AMBRE – a Mathematica package for the construction of Mellin-Barnes representations for Feynman integrals

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Abstract

The Mathematica toolkit AMBRE derives Mellin-Barnes (MB) representations for Feynman integrals in $d = 4 - 2\varepsilon$ dimensions. It may be applied for tadpoles as well as for multileg and multiloop scalar and tensor integrals. AMBRE uses a loop-by-loop approach and aims at lowest dimensions of the final MB representations. It integrates the package MB for the determination of the singularity structure in $\varepsilon$. The correctness of the representations has to be checked numerically. AMBRE works fine for planar Feynman diagrams. The package contains various sample applications for Feynman integrals with up to six external particles and up to four loops.
# Contents

1. Introduction
2. Construction of Mellin-Barnes representations
3. Using AMBRE
4. One-loop integrals
   4.1 Example: the pentagon diagram of massive QED
4.2 Numerators
4.3 More masses
4.4 More legs
5. Multi-loop integrals: loop-by-loop integrations
   5.1 Example: two-loop planar box in massive QED
5.2 Special numerators
5.3 Further examples: A three-loop planar box, a four-loop self-energy, and a two-loop pentagon
6. Tadpoles
7. Non-planar topologies
8. Conclusions
9. Acknowledgments
References
# List of Figures

1. Pentagon diagram of massive QED (example1a.nb, example1b.nb) 12
2. The one-loop massive QED box (example10.nb) 14
3. General one-loop vertex (example8.nb) 16
4. Six-point scalar functions; left: massless case (example9.nb), right: massive case 16
5. Massive two-loop planar QED box (example2.nb, section 5.2: example7.nb) 17
6. The loop-by-loop iterative procedure for planar topologies (example3.nb) 20
7. Massless topologies; left: four-loop two-point diagram (example4.nb), right: two-loop five-point diagram (example5.nb) 21
8. Two four-loop tadpoles with three massive lines; left: one-dimensional MB-representation (example6a.nb), right: six-dimensional MB-representation (example6b.nb) 22
9. Non-planar massless vertex 23
1 Introduction

Recently, Mellin-Barnes (MB) representations of Feynman integrals have been used extensively in various phenomenological and theoretical studies of quantum field theory. In many applications, sometimes in quite sophisticated ones [1–3], the MB-integrals can be solved analytically. One also may merge knowledge of some analytical solutions given by MB-integrals with other methods, e.g. the differential equations approach, as demonstrated in [4]. An introduction to the subject with many examples may be found in the monographs [5,6]. A systematic derivation and numerical evaluation of MB-representations for Feynman integrals with a (unpublished) Maple package was described in [7]. At the same time, the Mathematica program MB for the automatized analytic continuation of MB-integrals was published in [8]. With AMBRE, we deliver a Mathematica tool for the derivation of MB-integrals and their subsequent analytic continuation and numerical evaluation with MB.

The article is organized as follows. In section 2 we introduce the formulae used for the MB-representation of a general Feynman integral, and in section 3 the basic features of AMBRE are described. One-loop examples are given in section 4, section 5 describes the implementation of the loop-by-loop approach to multi-loop integrals, and tadpoles are discussed in section 6. Comments on limitations of AMBRE and conclusions follow in sections 7 and 8.

2 Construction of Mellin-Barnes representations

The backbone of the procedure to build up MB-representations is the relation

\[
\frac{1}{(A + B)^\nu} = \frac{B^{-\nu}}{2\pi i \Gamma(\nu)} \int_{-\infty}^{\infty} d\sigma A^\sigma B^{-\sigma} \Gamma(-\sigma) \Gamma(\nu + \sigma),
\]

(1)

where the integration contour separates the poles of the \( \Gamma \)-functions.

The object to be evaluated by AMBRE is an \( L \)-loop Feynman integral\(^1\) in \( d = 4 - 2\varepsilon \) dimensions with \( N \) internal lines with momenta \( q_i \) and masses \( m_i \), and \( E \) external legs with momenta \( p_e \):

\[
G_L[T(k)] = \frac{1}{(i\pi^{d/2})^L} \int \frac{d^d k_1 \cdots d^d k_L}{(q_1^2 - m_1^2)^{\nu_1} \cdots (q_i^2 - m_i^2)^{\nu_i} \cdots (q_N^2 - m_N^2)^{\nu_N}} T(k).
\]

(2)

\(^1\) Often one uses the additional normalization \( e^{\varepsilon \gamma E} \); we leave this to the later evaluation with the package MB [8].
The numerator $T(k)$ is a tensor in the integration variables:

$$T(k) = 1, k_i^\mu, k_i^\mu k_i^\nu, \ldots$$  \hspace{1cm} (3)

The momenta of the denominator functions $d_i$ may be expressed by external and loop momenta:

$$d_i = q_i^2 - m_i^2;$$  \hspace{1cm} (4)

$$q_i = \sum_{l=1}^{L} \alpha_i l k_l - P_i;$$  \hspace{1cm} (5)

$$P_i = \sum_{e=1}^{E} \beta_{ie} p_e.$$  \hspace{1cm} (6)

In the package AMBRE, in a first step the momentum integrals are replaced by Feynman parameter integrals:

$$G_L[T(k)] = (-1)^{N_v} \frac{\Gamma\left(\nu - \frac{d}{2}\right)}{\Gamma(\nu_1) \cdots \Gamma(\nu_N)} \int_0^1 \prod_{j=1}^{N} dx_j x_j^{\nu_j - 1} \delta \left(1 - \sum_{i=1}^{N} x_i\right) \frac{U(x)_{N_v-d(L+1)/2}}{F(x)_{N_v-dL/2}} P_L[T]$$  \hspace{1cm} (7)

with

$$N_v = \sum_{i=1}^{N} \nu_i.$$  \hspace{1cm} (8)

The two functions $U$ and $F$ are polynomials in $x$. They are characteristics of the topology of the underlying Feynman diagram; one may derive them from

$$N = \sum_{i=1}^{N} x_i(q_i^2 - m_i^2) \equiv Mk - 2kQ + J,$$  \hspace{1cm} (9)

where $M_{ll'} = \sum_{i=1}^{N} \alpha_i l l', \alpha_i l; Q_l = \sum_{i=1}^{N} \alpha_i l P_i x_i; J = \sum_{i=1}^{N} (P_i^2 - m_i^2) x_i.$ Namely:

$$U(x) = \det(M),$$  \hspace{1cm} (10)

$$F(x) = -\det(M) \cdot J + Q\tilde{M}Q.$$  \hspace{1cm} (11)

The $U$ and $F$ as well as $\tilde{M} = \det(M) \cdot M^{-1}$ are polynomials in $x$. Further, the numerator functions $P(T)$ for scalar and vector integrals are:
\[ P_L(1) = 1, \]
\[ P_L(k_i^a) = \sum_{l'=1}^{L} \tilde{M}_{ll'} Q_i^{l'}. \]

Tensors of higher degree depend additionally on the diagonalizing rotation \( V \) for \( \mathcal{N} \),
\[ \mathcal{N}_{\text{diag}} = (\alpha_1, \ldots, \alpha_L) = (V^{-1})^T MV^{-1}. \]

As an example, we quote here the case of a tensor of degree two:
\[ P_L(k^a\kappa^\beta) = \sum_{i=1}^{L} \left[ \tilde{M}_{ii} Q_i^a [\tilde{M}_{ii'} Q_i^\beta] - \frac{\Gamma(N_\nu - \frac{d}{2}L - 1)}{\Gamma(N_\nu - \frac{d}{2}L)} UF(V_i^{(1)} + V_i^{(1)}^{-1}) \frac{g^{\alpha\beta}}{2} \right]. \]

The formulae simplify considerably for one-loop integrals:
\[ U = M = \tilde{M} = \det(M) = V = \sum_i x_i = 1, \]
\[ F = -UJ + Q^2 = \sum_{i,j} [P_i P_j - P_i^2 + m_i^2] x_i x_j \equiv \sum_{i,j} f_{ij} x_i x_j. \]

Then, the tensor factors \( P(T) \) will become:
\[ P_1(1) = 1, \]
\[ P_1(k^a) = \sum_{i=1}^{N} x_i P_i, \]
\[ P_1(k^a\kappa^\beta) = \sum_{i=1}^{N} x_i P_i^a \sum_{j=1}^{N} x_j P_j^\beta - \frac{\Gamma(N_\nu - \frac{d}{2} - 1)}{\Gamma(N_\nu - \frac{d}{2})} F \frac{g^{\alpha\beta}}{2}, \text{ etc.} \]

For the general case \( P_1(T) \) see section 4.2.

One now has to perform the \( x \)-integrations. In AMBRE, we will do this by the following simple formula:
\[ \int_0^1 \prod_{i=1}^{N} dx_i x_i^{q_i - 1} \delta \left( 1 - \sum_j x_j \right) = \frac{\Gamma(q_1) \cdots \Gamma(q_N)}{\Gamma(q_1 + \cdots + q_N)}. \]

From the above text it is evident that the integrand of (7) contains besides simple sums of monomials \( \prod_i x_i^{n_i} \) also different structures. This is due to the appearance of the factors \( U(x) \) and \( F(x) \). Beginning with twoloop integrals, one faces additionally a dependence
of $P(T)$ on $x$ for higher rank tensors $T$ due to the appearance of $V$ and $\alpha$, see (15). For this reason, \textsc{Ambre} is restricted to scalar and vector integrals and/or to one-loop integrals. Then we have to rewrite $F(x)$ and $U(x)$ so that (21) becomes applicable, for the one-loop case even only the $F(x)$. We discuss here only the $F(x)$. The $F(x)$ may be written as a sum of $N_F \leq \frac{1}{2}N(N + 1)$ non-vanishing, bilinear terms in $x_i$:

$$F(x) = \sum_{n=1}^{N_F} f_n(i, j)x_i x_j$$

$$= \frac{1}{\Gamma(N_\nu - dL/2)} \left( \frac{1}{(2\pi i)^N} \prod_{i=1}^{N_F} \int_{i\infty+u_i}^{\infty+u_i} dz_i \prod_{n=2}^{N_F} [f_n(ij)x_i x_j]^{z_n} \right)$$

$$\times \left( \frac{1}{\Gamma(N_\nu - dL/2)} \sum_{j=2}^{N_F} z_j \right) \left( \frac{N_\nu - dL/2}{2} + \sum_{j=2}^{N_F} z_j \right) \prod_{j=2}^{N_F} \Gamma(-z_j).$$

(22)

Here, $f_n(i, j) = f_{ij}$ if $f_{ij} \neq 0$. Inserting (22) (and if needed a similar representation for the $U(x)$) and one of the tensor functions $P(T)$ into (2) allows to apply (21) for an evaluation of the $x$-integrations.

As a result, any scalar Feynman integral may be represented by a single multi-dimensional MB-integral and all the one-loop tensor integrals (and the vector $L$-loop integrals) by finite sums of MB-integrals. In practice, with \textsc{Ambre} we will evaluate the $L$-loop integrals by a loop-by-loop technique, which essentially allows us to restrict the formalism to the one-loop case. By the examples it will be seen that this is a powerful ansatz for many applications.

In subsequent steps, the package \textsc{MB} may be called. This package needs as input some MB-integral(s) as being prepared by \textsc{Ambre}. As described in detail in [8], \textsc{MB} allows to analytically expand a Feynman integral in $\varepsilon$ and to evaluate the resulting sequence of MB-integrals by one or the other method.

3 Using \textsc{Ambre}

In this section we describe the use of the Mathematica package \textsc{Ambre} (\textsc{Ambre} stands for Automatic Mellin-Barnes Representation). It is a semi-automatic procedure written for multiloop calculations. The package works with Mathematica 5.0 and later versions of it.

The algorithm to build up MB-representations for Feynman integrals as described in the last section consists of the following parts:

(i) define kinematical invariants which depend on the external momenta;
(ii) make a decision about the order in which $L$ 1-loop subloops ($L \geq 1$) will be worked out sequentially;
(iii) construct a Feynman integral for the chosen subloop and perform manipulations on the corresponding $F$-polynomial to make it optimal for later use of the MB representations;
(iv) use equation (22);
(v) perform the integrations over Feynman parameters with equation (21);
(vi) go back to step (iii) and repeat the steps for the next subloop until $F$ in the last, $L^{th}$ subloop will be changed into an MB-integral.

(23)

The steps (ii) and (iii) must be analyzed carefully, because there exists some freedom of choice on the order of loop integrations in step (ii) and also on the order of MB integrations in step (iii). Different choices may lead to different forms of MB-representations.

The present version 1.0 of AMBRE can be used to construct planar Mellin-Barnes representation for:

- scalar multi-loop, multi-leg integrals
- tensor one-loop integrals
- integrals with specific higher-rank numerators ending up with a single MB-integral

In the next sections several examples will be used for an introduction to specific features of the package.

Here, we describe basic functions of the package. The starting point of all calculations is a proper definition of the integral (2) and of the kinematical invariants to be used. Formally, it has to be done in the following way:

\[
\text{Fullintegral}\{\text{numerator}, \text{denominator}, \text{internal momenta}\};
\text{invariants} = \{\text{invariants as a rule}\};
\]

(24)

We recommend to use $k_i$ and $p_i$ as symbols for internal and external momenta, respectively. Also masses should appear as symbols; a numeric value may cause problems in multi-loop calculations.

The command Fullintegral defines a given integral. For example:

\[
\text{Fullintegral}\{\{1\}, \{\text{PR}[k1, 0, n1]*\text{PR}[k1 + p1, m, n2]\}, \{k1\}];
\]

(25)

corresponds to:

\[
\int \frac{1}{i\pi^{d/2}} \frac{d^d k_1}{(k_1^2)^{n1}[(k_1 + p_1)^2 - m^2]^{n2}}.
\]

(26)
The last argument in the \texttt{Fullintegral} function is a list of internal momenta. The order of internal momenta in this list controls the ordering of integrations (if iterated). For example \{k3,k2,k1\} defines the first integration to be over \(k_3\), the second over \(k_2\) and the third over \(k_1\). The next step is to prepare a subloop of the full integral by collecting all propagators which carry a given loop momentum \(k_i\). We do this by initiating the consecutive functions:

\begin{equation}
\text{IntPart}[\text{iteration}]
\end{equation}

Each iteration, \(i = 1, 2, \ldots, L\), prepares the appropriate subloop for the integration over the corresponding internal momentum. It will display a piece of the \texttt{Fullintegral} with:

- the numerator associated with the given subloop
- subloop for a given internal momentum
- internal momentum for which \texttt{AMBRE} will integrate the subloop

The execution of \texttt{IntPart}[\text{iteration}] proceeds as follows: \texttt{IntPart[1]}, then \texttt{IntPart[2]}, then \texttt{IntPart[3]}, and so on. If there is a need to change the ordering of integrations, one has to change the order in the starting list of internal momenta (24). Inserting \texttt{IntPart[2]} before \texttt{IntPart[1]} is not a proper way to do this. In the output of \texttt{IntPart[iteration]} a tag message will be displayed:

\begin{verbatim}
Fauto::mode: U and F polynomials will be calculated in AUTO mode. In order to use MANUAL mode execute Fauto[0].
\end{verbatim}

(28)

By running \texttt{Fauto[0]}, \texttt{AMBRE} will calculate the \(F\)-polynomial (with name \texttt{fupc}) for a given subloop. At this stage, a user may wish to modify \texttt{fupc} manually, e.g. by applying some changes in kinematics.

During the calculations, the \texttt{FX} function of \texttt{AMBRE} may appear in the \(F\)-polynomial. This function collects full squares of sums of Feynman parameters, e.g.:

\begin{equation}
\text{FX}[x[1]+x[3]]^2 \equiv (x_1 + x_3)^2.
\end{equation}

(29)

Such terms appear in the \(F\)-polynomials if some masses in the loops are equal. They will later allow to apply Barnes’ lemma leading to lower dimensional \(MB\)-representations. At the other hand, the exponent two of the square may lead to arguments of \(\Gamma\)-functions in 22 with doubled integration variables, with far-reaching consequences for a later analytical evaluation when a sum over an infinite series of residua is tried.

The basic function for deriving the Mellin-Barnes representation is:

\begin{equation}
\text{SubLoop[integral]}
\end{equation}

(30)
This function takes output generated by \texttt{IntPart[iteration]} and performs the following calculations:

- calculate the $F$-polynomial for the subloop (only if \texttt{Fauto[1]} is set)
- determine the MB-representation for the $F$-polynomial
- integrate over Feynman parameters $x_i$

As a result, the MB-representation for a given subloop integral will be displayed. In multiloop calculations one will notice additional propagators (marked in red in the output of \texttt{AMBRE}) which appear from the intermediate $F$-polynomial (see section 5.1 for an instructive example).

As mentioned, \texttt{AMBRE} can construct Mellin-Barnes representations for general one loop tensor integrals. The procedure of calculating such cases is basically the same, with few minor differences. First of all, the numerator input must be defined. A one-loop box with numerators $(k_1p_1)(k_1p_2)(k_1p_3)$ might look like this:

\begin{verbatim}
Fullintegral[
{k1*p1,k1+p1,p2,k1+p3},
{PR[k1,m,n1]PR[k1+p1,0,n2]PR[k1+p1+p2,m,n3]PR[k1+p3,0,n4]},{k1}];
\end{verbatim}  

We have written this procedure such that numerators consist of scalar products of internal and external momenta. In the calculations with tensors, the definitions of momentum flows in the subloops play a crucial role for the results and have to be controlled carefully. In one-loop subloops, the internal momentum $k_i$ is assumed to have positive sign in (5). Another difference to scalar cases is the way how \texttt{AMBRE} displays results. Because they can be long, we decided to use a short notation. For example, the following list is likely to be obtained:

\begin{verbatim}
{ARint[1],ARint[2],ARint[3]}
\end{verbatim}  

The result of the evaluation has to be understood as the sum of the elements, $\text{ARint[1]}+\text{ARint[2]}+\text{ARint[3]}$, where each $\text{ARint[i]}$ is one of the resulting MB-integrals. By executing

\begin{verbatim}
ARint[name of representation result,i]
\end{verbatim}

one may display the appropriate $\text{ARint[i]}$. The procedure uses the short notation by default, but it is also possible to use the option \texttt{Result->True} in order to force \texttt{SubLoop} to display the full result:

\begin{verbatim}
SubLoop[integral,Result->True];
\end{verbatim}  

Finally, we have also implemented Barnes’ lemmas:

1st Barnes’ lemma
\[ \int_{-i\infty}^{i\infty} dz \Gamma(a+z)\Gamma(b+z)\Gamma(c-z)\Gamma(d-z) = \frac{\Gamma(a+c)\Gamma(a+d)\Gamma(b+c)\Gamma(b+d)}{\Gamma(a+b+c+d)}. \] (34)

2nd Barnes’ lemma

\[ \int_{-i\infty}^{i\infty} dz \frac{\Gamma(a+z)\Gamma(b+z)\Gamma(c+z)\Gamma(d-z)\Gamma(e-z)}{\Gamma(a+b+c+d+e+z)} = \frac{\Gamma(a+d)\Gamma(a+e)\Gamma(b+d)\Gamma(b+e)\Gamma(c+d)\Gamma(c+e)}{\Gamma(a+b+d+e)\Gamma(a+c+d+e)\Gamma(b+c+d+e)}. \] (35)

The usage of Barnes’ lemmas is simple; one has to execute:

\[ \text{BarnesLemma[representation,i]} \] (36)

where \( i \) is 1 or 2 for the first or second Barnes’ lemma, respectively. This function tries to apply the lemma on all integration variables \( z_i \) which do not appear in powers of kinematical invariants. A comment will be displayed if the lemma was successfully applied.

4 One-loop integrals

We will give a couple of examples starting with construction of MB-representations for the 1-loop Feynman integrals which are an important ingredient of the algorithm Eq. 23. Most of the cases considered in subsequent sections are connected with massless gauge theories or massive QED.

4.1 Example: the pentagon diagram of massive QED

Let us consider the one-loop five-point function shown in figure 1. If we use the FUPolynomial function of the MB package, we will get:

\[ U = x_1 + x_2 + x_3 + x_4 + x_5, \] (37)

\[ F = m^2x_1^2 + 2m^2x_1x_3 - s_{15}x_1x_3 + m^2x_3^2 + 2m^2x_1x_4 - s_{23}x_1x_4 + m^2x_2x_4 \]
\[ - s_{45}x_2x_4 + 2m^2x_3x_4 + m^2x_4^2 - s_{12}x_2x_5 + m^2x_3x_5 - s_{45}x_3x_5. \] (38)

The external momenta fulfill \( p_5^2 = 0, p_i^2 = m^2 \) for the other particles, and the \( s_{ij} = (p_i + p_j)^2 \) are kinematical invariants of the process. A simple counting of terms in the \( F \)-polynomial
Fig. 1. Pentagon diagram of massive QED (example1a.nb, example1b.nb)

would prove that we get a twelve-dimensional MB-integral. Of course the terms in $F$ can be grouped from the beginning and we will see in a minute that a five-fold MB-integral may be obtained; see also the sample file example1a.nb\(^2\).

First, propagators and kinematical invariants are defined:

$$\text{Fullintegral}[
{1},
\{\text{PR}[k1 + p1, 1, n1] \ast \text{PR}[k1 + p1 + p5, 0, n2] \ast \text{PR}[k1 + p1 + p4 + p5, 1, n3] \ast \text{PR}[k1 + p1 + p3 + p4 + p5, 1, n4] \ast \text{PR}[k1 + p1 + p2 + p3 + p4 + p5, 0, n5]\},\{k1\};$$

The kinematics is defined in a cyclic way (see the set Elinv in kinematics.m)

$$p_i^2 = m_i^2; \quad s_{i,i+1} = (p_i + p_{i+1})^2, \quad i = 1, ..., 5.$$  \(40\)

Then, using the \texttt{IntPart} and \texttt{SubLoop} functions the steps (ii)-(v) of the algorithm (23) are worked out and we end up with a nine-fold MB-representation. This representation is due to the following $F$-polynomial, constructed in the automatic way by AMBRE:

$$F' = m^2(x_1 + x_3 + x_4)^2 - s_{15}x_1x_3 - s_{23}x_1x_4 + m^2x_2x_4 - s_{45}x_2x_4 - s_{12}x_2x_5$$
$$+ m^2x_3x_5 - s_{34}x_3x_5.$$  \(41\)

Some mass terms have been collected here, but the $F$-polynomial can be further simplified by redefining $s_{34} \rightarrow \bar{s}_{34} + m^2$ and $s_{45} \rightarrow \bar{s}_{45} + m^2$, so that each term $x_ix_j$ appears only once. The $F'$ polynomial becomes finally:

$$F'' = m^2(x_1 + x_3 + x_4)^2 - s_{15}x_1x_3 - s_{23}x_1x_4 - \bar{s}_{45}x_2x_4 - s_{12}x_2x_5 - \bar{s}_{34}x_3x_5,$$  \(42\)

which gives a seven-fold MB-representation. In certain cases, some of the MB-integrations do not depend on the kinematics and Barnes lemmas may be applied. Here, due to the term

\(^2\) The sample Mathematica files are part of the package AMBRE. They are also available at [9,10]
one may twice use Barnes’ first lemma \[34\] and thus the MB-representation can be further reduced to a five-fold integral, see \texttt{example1a.nb} for details \[10\]. A \(2 \to 3\) scattering process depends on five variables (plus a mass in Bhabha scattering), so a further simplification is impossible.

In sample file \texttt{example1b.nb}, we use another definition of kinematical variables, namely

\[
\begin{align*}
p_i^2 &= m^2, & p_1p_2 &= \frac{1}{2}(t' - 2m^2), & p_1p_3 &= \frac{1}{2}(t - t' - v), \\
p_1p_4 &= m^2 + \frac{1}{2}(v_1 - s - t), & p_1p_5 &= \frac{1}{2}(s - 2m^2), & p_2p_3 &= \frac{1}{2}v, \\
p_2p_4 &= \frac{1}{2}(s - v_1 - v_2 - 2m^2), & p_2p_5 &= \frac{1}{2}(v_2 - s - t' + 2m^2), \\
p_3p_4 &= \frac{1}{2}v_2, & p_3p_5 &= \frac{1}{2}(t' - t - v_2), & p_4p_5 &= \frac{1}{2}(t - 2m^2).
\end{align*}
\] (43)

We get directly \(F\) in the form:

\[
F''' = (x_1 + x_3 + x_4)^2 - tx_1x_3 - t'x_1x_4 - v_2x_2x_4 - sx_2x_5 - v_1x_3x_5.
\] (44)

No wonder, that using function \texttt{SubLoop} we obtain directly the smallest, seven-dimensional integral, which then again reduces to the five-fold integral. The resulting MB-representation for the scalar Feynman integral is:

\[
G[1] = \frac{-e^{\gamma E}}{(2\pi i)^5} \prod_{i=1}^{5} \int_{-i\infty + u_i}^{+i\infty + u_i} dr_i (-s)^{-3-\varepsilon-r_1}(-t)^{r_2}(-t')^{r_3} \left(\frac{v_1}{s}\right)^{r_4} \left(\frac{v_2}{s}\right)^{r_5} \prod_{j=1..12} \Gamma_j \frac{\Gamma_0 \Gamma_{13}}{\Gamma_0 \Gamma_{13}},
\] (45)

with a normalization

\[
\Gamma_0 = \Gamma[-1 - 2\varepsilon],
\] (46)

and the other \(\Gamma\)-functions are:

\[
\begin{align*}
\Gamma_1 &= \Gamma[-r_2], \\
\Gamma_2 &= \Gamma[-r_3], \\
\Gamma_3 &= \Gamma[1 + r_2 + r_3], \\
\Gamma_4 &= \Gamma[-r_1 + r_2 + r_3], \\
\Gamma_5 &= \Gamma[-2 - \varepsilon - r_1 - r_4], \\
\Gamma_6 &= \Gamma[-r_4], \\
\Gamma_7 &= \Gamma[1 + r_2 + r_4], \\
\Gamma_8 &= \Gamma[-2 - \varepsilon - r_1 - r_5], \\
\Gamma_9 &= \Gamma[-r_5],
\end{align*}
\]
Fig. 2. The one-loop massive QED box (example10.nb)

\[ \Gamma_{10} = \Gamma[1 + r_3 + r_5], \]
\[ \Gamma_{11} = \Gamma[3 + \varepsilon + r_1 + r_4 + r_5], \]
\[ \Gamma_{12} = \Gamma[3 + 2r_1 + r_4 + r_5], \]
\[ \Gamma_{13} = \Gamma[3 + 2(r_2 + r_3) + r_4 + r_5]. \] (47)

and

\[ \Gamma_{13} = \Gamma[3 + 2(r_2 + r_3) + r_4 + r_5]. \] (48)

The real parts of the integration strips are
\(-2 < u_1 < -1\) and \(-1/2 < u_i < 0, i = 2 \ldots 5\).

A subsequent application of MB shows that up to constant terms in \(\varepsilon\), needed for an evaluation of two-loop massive Bhabha scattering [11], there are maximally three-dimensional finite contributions to be evaluated further.

4.2 Numerators

AMBRE may handle arbitrary one-loop tensor integrals with higher powers of propagators. The corresponding Feynman parameter integral is the generalization of equation (20):

\[ G_1(T) = \frac{(-1)^N \nu}{\prod_{i=1}^{N} \Gamma(\nu_i)} \int \prod_{i=1}^{N} dx_i x_i^{\nu_i - 1} \delta(1 - \sum_{j=1}^{N} x_j) \sum_{r \leq m} \frac{\Gamma(n - \frac{d}{2} - \frac{r}{2})}{(-2)^{\frac{d}{2} - \frac{r}{2}}} \left\{ A_r P^{m-r} \right\}^{[\mu_1, \ldots, \mu_m]} . \] (49)

Here \(F\) and \(P = P_1(k_\alpha)\) were introduced in equations (11) and (19). The \(r\) starts from zero (with \(A_0 = 1\)), and it is \(A_r = 0\) for \(r\) odd, and \(A_r = g_{[\mu_1 \mu_2 \ldots g_{\mu_r-1 \mu_r]}^r}\) for \(r\) even. The convention \([\mu_1 \ldots]\) means the totally symmetric combination of the arguments.

The momenta flow in the loop must be chosen such that the internal momentum \(k_i\) is non-negative.

In AMBRE tensorial numerators are assumed to be contracted with the external momenta \(p_i\).

As an example, we have prepared the massive QED one-loop box of figure 2 in sample file example10.nb with the numerator \((k_1 \cdot p_1)(k_1 \cdot p_2)(k_1 \cdot p_3)\). The corresponding definition used in AMBRE is:
Fullintegral[
{k1*p1,k1*p2,k1*p3},
{PR[k1,m,n1]PR[k1+p1,0,n2]PR[k1+p1+p2,m,n3]PR[k1+p3,0,n4]},{k1}];

Obviously, when working with tensor integrals we expect the result to be a sum of several MB-integrals (the higher the rank is, the more integrals will be obtained). Results for tensor integrals can be quite long, so that AMBRE displays the result in short notation, \{ARint[1],ARint[2]\}, see for that section 3. Because the result is here a sum of MB-integrals, subsequent calculations using the MB package [8] are performed separately for each of the integrals We have cross checked numerically results for two-, three- and four-point functions by comparing our results with decompositions of integrals into master integrals using the Integration-by-parts method implemented in the package IdSolver. Cross checks were done for numerators with up to six scalar products.

Finally we refer to section 5.2 for the interesting special case of irreducible numerators arising in intermediate subloops. In certain cases, the result for a tensor integral may remain as compact as it is for scalar integrals.

4.3 More masses

\(N\)-point functions with arbitrary internal masses and off-shell external legs give complicated multi-dimensional MB-integrals. Let us consider here and in example8.nb a general one-loop scalar vertex, Fig. 3. In this case we get a five-dimensional MB-integral:

\[
V_{\text{general}} = \frac{(-1)^{n_{132}}}{\Gamma[n_1]\Gamma[n_2]\Gamma[n_3]\Gamma[4-2\varepsilon-n_{132}]} \frac{1}{(2\pi i)^5} \int_{-i\infty}^{+i\infty} dz_1 \int_{-i\infty}^{+i\infty} dz_2 \int_{-i\infty}^{+i\infty} dz_3 \int_{-i\infty}^{+i\infty} dz_4 \int_{-i\infty}^{+i\infty} dz_5 \prod_{i=1}^{5} [\Gamma[-z_i]] \\
(m_1^2)^{z_1} (m_2^2)^{z_2} (m_3^2)^{z_3} (2-\varepsilon-n_{132}-n_{12345}) (m_1^2 + m_2^2 - M_1^2)^{z_2} (m_1^2 + m_3^2 - M_2^2)^{z_4} \\
(m_2^2 + m_3^2 - M_1^2 - M_2^2)^{z_5} \Gamma[n_1 + z_{1124}] \Gamma[4-2\varepsilon-n_{11233} - z_{12345}] \\
\Gamma[n_2 + z_{2335}] \Gamma[-2 + \varepsilon + n_{132} + z_{12345}],
\]

where we abbreviated \(z_{1124} = 2z_1 + z_2 + z_4\) and \(n_{132} = n_1 + n_2 + n_3\), etc.

For some physical case, \(M_1 = m_2\) and \(M_2 = m_3\), we get a compact three-dimensional MB-representation:
\[ V_{simpler} = \frac{(-1)^{n_{123}}}{\Gamma[n_1]\Gamma[n_2]\Gamma[n_3]} \left( \frac{1}{(2\pi i)^3} \int_{-i\infty}^{+i\infty} dz_1 \int_{-i\infty}^{+i\infty} dz_2 \int_{-i\infty}^{+i\infty} dz_3 \prod_{i=1}^{3} \frac{\Gamma[-z_i]}{\Gamma[4 - 2\varepsilon - n_{123} - z_1]} \right) \]

\[ (m_1^2)^{z_1}(m_2^2)^{z_2}(m_3^2)^{2-\varepsilon-n_{123}-z_{123}}(-s)^{z_3} \Gamma[n_1 + z_1]\Gamma[4 - 2\varepsilon - n_{11223} - z_{11223}]\Gamma[n_2 + z_{223}]\Gamma[-2 + \varepsilon + n_{123} + z_{123}] \]

\[ \text{(52)} \]

4.4 More legs

For topologies with a higher number of legs, there is an increasing number of kinematical invariants and so the dimension of MB-representations increases. The number of dimensions may become smaller after analytical continuation in \( \varepsilon \). For a scalar or vector Bhabha massive five-point function, Fig. 1, up to constant terms in \( \varepsilon \), it includes at most three-dimensional integrals, which hopefully can be solved even analytically [11]. In general, the MB-representation for that case is five-dimensional, see section 1.

In example9.nb we derive MB-representations for a massless and a massive one-loop hexagon scalar diagram, see figure 4. In general, it is an eight-fold integral, but the constant term in \( \varepsilon \) includes again only up to three-dimensional MB-integrals.

If all internal lines are massive, one has to deal with a nine-dimensional MB-integral. Again, the numerical results have been checked for both cases in the Euclidean region against sector decomposition. The package contains the auxiliary file KinematicsGen.m which generates the kinematics for six-point functions with arbitrary external legs.
5 Multi-loop integrals: loop-by-loop integrations

The Feynman integral (7) includes a delta-function which makes $U = 1$ for one-loop diagrams there so that the MB-relation (1) acts only on $F$. This simplification can be made also useful in multi-loop integrals by performing loop-by-loop integrations. We collected few examples which will exhibit several specific features.

5.1 Example: two-loop planar box in massive QED

Let us take the massive two-loop planar box topology\footnote{In fact there are three double-box diagrams in massive QED. One of them is non-planar, and we discuss here the so-called first planar diagram [3].} with seven internal lines as introduced in example2.nb. The momentum flow is defined in the following way, with all momenta being incoming:

\[
\text{Fullintegral[}
\{1\},
\{PR[k1, m, n1]PR[k1 + p1, 0, n2]PR[k1 + p1 + p2, m, n3]
PR[k1 - k2, 0, n4]PR[k2, m, n5]PR[k2 + p1 + p2, m, n6]
PR[k2 - p3, 0, n7]\}, \{k2, k1\}\].
\]

First, the momentum integration over $k_2$ is taken. The $k_2$ flow in the first subloop is defined by the function IntPart[1], which contains all propagators with momentum $k_2$:

\[
\text{integral} = PR[k1 - k2, 0, n4]\text{\*PR[k2, m, n5]\*}
PR[k2 + p1 + p2, m, n6]PR[k2 - p3, 0, n7].
\]

We just mention that generally it is preferred to choose the order of iteration such that first the loops with lowest number of lines are executed. Then their $F$-polynomials have a minimal number of terms. The first loop’s $F$-polynomial is the SubLoop[integral] function:

---

Fig. 5. Massive two-loop planar QED box (example2.nb, section 5.2: example7.nb)
\[ F[k_2] \equiv fupc = \]
\[
\begin{align*}
\end{align*}
\]  \hspace{1cm} (55)

It is reproduced here as derived without interactions by the user. The \( F \)-polynomial contains a mass term with the \( FX \) function which later will allow to apply Barnes’ first lemma successfully, and also a redundancy in \( X[3] \cdot X[4] \). The following nine-fold MB-representation after integrating over \( k_2 \) is obtained:

\[
\text{SubLoop1} \left[ \((-1)\)^{n_4 + n_5 + n_6 + n_7 + z_2 + z_3 + z_5} \cdot 4^{z_6} \right. \\
\left. (m^2(z_1 + z_6) \cdot (-s)^{z_4 + z_7} \cdot (-t)^{z_8} \right. \\
\left. (-u)^{(2 - ep - n_4 - n_5 - n_6 - n_7 - z_1 - z_2 - z_3 - z_4 - z_5 - z_6 - z_7 - z_8) \right. \\
\Gamma[-z_1] \Gamma[(-z_2)] \Gamma[(-z_3)] \Gamma[2 - ep - n_4 - n_5 - z_1 - z_2 - z_3 - z_4] \Gamma[(-z_4)] \\
\Gamma[(-z_5)] \Gamma[n_4 + z_2 + z_3 + z_5] \Gamma[(-z_6)] \Gamma[(-z_7)] \\
\Gamma[(-z_8)] \Gamma[-z_9] \Gamma[(-2) \cdot z_1 + z_9] \Gamma[n_5 + z_2 + z_4 + z_9]) / \\
\left. \Gamma[4 - 2 \cdot ep - n_4 - n_5 - n_6 - n_7] \Gamma[n_7] \Gamma[(-2) \cdot z_1] \right]
\]  \hspace{1cm} (56)

It is clear that the factors in front of the \( X[3] \cdot X[4] \) coefficient sum up to zero, due to \( s + t + u = 4m^2 \). To remove them from the beginning, the \texttt{Fauto[0]} option must be executed, followed by a modification of \( F \):

\[ fupc = fupc /. \ u -> 4 \cdot m^2 \cdot 2 \cdot s - t. \]  \hspace{1cm} (57)

In this way, executing the \texttt{SubLoop[integral]} function again, the MB-representation becomes five-dimensional, and also the unpleasant term \( 4^{z_6} \) is absent now.

The same situation appears in the second iteration, when integrating over \( k_1 \). We can switch to the \texttt{Fauto[0]} mode and again modify \( F \). After again applying Barnes’ first lemma, we end up with a six-dimensional integral.

Of course, by writing from the very beginning the kinematical invariants without the invariant \( u \), one can work out the whole case fully automatic with mode \texttt{Fauto[1]}.  

18
5.2 Special numerators

The example is interesting in yet another respect. After the first integration, the propagators for the second one contain four propagators, some of them with shifted indices compared to the input:

\[ \text{PR}[k_1, m, n_1+z_2] \text{PR}[k_1 + p_1, 0, n_2] \text{PR}[k_1 + p_1 + p_2, m, n_3+z_3] \]
\[ \text{PR}[k_1 - p_3, 0, z_5]. \quad (58) \]

This corresponds to the one-loop box of example `example10.nb` discussed in section 4.2, but with shifted indices. It includes the one new propagator with momentum \( q_5 = k_1 - p_3 \).

If we would have been evaluating an integral with numerator \( (q_5^2)^{-ns} \) we repeat the calculation, and get after the second integral an \( F \)-polynomial with one of the terms including the propagator \( \text{PR}[k_1 + p_1 + p_2 + p_4, 0, 1] \); see `SubLoop[integral]` in `example7.nb` [5]. It will sum up with \( \text{PR}[k_1 + p_1 + p_2 + p_4, 0, -n_8] \) resulting in the following integral:

\[
\text{integral} = \text{PR}[k_1, m, n_1 - z_2] \text{PR}[k_1 + p_1, 0, n_2] \\
\quad \text{PR}[k_1 + p_1 + p_2, m, n_3 - z_3] \\
\quad \text{PR}[k_1 + p_1 + p_2 + p_4, 0, -2 + e \pm n_4 + n_5 + n_6 + n_7 - n_8 + z_1 + z_2 + z_3 + z_4].
\quad (59)
\]

which has the following well-known \( F \)-form of the one-loop box:

\[
\]

What is essential here, no additional momentum structure appears.

Analyzing the irreducible numerators of the topology for the given momentum choice, one finds that there are two scalar products which may not be represented by linear combinations of the propagators (and thus are called irreducible): \( k_1 p_3 \) together with \( k_2 p_1 \) or \( k_1 p_3 \) together with \( k_2 p_2 \). So, \( q_5^2 \) represents one of two existing irreducible numerators and it is quite useful to have a simple MB-representation for that case. We see that there are integrals with (selected) numerators which may be represented by a single MB-representation as if a scalar integral would have been studied. This was used several times in examples given in [5,6], and was also used e.g. in [12] for a study of massive two-loop box master integrals, and for more sophisticated four-loop cases in [13].

Finally, a six-dimensional MB-integral emerges like in the scalar case. To check this integral numerically with the MB package, two analytical continuations, one in \( \epsilon \) and one in one of the powers of propagators must be done. We have checked the numerical result also against the results we got from a sector decomposition calculation (unpublished) and from a small-mass expanded version [14] or [12].

19
5.3 Further examples: A three-loop planar box, a four-loop self-energy, and a two-loop pentagon

A three-loop planar integral, shown in figure 6, is treated in example3.nb. The result is a 10-fold MB-representation. With the MB package it was shown that the numerical result agrees with [6].

The dimensions of some MB-representations for several massless and massive ladder topologies are summarized in Table 1. We apply an iterative procedure. For planar topologies the loop-by-loop iteration gives always proper topologies which obey momentum conservation. Only some powers of propagators change into non-integer (complex) numbers.

<table>
<thead>
<tr>
<th></th>
<th>Massless</th>
<th>Massive</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-loop</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>2-loop</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>3-loop</td>
<td>7</td>
<td>13</td>
</tr>
<tr>
<td>4-loop</td>
<td>10</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>14</td>
</tr>
</tbody>
</table>

Table 1
Dimensions of ladder topologies before and after Barnes first Lemma.

A similar procedure can be applied to more complicated topologies which obey the same rule: integrating over an internal momentum leads to a topology with propagators and momentum flow obeying momentum conservation in the remaining parts, i.e. we get regular subtopologies.

In this procedure, the choice of momenta flowing and the order of iterations are very important. Look at the two-loop ladder example shown. If we allow for the momentum flow \( k_1 \) through all the outer lines, and take first the integration over \( k_1 \) and then that over \( k_2 \), the final representation is not optimal (and Barnes' lemmas do not help). Starting instead with the \( k_2 \) integration, we will again end up with a six-dimensional representation.

In files example4.nb and example5.nb, massless MB-representations are constructed for a four-loop two-point topology and for a two-loop five-point massless topology, see figure 7. The six-dimensional four-loop self-energy has been checked numerically against sector decomposition. In example5.nb, there are three different derivations of MB-representations for the same kinematics, defined by equation (40). In each case we got another dimen-
Fig. 7. Massless topologies; left: four-loop two-point diagram (example4.nb), right: two-loop five-point diagram (example5.nb)

sion of MB-integrals (though the numerical results agree). The minimal dimension of the integral is seven when we integrate first over internal momenta of the box and then over that of the pentagon. We checked that numerically this is the same as in [15] where also a seven-dimensional MB-integral has been obtained. If we integrate first over the internal momentum running in the pentagon and next over that in the box, then a nine-dimensional MB-integral is obtained. In the third derivation, the momentum flow in the propagators is chosen in a different way. Then a 13-dimensional MB-integral results.

6 Tadpoles

The loop-by-loop approach can also be applied to planar tadpoles. Attention must be paid to keep the right order of integrations. Making iterations with the \texttt{Fauto[1]} option (i.e. automatic), we may end up with three different forms of propagators in the last iteration: one massive propagator, massive and massless propagators, or one massless propagator. For the first situation the well known formula is used:

\[
\int \frac{d^d k}{(k^2 - q^2)\nu} = i\pi^{d/2} (-1)^\nu \frac{\Gamma[\nu + \epsilon - 2]}{\Gamma[\nu]} \frac{1}{(q^2)^{\nu + \epsilon - 2}}. \tag{60}
\]

We found that for some massive tadpoles a term \((-m)^\alpha\) can appear which would lead to an oscillatory error while doing numerical calculations with \texttt{MB}. In such a situation one has to go back to the previous subloop and modify the \texttt{F}-polynomial with \texttt{Fauto[0]} so that two propagators with equal momenta appear: a massive and a massless one. The same procedure must be applied when a single massless propagator appears in the last integral.

We give as an example \texttt{example6a.nb}, for the diagram also shown left in figure 8. Using \texttt{AMBRE}, we have constructed a one-dimensional Mellin-Barnes representation:

\[
T(n_1, ..., n_5) = (-1)^{2+n_{12345}} (m^2)^{8-4\epsilon-n_{12345}} \prod_{i=1}^{6} \frac{\Gamma[n_i] \Gamma[2 - \epsilon - n_4] \Gamma[2 - \epsilon - n_5]}{\Gamma[2 - \epsilon]} \int_{-\infty}^{+\infty} dz_1 \\
\times \frac{\Gamma[-z_1] \Gamma[2 - \epsilon - n_1 - z_1] \Gamma[2 - \epsilon - n_2 - z_1] \Gamma[4 - 2\epsilon - n_{12} - z_1]}{\Gamma[2 - \epsilon] \Gamma[2 - \epsilon - n_1] \Gamma[2 - \epsilon - n_2] \Gamma[2 - \epsilon - n_3] \Gamma[2 - \epsilon - n_4] \Gamma[2 - \epsilon - n_5]}
\]
At this point the proper order of integrations is very important. A different choice can lead to two- or even higher-dimensional representations.

Using MB we got for the basic integral numerically:

\[
T(1, 1, 1, 1, 1) = 0.25 \frac{1}{4\epsilon^4} + \frac{1}{4\epsilon^3} + 2.843300366757447 \frac{1}{4\epsilon^2} + 5.781543610421033 \frac{1}{4\epsilon} \\
+ 22.955621881705923 + 80.89550616785341 \epsilon + 1085.2836587072804 \epsilon^2 \\
+ 4545.303884134432 \epsilon^3 + 35998.99383263255 \epsilon^4,
\]

(62)

This is in agreement with [16].

However, it appears that MB-representations for four-loop tadpoles can be more complicated. In example6b.nb, treating the diagram in the right of figure 8, we give an example where a six-dimensional MB-integral appears. Taking into account other approaches [16,17], one may see that the MB-approach to multi-loop calculations has natural limits, especially in the massive cases.

7 Non-planar topologies

The loop-by-loop iterative procedure described in this paper seems to be not the most efficient approach in the case of non-planar topologies. It is known from [18] that the massless non-planar vertex is described by a two-dimensional Feynman parameter integral. If we consider the loop-by-loop procedure for this case, we can divide the two-loop topology in figure 9 into two parts (follow the vertical line). The hourglass topology on the right-hand side, with two off-shell legs, gives a three-dimensional MB-representation [6], and adding the second part on the left-hand side we end up with a four-dimensional integral. No matter how we arrange the momenta flows in the diagram, it cannot become better. To get the minimal, two-dimensional integral, another approach must be realized. It is an
Fig. 9. Non-planar massless vertex

open question to us if the representation of non-planar diagrams can be automatized in a way like that for planar cases.

8 Conclusions

We have described the package AMBRE for the construction of MB-representations for planar Feynman integrals and gave a variety of applications. For these cases, the iterative loop-by-loop approach gives a possibility to construct MB-integrals of minimal dimension. Usually Barnes’ first and second lemmas help to get the minimal dimension of MB-integrals independently both of the flow of momenta in diagrams and of the order of iterations. However, for more complicated kinematics, starting with five legs, the order of iterations and the choice of momenta flows matters. As is shown in the case of tadpoles, MB-representations for massive topologies are not always the best way of choice. For some topologies quite simple representations are found, however, also multi-dimensional MB-integrals may arise from which it is hard to get stable, accurate numerical or analytical results.

Constructing useful MB-representations for a given Feynman integral is a kind of an art. As an example, in [12] it was found that by contracting directly two lines in the massive Bhabha two-loop planar integral \( B7l4m1 \) (notations due to [19,12]), the integral \( B5l2m2 \) can be obtained. After expansion in \( \epsilon \), \( B5l2m2 \) consists of eleven integrals, one being four-dimensional. On the other hand, constructing \( B5l2m2 \) from the scratch, loop-by-loop, after expansion in \( \epsilon \), we are left with four integrals, the highest is three-dimensional. This can be checked easily by the reader using the MB package and both the representation \( B7l4m1 \) given in [3] (the contraction of two lines must be done there) and the representation \( B5l2m2 \) given in [12]. There is no simple relation between both representations. Of course, the difference is due to our lack of knowledge about more complicated relations between integrals of different dimensionality.

Certainly, the number of integration variables is of importance for the final evaluation of MB-integrals, both in a fully analytical form or using approximations in some kinematical limits. In many cases some package like XSUMMER [20,21] can be used after deriving sums over residua. This again might become non-effective if the number of the nested sums

\footnote{The non-planar examples in the study [7] do not go beyond our observations stated here.}
– connected with dimension of the MB-integral – is too large or if the result is not in the class of functions covered by (e.g.) \texttt{XSUMMER}. This is also true for the case of a fully numerical evaluation of MB-integrals.

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