
K[241] = HoldForm[K[235]]*MBM[0, 1] + HoldForm[K[236]]*MBM[0, 2] +
  HoldForm[K[237]]*MBM[1, 1] + HoldForm[K[238]]*MBM[1, 2] +
  HoldForm[K[239]]*MBM[2, 1] + HoldForm[K[240]]*MBM[2, 2]
K[242] = 2*ALP4PI*FLUFAC*HoldForm[K[241]]
EEZHBOXES = HoldForm[K[242]]

```

## 4 Miscellaneous Functions

### 4.1 Functions for Polynomial Manipulations

Unfortunately the built-in *Mathematica* 2.0 functions `Factor`, `Collect` and `Together` are either not general enough for the purposes needed in *FeynCalc* or consume too much CPU time for certain polynomials with rational coefficients. The *FeynCalc* extensions `Factor2`, `Collect2` and `Combine` should be used instead when manipulating the rational polynomials emerging from `OneLoop` or `PaVeReduce`.

<code>Collect2[expr, x]</code>	group together powers of terms containing $x$
<code>Collect2[expr, {x<sub>1</sub>, x<sub>2</sub>, ...}]</code>	group together powers of terms containing $x_1, x_2, \dots$
<code>Combine[expr]</code>	put terms in a sum over a common denominator
<code>Factor2[expr]</code>	factor a polynomial in a canonical way

Modifications of `Collect`, `Together` and `Factor`.

This collects  $f[x]$  and  $p[y]$ . The default setting is not to expand terms like  $(b-2)d$ .

```
In[1]:= Collect2[(b - 2) d f[x] + f[x] + c p[y] + p[y], {f, y}]
Out[1]= (1 + (-2 + b) d) f[x] + (1 + c) p[y]
```

With the setting `ProductExpand -> True` the product  $3(s + m^2)$  gets expanded.

```
In[2]:= Collect2[3 B0[pp,m1,m2] (m^2 + s) + B0[pp,m1,m2] s,
                B0, ProductExpand -> True]
Out[2]= (3 m^2 + 4 s) B0[pp, m1, m2]
```

This puts terms over a common denominator without expanding the numerator.

```
In[3]:= Combine[(a - b) (c - d)/e + g]
Out[3]= 
$$\frac{(a - b) (c - d) + e g}{e}$$

```

Consider a generic polynomial.

```
In[4]:= test = (a - b) x + (b - a) y
Out[4]= (a - b) x + (-a + b) y
```

Mapping `Factor` on the summands shows that no canonical factorization of integers in sums takes place.

```
In[5]:= Map[Factor, test]
Out[5]= (a - b) x + (-a + b) y
```

If a canonical factorization is desired, you may use `Factor2` instead.

```
In[6]:= Map[Factor2, test]
Out[6]= (a - b) x - (a - b) y
```

This is the overall factorization of `Factor`.

```
In[7]:= Factor[test]
Out[7]= (-a + b) (-x + y)
```

Here the convention used by `Factor2` becomes clear: the first term in a subsum gets a positive prefactor.

```
In[8]:= Factor2[test]
Out[8]= (a - b) (x - y)
```

<i>option name</i>	<i>default value</i>	
<code>ProductExpand</code>	<code>False</code>	whether to expand products

An expansion controlling option for `Collect2` and `Combine`.

## ■ 4.2 An Isolating Function for Automatically Introducing Abbreviations

<code>Isolate[expr]</code>	substitute an abbreviation $K[i]$ for $expr$ , if $\text{Length}[expr] > 0$
<code>Isolate[expr, {x<sub>1</sub>, x<sub>2</sub>, ...}]</code>	substitute abbreviations for subsums free of $x_1, x_2, \dots$

A function for isolating variables by introducing abbreviations for common subexpressions.

<i>option name</i>	<i>default value</i>	
<code>IsolateHead</code>	<code>K</code>	the head of the abbreviations
<code>IsolateSplit</code>	<code>442</code>	a limit beyond which <code>Isolate</code> splits the expression into two sums

Options for `Isolate` and `Collect2`.

`Isolate` with one argument introduces a single  $K[i]$  as abbreviation.

```
In[1]:= Isolate[a + b]
Out[1]= K[1]
```

Here  $f$  gets isolated with  $K[1]$  and  $K[2]$  replacing the bracketed subsums.

```
In[2]:= test = Isolate[(a + b) f + (c + d) f + e, f]
Out[2]= e + f K[1] + f K[2]
```

Looking at the `FullForm` reveals that the  $K[i]$  are given in `HoldForm`.

```
In[3]:= FullForm[test]
Out[3]//FullForm=
Plus[e, Times[f, HoldForm[K[1]]], Times[f, HoldForm[K[2]]]]
```

Asking for  $K[1]$  returns its value, but in `test K[1]` is held.

```
In[4]:= { K[1], test, ReleaseHold[test] }
Out[4]= {a + b, e + f K[1] + f K[2],
e + (a + b) f + (c + d) f}
```

For the term  $(b + c (y + z))$  a single abbreviation  $F[2]$  is returned.

```
In[5]:= Isolate[a[z] (b + c (y + z)) + d[z] (y + z), {a, d},
IsolateHead -> F]
Out[5]= d[z] F[1] + a[z] F[2]
```

With the help of an additional function it is easy to introduce identical abbreviations for each subsum.

```
In[6]:= ( und[x...] := Isolate[Plus[x], IsolateHead -> H] /;
FreeQ2[{x}, {a, d}];
(a[z] (b + c (y + z)) + d[z] (y + z)) / .
Plus -> und /. und -> Plus
)
```

This trick of selectively substituting functions for sums is also quite useful in special reordering of polynomials.

```
Out[6]= d[z] H[1] + a[z] H[2]
```

Here it is clear that  $H[1] = (y + z)$  is part of  $H[2]$ .

```
In[7]:= ReleaseHold[%]
Out[7]= (y + z) d[z] + a[z] (b + c H[1])
```

Decreasing the option `IsolateSplit` significantly forces `Isolate` to use more than one  $L$  for  $a - b - c - d - e$ .

```
In[8]:= Isolate[ a - b - c - d - e, IsolateHead -> L,
IsolateSplit -> 15 ]
Out[8]= L[2]
```

This shows the values of the  $L$ .

```
In[9]:= {L[2], L[1]}
Out[9]= {-c - d - e + L[1], a - b}
```

The importance of `Isolate` is significant, since it gives you a means to handle very big expressions. The use of `Isolate` on big polynomials before writing them to a file is especially recommended.

A subtle issue of the option `IsolateSplit`, whose setting refers to the `Length[Characters[ToString[FortranForm[expr]]]]`, is that it inhibits too many continuation lines in the Fortran file when writing out expressions involving `K[i]` with `Write2`. See Section 4.4.4 for the usage of `Write2`.

You may want to change `IsolateSplit` when working with big rational polynomials in order to optimize the successive (`FixedPoint`) application of functions like `Combine[ReleaseHold[#]]&`.

### ■ 4.3 An Extension of `FreeQ` and Two Other Useful Functions

<code>FreeQ2[expr, {f<sub>1</sub>, f<sub>2</sub>, ...}]</code>	yields <code>True</code> if <code>expr</code> does not contain any occurrence of <code>f<sub>1</sub>, f<sub>2</sub>, ...</code>
<code>NumericalFactor[expr]</code>	gives the numerical factor of <code>expr</code>
<code>PartitHead[expr, h]</code>	returns a list <code>{a, h[b]}</code> with <code>a</code> free of expressions with head <code>h</code> , and <code>h[b]</code> having head <code>h</code>

Three useful functions.

`FreeQ2` is an extension of `FreeQ`, allowing a list as second argument.

```
In[1]:= FreeQ2[M^2 + m^2 B0[pp, m1, m2], {M, B0}]
Out[1]= False
```

This gives the numerical factor of the expression.

```
In[2]:= NumericalFactor[-137 x]
Out[2]= -137
```

The action of `PartitHead` on a product is to split the product apart.

```
In[3]:= PartitHead[f[m] (s - u), f]
Out[3]= {s - u, f[m]}
```

A sum gets separated into subsums.

```
In[4]:= PartitHead[s^2 + M^2 - f[m], f]
Out[4]= {M^2 + s^2, -f[m]}
```

### ■ 4.4 Writing Out to *Mathematica*, Fortran, Macsyma and Maple

The *Mathematica* functions `Write` and `Save` may on rare occasions create files that cannot be read in again correctly. What sometimes happens is that, for example, a division operator is written out as the first character of a line. When the file is loaded again, *Mathematica* may simply ignore that part of the file before this character. You can easily work around this bug by editing the file and wrapping the expressions with brackets “()”. Since this is a cumbersome procedure for lots of expressions, it has been automatized in `Write2` for writing out expressions. If you want to use `Save`, you have to check the output file for correctness.

An option can be given to `Write2` allowing the output to be written in `FortranForm`, `MacsymaForm` or `MapleForm`. These facilities are elementary and mostly limited to the type of polynomials usually encountered in *FeynCalc*.

The `FortranForm` option should not be used if the expression to be written out contains a term  $(-x)^n$ , where  $n$  is a symbol, (which is never produced by *FeynCalc* unless you enter it). The *Mathematica* `FortranForm` is also incapable of recognizing some elementary functions like `Exp[x]`. Therefore if you want to generate more elaborate Fortran code you may want to use *Maple*, which provides an optimization option when translating *Maple* code to Fortran, or the most sophisticated package for Fortran code generation from computer algebra systems: *Gentran* (by B. Gates and H. van Hulzen), which is available in some *Macsyma* versions. To this end the `Write2` option `FormatType` may be set to `MacsymaForm` or `MapleForm`.

You may achieve a rudimentary optimization by applying `Isolate` on the expression to be written out. With the option setting `FortranForm` and `InputForm` the function `Write2` recognizes variables in `HoldForm` and writes these out with their definitions first.

<code>Write2[fi, x = y]</code>	write the setting $x = y$ into a file $fi$
<code>Write2[fi, x = y, a = b, ...]</code>	write the setting $x = y$ into a file $fi$

A modification of `Write`.

<i>option name</i>	<i>default value</i>	
<code>FormatType</code>	<code>InputForm</code>	language in which to write the output
<code>D0Convention</code>	0	which convention to use for the scalar integrals

Options for `Write2`.

With the default setting 0 of `D0Convention` the arguments of the scalar integrals are not changed when written into a Fortran program. Another possible setting is 1, which interchanges the fifth and sixth arguments of `D0` and writes the mass arguments of `A0`, `B0`, `C0` and `D0` without squares.

<code>InputForm</code>	write out in <i>Mathematica</i> form
<code>FortranForm</code>	write out in Fortran form
<code>MacsymaForm</code>	write out in <i>Macsyma</i> form
<code>MapleForm</code>	write out in <i>Maple</i> form

The four possible settings of the option `FormatType` for `Write2`.

The output possibilities for the two other computer algebra systems *Macsyma* and *Maple* are very limited: square brackets in *Mathematica* are changed to round brackets, additional replacements are for `MacsymaForm` { `Pi` -> `%pi`, `I` -> `%i`, `=` -> `:` } and for `MapleForm` { `=` -> `:=` }.

The reasons to include these interface abilities are to utilize the Fortran optimization possibility of *Maple* and the excellent package *Gentran* for Fortran code generation.

Create a test polynomial.

```
In[1]:= tpol = z + Isolate[Isolate[2 Pi I + f[x] (a - b), f]]
Out[1]= z + K[2]
```

The default writes out in *Mathematica* `InputForm`.

```
In[2]:= Write2["test.m", test = tpol];
```

Show the content of the file.

```
In[3]:= !!test.m
K[1] = a - b
K[2] = 2*I*Pi + f[x]*HoldForm[K[1]]
test = z + HoldForm[K[2]]
```

This writes a file in Fortran format.

```
In[3]:= Write2["test.for", test = tpol,
             FormatType -> FortranForm];
```



Show the content of the Fortran file.

```
In[4]:= !!test.for
      K(1) = a - b
      K(2) = (0,2)*Pi + f(x)*K(1)
      test = z + K(2)
```

This writes a file in *Macysma* format.

```
In[4]:= Write2["test.mac", test = tpol,
      FormatType -> MacsmaForm];
```

Here the *Macysma* format can be seen.

```
In[5]:= !!test.mac
test : ( 2*i*%pi + z + (a - b)*f(x) )$
```

Here we get a file with *Maple* conventions.

```
In[5]:= Write2["test.map", test = tpol,
      FormatType -> MapleForm];
```

The *Maple* conventions are only slightly different.

```
In[6]:= !!test.map
test := ( 2*I*Pi + z + (a - b)*f(x) );
```

## ■ 4.5 More on Levi-Civita Tensors

**EpsEvaluate** [*expr*]    evaluate Levi-Civita Eps

A function for total antisymmetry and linearity with respect to Momentum.

This is  $\varepsilon^{\mu\nu\rho\sigma}(p+q)_\sigma = \varepsilon^{\mu\nu\rho(p+q)}$ .

```
In[1]:= Contract[ LeviCivita[m, n, r, s] FourVector[p + q, s] ]
Out[1]= eps[m, n, r, p + q]
```

In this way linearity is enforced:  
 $\varepsilon^{\mu\nu\rho(p+q)} = \varepsilon^{\mu\nu\rho p} + \varepsilon^{\mu\nu\rho q}$ .

```
In[2]:= EpsEvaluate[ % ]
Out[2]= eps[m, n, r, p] + eps[m, n, r, q]
```

This does not evaluate directly to 0.

```
In[3]:= LeviCivita[a, b, c, d] /. d -> c
Out[3]= eps[a, b, c, c]
```

Like this you can always evaluate  $\varepsilon$ 's.

```
In[4]:= EpsEvaluate[ % ]
Out[4]= 0
```

**EpsChisholm** [*expr*]    utilize  $\gamma_\mu \varepsilon^{\mu\nu\rho\sigma} = +i(\gamma^\nu \gamma^\rho \gamma^\sigma - g^{\nu\rho} \gamma^\sigma - g^{\rho\sigma} \gamma^\nu + g^{\nu\sigma} \gamma^\rho) \gamma^5$

A function for the Chisholm identity.

<i>option name</i>	<i>default value</i>	
LeviCivitaSign	-1	sign convention for the $\varepsilon$ -tensor

An option for **EpsChisholm**

Use the Chisholm identity for  $\varepsilon^{abcd} \gamma^\mu \gamma^\nu$ .

```
In[5]:= EpsChisholm[ LeviCivita[a, b, c, d] DiracMatrix[mu, a] ]
Out[5]= +I (-g[c, d] ga[mu] ga[b] ga[5] +
      g[b, d] ga[mu] ga[c] ga[5] -
      g[b, c] ga[mu] ga[d] ga[5] +
      ga[mu] ga[b] ga[c] ga[d] ga[5])
```

`Eps[a, b, c, d]` internal representation of Levi-Civita tensors, where the arguments must have head `LorentzIndex` or `Momentum`

The internal function for Levi-Civita tensors, see also section 4.4.9.

This clears `$PrePrint`.

```
In[6]:= $PrePrint=.
```

This shows the internal structure.

```
In[7]:= LeviCivita[m, n, r, s]
Out[7]= Eps[LorentzIndex[m], LorentzIndex[n],
LorentzIndex[r], LorentzIndex[s]]
```

Contracting  $\varepsilon^{\nu\rho\sigma} p^\sigma$  yields  $\varepsilon^{\mu\nu\rho p}$ .

```
In[8]:= Contract[% FourVector[p, s]]
```

In the internal representation the  $p$  has the head `Momentum` wrapped around it.

```
Out[8]= Eps[LorentzIndex[m], LorentzIndex[n],
LorentzIndex[r], Momentum[p]]
```

## ■ 4.6 Polarization Sums

<code>PolarizationSum[mu, nu]</code>	$-g^{\mu\nu}$
<code>PolarizationSum[mu, nu, k]</code>	$-g^{\mu\nu} + k^\mu k^\nu / k^2$
<code>PolarizationSum[mu, nu, k, n]</code>	$-g^{\mu\nu} - k_\mu k_\nu n^2 / (k \cdot n)^2 + (n_\mu k_\nu + n_\nu k_\mu) / (k \cdot n)$

Three polarization sums.

This is the polarization sum for massive bosons:  $\Sigma \varepsilon_\mu \varepsilon_\nu^* = -g^{\mu\nu} + k^\mu k^\nu / k^2$ .

```
In[1]:= PolarizationSum[mu, nu, k]
```

```
Out[1]= -g[mu, nu] + (k[mu] k[nu]) / (k.k)
```

Here the polarization sum for gluons is given; with external momentum  $n = p_1 - p_2$ .

```
In[2]:= PolarizationSum[mu, nu, k, p1 - p2]
```

```
Out[2]= -g[mu, nu] - ((p1.p1 - 2 p1.p2 + p2.p2) k[mu] k[nu]) / ((k.p1 - k.p2)^2) +
(k[nu] (p1 - p2)[mu] + k[mu] (p1 - p2)[nu]) / (k.p1 - k.p2)
```

## ■ 4.7 Simplifications of Expressions with Mandelstam Variables

`TrickMandelstam[expr, {s, t, u, m1^2 + m2^2 + m3^2 + m4^2}]`  
simplifies `expr` to the shortest form removing `s, t` or `u` in each sum using  $s + t + u = m_1^2 + m_2^2 + m_3^2 + m_4^2$

For tricky Mandelstam variable substitution.

This is an easy example of simplifications done by `TrickMandelstam`.

```
In[1]:= TrickMandelstam[(s + t - u) (2 mw2 - t - u),
{s, t, u, 2 mw2}]
```

```
Out[1]= 2 s (mw2 - u)
```

The result is always given in a factorized form.

```
In[2]:= TrickMandelstam[M^2 s - s^2 + M^2 t - s t + M^2 u - s u,
                        {s, t, u, 2 M^2}]
```

```
Out[2]= 2 M^2 (M^2 - s)
```

## ■ 4.8 Permuting the Arguments of the Four-Point Function

The arguments of the Passarino-Veltman four-point function  $D_0$  are permuted by *FeynCalc* into an alphabetical order. If you want a specific permutation of the 24 possible ones, you can use the function `PaVeOrder`.

`PaVeOrder[expr]` order the arguments of  $C_0$  and  $D_0$  in *expr* canonically.

An ordering function for  $C_0$  and  $D_0$ .

option name	default value	
<code>PaVeOrderList</code>	<code>{}</code>	order $D_0$ according to <code>{..., a, b, ...}</code>

An ordering option for `PaVeOrder`.

With the default setting of `PaVeOrder` a canonical ordering is chosen.

It is sufficient to supply a subset of the arguments.

```
In[1]:= PaVeOrder[D0[me2, me2, mw2, mw2, t, s, me2, 0, me2, 0],
                  PaVeOrderList -> {me2, me2, 0, 0}]
```

```
Out[1]= D0[me2, s, mw2, t, mw2, me2, me2, 0, 0, me2]
```

This interchanges the  $f$  and  $e$ .

```
In[2]:= PaVeOrder[D0[a, b, c, d, e, f, m12, m22, m32, m42],
                  PaVeOrderList -> {f, e}]
```

```
Out[2]= D0[a, d, c, b, f, e, m22, m12, m42, m32]
```

This shows how to permute several  $D_0$ .

```
In[3]:= PaVeOrder[ D0[a, b, c, d, e, f, m12, m22, m32, m42] +
                  D0[me2, me2, mw2, mw2, t, s, me2, 0, me2, 0],
                  PaVeOrderList-> { {me2, me2, 0, 0}, {f, e} }
                  ]
```

```
Out[3]= D0[a, d, c, b, f, e, m22, m12, m42, m32] +
        D0[me2, s, mw2, t, mw2, me2, me2, 0, 0, me2]
```

## ■ 4.9 On the Internal Representation

With the default setting of `$PrePrint = FeynCalcForm` the internal representation is hidden. However for special purposes it may be useful to know how the various objects are represented in *FeynCalc*. The basic idea is to wrap a special head around each special input variable. For example, the arguments of `MetricTensor[μ, ν]` each get a head `LorentzIndex`. The data-type heads usually have only a few functional definitions. A  $D$ -dimensional `LorentzIndex[μ, D]`, for example, simplifies to `LorentzIndex[μ]` when  $D$  is replaced by 4.

Inside *FeynCalc* the functions and rules are specified in such a way that they only apply to those objects with the right head. This programming style, which makes strong use of the pattern matching capabilities of *Mathematica*, especially the so-called upvalue function definitions, has proven to be very useful and reliable for the purpose of calculations in physics.

The exported heads, which often also have simple functional definitions, are listed briefly in this section.

<code>DiracGamma[x]</code>	head of a four-dimensional $\gamma^\mu$ or $\not{p}$
<code>DiracGamma[x, d]</code>	head of a $d$ -dimensional $\gamma^\mu$ or $\not{p}$
<code>LorentzIndex[mu]</code>	head of a four-dimensional Lorentz index $\mu$
<code>LorentzIndex[mu, d]</code>	head of a $d$ -dimensional Lorentz index $\mu$
<code>Momentum[p]</code>	head of a four-dimensional momentum $p$
<code>Momentum[p, d]</code>	head of a $d$ -dimensional momentum $p$

Heads of the internal representation.

Clear `$PrePrint`.

```
In[1]:= $PrePrint=.
```

This is the internal representation of a  $\gamma^\mu$  in four dimensions.

```
In[2]:= DiracMatrix[mu]
Out[2]= DiracGamma[LorentzIndex[mu]]
```

Here is a  $\gamma^\alpha$  in  $D$  dimensions.

```
In[3]:= DiracMatrix[al, Dimension -> D]
Out[3]= DiracGamma[LorentzIndex[al, D], D]
```

This a product of a four-dimensional  $\not{q}$  and a  $D$ -dimensional  $\not{q}$ .

```
In[4]:= DiracSlash[q] . DiracSlash[q, Dimension -> D]
Out[4]= DiracGamma[Momentum[q]] .
        DiracGamma[Momentum[q, D], D]
```

This shows the convention for a four-dimensional scalar product.

```
In[5]:= DiracSimplify[%]
Out[5]= Pair[Momentum[q], Momentum[q]]
```

The head `Pair` is also used for metric tensors.

```
In[6]:= MetricTensor[mu, nu]
Out[6]= Pair[LorentzIndex[mu], LorentzIndex[nu]]
```

Clearly a four-vector is also represented as a pair internally.

```
In[7]:= FourVector[l, mu]
Out[7]= Pair[LorentzIndex[mu], Momentum[l]]
```

The dimension is provided as an argument.

```
In[8]:= MetricTensor[a, b, Dimension -> d]
Out[8]= Pair[LorentzIndex[a, d], LorentzIndex[b, d]]
```

A  $(D - 4)$ -dimensional four-vector.

```
In[9]:= FourVector[p, mu, Dimension -> D - 4]
Out[9]= Pair[LorentzIndex[mu, -4 + D], Momentum[p, -4 + D]]
```

Obviously it vanishes after changing the dimension.

```
In[10]:= % /. D -> 4
Out[10]= 0
```

This shows the internal representations of polarization vectors.

```
In[11]:= PolarizationVector[k, mu]
Out[11]= Pair[LorentzIndex[mu], Momentum[Polarization[k]]]
```

<code>Pair[a, b]</code>	the head of $g^{\mu\nu}$ , $p^\mu$ and $(p \cdot q)$
-------------------------	--

The head for a pairing of `LorentzIndex` and `Momentum`

## ■ 4.10 FeynCalcForm

<code>FeynCalcForm[expr]</code> displays <i>expr</i> in an easy to read form
--

The print form of *FeynCalc*.

`FeynCalcForm` changes the internal representation of *FeynCalc* to a shorter and more readable form. The effect of `FeynCalcForm` can be investigated by clearing `$PrePrint` (`$PrePrint=.`) and applying `FeynCalcForm` on the various objects.

The default setting of `$PrePrint` is `FeynCalcForm`.

The effect of `FeynCalcForm` can be seen in the examples.

## ■ 4.11 Three New Global Variables

<i>option name</i>	<i>default value</i>	
<code>\$BreitMaison</code>	False	whether to use the Breitenlohner-Maison scheme
<code>\$VeryVerbose</code>	0	display intermediate messages from <i>FeynCalc</i>
<code>\$MemoryAvailable</code>	8	the amount of available main memory in MB

The three global variables of *FeynCalc*.

The variable `$VeryVerbose` may be set to 0, 1, 2 or 3. The higher the setting the more intermediate information is printed during the calculations.

*FeynCalc* is designed in such a way that certain intermediate results are stored in order to speed up the computations. This of course may become too memory consuming. The storing stops, when the value of  $10^{(-6)} * \text{MemoryInUse}[]$  gets bigger than `$MemoryAvailable-1`.

The default of *FeynCalc* is to use the naive  $\gamma^5$  prescription, i.e.,  $\gamma^5$  anticommutes with  $D$  - dimensional Dirac matrices. Setting `$BreitMaison` to `True` changes the commutation behavior of  $\gamma^5$ : It anti-commutes with Dirac matrices in four dimensions, but commutes with the  $(D-4)$ -dimensional part. The basic features of the Breitenlohner-Maison scheme are implemented in *FeynCalc* but they have not very thoroughly been tested. Therefore one should check the results for correctness.

If you want to change the default setting of `BreitMaison` you must set it *before* loading *FeynCalc*.

In the following example the automatic loading of *FeynCalc* from the `init.m` file has been turned off.

Set the variable <code>\$BreitMaison</code> before loading <i>FeynCalc</i> .	<code>In[1]:= \$BreitMaison = True</code> <code>Out[1]= True</code>
Loading <i>FeynCalc</i> .	<code>In[2]:= &lt;&lt;FeynCalc.m</code>
Entering $\gamma^5 \gamma^\mu$ , with $\gamma^\mu$ $D$ -dimensional.	<code>In[3]:= DiracMatrix[5] . DiracMatrix[mu, Dimension -&gt;D]</code> <code>Out[3]= ga[5] ga [mu]</code> <div style="text-align: center;"><math>\mu</math></div>

Now only the 4-dimensional part of  $\gamma^\mu$  anticommutes, while the  $D - 4$  dimensional one commutes.

```
In[4]:= DiracSimplify[%]
Out[4]= -ga[mu] ga[5] + ga[mu] ga[5]
          D-4
```

The chirality projectors  $\omega_+$  and  $\omega_-$  are inserted automatically.

```
In[5]:= ChiralityProjector[+1] . DiracMatrix[mu, Dimension ->D]
Out[5]= (- + ga[5]) ga[mu]
          2 2 D
```

The expression is expanded and  $\gamma^5$  is moved to the right.

```
In[6]:= DiracSimplify[%]
Out[6]= -ga[mu] ga[5] ga[mu] ga[5] ga[mu]
          2 D-4 D 2
```

This is *FeynCalc* does not simplify this sum directly, but you can easily specify which dimension to "eliminate".

```
In[7]:= test = 2 DiracMatrix[mu] +
            DiracMatrix[mu, Dimension -> D] +
            DiracMatrix[mu, Dimension -> D-4]
Out[7]= 2 ga[mu] + ga[mu] + ga[mu]
          D-4 D
```

Switch to the internal representation.

```
In[8]:= ($PrePrint=. ; test)
Out[8]= 2 DiracGamma[LorentzIndex[mu]] +
            DiracGamma[LorentzIndex[mu, -4 + D], -4 + D] +
            DiracGamma[LorentzIndex[mu, D], D]
```

This eliminates each Dirac matrix in  $D - 4$  dimension.

```
In[9]:= DiracGamma[x_[y_., d_Symbol - 4], d_Symbol - 4] :=
            DiracGamma[x[y, d], d] - DiracGamma[x[y]]
```

Only  $D$ - and 4-dimensional objects are left.

```
In[10]:= test
Out[10]= DiracGamma[LorentzIndex[mu]] +
            2 DiracGamma[LorentzIndex[mu, D], D]
```

## References

- [1] J. Küblbeck, M. Böhm and A. Denner, *Comp. Phys. Comm.* **60** (1990) 165.
- [2] A. Denner, Techniques for the Calculation of Electroweak Radiative Corrections at the One-Loop Level and Results for  $W$ -Physics at LEP200, to appear in *Fortschritte der Physik 1992, 41* and references therein.
- [3] R. Mertig, M. Böhm, and A. Denner, *Comp. Phys. Comm.* **64** (1991) 345.
- [4] E. Yehudai and A. C. K. Hsieh, *HIP* — Symbolic High-Energy Physics Calculations, *SLAC-PUB- 5576* July 1991.
- [5] G. J. van Oldenborgh, FF – a package to evaluate one-loop Feynman diagrams, *Comp. Phys. Comm.* **66** (1991) 1.
- [6] P. Cvitanovic, *Phys. Rev. D* **14** (1976) 1536.
- [7] A.P. Kryukov and A.Ya. Rodionov, *Comp. Phys. Comm.* **48** (1988) 327.
- [8] Matthias Jamin and E. Lautenbacher, TRACER, A Mathematica Package for  $\gamma$ -Algebra in Arbitrary Dimensions, preprint Technische Universität München, TUM-T31-20/91.
- [9] S. Wolfram, MACSYMA Tools for Feynman Diagram Calculations, *Proceedings of the 1979 MACSYMA Users Conference*.

## 5 Reference Guide for FeynCalc

### ■ A0

`A0[m^2]` is the scalar Passarino-Veltman one-point function.

`A0[m^2]` is an abbreviation for  $A_0(m^2) = -i\pi^{-2} \int d^D q (2\mu\pi)^{4-D} [q^2 - m^2]^{-1}$ . ■ `A0[0]` and `A0[Small[mass^2]]` give 0.

■ The following option can be given:

`A0ToB0` `True` whether to replace  $A_0(m^2)$  by  $m^2(1 + B_0(0, m^2, m^2))$

■ See page 21.

### ■ A0ToB0

`A0ToB0` is an option for `A0`. If set to `True`,  $A_0(m^2)$  is replaced by  $m^2(1 + B_0(0, m^2, m^2))$ .

■ See page 23.

### ■ B0

`B0[pp, m1^2, m2^2]` is the scalar Passarino-Veltman two-point function.

`B0[pp, m1^2, m2^2]` is an abbreviation for  $B_0(p^2, m_1^2, m_2^2) = -i\pi^{-2} \int d^D q (2\mu\pi)^{4-D} ([q^2 - m_1^2][(q+p)^2 - m_2^2])^{-1}$ . ■ `B0` is symmetric in its second and third argument. ■ The following options can be given:

`BReduce` `False` whether to reduce `B0` in special cases to `A0`

`B0Unique` `False` whether to replace  $B_0(a, 0, a)$  by  $(2 + B_0(0, a, a))$  and  $B_0(0, 0, a)$  by  $(1 + B_0(0, a, a))$ .

■ See page 21.

### ■ B00

`B00[pp, m1^2, m2^2]` is the coefficient of  $g^{\mu\nu}$  of the tensor integral decomposition of  $B^{\mu\nu}$ .

The following option can be given:

`BReduce` `True` whether to reduce `B00` to `B1` and `A0`

■ If `BReduce` is set to `True` the following simplification holds:

■  $B_{00}(a, b, c) = 1/6(A_0(b) + (B_1(a, b, c)(a - c + b) + a/3B_0(a, b, c) + 1/6(a + b - c/3)))$ . ■ See page 23.

### ■ B0Unique

`B0Unique` is an option for `B0`.

Sometimes it is useful to set this option to `True`, since only then all simplifications between different  $B_0$  occur. ■ See page 23.

### ■ B1

`B1[pp, m1^2, m2^2]` is the coefficient of  $p^\mu$  of the tensor integral decomposition of  $B^\mu$ .

The following option can be given:

`BReduce` `True` whether to reduce `B1` to `B0` and `A0`

■ If a variable of  $B_1$  has head `Small` and at least one other variable does not (and is different from 0), the variable with head `Small` is set to 0. ■ If the option `BReduce` is set to `True`, `B1` simplifies in the following way, where  $a, b, c$  are  $m^2$  with no head `Small`. The same simplifications are performed if instead of 0 a `Small` variable is supplied as an argument.

■  $B_1(a, b, c) = 1/2(A_0(b) - A_0(b) - (a - c + b)B_0(a, b, c))$ . ■  $B_1(a, b, b) = -1/2B_0(a, b, b)$ .

■  $B_1(a, a, 0) = -1/2B_0(a, a, 0) - 1/2$ . ■  $B_1(a, 0, a) = 1/2 - 1/2B_0(a, 0, m)$ . ■  $B_1(0, 0, a) = -1/2B_0(0, 0, a) + 1/4$ .

■  $B_1(0, a, 0) = -1/2B_0(0, a, 0) - 1/4$ . ■ See page 23.

### ■ B11

`B11[pp, m1^2, m2^2]` is the coefficient of  $p^\mu p^\nu$  of the tensor integral decomposition of  $B^{\mu\nu}$ .

The following option can be given:

`BReduce` `True` whether to reduce `B11` to `B1` and `A0`

If the option `BReduce` is set to `True`, `B11` simplifies in the following way, where  $a, b, c$  are  $m^2$  with no head `Small`.

■  $B_{11}(a, b, c) = 1/(3a)(A_0(b) - 2(a - c + b)B_1(a, b, c) - aB_0(a, b, c) - 1/2(a + b - c/3))$ . ■  $B_{11}(0, a, a) = 1/3B_0(0, a, a)$ . ■ See page 23.

## ■ BReduce

**BReduce** is an option for **B0**, **B00**, **B1** and **B11**, determining whether reductions to lower-order **A** and **B** are done.

The default setting of **BReduce** is **True**. ■ See page 23.

## ■ CancelQ2

**CancelQ2** is an option for **OneLoop**. If set to **True**, cancellation of all  $q^2$  with the first propagator via  $q^2 \rightarrow ((q^2 - m^2) + m^2)$  is performed, where  $q$  denotes the integration momentum.

With the default **True** the translation of the integration momentum in the lower order Passarino Veltman functions is done such that the third mass argument of the 4-point integral is put at position 1. ■ See page 30.

## ■ CancelQP

**CancelQP** is an option for **OneLoop**. If set to **True**, cancellation of  $q \cdot p$  with propagators is performed, where  $q$  denotes the integration momentum.

■ See page 30.

## ■ ChiralityProjector

**ChiralityProjector**[+1] is an alternative input for  $\omega_+ = \frac{1}{2}(1 + \gamma^5)$ . **ChiralityProjector**[-1] denotes  $\omega_- = \frac{1}{2}(1 - \gamma^5)$ .

**ChiralityProjector**[1] is identical to **DiracMatrix**[6]. **ChiralityProjector**[-1] is identical to **DiracMatrix**[7]. The internal representation is **DiracGamma**[6] and **DiracGamma**[7]. ■ See page 9. ■ See also: **DiracMatrix**, **DiracGamma**.

## ■ C0

**C0**[ $p_{10}, p_{12}, p_{20}, m_1^2, m_2^2, m_3^2$ ] is the scalar Passarino-Veltman three-point function. The first three arguments of **C0** are the scalar products  $p_{10} = p_1^2$ ,  $p_{12} = (p_1 - p_2)^2$ ,  $p_{20} = p_2^2$ .

**C0**[ $p_{10}, p_{12}, p_{20}, m_1^2, m_2^2, m_3^2$ ] is an abbreviation for

$C_0 = -i\pi^{-2} \int d^D q (2\mu\pi)^{4-D} ([q^2 - m_1^2][(q + p_1)^2 - m_2^2][(q + p_2)^2 - m_3^2])^{-1}$ . ■ A standard representative of the six equivalent argument permutations is chosen. ■ See page 21. ■ See also: **PaVeOrder**.

## ■ Collect2

**Collect2**[*expr*, *x*] collects together terms which are not free of any occurrence of *x*.

**Collect2**[*expr*, {*x*<sub>1</sub>, *x*<sub>2</sub>, ...}] collects together terms which are not free of *x*<sub>1</sub>, *x*<sub>2</sub>, ...

The following option can be given.

<b>ProductExpand</b>	<b>False</b>	whether to expand products in <i>expr</i> free of <i>x</i> ( <i>x</i> <sub>1</sub> , <i>x</i> <sub>2</sub> , ...)
<b>IsolateHead</b>	<b>False</b>	whether to isolate with respect to { <i>x</i> <sub>1</sub> , <i>x</i> <sub>2</sub> , ...}
<b>IsolateSplit</b>	<b>442</b>	the limit before <b>Isolate</b> splits

■ See page 37. ■ See also: **Isolate**.

## ■ Combine

**Combine**[*expr*] puts terms in a sum over a common denominator. **Combine** is similar to **Together**, but works better on certain polynomials with rational coefficients.

The following option can be given:

<b>ProductExpand</b>	<b>False</b>	whether to expand products
----------------------	--------------	----------------------------

■ See page 37.

## ■ CombineGraphs

**CombineGraphs** is an option for **OneLoopSum**.

The utilization of this option may speed up *FeynCalc*. But depending on the available memory there is a turning point where *FeynCalc* becomes quite inefficient when forced to calculate too many complicated diagrams at once. ■ See page 31.



## Contract

`Contract[expr]` contracts pairs of Lorentz indices in `expr`.

For the contraction of two Dirac matrices `DiracSimplify` has to be used. ■ The result of `Contract` is not fully expanded.

■ The following options can be given:

`EpsContract` `False` whether to contract products of Levi-Civita (Eps) tensors via the determinant formula

`Expanding` `True` expand all sums containing Lorentz indices

`Factoring` `False` whether to factor the result canonically

■ Examples: `Contract[FourVector[p, mu]^2]` → `p.p.`

■ `Contract[FourVector[p + q, mu] FourVector[p' + q', mu]]//ExpandScalarProduct` → `p.p' + p.q' + q.p' + q.q'`.

■ `Contract[LeviCivita[m, n, r, a] LeviCivita[m, n, r, b], EpsContract -> True]` → `-6 g[a, b]`.

■ `Contract[Pair[LorentzIndex[mu, Dim], LorentzIndex[mu, Dim]]]` → `Dim`.

■ `Contract[Pair[LorentzIndex[mu, D - 4], LorentzIndex[mu]]]` → `4`.

■ `Contract[Pair[LorentzIndex[mu, D - 4], LorentzIndex[mu]]]` → `0`.

■ `Contract[Pair[LorentzIndex[mu, D - 4], LorentzIndex[mu, D - 4]]]` → `-4 + D`.

■ `Contract[f[___, LorentzIndex[mu], ___] MetricTensor[mu, al]]` →

`f[___, LorentzIndex[al], ___]`. ■ `Contract[f[___, LorentzIndex[mu], ___] FourVector[p, mu]]` → `f[___, FourVector[p], ___]`.

■ `Contract[DiracMatrix[mu, Dimension -> D] MetricTensor[mu, al]]` →

`DiracGamma[LorentzIndex[nu]]`.

■ `Contract[DiracGamma[LorentzIndex[mu, D-4], D-4] MetricTensor[mu, al]]` → `0`. ■ If big expressions are contracted and substitutions for the resulting scalar products are made, it is best not to do these replacements after contraction, but to set the values of the relevant scalar products *before* invoking `Contract`; in this way the intermediate expression swell is minimized. ■ See page 13. ■ See also: `ExpandScalarProduct`.

## D0

`D0[p10, p12, p23, p30, p20, p13, m12, m22, m32, m42]` is the scalar Passarino-Veltman four-point function. The first six arguments of `D0` are the scalar products

$$p_{10} = p_1^2, p_{12} = (p_1 - p_2)^2, p_{23} = (p_2 - p_3)^2, p_{30} = p_3^2, p_{20} = p_2^2, p_{13} = (p_1 - p_3)^2.$$

`D0[p10, p12, p23, p30, p20, p13, m12, m22, m32, m42]` is an abbreviation for

$$D_0 = -i\pi^{-2} \int d^4q [(q^2 - m_1^2)[(q + p_1)^2 - m_2^2][(q + p_2)^2 - m_3^2][(q + p_3)^2 - m_4^2]^{-1}. \quad \blacksquare \text{ See page 21. } \blacksquare \text{ See also: PaVe, PaVeOrder.}$$

## D0Convention

`D0Convention` is an option for `Write2`. Possible settings are 0 or 1. With the last setting the fifth and sixth arguments of `D0` are interchanged and all (internal) mass arguments of the scalar Passarino Veltman integrals are given square free.

■ See page 40. ■ See also: `Write2`, `D0`.

## DB0

`DB0[p10, m02, m12]` is the derivative  $\partial B_0(p^2, m_0^2, m_1^2)/\partial p^2$  of the two-point function `B0`.

■ See page 21. ■ See also: `B0`.

## DenominatorOrder

`DenominatorOrder` is an option for `OneLoop`, if set to `True` the `PropagatorDenominator` in `FeynAmpDenominator` will be ordered in a standard way.

You may want to set this option to `False` when checking hand calculations. ■ See page 30.

## Dimension

`Dimension` is an option for `DiracMatrix`, `DiracSlash`, `FourVector`, `MetricTensor`, `OneLoop` and `ScalarProduct`.

The setting of `Dimension` may be 4, `dim` or `dim-4`, where `dim` must be a *Mathematica* Symbol. ■ See page 30.

## ■ DiracGamma

`DiracGamma[x, optdim]` is the head of all Dirac matrices and Feynman slashes  $\not{x}$  ( $= \gamma_\mu p^\mu$ ) in the internal representation.

A four-dimensional Dirac matrix  $\gamma_\mu$  is `DiracGamma[ LorentzIndex[mu] ]`, a four-dimensional Feynman slash is `DiracGamma[ Momentum[p] ]`.

$\gamma_5$  is represented as `DiracGamma[5]`, the helicity projectors  $\gamma_6 = (1 + \gamma_5)/2$  and  $\gamma_7 = (1 - \gamma_5)/2$  as `DiracGamma[6]` and `DiracGamma[7]` respectively.

For other than four dimensions an additional argument is necessary:

`DiracGamma[ LorentzIndex[mu, Dim], Dim ]` and

`DiracGamma[ Momentum[q, Dim], Dim ]`.

For standard input `DiracMatrix` and `DiracSlash` are more convenient. ■ Note that `DiracGamma[exp, Dim]` projects out the smaller dimension of the objects `exp` and `Dim`. There are special relationships if `Dim` takes the form `D-4`, e.g., `DiracGamma[ LorentzIndex[mu, D-4], 4 ]`  $\rightarrow 0$ . ■ See page 43.

## ■ DiracMatrix

`DiracMatrix[mu]` is an input function for a Dirac matrix. A product of Dirac matrices  $\gamma^\mu \gamma^\nu \gamma^\rho \dots$  is entered as `DiracMatrix[ mu, nu, ro, ... ]` or equivalently as `DiracMatrix[mu] . DiracMatrix[nu] . DiracMatrix[ro] . ...`

$\gamma^5$  may be entered as `DiracMatrix[5]`,  $\gamma^6 = (1 + \gamma^5)/2$  as `DiracMatrix[6]` and  $\gamma^7 = (1 - \gamma^5)/2$  as `DiracMatrix[7]`

The following option can be given:

`Dimension 4` space-time dimension

■  $\gamma^5, \gamma^6$  and  $\gamma^7$  are defined purely in four dimensions. ■ See page 9. ■ See also: `DiracGamma`, `DiracSlash`, `ChiralityProjector`, `BreitMaison`.

## ■ DiracOrder

`DiracOrder[expr, orderlist]` orders the Dirac matrices in `expr` according to `orderlist`.

`DiracOrder[expr]` orders the Dirac matrices in `expr` alphabetically.

$\gamma_5, \gamma_6$  and  $\gamma_7$  are not ordered; use `DiracSimplify` to push them all to the right. ■ Example:

`DiracOrder[ DiracSlash[a, q, a, p] ]`  $\rightarrow -2 \text{ a.a.p.q} + 2 \text{ a.q.gs[a] . gs[p]} + \text{a.a.gs[p] . gs[q]}$ .

■ `DiracOrder[ DiracSlash[p], DiracSlash[q], {q,p} ]`  $\rightarrow 2 \text{ p.q} - \text{gs[q] . gs[p]}$ . ■ This function is just the implementation of the anticommutator relation for Dirac matrices. ■ See page 14. ■ See also: `DiracSimplify`.

## ■ DiracSimplify

`DiracSimplify[expr]` simplifies products of Dirac matrices in `expr`. Double Lorentz indices and four vectors are contracted. The Dirac equation is applied. All `DiracMatrix[5]`, `DiracMatrix[6]` and `DiracMatrix[7]` are moved to the right. The order of the Dirac matrices is not changed.

$(\not{p} - m)u(p) = 0$ ,  $(\not{p} + m)v(p) = 0$  and  $\bar{u}(p)(\not{p} - m) = 0$ ,  $\bar{v}(p)(\not{p} + m) = 0$  are represented by:

`DiracSimplify[ (DiracSlash[p] - m) . Spinor[p, m] ]`  $\rightarrow 0$ ,

`DiracSimplify[ (DiracSlash[p] + m) . Spinor[-p, m] ]`  $\rightarrow 0$ ,

`DiracSimplify[ Spinor[p, m] . (DiracSlash[p] - m) ]`  $\rightarrow 0$ ,

`DiracSimplify[ Spinor[-p, m] . (DiracSlash[p] + m) ]`  $\rightarrow 0$  ■ Examples:

`DiracSimplify[ DiracMatrix[mu, mu] ]`  $\rightarrow 4$ . ■ `DiracSimplify[ DiracSlash[p, p] ]`  $\rightarrow \text{p.p.}$

■ `DiracSimplify[ DiracMatrix[al, be, al] ]`  $\rightarrow -2 \text{ ga[be]}$ .

■ `DiracSimplify[ DiracMatrix[al, be, al, Dimension->D] ]//Factor`  $\rightarrow (2-D) \text{ ga[be]}$ .

■ `DiracSimplify[ DiracSlash[p], (DiracSlash[-q]+m), DiracSlash[p] ]`  $\rightarrow$

`gs[q] p.p - 2 gs[p] p.q + p.p.m.` ■ `DiracSimplify[ DiracMatrix[5, mu] ]`  $\rightarrow -\text{ga[mu] . ga[5]}$ .

■ `DiracSimplify[ DiracMatrix[6, mu] ]`  $\rightarrow \text{ga[mu] . ga[7]}$ . ■ See page 14. ■ See also: `DiracOrder`.

## ■ DiracSlash

`DiracSlash[p]` is an input function for a Feynman slash  $\not{p} = \gamma^\mu p_\mu$ . A product of slashes may be entered by `DiracSlash[p, q, ...]` or `DiracSlash[p], DiracSlash[q], ...`

The following option can be given:

`Dimension 4` space-time dimension

■ The internal representation of a four-dimensional `DiracSlash[p]` is `DiracGamma[ Momentum[p] ]`, a D-dimensional `DiracSlash[ p, Dimension -> D ]` is transformed into `DiracGamma[ Momentum[p, D], D]`. ■ See page 9. ■ See also: `DiracGamma`, `DiracMatrix`.

## ■ DiracTrace

`DiracTrace[expr]` is the head of a Dirac trace. Whether the trace is evaluated depends on the option `DiracTraceEvaluate`. The argument `expr` may be a product of Dirac matrices or slashes separated by “.”.

The following options can be given:

<code>DiracTraceEvaluate</code>	<code>False</code>	evaluating the trace
<code>LeviCivitaSign</code>	<code>-1</code>	sign convention for the $\varepsilon$ -tensor
<code>Factoring</code>	<code>False</code>	whether to factor the result
<code>Mandelstam</code>	<code>{}</code>	if set to $\{s, t, u, m_1^2 + m_2^2 + m_3^2 + m_4^2\}$ , <code>TrickMandelstam</code> will be used
<code>PairCollect</code>	<code>True</code>	whether to collect the results with respect to products of <code>Pair</code>

■ Examples: `DiracTrace[ DiracMatrix[a1, be] ]`  $\rightarrow 4 g[a1, be]$ . ■ `DiracTrace[ DiracSlash[p, q] ]`  $\rightarrow 4 p \cdot q$ . ■ `DiracTrace[ DiracMatrix[mu, al, be, mu, Dimension->D] ]`  $\rightarrow 4 D g[a1, be]$ .  
 ■ `DiracTrace[ DiracMatrix[a, b, c, d, 5] ]`  $\rightarrow -4 I Eps[a, b, c, d]$ .  
 ■ `DiracTrace[ MetricTensor[a1, be] DiracMatrix[si, al, ro, si] ]`  $\rightarrow 16 g[be, ro]$ .  
 ■ `DiracTrace[ DiracSlash[p - q] . (DiracSlash[q] + m) + DiracSlash[k, 2 p] ]`  $\rightarrow 8 k \cdot p + 4 p \cdot q - 4 q \cdot q$ . ■ With `PP=DiracSlash[p']`; `P=DiracSlash[p]`; `MU=DiracMatrix[mu]`; `K=DiracSlash[k]`; `NU=DiracMatrix[nu]`: `DiracTrace[ (PP+m) . MU . (P+K+m) . NU . (P+m) . NU . (P+K+m) . MU/16 ]`  $\rightarrow 4 m^4 + k \cdot p (4 m^2 + 2 k \cdot p') + k \cdot p' (-4 m^2 + 2 p \cdot p) + k \cdot k (4 m^2 - p \cdot p') - 3 m^2 p \cdot p' + p \cdot p \cdot p'$ .  
 ■ To replace scalar products two possibilities exist: either set the corresponding `ScalarProduct` before calculating the trace: `ScalarProduct[p, q] = t/2`; `(DiracTrace[p . q])`  $\rightarrow 2 t$ , which is preferable, or substitute afterwards: `(DiracTrace[ DiracSlash[a, b] ] / . ScalarProduct[a, b]->s/2)`  $\rightarrow 2 s$ . ■ See page 16. ■ See also: `DiracMatrix`, `DiracSlash`, `ScalarProduct`, `TrickMandelstam`.

## ■ DiracTraceEvaluate

`DiracTraceEvaluate` is an option for `DiracTrace`. If set to `False`, `DiracTrace` remains unevaluated.

The reason for this option is that `OneLoop` needs traces in unevaluated form. ■ See page 17.

## ■ Eps

`Eps[a, b, c, d]` is the head of the totally antisymmetric four-dimensional epsilon (Levi-Civita) tensor. The  $a, b, c, d$  must have head `LorentzIndex` or `Momentum`.

All entries are transformed to four dimensions. ■ For user-friendly input of an `Eps` just with `LorentzIndex'` use

`LeviCivita`. ■ For  $\varepsilon^{\mu\nu\rho\sigma} q_\sigma$  the compact notation  $\varepsilon^{\mu\nu\rho\eta}$  is used, i.e.:

`Eps[LorentzIndex[mu], LorentzIndex[nu], LorentzIndex[ro], Momentum[q]]`. ■ `Eps` is just a head not having any functional properties. In order to exploit linearity ( $\varepsilon^{\mu\nu\rho(p+q)} \rightarrow \varepsilon^{\mu\nu\rho p} + \varepsilon^{\mu\nu\rho q}$ ) and the total antisymmetric property of the Levi-Civita tensor use `EpsEvaluate`. ■ See page 42.

## ■ EpsChisholm

`EpsChisholm[expr]` substitutes for a gamma matrix contracted with a Levi Civita tensor (`Eps`) the Chisholm identity:  $\gamma_\mu \varepsilon^{\mu\nu\rho\sigma} = +i (\gamma^\nu \gamma^\rho \gamma^\sigma - g^{\nu\rho} \gamma^\sigma - g^{\rho\sigma} \gamma^\nu + g^{\nu\sigma} \gamma^\rho) \gamma^5$ .

With the option `LeviCivitaSign` the sign of the right hand side of the equation above can be altered. ■ The following option can be given:

`LeviCivitaSign` `-1` sign convention

■ See page 41.

## ■ EpsContract

`EpsContract` is an option for `Contract` specifying whether Levi-Civita tensors `Eps` will be contracted, i.e., products of two `Eps` are replaced via the determinant formula.

■ See page 13.

## ■ EpsEvaluate

`EpsEvaluate[expr]` applies total antisymmetry and linearity (with respect to `Momentum`) to all Levi-Civita tensors (`Eps`) in `expr`.

■ See page 41.

## ■ EvaluateDiracTrace

`EvaluateDiracTrace[expr]` evaluates `DiracTrace` in *expr*.

- See page 17.

## ■ Expanding

`Expanding` is an option for `Contract` specifying whether expansion will be done in `Contract`. If set to `False`, not all Lorentz indices might get contracted.

- See page 13.

## ■ ExpandScalarProduct

`ExpandScalarProduct[expr]` expands scalar products in *expr*.

Examples: `ExpandScalarProduct[ScalarProduct[p-q,r+2 s]]` → `p.r + 2 p.s - q.r - 2 q.s`. ■ At the internal level `ExpandScalarProduct` expands actually everything with head `Pair`. ■ Since a four-vector has head `Pair` internally also, `ExpandScalarProduct[FourVector[p-2 q, mu]]` → `p[mu] - 2 q[mu]`. ■ See page 14.

## ■ Factor2

`Factor2[expr]` factors a polynomial in a standard way. `Factor2` works better than `Factor` on polynomials involving rationals with sums in the denominator.

In general it is better to use `Factor2` in *Mathematica* 2.0. ■ See page 37.

## ■ Factoring

`Factoring` is an option for `Contract`, `DiracTrace`, `DiracSimplify` and `OneLoop`. If set to `True` the result will be factored, using `Factor2`.

- See page 30.

## ■ FeynAmp

`FeynAmp[name, amp, q]` is the head of the Feynman amplitude given by *FeynArts*. The first argument *name* is for bookkeeping, *amp* is the analytical expression for the amplitude, and *q* is the integration variable. In order to calculate the amplitude replace `FeynAmp` by `OneLoop`. In the output of *FeynArts* *name* has the head `GraphName`.

- See page 31. ■ See also: `GraphName`, Guide to *FeynArts*.

## ■ FeynAmpDenominator

`FeynAmpDenominator[PropagatorDenominator[...], PropagatorDenominator[...], ...]` is the head of the denominators of the propagators, *i.e.*, `FeynAmpDenominator[x]` is the representation of  $1/x$ .

Example:  $1/([q^2 - m_1^2][(q + p_1)^2 - m_2^2])$  is represented as

`FeynAmpDenominator[PropagatorDenominator[q,m1], PropagatorDenominator[q+p1,m2]]`. ■ See page 11.

- See also: `PropagatorDenominator`.

## ■ FeynAmpList

`FeynAmpList[info ... ]FeynAmp[... ], FeynAmp[ ... ], ...]` is the head of a list of `FeynAmp` in the result of *FeynArts*.

- See page 31. ■ See also: Guide to *FeynArts*.

## ■ FeynCalcForm

`FeynCalcForm[expr]` changes the printed output of *expr* to an easy to read form. The default setting of `$PreRead` is `$PreRead = FeynCalcForm`, which forces to display everything after applying `FeynCalcForm`.

Small, Momentum and LorentzIndex are set to Identity by `FeynCalcForm`. ■ PaVe are abbreviated. ■ The action of `FeynCalcForm` is: `DiracMatrix[a] → ga[a]`. ■ `DiracSlash[p] → gs[p]`.  
 ■ `FeynAmpDenominator[ PropagatorDenominator[q,m1], PropagatorDenominator[q+p,m2] ] → 1/(q^2 - m1^2) ((p + q)^2 - m2^2)`. ■ `FourVector[p, mu] → p[mu]`. ■ `FourVector[p-q, mu] → (p - q)[mu]`. ■ `GellMannMatrix[a] → la[a]`. ■ `GellMannTrace[x] → tr[x]`. ■ `DiracTrace[x] → tr[x]`.  
 ■ `LeviCivita[a, b, c, d] → eps[a, b, c, d]`. ■ `MetricTensor[mu, nu] → g[mu nu]`.  
 ■ `Momentum[ Polarization[p] ] → ep[p]`. ■ `Conjugate[PolarizationVector[k, mu]] → ep(*)[k, mu]`.  
 ■ `PolarizationVector[p, mu] → ep[p][mu]`. ■ `ScalarProduct[p, q] → p.q`. ■ `Spinor[p, m] → u[p, m]`.  
 ■ `Spinor[-p, m] → v[p, m]`. ■ `QuarkSpinor[p, m] → u[p, m]`. ■ `QuarkSpinor[-p, m] → v[p, m]`.  
 ■ `SU3F[i, j, k] → f[i, j, k]`. ■ See page 45.

## ■ FinalSubstitutions

`FinalSubstitutions` is an option for `OneLoop` and `OneLoopSum`. All substitutions given to this option will be performed at the end of the calculation.

Example: `FinalSubstitutions → {mw^2 -> mw2, B0 -> B0R}`. ■ See page 30.

## ■ FourVector

`FourVector[p, mu]` is the input for a four vector  $p_\mu$ .

`FourVector[p, mu]` is directly transformed to the internal representation: `Pair[Momentum[p], LorentzIndex[mu]]`.

■ The following option can be given:

`Dimension 4` space-time dimension

■ See page 7. ■ See also: `LorentzIndex`, `Momentum`, `Pair`.

## ■ FreeQ2

`FreeQ2[expr, {form1, form2, ...}]` yields `True` if *expr* does not contain any occurrence of *form1*, *form2*, ... `FreeQ2[expr, form]` is the same as `FreeQ[expr, {form1, form2, ...}]`.

Stephen Wolfram pointed out that you can use alternatively `FreeQ[expr, form1 || form2 || ...]`. ■ See page 39.

## ■ GellMannMatrix

`GellMannMatrix[a]` is the Gell-Mann matrix  $\lambda_a$ . A product of Gell-Mann matrices may be entered as `GellMannMatrix[a, b, ...]` or as `GellmannMatrix[a] . GellMannMatrix[b] . ....`  
`GellMannMatrix[1]` denotes the unit-matrix in color space.

■ See page 10. ■ See also: `SU3Delta`, `SU3F`, `GellMannTrace`.

## ■ GellMannTrace

`GellMannTrace[expr]` calculates the trace of *expr*. All color indices should occur twice and *expr* must be a product of `SU3F`, `SU3Delta` and `GellMannMatrix`.

The Cvitanovic algorithm is used. ■ Examples: `[GellMannTrace[GellMannMatrix[i,i]] → 16`.

`GellMannTrace[GellMannMatrix[a . b . c] SU3F[a, b, c]] → 48 I`.

`GellMannTrace[GellMannMatrix[a . c . e . d] SU3F[a, b, e] SU3F[b, c, d]] → 0`.

`GellMannTrace[GellMannMatrix[1]] → 3`. ■ See page 20.

## ■ GraphName

`GraphName[a, b, c, d]` is the first argument of `FeynAmp` given by *FeynArts*. It may be used also as first argument of `OneLoop`. The arguments *a*, *b*, *c*, *d* indicate information of the graph under consideration.

■ See page 31.

## ■ InitialSubstitutions

`InitialSubstitutions` is an option for `PaVeReduce` and `OneLoop`. All substitutions hereby indicated will be performed at the beginning of the calculation. Energy momentum conservation may be especially indicated in the setting.

Example: `InitialSubstitutions → {CW^2 -> 1 - SW^2, k2 -> - k1 + p1 + p2 }`. ■ See page 30.

## ■ Isolate

`Isolate[expr, {x1, x2, ...}]` substitutes `K[i]` for all subsums in `expr` which are free of any occurrence of `x1, x2, ...`, if `Length[expr]>0`.

`Isolate[expr]` substitutes an abbreviation `K[i]` in `HoldForm` for `expr`, if `Length[expr]>0`.

The following options can be given:

`IsolateHead` `K` the head of the abbreviations

`IsolateSplit` `442` a limit beyond which `Isolate` splits the expression in two sums

■ `IsolateSplit` is the maximum of the characters of the `FortranForm` of the expression being isolated by `Isolate`. The default setting inhibits `Write2` from producing too many continuation lines when writing out in `FortranForm`. ■ The result of `Isolate` can always be recovered by `MapAll[ReleaseHold, result]`. ■ Example:

`Isolate[a[z] (b+c) + d[f] (x+y), {a,d}]` → `a[z] K[1] + d[f] K[2]`. ■ See page 38. ■ See also: `Write2`.

## ■ IsolateHead

`IsolateHead` is an option for `Isolate`.

■ See page 38.

## ■ IsolateSplit

`IsolateSplit` is an option for `Isolate`.

■ See page 38.

## ■ K

`K[i]` are abbreviations which may result from `PaVeReduce`, depending on the option `IsolateHead`. The `K[i]` are returned in `HoldForm` and may be recovered by `ReleaseHold`.

■ See page 24.

## ■ LeptonSpinor

`LeptonSpinor[p, mass]` specifies a Dirac spinor. Which of the spinors  $u, v, \bar{u}$  or  $\bar{v}$  is understood, depends on the sign of the mass argument and the relative position of `DiracSlash[p]`:

`LeptonSpinor[p, mass]` is that spinor which yields `mass*LeptonSpinor[p, mass]` if the Dirac equation is applied to `DiracSlash[p] . LeptonSpinor[p, mass]` or `LeptonSpinor[p, mass] . DiracSlash[p]`.

If a spinor is multiplied by a Dirac matrix or another spinor, the multiplication operator "." must be used.

■ See page 10. ■ See also: `DiracSimplify`.

## ■ LeviCivita

`LeviCivita[mu, nu, ro, si]` is an input function for the totally antisymmetric Levi-Civita tensor.

`LeviCivita[mu, nu, ro, si]` transforms to the internal representation

`Eps[LorentzIndex[mu], LorentzIndex[nu], LorentzIndex[ro], LorentzIndex[si]]`. ■ For simplification of Levi-Civita tensors use `EpsEvaluate`. ■ See page 7. ■ See also: `EpsEvaluate`.

## ■ LeviCivitaSign

`LeviCivitaSign` is an option for `DiracTrace`. The possible settings are (+1) or (-1). This option determines the sign convention of the result of  $tr(\gamma^a \gamma^b \gamma^c \gamma^d \gamma^5)$ .

■ See page 41.

## ■ LorentzIndex

`LorentzIndex[mu, optdim]` is the head of Lorentz indices. The internal representation of a four-dimensional  $\mu$  is `LorentzIndex[mu]`. For other than four dimensions enter `LorentzIndex[mu, dim]`.

`LorentzIndex[mu, 4]` simplifies to `LorentzIndex[mu]`. ■ See page 43.

### ■ MacsymaForm

`MacsymaForm` is an option for `FormatType` in `Write2`.

- See page 40. ■ See also: `Write2`, `MapleForm`.

### ■ Mandelstam

`Mandelstam` is an option for `DiracTrace`, `OneLoop` and `TrickMandelstam`. A typical setting is `Mandelstam -> {s, t, u, m1^2 + m2^2 + m3^2 + m4^2}`, which stands for  $s + t + u = m_1^2 + m_2^2 + m_3^2 + m_4^2$ .

- See page 30.

### ■ MapleForm

`MapleForm` is an option for `Write2`.

- See page 40. ■ See also: `Write2`, `MacsymaForm`.

### ■ MetricTensor

`MetricTensor[mu, nu]` is the input for a metric tensor  $g^{\mu\nu}$ .

The following option can be given:

`Dimension 4` space-time dimension

- See page 7. ■ See also: `Contract`, `LorentzIndex`, `Pair`.

### ■ Momentum

`Momentum[p, optdim]` is the head of a momentum in the internal representation. A four-dimensional momentum  $p$  is represented by `Momentum[p]`. For other than four dimensions an extra argument must be given: `Momentum[q, dim]`.

- See page 43. ■ `Momentum[p, 4]` is automatically transformed to `Momentum[p]`.

### ■ NumericalFactor

`NumericalFactor[expr]` gives the numerical factor of *expr*.

- See page 39.

### ■ OneLoop

`OneLoop[name, q, amplitude]` calculates the one-loop Feynman diagram amplitude. The argument *q* denotes the integration variable, *i.e.*, the loop momentum.

The following options can be given:

<code>ReduceToScalars</code>	<code>False</code>	reduce to $B_0, C_0, D_0$
<code>DenominatorOrder</code>	<code>True</code>	order the entries of <code>FeynAmpDenominator</code>
<code>Dimension</code>	<code>True</code>	dimension of integration
<code>FinalSubstitutions</code>	<code>{}</code>	substitutions done at the end of the calculation
<code>Factoring</code>	<code>False</code>	whether to factor the result
<code>InitialSubstitutions</code>	<code>{}</code>	substitutions done at the beginning of the calculation
<code>Mandelstam</code>	<code>{}</code>	indicate the Mandelstam relation
<code>Prefactor</code>	<code>1</code>	additional prefactor of the amplitude
<code>CancelQ2</code>	<code>True</code>	whether to cancel $q^2$
<code>CancelQP</code>	<code>True</code>	whether to cancel $q \cdot p$
<code>ReduceGamma</code>	<code>False</code>	whether to eliminate $\gamma_6$ and $\gamma_7$
<code>SmallVariables</code>	<code>{}</code>	a list of masses, which will get wrapped around the head <code>Small</code>
<code>WriteOut</code>	<code>True</code>	write out a result file <code>name.m</code>

- Energy momentum conservation may be given as rule of `InitialSubstitutions`. ■ See page 30.

### ■ OneLoopResult

`OneLoopResult[name]` is the variable in the result file written out by `OneLoop` to which the corresponding result is assigned. *name* is constructed from the first argument of `OneLoop`.

- See page 29.

## ■ OneLoopSum

`OneLoopSum[ FeynAmp[...], FeynAmp[...], ... ]` calculates a list of Feynman amplitudes by replacing `FeynAmp` step by step by `OneLoop` and sums the result.

The following options can be given:

<code>CombineGraphs</code>	<code>{}</code>	which amplitudes to sum before invoking <code>OneLoop</code>
<code>FinalSubstitutions</code>	<code>{}</code>	substitutions done at the end of the calculation
<code>IsolateHead</code>	<code>K</code>	whether to <code>Isolate</code> the result
<code>Mandelstam</code>	<code>{}</code>	whether to use the Mandelstam relation
<code>Prefactor</code>	<code>1</code>	multiply the result by a pre-factor
<code>ReduceToScalars</code>	<code>True</code>	reduce the summed result to scalar integrals
<code>SelectGraphs</code>	<code>All</code>	which amplitudes to select
<code>WriteOutPaVe</code>	<code>False</code>	whether to write out the reduced <code>PaVe</code>

Possible settings for `CombineGraphs` and `SelectGraphs` are lists of integers. For indicating a range of graphs also a list `{i, j}` instead of a single integer may be provided. ■ See page 31.

## ■ Pair

`Pair[a, b]` is the head of a special pairing used in the internal representation. The arguments  $a$  and  $b$  may have heads `LorentzIndex` or `Momentum`. If both  $a$  and  $b$  have head `LorentzIndex`, the metric tensor is understood. If  $a$  and  $b$  have head `Momentum`, a scalar product is meant. If one of  $a$  and  $b$  has head `LorentzIndex` and the other head `Momentum`, a Lorentz vector  $p^\mu$  is understood.

`Pair` has only one functional definition: any integers multiplied with  $a$  or  $b$  will be pulled out. ■ See page 44. ■ See also: `FourVector`, `LorentzIndex`, `MetricTensor`, `ExpandScalarProduct`, `ScalarProduct`.

## ■ PairCollect

`PairCollect` is an option for `Contract`.

■ See page 17. ■ See also: `Pair`, `ScalarProduct`, `Momentum`

## ■ PaVe

`PaVe[i, j, ..., plist, mlist]` denotes the Passarino-Veltman integrals. The length of the mass list `mlist` indicates if a one-, two-, three- or four-point integral is understood. The first set of arguments  $i, j, \dots$  signifies that the coefficient of  $p_i^\mu p_j^\nu, \dots$  of the tensor integral decomposition is meant, where  $p_0^\mu p_0^\nu = g^{\mu\nu}$ . Joining `plist` and `mlist` gives the same conventions as for `A0`, `B0`, `C0` and `D0`.

For the corresponding arguments of `PaVe` the special cases `A0`, `B0`, `C0`, `D0`, `B1`, `B00`, `B11` are returned. ■ See page 22.

## ■ PaVeOrder

`PaVeOrder[expr]` brings all arguments of `C0` and `D0` into a canical order.

The following option can be given:

`PaVeOrderList` `{}` whether to order according to a list of arguments

■ See page 43.

## ■ PaVeOrderList

`PaVeOrderList` is an option for `PaVeOrder` allowing to specify a specific order of the `D0` functions.

Possible settings are a sublist of the arguments of a `D0`, or a list of such lists. ■ See page 43.

## ■ PaVeReduce

`PaVeReduce[expr]` reduces Passarino-Veltman integrals `PaVe` to scalar integrals `B0`, `C0` and `D0`, depending on the option `BReduce` eventually also `A0`, `B1`, `B00` and `B11`.

The class of invariant Passarino-Veltman integrals which can be currently reduced consists of all coefficients of the Lorentz invariant decomposition of  $B^\mu, B^{\mu\nu}, C^\mu, C^{\mu\nu}, C^{\mu\nu\rho}, D^\mu, D^{\mu\nu}, D^{\mu\nu\rho}, D^{\mu\nu\rho\sigma}$ . ■ The following options can be given:

`IsolateHead` `False` whether to use `Isolate`  
`Mandelstam` `{}` Mandelstam relation, e.g., `{s, t, u, 2 mw^2}`

■ See page 24. ■ See also: `BReduce`, `K`, `PaVe`.



## ■ Polarization

`Polarization[p, optarg]` is the head of a polarization momentum  $\varepsilon(p)$ . `Polarization` must always occur inside `Momentum`. The full internal representation of  $\varepsilon(p)$  is `Momentum[Polarization[p]]`. With this notation transversality of polarization vectors is provided.

`Polarization[p, -1]` stands for the complex conjugate  $\varepsilon(p)^*$ . ■ See page 8. ■ See also: `PolarizationVector`.

## ■ PolarizationSum

`PolarizationSum[mu, nu, ...]` defines different polarization sums.

`PolarizationSum` does not calculate any polarization sum, it is just an abbreviation function.

■ `PolarizationSum[mu, nu] = -gμν`. ■ `PolarizationSum[mu, nu, k] = -gμν + kμkν/k2`.

■ `PolarizationSum[mu, nu, k, n] = -gμν - kμkνn2/(k·n)2 + (nμkν + nνkμ)/(k·n)`, where  $n_\mu$  denotes an external four vector. ■ See page 42.

## ■ PolarizationVector

`PolarizationVector[p, mu]` is an input function for a polarization vector  $\varepsilon(k)^\mu$ .

`Conjugate{PolarizationVector[p, mu]}` is the input for  $\varepsilon_\mu^*(k)$ . ■ The internal representation of `PolarizationVector[p, mu]` is `Pair[Momentum[Polarization[p]], LorentzIndex[mu]]`. ■ The internal representation of `Conjugate{PolarizationVector[p, mu]}` is `Pair[Momentum[Polarization[p, -1]], LorentzIndex[mu]]`. ■ See page 8. ■ See also: `Polarization`.

## ■ Prefactor

`Prefactor` is an option for `OneLoop` and `OneLoopSum`. If set as option of `OneLoop`, the amplitude is multiplied by `Prefactor` before calculation; if specified as option of `OneLoopSum`, it appears in the final result as a global factor.

A possible setting is  $1/(1 - D)$  for calculating the transverse part of self energies. The option `Dimension` of `OneLoop` must then be set to  $D$ . ■ See page 30.

## ■ ProductExpand

`ProductExpand` is an option for `Collect2` and `Combine`.

■ See page 37.

## ■ PropagatorDenominator

`PropagatorDenominator[q, m]` is the denominator of a propagator, *i.e.*,  $(q^2 - m^2)$ .

`PropagatorDenominator[q]` evaluates to `PropagatorDenominator[q, 0]`.

If  $q$  is supposed to be  $D$ -dimensional enter: `PropagatorDenominator[Momentum[q, D], m]`.

■ `PropagatorDenominator` must always occur only as an argument of `FeynAmpDenominator`. ■ See page 11.

## ■ QuarkSpinor

`QuarkSpinor[p, mass]` specifies a Dirac spinor. Which of the spinors  $u, v, \bar{u}$  or  $\bar{v}$  is understood, depends on the sign of the mass argument and the relative position of `DiracSlash[p]`:

`QuarkSpinor[p, mass]` is that spinor which yields `mass*QuarkSpinor[p, mass]` if the Dirac equation is applied to `DiracSlash[p]`. `QuarkSpinor[p, mass]` or `QuarkSpinor[p, mass]`. `DiracSlash[p]`.

If a spinor is multiplied by a Dirac matrix or another spinor, the multiplication operator "." must be used.

■ See page 10. ■ See also: `DiracSimplify`.

## ■ ReduceGamma

`ReduceGamma` is an option for `OneLoop` determining whether  $\gamma_6$  and  $\gamma_7$  are removed by their definitions  $1/2(1 + \gamma_5)$  and  $1/2(1 - \gamma_5)$ .

This option may be needed for certain standard matrixelements. ■ See page 30.

## ■ ReduceToScalars

`ReduceToScalars` is an option for `OneLoop` and `OneLoopSum` that specifies whether the result is reduced to scalar integrals.

Depending on the option `BReduce` the Passarino-Veltman functions `B1`, `B11`, `B00` and `B11` may also remain. ■ See page 31.  
 ■ See also: `PaVeReduce`.

## ■ ScalarProduct

`ScalarProduct` [ $p$ ,  $q$ ] is the input for a scalar product.

The following option can be given:

`Dimension` 4 space-time dimension

■ The internal representation is: `Pair[ Momentum[p], Momentum[q] ]`. ■ Scalar products may be set, e.g., `ScalarProduct[a, b] = c`; but `a` and `b` must not contain sums. ■ See page 7. ■ See also `ExpandScalarProduct`.

## ■ SelectGraphs

`SelectGraphs` is an option for `OneLoopSum`. The default setting is `All`. It may be set to a list indicating that only a subclass of all graphs supplied to `OneLoopSum` should be calculated.

■ See page 31.

## ■ SetMandelstam

`SetMandelstam` [ $s$ ,  $t$ ,  $u$ ,  $p_1$ ,  $p_2$ ,  $p_3$ ,  $p_4$ ,  $m_1$ ,  $m_2$ ,  $m_3$ ,  $m_4$ ] defines the Mandelstam variables  $s = (p_1 + p_2)^2$ ,  $t = (p_1 + p_3)^2$ ,  $u = (p_1 + p_4)^2$  and sets the  $p_i$  on-shell:  $p_i^2 = m_i^2$ , where the  $p_i$  satisfy  $p_1 + p_2 + p_3 + p_4 = 0$ .

If  $p_3 = -k_1$  and  $p_4 = -k_2$ , i.e.,  $p_1 + p_2 = k_1 + k_2$ , the input is:

`SetMandelstam` [ $s$ ,  $t$ ,  $u$ ,  $p_1$ ,  $p_2$ ,  $-k_1$ ,  $-k_2$ ,  $m_1$ ,  $m_2$ ,  $m_3$ ,  $m_4$ ] ■ See page 17.

## ■ SetStandardMatrixElements

`SetStandardMatrixElements` [{ $sma1 \rightarrow abb1$ }, { $sma2 \rightarrow abb2$ }, ... ,  $enmoconrule$  ] defines the standard matrix elements  $sma1$ ,  $sma2$ , .. (e.g., `Spinor` [ $p1$ ] . `DiracSlash` [ $k$ ] . `Spinor` [ $p2$ ]), as `StandardMatrixelement` [ $abb1$ ], `StandardMatrixelement` [ $abb2$ ], ... . The last argument  $enmomcon$  defines energy momentum conservation; e.g.,  $enmomcon = \{k2 \rightarrow p1 + p2 - k1\}$ .

`SetStandardMatrixElements` should be invoked only once for a whole process. It is most conveniently used in a separate specification batch file. In this file also the settings of the scalar products should be done either directly and/or with `SetMandelstam` before applying `SetStandardMatrixElements`. ■ It is not necessary to predefine standard matrix elements. ■ See page 28. ■ See also: `SetMandelstam`, `StandardMatrixelement`.

## ■ Small

`Small` [ $me$ ] is the head of a small mass  $me$ . The effect is that masses with this head are set to zero, if they occur outside a Passarino-Veltman function.

■ See page 12.

## ■ SmallVariables

`SmallVariables` is an option for `OneLoop`.

The setting of `SmallVariables` is a list containing masses which are small compared to others. If present the photon mass should always be listed. ■ See page 30.

## ■ Spinor

`Spinor` [ $p$ ,  $mass$ ] specifies a Dirac spinor. Which of the spinors  $u$ ,  $v$ ,  $\bar{u}$  or  $\bar{v}$  is understood, depends on the sign of the mass argument and the relative position of `DiracSlash` [ $p$ ]: `Spinor` [ $p$ ,  $mass$ ] is that spinor which yields  $mass * \text{Spinor}[p, mass]$  if the Dirac equation is applied to `DiracSlash` [ $p$ ] . `Spinor` [ $p$ ,  $mass$ ] or `Spinor` [ $p$ ,  $mass$ ] . `DiracSlash` [ $p$ ].

If a spinor is multiplied by a Dirac matrix or another spinor, the multiplication operator "." must be used.

■ See page 10. ■ See also: `DiracSimplify`.

## ■ StandardMatrixElement

`StandardMatrixElement[...]` is the head of the standard matrix elements. The standard matrix elements are a basis for the Feynman amplitude which contain the spinor structure and, eventually, the dependence on the polarization vectors.

If `SetStandardMatrixElements` has been used to define abbreviations for the non commutative and/or scalar-product structure, the arguments of `StandardMatrixElement` are these abbreviations itself. Without invoking `SetStandardMatrixElements` the arguments of `StandardMatrixElement` contain the basis for the amplitude, *i.e.*, the non commutative structure and/or scalar-product structure.

- See page 28.

## ■ SU3Delta

`SU3Delta[a, b]` is the Kronecker  $\delta$  with color indices  $a$  and  $b$ .

`SU3Delta[i, i] → 8`. ■ See page 10. ■ See also: `SU3F`, `GellMannMatrix`, `GellMannTrace`.

## ■ SU3F

`SU3F[a, b, c]` are the structure constants  $f_{abc}$  of  $SU(3)$ .

Only algebraic properties are implemented. ■ The following option can be given:

`SU3FToTraces True` whether to replace  $f_{abc}$  by  $\frac{i}{4}(tr(\lambda_a \lambda_c \lambda_b) - tr(\lambda_a \lambda_b \lambda_c))$ .

- See page 10. ■ See also: `SU3Delta`, `GellMannMatrix`, `GellMannTrace`.

## ■ SU3FToTraces

`SU3FToTraces` is an option for `SU3F`.

- See page 10.

## ■ Tr

`Tr[expr]` calculates the Dirac trace of  $expr$  directly. `Tr` is identical to `DiracTrace` up to the default setting of `DiracTraceEvaluate`.

The following options can be given:

<code>DiracTraceEvaluate</code>	<code>True</code>	evaluating the trace
<code>LeviCivitaSign</code>	<code>-1</code>	sign convention for the $\epsilon$ -tensor
<code>Factoring</code>	<code>False</code>	whether to factor the result
<code>Mandelstam</code>	<code>{}</code>	if set to $\{s, t, u, m_1^2 + m_2^2 + m_3^2 + m_4^2\}$ , <code>TrickMandelstam</code> will be used
<code>PairCollect</code>	<code>True</code>	whether to collect the results with respect to products of <code>Pair</code>

- See also: `DiracTrace`. ■ See page 16.

## ■ TrickMandelstam

`TrickMandelstam[expr, {s, t, u,  $m_1^2 + m_2^2 + m_3^2 + m_4^2$ }]` simplifies all sums in  $expr$  in such a way that one of the Mandelstam variables  $s, t$  or  $u$  is eliminated by the relation  $s + t + u = m_1^2 + m_2^2 + m_3^2 + m_4^2$ . The trick is that the resulting sum has the shortest number of terms.

Example: `TrickMandelstam[s + t + u, {s, t, u, 2 mw^2}] → 2 mw^2`. ■ See page 42.

## ■ Write2

`Write2[channel, val1 = expr1, val2 = expr2, ...]` writes the settings  $val_1 = expr_1, val_2 = expr_2$  in sequence followed by a newline, to the specified output channel.

The following options can be given:

<code>FormatType</code>	<code>InputForm</code>	in which language to write out to the result file
<code>D0Convention</code>	<code>0</code>	convention for scalar Passarino Veltman integrals

Other possible settings for `FormatForm` are `FortranForm`, `MacsymaForm` and `MapleForm`. ■ Be careful on the output when generating Fortran files. There might be problems like powers of ratios of integers which you have to correct by hand. ■ If the expressions contain variables in `HoldForm`, their values are written out first, if the output language is *Mathematica* or Fortran. ■ See page 40.

## ■ WriteOut

`WriteOut` is an option for `OneLoop`, `OneLoopSum`.

Possible settings are:

<code>True</code>	write output into a file <i>name</i> , where <i>name</i> is the first argument of <code>OneLoop</code>
<code>False</code>	write no output
<code>"/usr/hep/weinberg/"</code>	write result files in your local directory

■ See page 30.

## ■ WriteOutPaVe

`WriteOutPaVe` is an option for `OneLoopSum`. If set to a string the reduced PaVe are written into the file indicated.

■ See page 31.

## ■ \$BreitMaison

`$BreitMaison` is a global variable determining whether the Breitenlohner-Maison scheme is used.

The default setting is `False`, which implies the "naive"  $\gamma_5$  prescription:  $\gamma_5$  is assumed to anticommute with Dirac matrices in all dimensions. ■ If `$BreitMaison` is set to `True`,  $\gamma_5$  will anticommute with the four-dimensional part of a Dirac matrix and commute with the (D-4)-dimensional part. Reset `$PrePrint` for experimenting with the Breitenlohner-Maison scheme. `$BreitMaison` must be set to `True` before loading *FeynCalc*. The command is: `FeynCalc`$BreitMaison = True`. ■ Not every operation has been tested thoroughly, therefore beware! ■ See page 45.

## ■ \$MemoryAvailable

`$MemoryAvailable` is a global variable which is set to an integer *n*, where *n* is the available amount of main memory in MB. The default is 8. It should be increased if possible. The higher `$MemoryAvailable` can be, the more intermediate steps do not have to be repeated by *FeynCalc*.

■ See page 45.

## ■ \$VeryVerbose

`$VeryVerbose` is a global variable with default setting 0. If set to 1, 2, . . . , more and more intermediate comments and informations are displayed during calculations.

■ See page 45.

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