NINJA: Automated Integrand Reduction via Laurent Expansion for One-Loop Amplitudes

Tiziano Peraro
Max-Planck Insitut für Physik,
Föhringer Ring, 6, D-80805 München, Germany
E-mail: peraro@mppmu.mpg.de

Abstract

We present the public C++ library NINJA, which implements the Integrand Reduction via Laurent Expansion method for the computation of one-loop integrals. The algorithm is suited for applications to complex one-loop processes.
The investigation of the factorization properties of scattering amplitudes at their singularities [1–5] lead to important results in Quantum Field Theory and Theoretical Particle Physics, such as the development of new methods for phenomenological computations. In particular, integrand reduction methods, developed for one-loop diagrams [6,7] and recently extended
to higher loops \([8–12]\), use the knowledge of the analytic and algebraic structure of loop integrands in order to rewrite scattering amplitudes as linear combinations of Master Integrals.

At one loop, integrand-reduction methods allow to express any integrand in dimensional regularization as a sum of contributions with at most five propagators in the loop, regardless of the number of external legs of the amplitude. The numerators of these contributions are polynomial residues which have a universal parametric form that does not depend on the process. This parametric form can be written as a sum of monomials in the components of the loop momentum, multiplied by unknown process-dependent coefficients. After integration, the amplitude becomes a linear combination of known integrals. The coefficients of this linear combination can be identified with a subset of the ones which parametrize the residues. Therefore, the problem of the computation of any one-loop amplitude can be reduced to the one of performing a polynomial fit of the coefficients of the residues.

The fit of the unknown coefficients can be efficiently performed by evaluating the numerator of the integrand on multiple cuts, i.e. on values of the loop momentum such that a subset of the loop denominators vanish. The multiple-cut conditions can be viewed as projectors which isolate the residue sitting on the cut denominators. A residue can be evaluated by putting on-shell the corresponding loop propagators and subtracting from the integrand the non-vanishing contributions coming from higher-point residues. This leads to a top-down algorithm which allows to compute any one-loop amplitude with any number of external legs.

Within semi-numerical computations, the algorithm is usually implemented by sampling the integrand on several solutions of the multiple cuts and subtracting at each step of the reduction all the non-vanishing contributions coming from higher-point residues. This yields a system of equations for the coefficients of each residue. The method is suited for automation and it has been implemented in several codes, some of which are public (e.g. CutTOOLS \([13]\) and SAMURAI \([14]\)). Its usage within several automated frameworks \([15–23]\) has been particularly successful and produced highly non-trivial phenomenological results.

In this paper we present a new public C++ library called Ninja, which implements an alternative integrand-reduction algorithm first proposed in Ref. \([24]\). This is based on the systematic application of the Laurent series expansion to an integrand on the multiple cuts. After performing a suitable Laurent expansion on a multiple cut, in the asymptotic limit both the integrand and the subtraction terms exhibit the same polynomial behavior as the residue. This allows one to directly identify the coefficients of the residues (and thus the ones of the Master Integrals) with the ones of the Laurent expansion of the integrand, corrected by subtractions terms which can be computed once and for all as functions of a subset of the higher-point coefficients. This leads to a diagonal system of equations for each residue and to a significant reduction of the number of subtraction terms which affect the computation of lower-point contributions.

Ninja takes as input the numerator and three parametric expansions of the same. Since the integrand of a one-loop amplitude is a rational function of the loop momentum, the Laurent expansions can be performed via a partial fraction decomposition. Ninja implements it semi-numerically via a simplified polynomial division algorithm between the expansions of the numerator and the ones of the denominators. The coefficients of the Laurent expansion are then corrected by the subtraction terms and multiplied by the Master Integrals. These are computed by interfacing Ninja with an external library which can be specified by the user. Interfaces for OneLoop \([16, 25]\) and LoopTools \([15]\) are already provided with the distribution.

The simplified subtractions and the diagonal systems of equations make the algorithm
implemented in Ninja significantly simpler and lighter than the traditional one. The library has been interfaced with the one-loop package GoSam [20] and has already been used to compute NLO corrections to Higgs boson production in association with a top-quark pair and a jet [26] and several six-, seven- and eight-point amplitudes involving both massive and massless particles as external states or propagating in the loop [27]. This applications showed that Ninja has better performance and numerical stability than implementations of more traditional integrand reduction algorithms. With this paper, we make Ninja publicly available as a standalone library which can be interfaced to other packages and frameworks for one-loop computations.

In order to simplify the generation of the numerator expansions and the corresponding source code needed by Ninja as input, we distribute with the library a small Python package called NinjaNumGen which uses FORM-4 [28–30] in order to compute the expansions and produce an optimized source code which can be used by the library.

In this paper, besides the description of the implementation of the algorithm and the usage of the library, we also propose a method for the extension of the integrand reduction via Laurent expansion for the computation of integrals whose rank is one unity higher than the number of loop denominators. This method is not present elsewhere in the literature and has been implemented in Ninja, allowing thus to use the library for computations in non-renormalizable and effective theories as well.

The paper is organized as follows. In Section 2 we give a review of the Laurent Expansion method for the integrand reduction of one-loop amplitudes. In Section 3 we discuss its semi-numerical implementation in the library Ninja. The usage of the library is explained, with the help of simple examples, in Section 4. In Section 5 we give a description of the examples which are distributed with the library, giving a better view of its usage and capabilities. Appendix A gives more details on the usage of the package NinjaNumGen. In Appendix B we present the extension of the algorithm to higher-rank integrals. Appendix C gives more information on the interface between Ninja and the libraries of Master Integrals.

Since this paper is rather technical, the reader which is mostly interested in the usage of the library might want to read Sections 4 and 5 first, referring to the previous sections or the appendices at a later time if needed.

2 Integrand Reduction via Laurent Expansion

In this section we review the Integrand Reduction via Laurent-expansion method [24] for the computation of one-loop integrals. The method is based on the systematic application of the Laurent series expansion on the universal integrand decomposition of one-loop amplitudes, which allows to reduce any amplitude as a linear combination of known Master Integrals.

2.1 Universal one-loop decomposition

A generic contribution $M$ to an $n$-point one-loop amplitude in dimensional regularization has the form

$$M = h(\mu_R^2, d) \int d^d \bar{q} \mathcal{I} = h(\mu_R^2, d) \int d^d \bar{q} \frac{\mathcal{N}(\bar{q})}{D_0 \cdots D_{n-1}}. \quad (1)$$

In the previous equation, the integrand $\mathcal{I}$ is a rational function of the components of the $d$-dimensional loop momentum $\bar{q}$, with $d = 4 - 2\epsilon$. The numerator $\mathcal{N}(\bar{q})$ is a process-dependent
polynomial in $\bar{q}$, while the denominators $D_i$ are quadratic polynomials in $\bar{q}$ and correspond to Feynman loop propagators,

$$D_i = (\bar{q} + p_i)^2 - m_i^2.$$

The function $h$ appearing in Eq. (1) is a conventional normalization factor given by [31]

$$h(\mu_R^2, d) = h(\mu_R^2, 4 - 2\epsilon) = \frac{\mu_R^{2\epsilon}}{i\pi^{d/2}} \frac{\Gamma(1 - 2\epsilon)}{\Gamma(1 - \epsilon)\Gamma(1 + \epsilon)},$$

as a function of the renormalization scale $\mu_R^2$ and the dimension $d = 4 - 2\epsilon$. The $d$-dimensional loop momentum $\bar{q}$ can be split into a four-dimensional part $q$ and a $(-2\epsilon)$-dimensional part $\vec{\mu}$ as

$$\bar{q} = q + \vec{\mu}, \quad \bar{q}^2 = q^2 - \mu^2.$$  

The numerator $N$ will therefore be a polynomial in the four components of $q$ and the extra-dimensional variable $\mu^2$.

Every one-loop integrand in dimensional regularization can be decomposed as sum of integrands having five or less loop denominators [6,7]

$$\mathcal{I} \equiv \frac{\mathcal{N}}{D_0 \cdots D_{n-1}} = \sum_{k=1}^{5} \sum_{\{i_1, \ldots, i_k\}} \frac{\Delta_{i_1\cdots i_k}}{D_{i_1} \cdots D_{i_k}},$$

where the second sum on the r.h.s. runs over all the subsets of the denominator indexes $\{0, \ldots, n-1\}$ containing $k$ elements. The residues $\Delta_{i_1\cdots i_k}$ appearing in Eq. (5) are irreducible polynomials, i.e. polynomials which do not contain terms proportional to the corresponding loop denominators $D_{i_1}, \ldots, D_{i_k}$. These residues have a universal, process-independent parametric form in terms of unknown, process-dependent coefficients.

For any set of denominators $D_{i_1}, \ldots, D_{i_k}$ with $k \leq 5$, one can build a 4-dimensional basis of massless momenta $\mathcal{E}^{(i_1\cdots i_k)} = \{e_1, e_2, e_3, e_4\}$ [6,14,32,33]. The first two elements of the basis are linear combinations of two external momenta $K_1, K_2$ of the sub-diagram identified by the considered set of loop denominators. More explicitly, we define

$$e_1^\mu = \frac{1}{1 - r_1 r_2} (K_1^\mu - r_1 K_2^\mu), \quad e_2^\mu = \frac{1}{1 - r_1 r_2} (K_2^\mu - r_2 K_1^\mu),$$

with

$$K_1^\mu = p_{i_1}^\mu - p_{i_k}^\mu, \quad K_2^\mu = p_{i_2}^\mu - p_{i_1}^\mu, \quad r_1 = \frac{K_1^2}{\gamma}, \quad r_2 = \frac{K_2^2}{\gamma},$$

$$\gamma = (K_1 \cdot K_2) \left(1 + \sqrt{1 - \frac{K_1^2 K_2^2}{(K_1 \cdot K_2)^2}}\right).$$

If the sub-diagram has less than two independent external momenta, the remaining ones are substituted by arbitrary reference vectors in the definition of $e_1$ and $e_2$. The momenta $e_3$ and $e_4$ are instead chosen to be orthogonal to the first two and can be defined using the spinor notation as

$$e_3^\mu = \frac{(e_1 \gamma^\mu e_2)}{2}, \quad e_4^\mu = \frac{(e_2 \gamma^\mu e_1)}{2}.\)
They satisfy the property \((e_3 \cdot e_4) = -(e_1 \cdot e_2)\). For subsets of \(k = 4\) denominators, we also define the vectors \(v\) and \(v_\perp\)

\[
v_\mu = (e_4 \cdot K_3) e_3^\mu + (e_3 \cdot K_3) e_4^\mu, \quad v_\perp = (e_4 \cdot K_3) e_3^\mu - (e_3 \cdot K_3) e_4^\mu,
\]

with \(K_3^\mu = p_{i_3}^\mu - p_{i_2}^\mu\). We observe that the vector \(v_\perp\) is orthogonal to all the external legs of the sub-diagram identified by the four denominators.

By expanding the four dimensional part \(q\) of the loop momentum \(\bar{q}\) in the basis \(\mathcal{E}^{(i_1\ldots i_k)}\), the numerator and the denominators can be written as polynomials in the coordinates \(z \equiv (z_1, z_2, z_3, z_4, z_5) = (x_1, x_2, x_3, x_4, \mu^2)\),

\[
\mathcal{N}(\bar{q}) = \mathcal{N}(q, \mu^2) = \mathcal{N}(x_1, x_2, x_3, x_4, \mu^2) = \mathcal{N}(z),
\]

with

\[
q^\nu = -p_{i_1}^\nu + \frac{1}{e_1 \cdot e_2} (x_1 e_1^\nu + x_2 e_2^\nu - x_3 e_3^\nu - x_4 e_4^\nu).
\]

The coordinates \(x_i\) can be written as scalar products

\[
x_1 = (l_{i_1} \cdot e_2), \quad x_2 = (l_{i_1} \cdot e_1), \quad x_3 = (l_{i_1} \cdot e_4), \quad x_4 = (l_{i_1} \cdot e_3),
\]

where \(l_{i_1} \equiv (q + p_{i_1})\). For \(k = 4\) we also consider the alternative expansion of the loop momentum

\[
q^\nu = -p_{i_1}^\nu + \frac{1}{e_1 \cdot e_2} (x_1 e_1^\nu + x_2 e_2^\nu) + \frac{1}{v^2} (x_{3,v} v_3^\nu - x_{4,v} v_4^\nu).
\]

with

\[
x_{3,v} = (l_{i_1} \cdot v), \quad x_{4,v} = (l_{i_1} \cdot v_\perp).
\]

The universal parametric form of the residues \(\Delta_{i_1\ldots i_k}\) in a renormalizable theory is \([6,7,11]\)

\[
\begin{align*}
\Delta_{i_1i_2i_3i_4i_5} &= c_0 \mu^2 \\
\Delta_{i_1i_2i_3i_4} &= c_0 + c_1 x_{4,v} + \mu^2 (c_2 + c_3 x_{4,v} + \mu^2 c_4) \\
\Delta_{i_1i_2i_3} &= c_0 + c_1 x_4 + c_2 x_4 + c_3 x_4 + c_4 x_3 + c_5 x_3^2 + c_6 x_3^3 + \mu^2 (c_7 + c_8 x_4 + c_9 x_3) \\
\Delta_{i_1i_2} &= c_0 + c_1 x_1 + c_2 x_2 + c_3 x_4 + c_4 x_1 x_4 + c_5 x_3 + c_6 x_3^2 + c_7 x_1 x_4 + c_8 x_1 x_3 + c_9 \mu^2 \\
\Delta_{i_1} &= c_0 + c_1 x_2 + c_2 x_1 + c_3 x_4 + c_4 x_3.
\end{align*}
\]

where we understand that the unknown coefficients \(c_j\) depend on the indexes of the residue (e.g. \(c_j = c_j^{(i_1\ldots i_k)}\)), while the scalar products \(x_i\) and \(x_{i,v}\) depend on the both the indexes of the residue and the loop momentum \(q\). The parametrization in Eq. (14) can be extended to effective and non-renormalizable theories where the rank of the numerator can be larger than the number of loop propagators \([24]\). More details on the higher-rank case are given in Appendix B.

Most of the terms appearing in Eq. (14) are spurious, i.e. they vanish after integration and do not contribute to the final result. The amplitude \(\mathcal{M}\) can thus be expressed as a linear combination of Master Integrals, corresponding to the non-spurious terms of the integrand.
decomposition, namely

\[
\mathcal{M} = \sum_{\{i_1, i_2, i_3, i_4\}} \left\{ c_0^{(i_1 i_2 i_3 i_4)} I_{i_1 i_2 i_3 i_4} + c_4^{(i_1 i_2 i_3 i_4)} I_{i_1 i_2 i_3 i_4} [\mu^4] \right\} \\
+ \sum_{\{i_1, i_2, i_3\}} \left\{ c_0^{(i_1 i_2 i_3)} I_{i_1 i_2 i_3} + c_7^{(i_1 i_2 i_3)} I_{i_1 i_2 i_3} [\mu^2] \right\} \\
+ \sum_{\{i_1, i_2\}} \left\{ c_0^{(i_1 i_2)} I_{i_1 i_2} + c_1^{(i_1 i_2)} I_{i_1 i_2} [(q + p_{i_1}) \cdot e_2] \\
+ c_2^{(i_1 i_2)} I_{i_1 i_2} [(q + p_{i_1}) \cdot e_2)^2] + c_0^{(i_1 i_2)} I_{i_1 i_2} [\mu^2] \right\} \\
+ \sum_{i_1} c_0^{(i_1)} I_{i_1},
\]

(15)

where

\[
I_{i_1 \ldots i_k}[\alpha] \equiv h(\mu_R^2, d) \int \frac{d^d q}{D_{i_1} \ldots D_{i_k}} \frac{\alpha}{D_{i_1} \ldots D_{i_k}}, \quad I_{i_1 \ldots i_k} \equiv I_{i_1 \ldots i_k}[1].
\]

The coefficients of this linear combination can be identified with a subset of the coefficients of the parametric residues in Eq. (14). Since all the Master Integrals appearing in Eq. (15) are known, the problem of the computation of an arbitrary one-loop amplitude can be reduced to the problem of the determination of the non-spurious coefficients appearing in the parametrization of the residues \(\Delta_{i_1 \ldots i_k}\).

### 2.2 The Laurent expansion method

The coefficients appearing in the integrand decomposition can be computed by evaluating the integrand on multiple cuts, i.e. on values of the loop momentum \(\vec{q}\) such that a subset of loop denominators vanish [13]. More in detail, the coefficients of a \(k\)-point residue \(\Delta_{i_1 \ldots i_k}\) can be determined by evaluating the integrand on the corresponding \(k\)-ple cut \(D_{i_1} = \cdots = D_{i_k} = 0\). For this values of the loop momentum, the only non-vanishing contributions of the integrand decomposition are the ones coming from the residue in consideration and from all the higher-point residues which have \(\{D_{i_1}, \ldots, D_{i_k}\}\) as a subset of their loop denominators.

Within the original integrand reduction algorithm [13, 14, 34], the coefficients are computed by sampling the numerator of the integrand on a finite subset of the on-shell solutions, subtracting all the non-vanishing contributions coming from higher-point residues, and finally solving the resulting linear system of equations. This is therefore a top-down approach, where higher-point residues are computed first, starting from \(k = 5\), and systematically subtracted from the integrand for the evaluation of lower-point contributions. These are referred to as subtractions at the integrand level.

The integrand reduction via Laurent expansion method, presented in Ref. [24], improves this reduction strategy by elaborating on techniques proposed in [32, 35]. Whenever the analytic dependence of the integrand on the loop momentum is known, this approach allows to compute the coefficients of a residue \(\Delta_{i_1 \ldots i_k}\) by performing a Laurent expansion of the integrand with respect to one of the components of the loop momentum which are not fixed by the on-shell conditions of the corresponding multiple cut \(D_{i_1} = \cdots = D_{i_k} = 0\). In the asymptotic limit, both the integrand and the higher-point subtractions exhibit the same polynomial.
behavior as the residue. Therefore one can directly identify the unknown coefficients with the ones of the Laurent expansion of the integrand, corrected by the contributions coming from higher-point residues.

Hence, by choosing a suitable Laurent expansion, one obtains a diagonal system of equations for the coefficients of the residues, while the subtractions of higher-point contributions can be implemented as corrections at the coefficient level which replace the subtractions at the integrand level of the original algorithm. Since the polynomial structure of the residues is universal and does not depend on the process, the parametric form of the coefficient-level corrections can be computed once and for all, in terms of a subset of the higher-point coefficients. More in detail, the corrections at the coefficient level are known functions of a subset of the coefficients of 3- and 2-point residues. In particular, no subtraction term coming from 4- and 5-point contributions is ever needed. This allows to skip the computation of the (spurious) 5-point contributions entirely, and to completely disentangle the determination of 4-point residues from the one of lower point contributions.

In the following, we address more in detail the computation of 5-, 4-, 3-, 2-, and 1-point residues, also commonly known as pentagons, boxes, triangles, bubbles and tadpoles respectively. For simplicity, we focus on renormalizable theories, where (up to a suitable choice of gauge) the maximum allowed rank of the integrand is equal to the number of loop denominators and the most general parametrization of the residues is the one given in Eq. (14). NINJA can also be used for the computation of integrals whose rank exceeds the number of denominators by one. The extension of the method to the higher-rank case is discussed in Appendix B.

5-point residues As mentioned above, pentagon contributions are spurious. Within the original integrand reduction algorithm, their computation is needed because they appear in the subtractions at the integrand level required for the evaluation of lower-point contributions. A 5-point residue only has one coefficient, which can easily be computed by evaluating the numerator of the integrand on the corresponding 5-ple cut. Within the Laurent expansion approach, the subtraction terms coming from five-point residues always vanish in the asymptotic limits we consider, therefore their computation can be skipped. For this reason, in the library NINJA the computation of pentagons is disabled by default, even though it can be enabled for debugging purposes, as explained in Section 4.3.

4-point residues The coefficient $c_0$ of a box contribution $\Delta_{i_1i_2i_3i_4}$ can be determined via four-dimensional quadruple cuts [5]. A quadruple cut $D_{i_1} = \cdots = D_{i_4} = 0$ in four dimensions (i.e. $q = q$, $\mu^2 = 0$) has two solutions, $q_+$ and $q_-$. The coefficient $c_0$ can be expressed in terms of these solutions as

$$c_0 = \frac{1}{2} \left( \frac{\mathcal{N}(q)}{\prod_{j \neq i_1,i_2,i_3,i_4} D_j\bigg|_{q=q_+}} + \frac{\mathcal{N}(q)}{\prod_{j \neq i_1,i_2,i_3,i_4} D_j\bigg|_{q=q_-}} \right).$$

Given the simplicity of Eq. (16), this is the only coefficient which NINJA computes in the same way as the traditional algorithm. The coefficient $c_4$ can instead be determined by evaluating the integrand on $d$-dimensional quadruple cuts in the asymptotic limit of large $\mu^2$ [35]. A $d$-dimensional quadruple cut has an infinite number of solutions which can be parametrized by the $(-2\epsilon)$-dimensional variable $\mu^2$. These solutions become simpler in the considered limit,


\[ q_+^\nu = -p_1^\nu + a^\nu + \beta \frac{\mu_2^2}{\beta + \mu_2^2} e_3^\nu, \quad q_- = -p_1^\nu + a^\nu + \beta \frac{\mu_2^2}{\beta + \mu_2^2} e_3^\nu + t e_4^\nu, \]  

where the vector \( a^\nu \) and the constant \( \beta \) are fixed by the cut conditions. The coefficient \( c_4 \) is non-vanishing only if the rank of the numerator is greater or equal to the number of loop denominators. In a renormalizable theory, it can be found in the \( \mu^2 \to \infty \) asymptotic limit as the leading term of the Laurent expansion of the integrand

\[
\frac{N(q, \mu^2)}{\prod_{j\neq i_1,i_2,i_3} D_j} \bigg|_{q=\sqrt{\mu^2} v_\perp + O(1)} = c_4 \mu^4 + O(\mu^3). \tag{18}
\]

The other coefficients of the boxes are spurious and, since they neither contribute to the final result nor to the subtraction terms, their computation can be skipped.

**3-point residues** The coefficients of the residues of a generic triangle contribution \( \Delta_{i_1i_2i_3} \) can be determined by evaluating the integrand on the solutions of the corresponding \( d \)-dimensional triple cut \[ [32]. These can be parametrized by the variable \( \mu^2 \) and the free parameter \( t \),

\[
q_+^\nu = -p_1^\nu + a^\nu + t e_3^\nu + \frac{\beta + \mu_2^2}{\beta + \mu_2^2} e_3^\nu, \quad q_- = -p_1^\nu + a^\nu + \frac{\beta + \mu_2^2}{\beta + \mu_2^2} e_3^\nu + t e_4^\nu, \tag{19}
\]

where the vector \( a^\nu \) and the constant \( \beta \) are fixed by the cut conditions \( D_{i_1} = D_{i_2} = D_{i_3} = 0 \). The momentum \( a^\nu \) is a linear combination of \( e_1 \) and \( e_2 \) and is therefore orthogonal to \( e_3 \) and \( e_4 \). On these solutions, the non-vanishing contributions to the integrand decomposition are the ones of the residue \( \Delta_{i_1i_2i_3} \), as well as the ones of the boxes and pentagons which share the three cut denominators. However, after performing a Laurent expansion for large \( t \) and dropping the terms which vanish in this limit, the pentagon contributions vanish, while the box contributions are constant in \( t \) but they also vanish when taking the average between the parametrizations \( q_+ \) and \( q_- \) of Eq. (19). More explicitly,

\[
\frac{N(q_\pm, \mu^2)}{\prod_{j\neq i_1,i_2,i_3} D_j} = \Delta_{i_1i_2i_3} + \sum_j \frac{\Delta_{i_1j_i_j}}{D_j} + \sum_{jk} \frac{\Delta_{i_1j_i_jk}}{D_j D_k} = \Delta_{i_1i_2i_3} + d_1^+ + d_2^- \mu^2 + O(1/t), \quad \text{with } d_1^+ + d_2^- = 0. \tag{20}
\]

Moreover, the expansion of the integrand is given by

\[
\frac{N(q_+, \mu^2)}{\prod_{j\neq i_1,i_2,i_3} D_j} = n_0^+ + n_7^+ \mu^2 + (n_4 + n_9 \mu^2) t + n_5 t^2 + n_6 t^3 + O(1/t) \\
\frac{N(q_-, \mu^2)}{\prod_{j\neq i_1,i_2,i_3} D_j} = n_0^- + n_7^- \mu^2 + (n_1 + n_8 \mu^2) t + n_2 t^2 + n_3 t^3 + O(1/t) \tag{21}
\]

and it has the same polynomial behavior as the expansion of the residue \( \Delta_{i_1i_2i_3} \),

\[
\Delta_{i_1i_2i_3}(q_+, \mu^2) = c_0 + c_7 \mu^2 + (c_4 + c_9 \mu^2) (e_3 \cdot e_4) t + c_5 (e_3 \cdot e_4)^2 t^2 + c_6 (e_3 \cdot e_4)^3 t^3 + O(1/t) \\
\Delta_{i_1i_2i_3}(q_-, \mu^2) = c_0 + c_7 \mu^2 + (c_1 + c_8 \mu^2) (e_3 \cdot e_4) t + c_2 (e_3 \cdot e_4)^2 t^2 + c_3 (e_3 \cdot e_4)^3 t^3 + O(1/t). \tag{22}
\]
By comparison of Eqs (20), (21) and (22) one can directly identify the ten triangle coefficients as the corresponding terms of the expansion of the integrand,

\[ c_{0,7} = \frac{1}{2}(n_0^++n_0^-), \quad c_{1,4,8,9} = \frac{n_{1,4,8,9}}{(e_3 \cdot e_4)}, \quad c_{2,5} = \frac{n_{2,5}}{(e_3 \cdot e_4)^2}, \quad c_{3,6} = \frac{n_{3,6}}{(e_3 \cdot e_4)^3}. \] (23)

Hence, with the Laurent expansion method, the determination of the 3-point residues does not require any subtraction of higher-point terms.

**2-point residues** The coefficients of a generic 2-point residue \( \Delta_{i_1i_2} \) can be evaluated on the on-shell solutions of the corresponding double cut \( D_{i_1} = D_{i_2} = 0 \), which can be parametrized as

\[
q_+ = -p_0 + x e_1' + (\alpha_0 + x \alpha_1) e_2' + t e_3' + \frac{\beta_0 + \beta_1 x + \beta_2 x^2 + \mu^2}{2t(e_3 \cdot e_4)} e_4',
\]

\[
q_- = -p_0 + x e_1' + (\alpha_0 + x \alpha_1) e_2' + \frac{\beta_0 + \beta_1 x + \beta_2 x^2 + \mu^2}{2t(e_3 \cdot e_4)} e_5' + t e_4',
\] (24)

in terms of the three free parameters \( x, t \) and \( \mu^2 \), while the constants \( \alpha_i \) and \( \beta_i \) are fixed by the on-shell conditions. After evaluating the integrand on these solutions and performing a Laurent expansion for \( t \to \infty \), the only non-vanishing subtraction terms come from the triangles,

\[
\frac{\mathcal{N}(q_+, \mu^2)}{\prod_{j \neq i_1, i_2} D_j} = \Delta_{i_1i_2} + \sum_j \frac{\Delta_{i_1i_2j}}{D_j} + \sum_{jk} \frac{\Delta_{i_1i_2jk}}{D_j D_k} + \sum_{jkl} \frac{\Delta_{i_1i_2jkl}}{D_j D_k D_l},
\]

\[
= \Delta_{i_1i_2} + \sum_j \frac{\Delta_{i_1i_2j}}{D_j} + \mathcal{O}(1/t).
\] (25)

Even though the integrand and the subtraction terms are rational functions, in the asymptotic limit they both have the same polynomial behavior as the residue, namely

\[
\frac{\mathcal{N}(q_+, \mu^2)}{\prod_{j \neq i_1, i_2} D_j} = n_0 + n_9 \mu^2 + n_1 x + n_2 x^2 - (n_5 + n_8x)t + n_6 t^2 + \mathcal{O}(1/t)
\]

\[
\frac{\mathcal{N}(q_-, \mu^2)}{\prod_{j \neq i_1, i_2} D_j} = n_0 + n_9 \mu^2 + n_1 x + n_2 x^2 - (n_3 + n_7x)t + n_4 t^2 + \mathcal{O}(1/t)
\] (26)

\[
\frac{\Delta_{i_1i_2j}(q_+, \mu^2)}{D_j} = c_{s_{3,0}}^{(j)} + c_{s_{3,9}}^{(j)} \mu^2 + c_{s_{3,1}}^{(j)} x + c_{s_{3,2}}^{(j)} x^2 - (c_{s_{3,5}}^{(j)} + c_{s_{3,8}}^{(j)}x)t + c_{s_{3,6}}^{(j)} t^2 + \mathcal{O}(1/t)
\]

\[
\frac{\Delta_{i_1i_2j}(q_-, \mu^2)}{D_j} = c_{s_{3,0}}^{(j)} + c_{s_{3,9}}^{(j)} \mu^2 + c_{s_{3,1}}^{(j)} x + c_{s_{3,2}}^{(j)} x^2 - (c_{s_{3,3}}^{(j)} + c_{s_{3,7}}^{(j)}x)t + c_{s_{3,4}}^{(j)} t^2 + \mathcal{O}(1/t)
\] (27)

\[
\Delta_{i_1i_2}(q_+, \mu^2) = c_0 + c_9 \mu^2 + c_1 (e_1 \cdot e_2) x + c_2 (e_1 \cdot e_2)^2 x^2 + (c_5 + c_8 (e_1 \cdot e_2)) (e_3 \cdot e_4) t + c_6 (e_3 \cdot e_4)^2 t^2 + \mathcal{O}(1/t)
\]

\[
\Delta_{i_1i_2}(q_-, \mu^2) = c_0 + c_9 \mu^2 + c_1 (e_1 \cdot e_2) x + c_2 (e_1 \cdot e_2)^2 x^2 + (c_3 + c_7 (e_1 \cdot e_2)) (e_3 \cdot e_4) t + c_4 (e_3 \cdot e_4)^2 t^2 + \mathcal{O}(1/t).
\] (28)
The coefficients $c_{s_3,4}^{(j)}$ of the expansion of the subtractions terms in Eq.s (27) are known parametric functions of the triangle coefficients. Hence, the subtraction of the triangle contributions can be implemented by applying coefficient-level corrections to the terms appearing in the expansion of the integrand. More explicitly, by inserting Eq.s (26), (27) and (28) in Eq. (25) one gets

$$c_{0,9} = n_{0,9} - \sum_j c_{s_1,0,9}^{(j)}$$

$$c_{1,3,5} = \frac{1}{(\epsilon_1 \cdot \epsilon_2)} \left( n_{1,3,5} - \sum_j c_{s_2,1,3,5}^{(j)} \right)$$

$$c_{2,4,6,7,8} = \frac{1}{(\epsilon_1 \cdot \epsilon_2)^2} \left( n_{2,4,6,7,8} - \sum_j c_{s_3,2,4,6,7,8}^{(j)} \right).$$

### 1-point residues

The only non-spurious coefficient $c_0$ of a tadpole residue $\Delta_{i \downarrow}$ can be computed by evaluating the integrand on solutions of the single cut $D_{i \downarrow} = 0$. For this purpose, one can consider 4-dimensional solutions of the form

$$q_+^\nu = -p_{i \downarrow}^\nu + t e_3^\nu + \frac{m_{i_1}^2}{2t(e_3 \cdot e_4)} e_4^\nu,$$

parametrized by the free variable $t$. In the asymptotic limit $t \to \infty$, only bubble and triangle subtraction terms are non-vanishing,

$$\frac{\mathcal{N}(q_+)}{\prod_{j \neq i_1} D_j} = \Delta_{i \downarrow} + \sum_j \frac{\Delta_{i_1,j}}{D_j} + \sum_{jk} \frac{\Delta_{i_1,jk}}{D_j D_k} + \sum_{jkl} \frac{\Delta_{i_1,jkl}}{D_j D_k D_l}$$

$$= \Delta_{i \downarrow} + \sum_j \frac{\Delta_{i_1,j}}{D_j} + \sum_{jk} \frac{\Delta_{i_1,jk}}{D_j D_k} + \mathcal{O}(1/t).$$

Similarly to the case of the 2-point residues, in this limit the integrand and the subtraction terms exhibit the same polynomial behavior as the residue, i.e.

$$\frac{\mathcal{N}(q_+)}{\prod_{j \neq i_1} D_j} = n_0 + n_4 (e_3 \cdot e_4) t + \mathcal{O}(1/t)$$

$$\frac{\Delta_{i_1,j}(q_+)}{D_j} = c_{s_2,0}^{(j)} + c_{s_2,4}^{(j)} (e_3 \cdot e_4) t + \mathcal{O}(1/t)$$

$$\frac{\Delta_{i_1,jk}(q_+)}{D_j D_k} = c_{s_3,0}^{(jk)} + c_{s_3,4}^{(jk)} (e_3 \cdot e_4) t + \mathcal{O}(1/t)$$

$$\Delta_{i_1}(q_+) = c_0^{(i_1)} + c_4^{(i_1)} t + \mathcal{O}(1/t).$$

Putting everything together, the coefficient of the tadpole integral can be identified with the corresponding one in the expansion of the integrand, corrected by coefficient-level subtractions from bubbles and triangles

$$c_0 = n_0 - \sum_j c_{s_2,0}^{(j)} - \sum_{jk} c_{s_3,0}^{(jk)}.$$

The subtraction terms $c_{s_2,0}^{(j)}$ and $c_{s_3,0}^{(jk)}$, coming from 2-point and 3-point contributions respectively, are known parametric functions of the coefficients of the corresponding higher-point residues.
3  Semi-numerical implementation

The C++ library Ninja provides a semi-numerical implementation of the Laurent expansion method described in Section 2. The Laurent series expansion is typically an analytic operation, but since a one-loop integrand is a rational function of the loop variables, its expansion can be obtained via a partial fraction decomposition between the numerator and the denominators. This is implemented in Ninja via a simplified polynomial-division algorithm, which takes as input the coefficients of a parametric expansion of the numerator \( \mathcal{N} \) and computes the leading terms of the quotient of the polynomial division with respect to the uncut denominators. In this section we describe the input needed for the reduction performed by Ninja and we give further details about the implementation of the reduction.

3.1  Input

The inputs needed from the reduction algorithm implemented in Ninja are the momenta \( p_i \) and the masses \( m_i \) of the loop denominators defined in Eq. (2), besides the numerator \( \mathcal{N}(q, \mu^2) \) of the integrand. The latter must be cast in four different forms, one of which is optional. The C++ implementation requires the numerator to be an instance of a class inherited from the abstract class Ninja::Numerator. The latter defined as

```
class Numerator {

public:

    virtual Complex evaluate(const ninja::ComplexMomentum & q,
        const ninja::Complex & \mu^2,
        int cut,
        const ninja::PartitionInt partition[]) = 0;

    virtual void muExpansion(const ninja::ComplexMomentum v[],
        const ninja::PartitionInt partition[],
        ninja::Complex c[]) {}

    virtual void t3Expansion(const ninja::ComplexMomentum & v_0,
        const ninja::ComplexMomentum & v_3,
        const ninja::ComplexMomentum & v_4,
        const ninja::Complex & \beta,
        int mindeg,
        int cut,
        const ninja::PartitionInt partition[],
        ninja::Complex c[]) = 0;

    virtual void t2Expansion(const ninja::ComplexMomentum & v_1,
        const ninja::ComplexMomentum & v_2,
        const ninja::ComplexMomentum & v_3,
        const ninja::ComplexMomentum & v_4,
        const ninja::Complex \beta[] ,
        int mindeg,
        int cut,
        const ninja::PartitionInt partition[],
```

12
The input parameters `cut` and `partition` are common to more methods and give information about the multiple cut where NINJA is currently evaluating the numerator. Although this information is not always necessary, there might be occasions where it could be useful for an efficient evaluation of the numerator. The integer `cut` is equal to `k` if the numerator is being evaluated on a `k`-ple cut. This parameter is not given in the method `muExpansion` because the latter is always evaluated on quadruple cuts. The parameter `partition` points to an array of integers (namely of integer type `ninja::PartitionInt`), with length equal to `cut`, containing the indexes of the cut numerators. If the user asks to perform a global test (see Section 4.3), the numerator will also be evaluated outside the solutions of the multiple cuts, in which case the parameter `cut` will be set to zero.

Here is a detailed description of each method of the class `ninja::Numerator`, for a generic numerator of an `n`-point integrand of rank `r`. If the analytic expression of the integrand is available, all these methods can be easily generated with the help of the simple Python package `NINJA_NUMGEN`, which is distributed with the library and whose usage is described in Section 4.2 and in Appendix A. The details which follow are only needed to those who prefer to provide an alternative implementation of the required methods without `NINJA_NUMGEN`.

**The method Numerator::evaluate(q, \(\mu^2\), cut, partition)**

It must return the value of the numerator \(N(q, \mu^2)\) evaluated at the (complex) values of \(q\) and \(\mu^2\) given as input.

**The method Numerator::muExpansion(v, partition, c)**

This is used for the computation of the Laurent expansion in \(\mu^2\) required to obtain the coefficient \(c_4\) of the boxes. In the renormalizable case, this method should compute the leading term of a parametric expansion in \(t\) of the integrand defined by

\[
qu' \rightarrow tv'_\perp, \quad \mu^2 \rightarrow t^2 v'^2_\perp
\]

(37)

where \(v_\perp\) is given by \(v[0]\), i.e. by the zeroth entry of the array of momenta \(v\). For renormalizable theories this array will therefore only contain at most one element. The method should write the leading coefficient of the expansion in the zeroth entry of the array pointed by the parameter \(c\), i.e.

\[
c[0] = N[t'];
\]

The generalization of this method to the higher-rank case is described in Appendix B. The implementation of this method is only required when \(r \geq n\). It is also not needed if the user chooses to disable the \(\mu^2\)-expansion method for the boxes, but in that case more evaluations of the numerator will be needed and the computation of the pentagons will not be skipped.
The method `Numerator::t3Expansion(v_0, v_3, v_4, \beta, \text{mindeg}, \text{partition}, c)`

This method is used for the computation of the coefficients of the residues of both the triangles and the tadpoles. It is supposed to compute the coefficients of the terms $t^j \mu^{2k}$ for $j \in \{r, r - 1, r - 2, \ldots, r - \text{mindeg}\}$, given by substituting into the numerator the parametric expansion of the loop momentum defined by

$$q^\nu \rightarrow v_0^\nu + t v_3^\nu + \frac{\beta + \mu^2}{t} v_4^\nu, \quad v_3^2 = v_4^2 = 0, \quad (v_3 \cdot v_4) = \frac{1}{2},$$

(38)
as a function of the momenta $v_i^\nu$ and the constant $\beta$ which are passed as parameters. The maximum value of the parameter \text{mindeg} is $r - n + 3$. Since in a renormalizable theory $r \leq n$, and by definition of rank we have $j + 2k \leq r$, in this case at most 6 terms can be non-vanishing in the specified range of $j$. The method should write these terms in the entries of the array pointed by $c$, ordered by decreasing powers of $t$. Terms with the same power of $t$ should be ordered by increasing powers of $\mu^2$. A pseudo-implementation will therefore look like

```c
int idx = 0;
for (int j=r; j>=r-mindeg; --j)
for (int k=0; 2*k<=r-j; ++k) {
    c[idx] = N[t^j \mu^{2k}];
    ++idx;
}
```

The method `Numerator::t2Expansion(v_1, v_2, v_3, v_4, \beta, \text{mindeg}, \text{partition}, c)`

In the current version of \texttt{NINJA}, this method is called during the computation of the coefficients of the bubbles. It is supposed to compute the coefficients of the terms $t^j x^l \mu^{2k}$ for $j \in \{r, r - 1, \ldots, r - \text{mindeg}\}$, given by the expansion

$$q^\nu \rightarrow v_1^\nu + x v_2^\nu + t v_3^\nu + \frac{\beta_0 + \beta_1 x + \beta_2 x^2 + \mu^2}{t} v_4^\nu, \quad v_2 \cdot v_3 = v_2 \cdot v_4 = v_3^2 = v_4^2 = 0, \quad (v_3 \cdot v_4) = \frac{1}{2},$$

(39)
as a function of the momenta $v_i^\nu$ and the constants $\beta_i \equiv \beta[i]$, which are passed as parameters to the method. The maximum value of \text{mindeg} is $r - n + 2$. In a renormalizable theory, this implies that one can have at most 7 non-vanishing terms in this range of $j$. It is worth observing that the expansion in Eq. (39) can be obtained from the previous one in Eq. (38) with the substitutions

$$v_0^\nu \rightarrow v_1^\nu + x v_2^\nu, \quad \beta \rightarrow \beta_0 + \beta_1 x + \beta_2 x^2, \quad v_2 \cdot v_3 = v_2 \cdot v_4 = 0.$$

The terms of the expansion must be stored in the entries of the array pointed by $c$, ordered by decreasing powers of $t$. Terms with the same power of $t$ should be ordered from the lowest to the highest with respect to the lexicographical order in the variables $(x, \mu^2)$. A pseudo-implementation will have the form

```c
int idx = 0;
for (int j=r; j>=r-mindeg; --j)
```
for (int l=0; l<=r-j; ++l)
    for (int k=0; 2*k<=r-j-l; ++k) {
        c[idx] = \mathcal{N}[t^j x^l \mu^{2k}];
        ++idx;
    }

3.2 Reduction via polynomial division

For every phase-space point, \textsc{Ninja} at run-time computes the parametric solutions of the multiple cuts corresponding to each residue. The Laurent expansion of the integrand on these solutions is performed via a simplified polynomial division between the expansion of the numerator and the set of the uncut denominators. The coefficients of this expansion are corrected by the coefficient-level subtractions appearing in Eq.s (29) and (36). The non-spurious coefficients are finally multiplied by the corresponding Master Integrals in order to obtain the integrated result as in Eq. (15).

The coefficients of the expansions of the numerator are written on a contiguous array by the numerator methods described in Section 3.1. The Laurent expansion is obtained via a simplified polynomial division. The latter is performed in-place on the same array, keeping only the elements which are needed for the final result. A possible implementation for an univariate expansion, with a numerator \( N = \text{num}[0] t^n + \text{num}[1] t^{n-1} + \ldots + \text{num}[\text{nterms}-1] t^{r-\text{nterms}+1} + \mathcal{O}(t^{r-\text{nterms}}) \) and denominator \( D = \text{d}[0] t + \text{d}[1] + \text{d}[2] \frac{1}{t} \), would have the form

```cpp
void division(Complex num[], int nterms, Complex den[3])
{
    for (int i=0; i<nterms; ++i) {
        num[i] /= den[0];
        if (i+1<nterms) {
            num[i+1] -= den[1]*num[i];
            if (i+2<nterms)
                num[i+2] -= den[2]*num[i];
        }
    }
}
```

One can check that this routine correctly replaces the first \text{nterms} elements of the array \text{num} with the first \text{nterms} leading elements of the Laurent expansion of \( \mathcal{N}/D \). The actual implementation in \textsc{Ninja}, having to deal with multivariate expansions, is significantly more involved than the division procedure presented here. Nevertheless, it qualitatively follows the same algorithm.

The coefficients obtained by the division are then corrected by the coefficient-level subtractions and thus identified with the corresponding coefficients of the residues, as explained in Section 2. Once the reduction is complete, the non-spurious coefficients are multiplied by the corresponding Master Integrals.
3.3 Master Integrals

**NINJA** calls the routines implementing the Master Integrals through a generic interface which, as in the case of the numerator, is defined in the C++ code via an abstract class (called `IntegralLibrary`). This allows one to use any integral library which can be specified at run-time. More details on the implementation of this interface are given in Appendix C. The current version of **NINJA** already implements it for two integral libraries.

The first built-in interface, `ninja::AvHOneLoop`, is a C++ wrapper of the routines of the OneLoop library [16, 25]. This wrapper caches every computed integral allowing constant-time lookup of their values from their arguments. The caching of the integrals can significantly speed up the computation, especially for complex processes. Every instance of the class `AvHOneLoop` has an independent cache of Master Integrals (hence, one may safely use it in multi-threaded applications by using one instance of the class per thread).

The second implementation of the interface, `ninja::LoopTools`, uses instead the LoopTools library [15], which already has an internal cache of computed integrals.

4 Basic usage

In this Section we explain how to use the library for the computation of a generic one-loop integral. **NINJA** can be interfaced to any one-loop generator capable of providing the input it needs, and in particular to packages which can reconstruct the analytic dependence of the numerators on the loop momentum. An interface for the one-loop package GoSam [20] is already built in the library, and has been used for highly non-trivial phenomenological computations [26, 27]. An interface with the package FORMCALC [15] is currently under development. The author is open to give his assistance in interfacing other packages as well.

In this paper we focus on the usage of **NINJA** as a standalone library. We will show how to generate the numerator methods needed as input, starting from an analytic expression of the numerator, with the help of the Python package **NINJA**NUMGEN which is distributed with the library. We will then explain how to perform the reduction, and how to set the main available options.

4.1 Installation

**NINJA** can be obtained at the url [http://ninja.hepforge.org](http://ninja.hepforge.org). The library is distributed with its source code using the GNU build system (also known as AUTOTOOLS). It can be compiled and installed with the shell commands.

```
./configure
make
make install
```

This will typically install the library and the header files in sub-directories of `/usr/local`. The `--prefix` option can be used in order to specify a different installation path. In this case, you might need to update your `LD_LIBRARY_PATH` (or `DYLD_LIBRARY_PATH` on Mac OS) environment variable accordingly. In order to use **NINJA** for the production of phenomenological results, one must interface it with a library of Master Integrals. As already mentioned, interfaces to the ONELOOP and LOOPTOOLS libraries are provided (see Appendix C for interfacing a different library). They can be enabled by passing to the `configure` script the
options --with-avholo[=FLAGS] and --with-looptools[=FLAGS]. For instance, the following commands

```
./configure --prefix=$HOME/ninja \
   --with-avholo='-L/path/to/avh_olo/lib -lavh_olo' \
   FCINCLUDE=-I/path/to/avh_olo/include
make install
```

will install all the files in sub-directories of $HOME/ninja and build the interface with the ONELOOP library, which will be supposed to be already installed and linkable with the flags specified with the --with-avholo option. We also specified the FCINCLUDE variable with the flags which are needed to find FORTRAN-90 modules when they are not installed in a default directory. A full list of optional arguments for the configure script can be obtained with the command ./configure --help. While most of them are common to every package distributed with the GNU build system, some are instead specific to the NINJA library and they are described in Table 1. In most of the cases, only the options for interfacing the integral libraries should be needed.

| Option                      | Description                                                                 |
|-----------------------------|-----------------------------------------------------------------------------|
| --with-avholo[=FLAGS]       | Include an interface with the ONELOOP integral library [16, 25], specifying the corresponding flags for dynamic linking. If the FORTRAN module avh_olo is not in a standard path, one should add its directory to the FCINCLUDE variable when using this option. |
| --with-looptools[=FLAGS]    | Include an interface with the LOOPTOOLS library [15] (needs LOOPTOOLS version 2.9 or higher), specifying the corresponding flags for static linking. If the header file cloopttools.h is not in a standard path, one should add its directory to the CPPFLAGS variable when using this option. |
| --with-quadruple[=FLAGS]    | Compile the library in quadruple precision. This requires the GCC libquadmath library and one can specify the corresponding flags for the linker (-lquadmath by default). With this option, the types ninja::Real and ninja::Complex will be quadruple precision floating point numbers, and including any public header file of the library will define the macro NINJA_QUADRUPLE to 1 (it will not be defined otherwise). The user should also make sure, when using this option, that the libraries of Master Integrals used by NINJA are compiled in quadruple precision, with floating point types compatible with the ones of GCC. |
| --enable-higher_rank        | Enable support for higher-rank numerators. This is not needed for renormalizable theories. |
| --disable-gosam             | Do not include GoSAM interface. |
| --disable-avholo_cache      | Do not include a cache of Master Integrals in the interface with the ONELOOP library. |

Table 1: Options and environment flags for the configure script. Only the options which modify the default behaviour of NINJA are listed.
The user can also optionally install the PYTHON package *NINJANUMGEN*, which allows one to easily generate the input needed by *NINJA*. The package can be used both as a script and as a PYTHON module, and it could also be useful for interfacing *NINJA* to existing one-loop packages. In order to install the package, move in the *utils* folder and type

```
python setup.py install
```

where, as usual, an installation prefix can be specified using *--prefix*. In this case one might need to update the *PATH* and *PYTHONPATH* environment variables accordingly. The package needs FORM-4 [29, 30] in order to compute the expansions which are needed and for the generation of the corresponding source code.

### 4.2 Writing the Integrand

The most important input that the *NINJA* library needs, for the computation of a one-loop integral, is the numerator of the corresponding integrand, cast in a suitable form. This form has been described in detail in Section 3.1 but, as already mentioned, it can also be generated with the help of the package *NINJANUMGEN*. In this section we describe its usage as a script, with a simple example. A more detailed list of options, allowing to fine-tune the output according to the user’s needs, as well as the usage of the package as a PYTHON module, are described in Appendix A.

Let us define, as an example, the following 4-point one-loop integrand, with kinematics $k_0, k_1 \rightarrow k_2, k_3$

\[
\mathcal{I} = \frac{N(q, \mu^2)}{D_0 D_1 D_2 D_3} = \frac{(q \cdot v_1)(q \cdot v_2)(q \cdot v_3)(q \cdot v_4) + \mu^4}{D_0 D_1 D_2 D_3}
\]

\[
\begin{align*}
D_0 &= q^2 - m_0^2 \\
D_1 &= (q + k_0)^2 - m_1^2 \\
D_2 &= (q + k_0 + k_1)^2 - m_2^2 \\
D_3 &= (q + k_3)^2 - m_3^2.
\end{align*}
\]

(40)

where $v_i$ are arbitrary reference vectors. In order to generate the methods declared in the *ninja::Numerator* class, we first create a file *mynum.frm* containing a FORM [28] expression for the numerator

```
* mynum.frm
V v1,v2,v3,v4;
V Q;
S Mu2;
L Diagram = (Q.v1)*(Q.v2)*(Q.v3)*(Q.v4) + Mu2^2;
```

and we run the script with the command

```
ninjanumgen mynum.frm --nlegs 4 --rank 4 -o mynum.cc
```

which creates the source file *mynum.cc* with the definition of the methods, optimized for fast numerical evaluation using the recent features of FORM-4. The command also creates an header file *mynum.hh*, unless already present, which is supposed to contain the declaration of
the numerator class. The latter will have the following form (where for brevity we replaced the parameters of each method with ellipses),

```cpp
class Diagram : public ninja::Numerator {

public:
  virtual ninja::Complex evaluate(...);
  virtual void muExpansion(...);
  virtual void t3Expansion(...);
  virtual void t2Expansion(...);

public:
  // Add other public methods and data here

private:
  // Add other private methods and data here
};
```

If the numerator expression depends on other momenta or parameters, these should be visible inside the definitions of the methods. In our example, the numerator depends on the reference vectors \( v_i \) which appear in its FORM expression. One possibility would be declaring these vectors as global variables, but a better alternative could be defining them as data members of the numerator class. In this example we will declare them as public data members by inserting the following code inside the class definition

```cpp
public:
  ninja::ComplexMomentum v1, v2, v3, v4;
```

which defines the vectors as complex Lorentz momenta. This completes the generation of the input, which will allow NINJA to compute the integral.

### 4.3 Running the reduction

In this subsection we describe the usage of NINJA for the reduction of a generated integrand, such as the one in the example of the previous subsection. With the help of simple examples, we show how to specify the input and how to control the run-time behavior of the procedures of the library.

All the types, classes and functions provided by the NINJA library are defined inside the `ninja` namespace. In particular, the types `Real` and `Complex` are aliases for `double` and `std::complex<double>`, unless the library was compiled in quadruple precision. Classes for real and complex momenta are defined as `RealMomentum` and `ComplexMomentum` respectively. They are wrappers of four-dimensional arrays of the corresponding floating-point types, which overload arithmetic and subscript operators. All the public header files of the library are installed in the sub-directory `ninja` of the `include` path in the installation directory.

#### 4.3.1 A simple example

Here we show the contents of a file `simple_test.cc` which illustrate the basic usage of NINJA for computing the integral defined in Eq. (40).
#include <iostream>
#include <ninja/ninja.hh>
#include <ninja/rambo.hh>
#include "mynum.hh"
using namespace ninja;

int main()
{
    // External legs of the loop
    const int N_LEGS = 4;

    // Center of mass energy
    const Real ENERGY_CM = 50;

    // Invariant s
    const Real S = ENERGY_CM*ENERGY_CM;

    // Rank of the numerator
    const int RANK = 4;

    // Declare an instance of the numerator
    Diagram num;

    // Assign numerical values to the reference vectors
    num.v1 = ComplexMomentum(1.0,1.1,1.2,1.3);
    num.v2 = ComplexMomentum(1.4,1.5,1.6,1.7);
    num.v3 = ComplexMomentum(1.8,1.9,2.0,2.1);
    num.v4 = ComplexMomentum(2.2,2.3,2.4,2.5);

    // Define external momenta
    RealMomentum k[N_LEGS];

    // Get a random phase-space point
    Rambo phase_space(S,N_LEGS);
    phase_space.getMomenta(k);

    // Define the internal momenta of the loop
    RealMomentum pi[N_LEGS];
    pi[0] = RealMomentum(0,0,0,0);
    pi[1] = k[0];
    pi[2] = k[0]+k[1];
    pi[3] = k[3];

    // Define the square of the internal masses
    Real msq[N_LEGS] = {1.,2.,3.,4.};

    // Create an amplitude object
    Amplitude<RealMasses> amp(N_LEGS,RANK,pi,msq);

    // Evaluate the integral
    amp.evaluate(num);
In the example above, after specifying some constants, we declare an instance num of the user-defined class Diagram, which we constructed in Section 4.2. Then we give numerical values to the reference vectors $v_i$ appearing in the numerator definition of Eq. (40) as well as in the analytic expression given in corresponding FORM file. This defines our numerator. Next we randomly generate a phase-space point, by creating a Rambo object and calling its method getMomenta which fills the array $k$ of external momenta. The phase space generation is a translation in C++ of the corresponding GOSam implementation, which in turn is based on the one of Ref. [36]. Every call of the method getMomenta on the same Rambo object randomly generates a different phase-space point. The code above assumes the external legs to be massless. If the external legs were massive, with masses \{MASS_0, MASS_1, MASS_2, MASS_3\}, we should have generated the phase-space point by passing an array of external masses as third argument to the constructor of the Rambo object, i.e.

\[
\text{Real external_masses[N_LEGS] = \{MASS_0, MASS_1, MASS_2, MASS_3\};}
\]

\[
\text{Rambo phase_space(S, N_LEGS, external_masses);}
\]

\[
\text{phase_space.getMomenta(k);}\]

The momenta $p_i$ are then defined according to Eq. (40). Arithmetic operations between momentum types work as one would expect. After specifying the square of the internal masses, we create an Amplitude<RealMasses> object amp, whose method evaluate computes the integral with the numerator specified in its argument. This adds the computed integral to the total result stored internally by amp, which can then be accessed either with the methods eps0, epsm1, epsm2 or with the subscript operator “[]” as the example shows.

4.3.2 The Amplitude class

The Amplitude template class is the main class of the Ninja library and its method evaluate computes a one-loop integral. The method can take as input an object of a class derived from ninja::Numerator, which provides a generic interface to the methods defined by each Laurent expansion. The template parameter of the Amplitude class is the type of the internal masses. Allowed types are: RealMasses, ComplexMasses and Massless. The methods needed for the evaluation of the amplitude are instantiated for all these three types in the compiled code of the library.

Instantiation In the example above, we showed how to instantiate an Amplitude object passing to its constructor the number of external legs, the rank of the numerator, the momenta $p_i$ and the squared masses $m_i^2$ of the loop denominators (cfr. Eq. (2)). There we assumed the internal masses to be real. For the complex-masses case and the massless case, the relevant part of the source would have looked like
// Complex masses
Complex msq[N_LEGS] = {...};
Amplitude<ComplexMasses> amp(N_LEGS,RANK,pi,msq);
amp.evaluate(num);

and

// Massless (msq does not need to be specified here)
Amplitude<Massless> amp(N_LEGS,RANK,pi);
amp.evaluate(num);

respectively. The Amplitude class also has a default constructor, as well as methods which allow to set or change the kinematics, the (squared) internal masses, the number of legs or the rank of the numerator to be evaluated, as in the following

Amplitude<RealMasses> amp;
amp.setN(N_LEGS).setRank(RANK);
amp.setKinematics(pi).setInternalMasses(msq);

Renormalization scale Another important setting is the renormalization scale $\mu_R^2$ to be used. This is equal to 1 by default, and it only affects the computation of the Master Integrals. It can be set as in the following example

// takes the square of the scale
amp.setRenormalizationScale(50*50);

The S-matrix An optional parameter which can be set by the user is the so-called S-matrix. This is defined in NINJA by

$$s_{ij} = (p_i - p_j)^2$$

where $p_i$ are the momenta appearing in Eq. (2). When this is specified by the user, the computation of the Master Integrals might be more accurate. This can be particularly useful in the presence of infrared singularities, which otherwise might not be detected by the integral library in use. The S-matrix can be declared either by an SMatrix object or by a simple $n^2$-dimensional array, where $n$ is the number of loop denominators. In the simple example given in Section 4.3.1, using the definitions of Eq. (40) in Eq. (41), for massless external momenta $k_i$ we could have specified the following S-matrix

$$
(s_{ij}) = 
\begin{pmatrix}
0 & 0 & 2(k_0 \cdot k_1) & 0 \\
0 & 0 & 0 & -2(k_0 \cdot k_3) \\
2(k_0 \cdot k_1) & 0 & 0 & 0 \\
0 & -2(k_0 \cdot k_3) & 0 & 0 \\
\end{pmatrix}
$$

(42)

either with

SMatrix s_mat;
s_mat.allocate(N_LEGS); // allocate the matrix
s_mat.fill(0); // fill the entries with zeros

22
s_mat(0,2) = s_mat(2,0) = 2*mp(k[0],k[1]);
s_mat(1,3) = s_mat(3,1) = -2*mp(k[0],k[3]);
amp.setSMatrix(s_mat);

or with

Real s_mat[N_LEGS*N_LEGS] = {0, 0, 2*mp(k[0],k[1]), 0,
                            0, 0, 0, -2*mp(k[0],k[3]),
                            2*mp(k[0],k[1]), 0, 0, 0,
                            0, -2*mp(k[0],k[3]), 0, 0};
amp.setSMatrix(s_mat);

We recommend to specify the S-Matrix whenever possible, and in particular when infrared singularities are present. As an alternative to writing it explicitly, one could use the method

SMatrix &
SMatrix::fillFromKinematics(const RealMomentum pi[],
Real ir_threshold = 0);

before each call of evaluate. This will automatically compute the matrix from the momenta $p_i$, but it will set to zero all the matrix elements which are smaller than $ir\_threshold$.

It can be worth pointing out that an Amplitude object only stores a pointer to the internal masses ($msq$), momenta ($pi$), and S-matrix ($s\_mat$) to be used, therefore the user should make sure these will exist in memory until the end of the execution of the evaluate method.

**Stopping the reduction earlier** There are cases where lower-point residues do not contribute to the final result for the integral (see e.g. the example in Section 5.3). If the user knows that $k$-point residues with $k < MIN\_CUT$ will not contribute to an amplitude, this information can be passed to NINJA through the setCutStop method, as in this example

amp.setCutStop(MIN_CUT);

which will tell NINJA to stop the reduction right after the evaluation of the residues of $k$-ple cuts with $k = MIN\_CUT$.

**Master Integrals** Each instance of an Amplitude object can in principle use a different library of Master Integrals. The library to be used can be specified with the method setIntegralLibrary as in the following example

Amplitude<RealMasses> amp;
amp.setIntegralLibrary(loop_tools);

If an integral library is not set explicitly for an amplitude object, the instance will use the one which is the default at the time of its creation. The default library will be ONELOOP if enabled during configuration, otherwise it will be LOOPTOOLS. If none of the two is enabled, NINJA can still be used, but a different library of Master Integrals should be specified at run-time (more details about the implementation of the corresponding interface are given in Appendix C). The function setDefaultIntegralLibrary can be used to change the default integral library. Assuming both ONELOOP and LOOPTOOLS have been enabled, the user can change the default as in the following example

// Set default integral library to LOOPTOOLS
setDefaultIntegralLibrary(LOOPTOOLS);

// Change default integral library to ONELOOP
setDefaultIntegralLibrary(ONELOOP);
```cpp
#include <ninja/ninja.hh>
#include <ninja/avholo.hh>
#include <ninja/looptools.hh>
using namespace ninja;

int main()
{
    setDefaultIntegralLibrary(loop_tools);
    // Amplitude objects defined here will use LoopTools
    setDefaultIntegralLibrary(avh_olo);
    // Amplitude objects defined here will use OneLoop
    return 0;
}
```

**Disabling the\(\mu^2\) expansion** The user can choose to avoid using the \(\mu^2\) expansion for the boxes with

```cpp
amp.useMuExpansion(false);
```

In this case the method `muExpansion` of the numerator class does not need to be provided. However, this would increase the number calls of the numerator method `evaluate` and it would also require the computation of pentagons. Given the simplicity of the \(\mu^2\) expansion, disabling it is therefore not recommended, if not for debugging purposes.

**Evaluation of the integrals** As already explained, integrals are computed by calling the method `evaluate`. This has the following prototype

```cpp
template <typename MassType>
int Amplitude<MassType>::evaluate(Numerator & num);
```

and takes as input the numerator of the integrand, which must be an instance of a class inherited from `ninja::Numerator`. It returns an integer value which depends on the results of the internal tests which Ninja can optionally perform during the computation (see Section 4.3.3). By default no test is performed and the return value can be safely ignored. In general, the return value will be equal to

- `Amplitude<MassType>::TEST_FAILED` if any of the performed tests failed
- `Amplitude<MassType>::SUCCESS` otherwise.

Each call of the method `evaluate` adds the computed integral to the total result stored internally by the instance. It can then be accessed either with the methods `eps0, epsm1, epsm2` or with the subscript operator “[]” as we illustrated in the simple test described in this section. The result can be quickly reset to zero, by calling the `reset` method,

```cpp
amp.reset();
```

The finite term is given by the sum of the `cut-constructible part` and the `rational part`, but the two can also be accessed separately with
4.3.3 Global settings

By default Ninja tries to compute a minimal set of coefficients during the reduction, i.e. those which are needed for the determination of the final integrated result. These are only a subset of the ones which are required for the full reconstruction of the integrand decomposition of Eq. (5). Indeed the computation of spurious coefficients is skipped whenever possible, i.e. whenever these do not enter the coefficient-level subtractions needed for lower-point residues, as in the case of spurious coefficients of pentagons, boxes and tadpoles. Ninja also entirely skips the computation of residues whose non-spurious coefficients would multiply scaleless integrals.

For debugging purposes, the user can however ask Ninja to perform some tests on the quality of the reconstruction of the integrand, or print some information about the ongoing computation. These operations might require the computation of a larger set of coefficients.

There are two kinds of tests which Ninja can perform: global tests and local tests. The global $N = N$ test [6, 37] checks that the following equality, which follows from Eq. (5), is valid

$$N(q, \mu^2) = \sum_{k=1}^{5} \Delta_{i_1 \cdots i_k} \prod_{h \neq i_1 \cdots i_k} D_h.$$  \hspace{1cm} (43)

Another global test, which can be performed when the rank $r$ of the numerator is equal to the number of external legs $n$, is the so-called power-test introduced in Ref. [14]. This consists in checking that the sum of all the spurious tadpole coefficients is vanishing,

$$\sum_{i_1=0}^{n-1} \sum_{k=0}^{4} c_k^{(i_1)} = 0.$$  \hspace{1cm} (44)

Finally, Ninja can perform local $N = N$ tests on quadruple, triple, double and single cuts. These will check the validity of Eq. (43) on values of the loop momenta corresponding to a given $k$-ple cut, and they can be useful in order to pinpoint the multiple cut where an error or an instability has occurred.

By default Ninja does not perform any internal test. The tests to be performed during the execution of the method evaluate can be set using the function

```c
void setTest(unsigned flag);
```

where the parameter flag can be any of the following:

- **Test::NONE** no test is performed
- **Test::ALL** all tests are performed
- **Test::GLOBAL** global tests are performed
- **Test::LOCAL** with $k \in \{1, 2, 3, 4\}$, local tests on $k$-ple cuts are performed
- **Test::LOCAL** all local tests are performed
or any combination of these. Different flags can be combined with the operator “|”. For instance, the following command will ask NINJA to perform global tests and local tests on double cuts

\[
\text{setTest} (\text{Test}::\text{GLOBAL} \mid \text{Test}::\text{LOCAL}_2);
\]

The \( \mathcal{N} = \mathcal{N} \) tests will check whether the numerator \( N_{\text{rec}} \) reconstructed by evaluating the r.h.s. of Eq. (43) is equal to the numerator \( N_{\text{eva}} \), obtained by a direct evaluation through the \textit{evaluate} method of the numerator class, up to a given tolerance. More explicitly, it checks if

\[
\left| \frac{N_{\text{rec}} - N_{\text{eva}}}{N_{\text{eva}}} \right| < \delta_{\text{tol}}
\]

(45)

where the threshold \( \delta_{\text{tol}} \) is \( 10^{-5} \) by default, but it can specified by the user through the function

\[
\text{void setTestTolerance(Real test\_tolerance);}
\]

As explained in Section 4.3.2, the return value of the method \textit{evaluate} of the \texttt{Amplitude} class can be used in order to check whether any performed test has failed. These tests have been implemented for debugging purposes, and they are not meant as an estimate of the accuracy of the total result. Indeed, there are cases where a numerical instability might cause a test to fail, while having negligible effects on the total amplitude. The accuracy of the result can be better estimated by means of the scaling test proposed in Ref. [22] or the rotation test described in Ref. [27].

Another global option which can be set is the \texttt{verbosity}, i.e. the amount of information printed during the evaluation of an integral. By default nothing is printed during a computation. The setting can be controlled by calling the function

\[
\text{void setVerbosity(unsigned flag);}\]

where possible values of the parameter \texttt{flag} can be

\begin{itemize}
  \item \texttt{Verbose::NONE} nothing is printed
  \item \texttt{Verbose::ALL} everything is printed (equivalent to the combination of all the other options)
  \item \texttt{Verbose::GLOBAL\_TEST} the result of global tests are printed, when performed
  \item \texttt{Verbose::LOCAL\_TEST\_k} with \( k \in \{1,2,3,4\} \), the result of local tests on \( k \)-ple cuts are printed when performed
  \item \texttt{Verbose::LOCAL\_TESTS} the result of all performed local tests is printed
  \item \texttt{Verbose::TESTS} the result of all performed tests is printed
  \item \texttt{Verbose::COEFFS} with \( k \in \{1,2,3,4,5\} \), the value of the coefficients of the computed \( k \)-point residues is printed
  \item \texttt{Verbose::RESULT} the value of all the computed coefficients is printed
  \item \texttt{Verbose::INTEGRALS} the value of the Master Integrals is printed
\end{itemize}
Similarly to the options controlling the performed tests, any combination of the flags above can be specified using the operator “|”. As an example, the following instruction will ask Ninja to print the value of the triangle coefficients, and the result of the current integral

```cpp
setVerbosity(Verbose::C3 | Verbose::RESULT);
```

When not specified otherwise, Ninja will print everything to standard output. Any other output stream can be set, as in the following example

```cpp
#include <fstream>
#include <ninja/ninja.hh>
using namespace ninja;

int main()
{
    std::ofstream f;
    f.open("my_file.out");
    setOutputStream(f);

    // from now on everything will be printed
    // by Ninja on the file "myfile.out"

    // ...

    return 0;
}
```

## 5 Examples

In this Section we give a description of the examples which are distributed with the library. In order to compile the corresponding executables, one can use the command

```bash
make examples
```

either in the root directory or in the examples directory. These examples are meant to provide a more detailed description of the usage of the library in several kinds of problems which involve the computation of one-loop integrals. More involved computations performed with the help of Ninja have been presented in Ref.s [26,27].

Every example presented in this paper has been generated with the help of the package NinjaNumGen, which is distributed with Ninja and is described in Appendix A. The package can be used both as a script and as a Python module. For each example, we include in the distribution

- the FORM file (with extension .frm) containing the analytic expression of the numerator which is used as input
- a SHELL script (with extension .sh) with the command we used for the generation of the numerator class methods
• a Python script (with extension .py) which achieves the same by importing the ninjanumgen module
• the C++ source files (with extension .cc) and headers files (with extension .hh) defining the numerator and its methods, as well as a test program.

5.1 Simple Test
The first simple example has already been extensively described in Section 4, in order to illustrate the basic usage of the library. The numerator class is defined in the header file mynum.hh, the source file generated by NINJANUMGEN is mynum.cc, while the source file with the main function is simple_test.cc.

5.2 Four-photon helicity amplitudes
In this example we consider a four-photon amplitude [38, 39] and we describe the usage of NINJA for the definition of polarization vectors and other spinor objects which are needed for the evaluation of the numerator.

The integrand of a diagram contributing to a four-photon amplitude is given by

\[ I = \frac{\mathcal{N}(q, \mu^2)}{D_0 D_1 D_2 D_3} \]

\[ \mathcal{N}(q, \mu^2) = - \text{Tr} \left( (\bar{l}_1 + m_f)\gamma^\mu (\bar{l}_2 + m_f)\gamma^\nu (\bar{l}_3 + m_f)\gamma^\rho (\bar{l}_0 + m_f)\gamma^\sigma \right) \]

\[ D_i = \bar{l}_i^2 - m_f^2 \]

(46)

where \( m_f \) is the mass of the fermion propagating in the loop, and the momenta \( \bar{l}_i \) are defined by

\[ \bar{l}_0 = \bar{q}, \quad \bar{l}_1 = \bar{q} + k_0, \quad \bar{l}_2 = \bar{q} + k_0 + k_1, \quad \bar{l}_3 = \bar{q} - k_4. \]

(47)

For simplicity we have assumed the four photons to be all incoming, i.e. \( k_0, k_1, k_2, k_3 \to 0 \).

The extra-dimensional components \( \vec{\mu} \) of the loop momentum satisfy the (anti-)commutation relations

\[ \{ \vec{p}, \vec{q} \} = 0, \quad \{ \vec{p}, \vec{p} \} = -\mu^2, \]

(48)

for any four-dimensional momentum \( p \). This allows to work out the extra-dimensional algebra and rewrite the numerator in terms of four-dimensional spinor products between the polarization vectors \( \epsilon_i \), such as \( \langle \epsilon_i, \epsilon_j \rangle \) and \( [\epsilon_i, \epsilon_j] \), and scalar products involving the four-dimensional momenta \( q, k_i \), and momenta \( e_{ij} \) defined by

\[ e_{ij} \equiv \frac{\langle \epsilon_i \gamma^\mu \epsilon_j \rangle}{2}. \]

(49)

The FORM package SPINNEY [40] can help in this kind of algebraic operations. The final expression can be found in the FORM file 4photons.frm of the directory examples of the distribution.

NINJA includes a small library for massless spinors, which is used internally for building the bases of momenta corresponding to each residue. This can also be useful for defining polarization vectors and other spinor-related objects. The spinor library can be used by including the header file ninja/spinors.hh in the source and linking the program with the NINJA library. Polarization vectors can be defined with
// Positive helicity (right-handed)
ComplexMomentum epsilon_r = polarizationVectorR(r,k);

// Negative helicity (left-handed)
ComplexMomentum epsilon_l = polarizationVectorL(r,k);

where \( r \) is an arbitrary reference momentum and \( k \) is the momentum of the corresponding photon (or gluon). These functions assume the momenta to be incoming, while for outgoing momenta the helicity should be reversed. Angle-bracket and square-bracket spinor products can be computed with the functions \texttt{spaa} and \texttt{spbb} respectively. If \( k \) is a (real or complex) massless momentum, the corresponding spinor \texttt{spinor\_k} can be defined as

\[
\texttt{Spinor spinor\_k = Spinor(k)};
\]

and can be used as input parameter for the functions described above. This turns out to be more efficient when several spinor-related operations need to be performed on the same momentum. Instances of the class \texttt{Spinor} can also be used in order to define vectors as in Eq. (49), using the following function

\[
// this returns \langle p\gamma^\mu q \rangle/2
\]

\[
\texttt{ComplexMomentum}
\texttt{momentumFromSpinors(const Spinor & p, const Spinor & q)};
\]

In the header file \texttt{4photons\_num.hh} we define a numerator class \texttt{FourPhotons} containing, as private data members, the values of all the momenta and spinor products appearing in the integrand. The numerator methods have been generated with \texttt{NINJA\_NUM\_GEN} and written in the file \texttt{4photons\_num.cc}. The file \texttt{4photons\_init.cc} contains the implementation of an \texttt{init} method which initializes the data members using the spinor-related operations described above, while \texttt{4photons.cc} contains a simple test. In this test, the mass of the fermion is complex, thus it can have a width. The results have been compared with the ones in Ref [39] as well as with a similar computation performed with \texttt{SAMURAI} for several choices of the external helicity states and the fermion mass. In Figure 1 we show a typical output for this example.

### 5.3 Six-photon helicity amplitudes

In this example we consider six incoming photons [37,41–46]. This is a non-trivial case where the \texttt{setCutStop} method of an \texttt{Amplitude} class can make the computation more efficient when lower point cuts do not contribute to the total result.

A generic six-photon diagram has an integrand of the form

\[
\mathcal{I} = \frac{\mathcal{N}(q, \mu^2)}{D_0 D_1 D_2 D_3 D_4 D_5}
\]

\[
\mathcal{N}(q, \mu^2) = - \text{Tr} \left( \bar{l}_i \epsilon_i \bar{l}_2 \epsilon_2 \bar{l}_3 \epsilon_3 \bar{l}_4 \epsilon_4 \bar{l}_5 \epsilon_5 \bar{l}_6 \epsilon_6 \right)
\]

\[
D_i = l_i^2
\]

where we assumed the fermion running in the loop to be massless. The momenta \( \bar{l}_i \) are defined
Finite: \((-0.184034, -0.16765)\)
Abs. val.: 0.248948
Single pole: \((-3.73035e-13, 3.98348e-13)\)
Double pole: (0,0)

Figure 1: Phase space point and output for the example in 4photons.cc. It shows the computed finite part and poles of an all-plus four-photon helicity amplitude, using a complex fermion mass \(m_f = 10.0 - 1.0i\).

by

\[
\begin{align*}
\bar{l}_0 &= \bar{q}, \\
\bar{l}_1 &= \bar{q} + k_0, \\
\bar{l}_2 &= \bar{q} - k_4 - k_5, \\
\bar{l}_3 &= \bar{q} - k_5.
\end{align*}
\]

One can work out the algebra, define the corresponding spinor products and vectors, and generate the input for \textsc{Ninja} in the same way as for the four-photons case. One can also check that the terms proportional to \(\mu^2\) in the final expression for the integral vanish upon integration. Therefore, we can perform the simplification \(\bar{l}_i \to l_i\), or equivalently \(\mu^2 \to 0\), in the numerator. Moreover, one can exploit the knowledge that only the cut-constructible contributions of boxes and triangles contribute to the total result, hence we can ask \textsc{Ninja} to stop the reduction at triple cuts with

\[
\text{amp.setCutStop}(3);
\]

and remove the rational part from the result with
which will make the computation more efficient (in the example implemented here, the run-
time is reduced by about 33%).

In the file `6photons.cc` we call the method `evaluate` on all the independent permutations
of the external legs, generated at run-time with the function `std::next_permutation` of the
C++ standard library. The results have been compared with the ones in Ref.s [45,46] as well
as with a similar computation performed with SAMURAI for several helicity choices.

5.4 Five-point diagram of $gg \rightarrow Ht\bar{t}$

With this example, we discuss a possible strategy for the generation of the input needed by
NINJA which can be suited for more complex computations where an efficient evaluation of
the numerator methods at run-time can be important.

We consider the one-loop integral defined by the diagram depicted in Figure 2, contributing
to the 5-point helicity amplitude $g(k_1, -), g(k_2, -) \rightarrow H(k_3), t(k_4, +), \bar{t}(k_5, -)$. The analytic

![Diagram](image)

Figure 2: Diagram contributing to $gg \rightarrow Ht\bar{t}$. This picture has been generated using GoSam.

expression for the integrand of this example, which can be worked out from the Feynman rules
of the Standard Model, has been generated with the help of the package GoSAM and can
be found in the FORM file `ttbarh.frm`. This already contains some abbreviations which are
independent from the loop momentum $\bar{q}$ of the diagram. At run-time, these $\bar{q}$-independent
abbreviations are computed only once per phase space point, making thus the evaluation of
the numerator and its expansions more efficient. This analytic expression is processed by
NINJA_NUMGEN which produces the numerator expansions. We also add to the numerator
class `TTbarHDiagram` an `init` method which uses the spinor library described in Section 5.2
in order to compute the relevant spinor products and polarization vectors, as well as the
abbreviations which do not depend on the loop momentum. These are stored as private data
members of the class. For simplicity, our result neglects the coupling constants and an overall
color factor.

Even though we considered a single diagram and a specific helicity choice, this example
illustrates a general strategy for the generation of an analytic numerator expression which is
suited for the numerical evaluations performed by integrand-reduction algorithms like the one implemented in the library Ninja. The full amplitude for this process has been computed in Ref.s [47–51], while an additional jet has recently been added to the final state in Ref.s [26,27] where Ninja has been used for the reduction of the corresponding integrands generated by GoSam.

5.5 Higher-rank example

In this example we show how Ninja can be used in order to compute integrals whose rank is higher than the number of loop denominators. This simple test is similar to the example presented in Ref. 4, hence we will describe each step as in the previous case. We define a 5-point amplitude of rank 6, with kinematics $k_0, k_1 \rightarrow k_2, k_3, k_4, k_5$ and integrand

$$I = \frac{\mathcal{N}(q, \mu^2)}{D_0 D_1 D_2 D_3 D_4}$$

$$\mathcal{N}(q, \mu^2) = \prod_{i=0}^{2} \left( (q \cdot v_{2i})(q \cdot v_{2i+1}) + \mu^2 (v_{2i} \cdot v_{2i+1}) \right)$$

$$D_i = l_i^2 - m_i^2$$

in terms of the reference vectors $v_i$ ($i = 0, \ldots, 5$) and the momenta $\bar{l}_i$ running into the loop

$$\bar{l}_0 = \bar{q}, \quad \bar{l}_1 = \bar{q} + k_0, \quad \bar{l}_2 = \bar{q} + k_0 + k_1, \quad \bar{l}_3 = \bar{q} + k_3 + k_4, \quad \bar{l}_4 = \bar{q} + k_4.$$  

We follow the same steps outlined in Section 4. With NinjaNumGen we generate the methods for Ninja. After writing the integrand in the FORM file mynumhr.frm we call the script with the command

```
ninjanumgen mynumhr.frm --nlegs 5 --rank 6 -o mynumhr.cc
```

which creates the file mynumhr.cc and a template for the header mynumhr.hh. Once again, we define the vectors $v_i$ as public members of the numerator class Diagram, by inserting

```
public:
    ninja::ComplexMomentum v0,v1,v2,v3,v4,v5;
```

in the class definition. A possible test program can be almost identical to the one we showed in Section 4.3.1, with obvious changes in the definition of the rank, the number of external legs and the reference vectors $v_i$. This is implemented in the file simple_higher_rank_test.cc. In order to run this example, the user must compile the library with the --enable-higher rank option, otherwise a call to the evaluate method of an Amplitude object will cause a run-time error.

As one can see, when NinjaNumGen is used for the generation of the expansions, the higher-rank case is handled automatically without any intervention by the user. Besides, the internal higher-rank routines of Ninja will be automatically called whenever the rank $r$ equals to $n + 1$ (where $n$ is the number of loop propagators), while in the public programming interface there is no difference with respect to the normal-rank case.
5.6 Usage in multi-threaded applications

In the last examples, we wish to illustrate the possibility of using Ninja in a multi-threaded application. These examples are implemented using POSIX threads, which are a standard in Unix-like operating systems, but adapting them to different programming interfaces for threads (such as OpenMP) should be straightforward.

In order to implement a thread-safe application, one should avoid race conditions which might occur if different threads try to write on the same variables. In particular, one should avoid accessing global variables for writing from different threads. The only global variables used directly by Ninja are those controlling the global options described in Section 4.3.3. As explained in that section, these options are only meant to change the general behavior of the library for debugging purposes (e.g. for checking that the provided numerator methods are correct), while in general the default options should not be changed during a phase-space integration, especially when performance is important. Hence, on the side of the Ninja library, there should be no issue and one can safely call the `evaluate` method from different `Amplitude` objects in different threads.

During a call of the `evaluate` method on an `Amplitude` object, issues might however arise from global variables used by the chosen library of Master Integrals or the numerator methods. As for the numerator methods, all the examples distributed with Ninja define a thread-safe numerator class (more specifically, one can safely call numerator methods from different instances of the class in different threads). This is simply done by using data members of the class instead of global variables, making thus different instances of the same class completely independent.

If the procedures implemented by libraries of Master Integrals are thread-safe, one can therefore use Ninja in multi-threaded applications. As an example, one can use the class `AvHOneLoop` which, as explained in Section 3.3, wraps routines of the `OneLoop` library and adds a cache of computed integrals. The cache is a non-static data member of the class. One can therefore create one instance of this class per thread and assign it accordingly to the `Amplitude` objects to be evaluated in the same thread. As an example, with

```c
avh_olo.init(1);
AvHOneLoop my_lib[N_THREADS];
Amplitude<RealMasses> amp[N_THREADS];
for (int i=0; i<N_THREADS; ++i)
    amp[i].setIntegralLibrary(my_lib[i]);
```

we create `N_THREADS` amplitude objects whose `evaluate` method can be safely called in a separate thread (in the first line, we called the `init` method on the global instance `avh_olo` defined in the library, in order to allow `OneLoop` to perform its global initialization). In this way, different threads will also have an independent cache of Master Integrals. This strategy allows to build a multi-threaded application which uses Ninja for the reduction of one-loop integrals.

In the following we discuss the possibility to build a multi-threaded application with Ninja and any other (not necessarily thread-safe) library of Master Integrals. Indeed, even though Ninja has obviously no control over possible issues arising from routines of external libraries, we offer an easy way to work around any potential problem. In this case, there is no general way to ensure that calling routines of the same integral library from different threads will not cause conflicts. However, one can avoid these conflicts by scheduling the calls of the external
procedures in such a way that they are never evaluated at the same time from two or more threads. If the computation of the Master Integrals takes only a small fraction of the total run time (which is usually the case when a cache of integrals is present), the effects of this on the performance will in general be reasonably small.

Within Ninja, implementing a scheduled access on the routines used by a library of Master Integrals is straightforward. As explained more in detail in Appendix C, the generic interface used by Ninja in order to call Master Integral procedures, has two methods called init and exit which are evaluated exactly once in each call of the evaluate method, immediately before the computation of the first Master Integral and after the computation of the last Master Integral respectively. Therefore we can use mutexes (such as the ones present the POSIX standard for threads) in order to lock the calls to the Master Integrals in the init method and unlock them in the exit method. This makes sure that, between the calls of the init and exit methods, no other thread will use the Master Integral routines, hence avoiding any possible conflict.

In order to make a library of Master Integrals thread-safe, we use the template class ThreadSafeIntegralLibrary, which is included in the distribution. This automatically wraps an existing class derived from IntegralLibrary and adds to it a mutex that schedules the calls to the Master Integrals as explained above. As an example, defining a thread-safe version of a generic library BaseLibrary can be simply achieved with

```cpp
#include <ninja/thread_safe_integral_library.hh>
using namespace ninja;
ThreadSafeIntegralLibrary<BaseLibrary> my_lib;
```

which defines a new interface `my_lib` that can be made the default by calling

```cpp
setDefaultIntegralLibrary(my_lib);
```

before any thread is created (alternatively, we could call the setIntegralLibrary method on each Amplitude object, either outside or inside the threads).

In the files thread_4photons.cc and thread_6photons.cc we repeat the examples of the four- and six-photons amplitudes, but this time we compute several phase-space points in parallel on different threads. As mentioned before, we do not need to implement other numerator classes, since the ones described in Sections 5.2 and 5.3 can be safely used in multi-threaded applications. In the source files, we implement both the approaches described in this section. The preprocessor will select the former if the ONELOOP interface has been enabled and the latter otherwise. The multi-threaded examples can be compiled with

```bash
make thread-examples
```

if at least one between the ONELOOP and LOOPTOOLS libraries was enabled during configuration and your system supports POSIX threads.

A complete discussion on the implementation of multi-threaded applications for doing phenomenology at one-loop is beyond the purposes of this paper. Moreover, a detailed assessment of possible advantages of this approach would generally depend on the generator of the integrands and the phase space integration. In these examples we showed that the methods implementing the reduction via Laurent expansion in Ninja can be safely used in multi-threaded programs.
6 Conclusions

We presented the public C++ library **NINJA** which implements the Integrand Reduction via Laurent Expansion method for the computation of one-loop amplitudes in Quantum Field Theories. The main procedures of the library take as input the numerator of the integrand and some parametric expansions of the same, which can be generated with the help of the simple **PYTHON** package **NINJANUMGEN** included with the distribution. The expansions of the integrand on the multiple cuts are computed semi-numerically at run-time, via a simplified polynomial-division algorithm. Some of the coefficients of the Laurent expansions are thus identified with the ones which multiply the Master Integrals. The algorithm is light and proved to have good performance and numerical stability, hence it is suited for applications to complex one-loop processes, characterized by either several external legs or several mass scales.

We described the usage of the library, in particular the generation of the input, the calls of the procedures for the reduction, and the interface to libraries of Master Integrals. This information can be used in order to interface the library with existing one-loop packages. We thus expect that **NINJA** will be useful for future computations in high-energy physics, especially for those involving more complex processes.

Acknowledgments

The author thanks all the other members of the GoSam collaboration for the common development of a one-loop package which could be interfaced with **NINJA**, and especially Pier-paolo Mastrolia, Edoardo Mirabella and Giovanni Ossola for innumerable discussions and exchanges. The author also thanks Thomas Hahn for his support with **LOOPTOOLS**. This work was supported by the Alexander von Humboldt Foundation, in the framework of the Sofja Kovalevskaja Award Project “Advanced Mathematical Methods for Particle Physics”, endowed by the German Federal Ministry of Education and Research.

Appendix A The **PYTHON** package **NINJANUMGEN**

The reduction procedures implemented in **NINJA** take as input a class derived from the abstract class **ninja::Numerator**. This is supposed to implement the required expansions in the corresponding methods. The source code for the methods can be automatically generated with the help of the simple **PYTHON** package **NINJANUMGEN** which is distributed with the library and can be installed as explained in Section 4.1. The package can be used both as a script or as a module within **PYTHON**.

In Section 4.2 we already gave a simple example of its usage as a script. As explained there, the user is supposed to create a file containing a **FORM** expression of the numerator of the integrand. The package uses **FORM-4** in order to generate the expansions which are needed and produce a C++ source file with the definitions of the corresponding methods. If not already present, an header file with a sketch of the definition of the class will also be created. The user can complete it by adding data members and methods which are specific of this class. **FORM** allows one to define symbols between square brackets (e.g. [[symbol name]]), containing characters which otherwise would not be permitted in a declaration. **NINJANUMGEN** also allows the usage of such symbols in the expression of the numerator, and it will
remove the brackets (which would produce illegal C++ code) when writing the final source files. This gives the user a wider range of possibilities, for instance using symbols which correspond to variable names containing underscores or data members of structures (e.g. with [structure_instance.data_member]).

We first give a few more details about the usage of the package as a script. It is invoked with the command

```
ninjanumgen --nlegs NLEGS optional-arguments file
```

where file is the name of the file which contains the numerator expression and NLEGS is the number of external legs of the loop, which is equal to the number of loop denominators. A description of all the allowed arguments can be obtained with the command

```
ninjanumgen --help
```

and the most important ones are:

--rank RANK, -r RANK rank of the numerator, by default it will be assumed to be equal to the number of external legs of the loop

--diagname DIAGNAME, -d DIAGNAME name of the numerator expression in the FORM file, by default it will be assumed to be Diagram

--cdiagname CDIAGNAME name of the numerator class in the generated C++ files, by default it will be the same as the FORM expression

--formexec FORMEXEC the FORM executable, the default is form

--qvar QVAR name of the loop momentum variable q defined in Eq. (4), the default is Q

--mu2var MU2VAR name of the loop variable µ^2 defined in Eq. (4), the default is Mu2

--output OUTPUT, -o OUTPUT name of the output source file, the default is ninjanumgen.cc

--header HEADER C++ header file containing the definition of the numerator class: if the file does not exists, one will be created. By default it will have the same name as the output but with .hh extension.

As mentioned, one can also use the package as a PYTHON module (ninjanumgen). This contains a class DiagramExpansion which can be used for the generation of the source code which implements the numerator methods. The input parameters of the constructor of this class roughly correspond to the arguments which can be used in the script. A detailed description can be obtained, after installation, by invoking PYTHON in interactive mode (usually done with the command python) and typing

```
import ninjanumgen
help(ninjanumgen.DiagramExpansion)
```

The method writeSource generates the source files. As a simple example, the source for the integrand we defined in Section 4.2 could have been generated within PYTHON with the commands
# import the module
import ninjanumgen

# define the mandatory arguments for the constructor
n_legs = 4
input_file = 'mynum.frm'
output_file = 'mynum.cc'

# define an instance of the class DiagramExpansion
mynum = ninjanumgen.DiagramExpansion(input_file,
                                      output_file,
                                      n_legs, rank=4)

# generate the source
mynum.writeSource()

We suggest to look at the PYTHON files in the examples directory for other basic examples.

**Appendix B Higher-rank numerators**

As pointed out in Ref. [24], the Laurent expansion method can be generalized to non-renormalizable and effective theories with higher-rank numerators. In a renormalizable theory, with a proper choice of gauge the rank \( r \) cannot be greater than the number \( n \) of loop propagators. Ninja, if configured with the `--enable-higher_rank` option, can also be used for the computation of integrals with \( r = n + 1 \). Here we describe the generalization of the method to the higher-rank case, underlining the points where it differs from the renormalizable case.

In Eq. (14), we gave the most general parametrization of the residues \( \Delta_{i_1 \cdots i_k} \) in a renormalizable theory. In the higher-rank case with \( r = n + 1 \), such parametrization is generalized as follows [24]

\[
\begin{align*}
\Delta_{i_1 i_2 i_3 i_4 i_5}^{(r=n+1)} &= \Delta_{i_1 i_2 i_3 i_4 i_5} \\
\Delta_{i_1 i_2 i_3 i_4}^{(r=n+1)} &= \Delta_{i_1 i_2 i_3 i_4} + c_5 \mu^4 x_{e,4} \\
\Delta_{i_1 i_2 i_3}^{(r=n+1)} &= \Delta_{i_1 i_2 i_3} + \mu^2 (c_{10} x_1^2 + c_{11} x_3^2) + c_{12} x_4^4 + c_{13} x_3^4 + c_{14} \mu^4 \\
\Delta_{i_1 i_2}^{(r=n+1)} &= \Delta_{i_1 i_2} + \mu^2 (c_{10} x_1 + c_{11} x_4 + c_{12} x_3) + c_{13} x_1^3 + c_{14} x_3^4 + c_{15} x_3^3 \\
&\quad + c_{16} x_1^2 x_4 + c_{17} x_1^2 x_3 + c_{18} x_1 x_3^2 + c_{19} x_1 x_3^2 \\
\Delta_{i_1}^{(r=n+1)} &= \Delta_{i_1} + c_5 x_2^2 + c_6 x_1^2 + c_7 x_4^2 + c_8 x_3^2 + c_{10} x_2 x_4 + c_{11} x_2 x_3 \\
&\quad + c_{12} x_1 x_4 + c_{13} x_1 x_3 + c_{14} \mu^2 + c_{15} x_3 x_4. \quad (B.1)
\end{align*}
\]
The generalized integral decomposition is thus

\[ M^{(r=n+1)} = M^{(r=n)} + \sum \left\{ c_{14}^{(i_1 i_2 i_3)} I_{i_1 i_2 i_3} [\mu^4] \right\} + \sum \left\{ c_{10}^{(i_1 i_2)} I_{i_1 i_2} [\mu^2 (q + p_{i_1}) \cdot e_2] + c_{13}^{(i_1 i_2)} I_{i_1 i_2} [(q + p_{i_1}) \cdot e_2)] \right\} + \sum \left\{ c_{14} I_{i_1} [\mu^2] + c_{15} I_{i_1} [(q + p_{i_1}) \cdot e_3)((q + p_{i_1}) \cdot e_4)] \right\} \]  

(B.2)

This higher-rank decomposition has been used for the computation of NLO corrections to Higgs-boson production in association with two [52] and three [27, 53] jets. Other libraries which implement the reduction of higher-rank integrals are XSAMURAI [54], which extends the more traditional integrand reduction algorithm of SAMURAI, and GLOEM95 [55–57].

### B.1 Reduction of higher-rank integrands

While the extension of the Laurent expansion method for the computation of higher-rank 3-point and 2-point residues is straightforward, for 4-point and 1-point residues some further observations are in order. Here we propose a generalization of the Laurent expansion method which allows to efficiently compute the non-spurious coefficients of 4- and 1-point residues without spoiling the nice features of the algorithm, such as the simplified subtractions of higher-point contributions and the diagonal systems of equations. This generalization is not present elsewhere in the literature and has been implemented in the NINJA library.

**4-point residues** The coefficient \( c_0 \) can be computed exactly as in the renormalizable case. For the coefficient \( c_4 \), one needs instead to keep also the next-to-leading term in the \( \mu^2 \) expansion described before, so that the \( d \)-dimensional solutions of a quadruple cut, given in Eq. (17), in the asymptotic limit become

\[ q_\pm^\nu = -p_\nu^\nu + a^\nu \pm \sqrt{\mu^2 + \beta v_{\perp}^\nu} \mu^2 \rightarrow \infty -p_\nu^\nu + a^\nu \pm \sqrt{\mu^2} v_{\perp}^\nu + O \left( \frac{1}{\sqrt{\mu^2}} \right), \]  

(B.3)

where it is worth noticing that \( a^\nu \) can be obtained as the average of the two solutions of the corresponding four-dimensional quadruple cut. In this limit, the expansion of the integrand reads

\[ \mathcal{N}(q, \mu^2) \prod_{j \neq i_1, i_2, i_3, i_4} D_j \bigg|_{q = \sqrt{\mu^2 v_{\perp} + a + O(\mu^{-1})}} = c_5 v_{\perp}^2 \mu^5 + c_4 \mu^4 + O(\mu^3), \]  

(B.4)

hence the leading term is now the spurious coefficient \( c_5 \), but \( c_4 \) can still be obtained as the next-to-leading term. This can be implemented semi-numerically, by keeping the two leading terms of the expansion of the numerator and performing a polynomial division with respect to the two leading terms in the expansion of the uncut denominators which have the form

\[ D_{h \neq i_1, i_2, i_3, i_4} \big|_{q = q_4} = d_{h, 0} \sqrt{\mu^2} + d_{h, 1} + O \left( \frac{1}{\sqrt{\mu^2}} \right). \]  

(B.5)

Given the very limited number of terms involved, the division can be implemented very efficiently in a small number of operations. More in detail, if \( \text{num} \) and \( \text{den} \) are arrays of length...
two containing the leading and next-to-leading terms in the expansion of the numerator and a denominator respectively, we can perform the division in place with the commands

```c
num[0] /= den[0];
num[1] -= den[1]*num[0];
num[1] /= den[0];
```

which will have the effect of replacing the entries of `num` with the ones of the expansion of \(N/D\). We also observe that the computation and the subtraction of pentagons is not needed in the higher-rank case either.

1-point residues On higher-rank 1-point residues \(\Delta_i\), we consider \(d\)-dimensional solutions of the corresponding single cut of the form

\[ q_+ = -p_i + t e_1^0 + \frac{m_{i1}^2 + \mu^2}{2t(e_1 \cdot e_2)} e_2^0, \quad q_- = -p_i + t e_3 + \frac{m_{i1}^2 + \mu^2}{2t(e_3 \cdot e_4)} e_4^0, \]  

in terms of the free variables \(t\) and \(\mu^2\). By taking the \(t \to \infty\) limit of the integrand and the subtraction terms evaluated on these solutions, we obtain an asymptotic polynomial expansion of the form

\[ \frac{N(q_\pm,\mu^2)}{\prod_{j \neq i} D_j} = n_0^+ + n_1^+ t + n_2^+ t^2 + n_3^+ \mu^2 + \mathcal{O}(1/t) \]  

\[ \frac{\Delta_{i+j}(q_\pm,\mu^2)}{D_j} = c_{s_{2,1}}^{\pm(j)} + c_{s_{2,2}}^{\pm(j)} t + c_{s_{2,3}}^{\pm(j)} \mu^2 + \mathcal{O}(1/t) \]  

\[ \frac{\Delta_{i+j+k}(q_\pm,\mu^2)}{D_j D_k} = c_{s_{3,0}}^{\pm(j,k)} + c_{s_{3,1}}^{\pm(j,k)} t + c_{s_{3,2}}^{\pm(j,k)} \mu^2 + \mathcal{O}(1/t). \]

One can check that the non-spurious coefficients of the tadpole are given in terms of the ones of the expansions above by

\[ c_0 = n_0^+ - \sum_j c_{s_{2,0}}^{+(j)} - \sum_{jk} c_{s_{3,0}}^{+(j,k)}, \]

\[ c_{14} = n_3^+ - \sum_j c_{s_{3,2}}^{+(j)} - \sum_{jk} c_{s_{3,3}}^{+(j,k)}, \]

\[ c_{15} = \frac{2}{(e_3 \cdot e_4)} \left( n_3^- - \sum_j c_{s_{2,3}}^{-(j)} - \sum_{jk} c_{s_{3,3}}^{-(j,k)} - c_{14} \right). \]

B.2 The input

In the higher-rank case, the `muExpansion` method of the numerator needs to compute both the leading and the next-to-leading term of the expansion in \(\mu^2\). The package `NinjaNumGen`, takes care of this automatically when the specified rank is higher than the number of external legs of the loop. The information in the next paragraph is only needed for a custom implementation of the method without `NinjaNumGen`.

The `muExpansion` method in the higher-rank case should compute the two leading terms of the expansion in \(t\) of the numerator, defined by

\[ q' \to t v_0' t + v_1', \quad \mu^2 \to t^2 v_0^2 \]  

(B.11)
with $v_i \equiv v[1]$, where $v$ is the array of momenta passed as input parameter. The leading and next-to-leading terms of the expansion should be written in the first two entry of the array pointed by the parameter $c$, i.e.

\[
\begin{align*}
    c[0] &= \mathcal{N}[t'] ; \\
    c[1] &= \mathcal{N}[t'^{-1}] ; 
\end{align*}
\]

All the other methods should have instead the same definition described in Section 3.1.

### B.3 Higher-rank Master Integrals

As one can see form Eq. (B.2), in the higher-rank case five new types of integral appear in the final decomposition. They are a 2-point integral of rank 3, a 1-point integral of rank 2, and three more integrals containing $\mu^2$ at the numerator which contribute to the rational part of the amplitude.

**Ninja** contains an implementation of all these higher-rank integrals in terms of lower-rank ones. This means that, should the user choose to interface a custom integral library (see Appendix C), these higher-rank integrals would not be needed, although specifying an alternative implementation would still be possible.

All the integrals of Eq. (B.2) which contribute to the rational part of the amplitude have already been computed in Ref. [24]. With our choice of the normalization factor given in Eq. (3), they read

\[
I_{i_1i_2i_3}[\mu^4] = \frac{1}{6} \left( \frac{s_{i_2i_1} + s_{i_3i_2} + s_{i_1i_3}}{4} - m^2_{i_1} - m^2_{i_2} - m^2_{i_3} \right) + \mathcal{O}(\epsilon) \quad (B.12)
\]

\[
I_{i_1i_2}[\mu^2 ((q + p_{i_1}) \cdot v)] = \frac{(p_{i_2} - p_{i_1}) \cdot v}{12} \left( s_{i_2i_1} - 2 m^2_{i_1} - 4 m^2_{i_2} \right) + \mathcal{O}(\epsilon) \quad (B.13)
\]

\[
I_{i_1}[\mu^2] = \frac{m^4_{i_1}}{2} + \mathcal{O}(\epsilon) \quad (B.14)
\]

where $s_{ij}$ were defined in Eq. (41) and $v$ is an arbitrary vector. The tadpole of rank 2 appearing in Eq. (B.2) can also be written as a function of the scalar tadpole integral $I_{i_1}$ as follows

\[
I_{i_1}[(q + p_{i_1}) \cdot e_3 ((q + p_{i_1}) \cdot e_4)] = m^2_{i_1} \left( \frac{e_3 \cdot e_4}{4} \right) \left( I_{i_1} + \frac{m^2_{i_1}}{2} \right) + \mathcal{O}(\epsilon). \quad (B.15)
\]

Since the vector $e_2$ in the bubble integral of rank 3 appearing in Eq. (B.2) is massless, the corresponding integral is simply proportional to the form factor $B_{111}$.

\[
I_{i_1i_2}[(q + p_{i_1}) \cdot e_2]^3 = ((p_{i_2} - p_{i_1}) \cdot e_2)^3 B_{111}(s_{i_2i_1}, m^2_{i_1}, m^2_{i_2}). \quad (B.16)
\]

The form factor can be computed using the formulas of Ref. [58], as a function of form factors of scalar integrals $B_0$. In the special case with $s_{i_2i_1} = 0$ we use Eqs (A.6.2) and (A.6.3) of
that reference. For the general case \( s_{i2i1} \neq 0 \) we implement instead the following formula

\[
B_{111}(s_{i2i1}, m_{i1}^2, m_{i2}^2) = \frac{1}{4 s_{i2i1}} \left\{ s_{i2i1} \left( m_{i1}^2 I_{i1} + I_{i1} [\mu^2] - m_{i2}^2 I_{i2} - I_{i2} [\mu^2] \right) \\
- 4 I_{i1i2} [\mu^2 ((q + p_{i1}) \cdot (p_{i2} - p_{i1}))] \\
- 4 m_{i1}^2 I_{i1i2} [(q + p_{i1}) \cdot (p_{i2} - p_{i1})] \\
+ 4 (m_{i2}^2 - m_{i1}^2 - s_{i2i1}) I_{i1i2} [(q + p_{i1}) \cdot (p_{i2} - p_{i1})]^2 \right\},
\]

(B.17)

Appendix C Interfaces to Integral Libraries

\textsc{Ninja} already implements interfaces for the \textsc{Oneloop} and the \textsc{LoopTools} integral libraries. These libraries have been used in a large number of computations and provide very reliable results, hence they should suffice for most purposes. However, \textsc{Ninja} has been designed considering the possibility of using any other library of Master Integrals.

The Master Integrals are computed by calling virtual methods of the abstract class \texttt{ninja::IntegralLibrary}, which is defined in the header file \texttt{ninja/integral_library.hh}. Therefore, any library of Master Integrals can be interfaced by implementing a class derived from \texttt{IntegralLibrary}. Each method of the library corresponds to a different Master Integral appearing in Eq. (15), which should be implemented for both real and complex internal masses (and optionally for the massless case). An implementation of higher-rank integrals can also be provided but it is not needed, since \textsc{Ninja} has a default implementation of them in terms of lower rank integrals. There are two further methods, namely \texttt{init} and \texttt{exit}. The former is called inside the method \texttt{Amplitude::evaluate} just before the computation of the first needed Master Integral, while the latter is called after the last Master Integral has been computed. The method \texttt{init}

\begin{verbatim}
  virtual void init(Real muRsq) = 0;
\end{verbatim}

takes as input the square of the renormalization scale to be used in the subsequent calls of the methods implementing the Master Integrals. It can also be used in order to perform any other initialization the library might need before computing the integrals. The \texttt{exit} method instead, does not need to be implemented and by default it will not perform any action. In Section 5.6 we gave an example of a case where a non-trivial implementation of the \texttt{exit} method could be useful.

The other methods should compute the finite part and the poles of the corresponding Master Integrals. As an example, the methods for the box integrals have the following declarations

\begin{verbatim}
// - real masses
  virtual void getBoxIntegralRM(Complex rslt[3],
      Real s21, Real s32, Real s43,
      Real s14, Real s31, Real s42,
      Real m1sq, Real m2sq,
      Real m3sq, Real m4sq) = 0;
\end{verbatim}
virtual void getBoxIntegralCM(Complex rslt[3],
    Real s21, Real s32, Real s43,
    Real s14, Real s31, Real s42,
    const Complex & m1sq,
    const Complex & m2sq,
    const Complex & m3sq,
    const Complex & m4sq) = 0;

and they are supposed to write the $O(\epsilon^{-i})$ term of the result in the $i$-th entry of the array rslt, for $i \in \{0,1,2\}$. The arguments are the invariants $s_{ij}$ and the squared masses $m^2_i$.

Similar methods need to be provided for 3-point, 2-point and 1-point Master Integrals, as described in detail in the comments inside the header file ninja/integral_library.hh.

C.1 Built-in interfaces

Examples of implementation of this interface for the libraries ONELOOP and LOOPTOOLS can be found in the source code. More in detail, we define the instances ninja::avh_olo and ninja::loop_tools of the classes ninja::AvHOneLoop and ninja::LoopTools respectively, which implement the methods described above as wrappers of the corresponding routines in each integral library.

The ONELOOP interface also implements a cache of Master Integrals on top these routines. The cache is implemented similarly to an hash table, which allows constant-time look-up of each computed integral from its arguments. Hence, the methods of the AvHOneLoop class will call the routines of the ONELOOP library only if a Master Integral is not found in the cache. The cache can be cleared with the class method AvHOneLoop::clearIntegralCache. During a phase-space integration, we suggest calling this method once per phase space point, especially for most complex processes. This method does not completely free the allocated memory, but keeps the buckets of the hash table available in order to store the integrals more efficiently in subsequent calls of the respective methods. If the user wishes to completely free the allocated memory, the method AvHOneLoop::freeIntegralCache can be used, although in general clearIntegralCache should be preferred. As already mentioned, every instance of AvHOneLoop has a cache of Master Integrals as data member. This can be useful for building multi-threaded applications, as discussed in the examples of Section 5.6.

Since LOOPTOOLS already has an internal cache of Master Integrals, the implementation of its interface is much simpler and only consists in wrapper of its routines. We implemented a clearIntegralCache method in the LoopTools class as well, which in this case simply calls the routine which clears the cache of integrals in LOOPTOOLS.

References

[1] F. Cachazo, P. Svrcek and E. Witten, JHEP 0409 (2004) 006 [hep-th/0403047].

[2] R. Britto, F. Cachazo and B. Feng, Nucl. Phys. B 715 (2005) 499 [hep-th/0412308].
[3] R. Britto, F. Cachazo, B. Feng and E. Witten, Phys. Rev. Lett. 94 (2005) 181602 [hep-th/0501052].

[4] Z. Bern, L. J. Dixon, D. C. Dunbar and D. A. Kosower, Nucl. Phys. B 425 (1994) 217 [hep-ph/9403226].

[5] R. Britto, F. Cachazo and B. Feng, Nucl. Phys. B 725 (2005) 275 [hep-th/0412103].

[6] G. Ossola, C. G. Papadopoulos and R. Pittau, Nucl. Phys. B 763 (2007) 147 [hep-ph/0609007].

[7] R. K. Ellis, W. T. Giele and Z. Kunszt, JHEP 0803 (2008) 003 [arXiv:0708.2398 [hep-ph]].

[8] P. Mastrolia and G. Ossola, JHEP 1111 (2011) 014 [arXiv:1107.6041 [hep-ph]].

[9] S. Badger, H. Frellesvig and Y. Zhang, JHEP 1204 (2012) 055 [arXiv:1202.2019 [hep-ph]].

[10] Y. Zhang, JHEP 1209 (2012) 042 [arXiv:1205.5707 [hep-ph]].

[11] P. Mastrolia, E. Mirabella, G. Ossola and T. Peraro, Phys. Lett. B 718 (2012) 173 [arXiv:1205.7087 [hep-ph]].

[12] P. Mastrolia, E. Mirabella, G. Ossola and T. Peraro, Phys. Lett. B 727 (2013) 532 [arXiv:1307.5832 [hep-ph]].

[13] G. Ossola, C. G. Papadopoulos and R. Pittau, JHEP 0803 (2008) 042 [arXiv:0711.3596 [hep-ph]].

[14] P. Mastrolia, G. Ossola, T. Reiter and F. Tramontano, JHEP 1008 (2010) 080 [arXiv:1006.0710 [hep-ph]].

[15] T. Hahn and M. Perez-Victoria, Comput. Phys. Commun. 118 (1999) 153 [hep-ph/9807565].

[16] A. van Hameren, C. G. Papadopoulos and R. Pittau, JHEP 0909 (2009) 106 [arXiv:0903.4665 [hep-ph]].

[17] G. Bevilacqua, M. Czakon, M. V. Garzelli, A. van Hameren, A. Kardos, C. G. Papadopoulos, R. Pittau and M. Worek, Comput. Phys. Commun. 184 (2013) 986 [arXiv:1110.1499 [hep-ph]].

[18] C. F. Berger, Z. Bern, L. J. Dixon, F. Febres Cordero, D. Forde, H. Ita, D. A. Kosower and D. Maitre, Phys. Rev. D 78 (2008) 036003 [arXiv:0803.4180 [hep-ph]].

[19] V. Hirschi, R. Frederix, S. Frixione, M. V. Garzelli, F. Maltoni and R. Pittau, JHEP 1105 (2011) 044 [arXiv:1103.0621 [hep-ph]].

[20] G. Cullen, N. Greiner, G. Heinrich, G. Luisoni, P. Mastrolia, G. Ossola, T. Reiter and F. Tramontano, Eur. Phys. J. C 72 (2012) 1889 [arXiv:1111.2034 [hep-ph]].

[21] F. Cascioli, P. Maierhofer and S. Pozzorini, Phys. Rev. Lett. 108 (2012) 111601 [arXiv:1111.5206 [hep-ph]].
[22] S. Badger, B. Biedermann and P. Uwer, Comput. Phys. Commun. 182 (2011) 1674 [arXiv:1011.2900 [hep-ph]].

[23] S. Badger, B. Biedermann, P. Uwer and V. Yundin, Comput. Phys. Commun. 184 (2013) 1981 [arXiv:1209.0100 [hep-ph]].

[24] P. Mastrolia, E. Mirabella and T. Peraro, JHEP 1206 (2012) 095 [Erratum-ibid. 1211 (2012) 128] [arXiv:1203.0291 [hep-ph]].

[25] A. van Hameren, Comput. Phys. Commun. 182 (2011) 2427 [arXiv:1007.4716 [hep-ph]].

[26] H. van Deurzen, G. Luisoni, P. Mastrolia, E. Mirabella, G. Ossola and T. Peraro, Phys. Rev. Lett. 111 (2013) 171801 [arXiv:1307.8437 [hep-ph]].

[27] H. van Deurzen, G. Luisoni, P. Mastrolia, E. Mirabella, G. Ossola and T. Peraro, arXiv:1312.6678 [hep-ph].

[28] J. A. M. Vermaseren, math-ph/0010025.

[29] J. Kuipers, T. Ueda, J. A. M. Vermaseren and J. Vollinga, Comput. Phys. Commun. 184 (2013) 1453 [arXiv:1203.6543 [cs.SC]].

[30] J. Kuipers, T. Ueda and J. A. M. Vermaseren, arXiv:1310.7007 [cs.SC].

[31] R. K. Ellis and G. Zanderighi, JHEP 0802 (2008) 002 [arXiv:0712.1851 [hep-ph]].

[32] D. Forde, Phys. Rev. D 75 (2007) 125019 [arXiv:0704.1835 [hep-ph]].

[33] F. del Aguila and R. Pittau, JHEP 0407 (2004) 017 [hep-ph/0404120].

[34] P. Mastrolia, G. Ossola, C. G. Papadopoulos and R. Pittau, JHEP 0806 (2008) 030 [arXiv:0803.3964 [hep-ph]].

[35] S. D. Badger, JHEP 0901 (2009) 049 [arXiv:0806.4600 [hep-ph]].

[36] R. Kleiss, W. J. Stirling and S. D. Ellis, Comput. Phys. Commun. 40 (1986) 359.

[37] G. Ossola, C. G. Papadopoulos and R. Pittau, JHEP 0707 (2007) 085 [arXiv:0704.1271 [hep-ph]].

[38] G. J. Gounaris, P. I. Porfyriadis and F. M. Renard, Eur. Phys. J. C 9 (1999) 673 [hep-ph/9902230].

[39] C. Bernicot, arXiv:0804.0749 [hep-ph].

[40] G. Cullen, M. Koch-Janusz and T. Reiter, Comput. Phys. Commun. 182 (2011) 2368 [arXiv:1008.0803 [hep-ph]].

[41] G. Mahlon, Phys. Rev. D 49 (1994) 2197 [hep-ph/9311213].

[42] Z. Nagy and D. E. Soper, Phys. Rev. D 74 (2006) 093006 [hep-ph/0610028].

[43] T. Binoth, G. Heinrich, T. Gehrmann and P. Mastrolia, Phys. Lett. B 649 (2007) 422 [hep-ph/0703311].
[44] W. Gong, Z. Nagy and D. E. Soper, Phys. Rev. D 79 (2009) 033005 [arXiv:0812.3686 [hep-ph]].

[45] C. Bernicot and J. -P. .Guillet, JHEP 0801 (2008) 059 [arXiv:0711.4713 [hep-ph]].

[46] C. Bernicot, arXiv:0804.1315 [hep-ph].

[47] W. Beenakker, S. Dittmaier, M. Kramer, B. Plumber, M. Spira and P. M. Zerwas, Phys. Rev. Lett. 87 (2001) 201805 [hep-ph/0107081].

[48] W. Beenakker, S. Dittmaier, M. Kramer, B. Plumber, M. Spira and P. M. Zerwas, Nucl. Phys. B 653 (2003) 151 [hep-ph/0211352].

[49] S. Dawson, L. H. Orr, L. Reina and D. Wackeroth, Phys. Rev. D 67 (2003) 071503 [hep-ph/0211438].

[50] S. Dawson, C. Jackson, L. H. Orr, L. Reina and D. Wackeroth, Phys. Rev. D 68 (2003) 034022 [hep-ph/0305087].

[51] S. Dittmaier, M. Kramer, 1 and M. Spira, Phys. Rev. D 70 (2004) 074010 [hep-ph/0309204].

[52] H. van Deurzen, N. Greiner, G. Luisoni, P. Mastrolia, E. Mirabella, G. Ossola, T. Peraro and J. F. von Soden-Fraunhofen et al., Phys. Lett. B 721 (2013) 74 [arXiv:1301.0493 [hep-ph]].

[53] G. Cullen, H. van Deurzen, N. Greiner, G. Luisoni, P. Mastrolia, E. Mirabella, G. Ossola and T. Peraro et al., Phys. Rev. Lett. 111 (2013) 131801 [arXiv:1307.4737 [hep-ph]].

[54] H. van Deurzen, Acta Phys. Polon. B 44 (2013) 11, 2223.

[55] T. Binoth, J. -P. .Guillet, G. Heinrich, E. Pilon and T. Reiter, Comput. Phys. Commun. 180 (2009) 2317 [arXiv:0810.0992 [hep-ph]].

[56] G. Cullen, J. P. .Guillet, G. Heinrich, T. Kleinschmidt, E. Pilon, T. Reiter and M. Rodgers, Comput. Phys. Commun. 182 (2011) 2276 [arXiv:1101.5595 [hep-ph]].

[57] J. P. .Guillet, G. Heinrich and J. F. von Soden-Fraunhofen, arXiv:1312.3887 [hep-ph].

[58] R. G. Stuart, Comput. Phys. Commun. 48 (1988) 367.