AutoDipole – Automated generation of dipole subtraction terms – ✩

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We present an automated generation of the subtraction terms for next-to-leading order QCD calculations in the Catani–Seymour dipole formalism. For a given scattering process with \( n \) external particles our Mathematica package generates all dipole terms, allowing for both massless and massive dipoles. The numerical evaluation of the subtraction terms proceeds with MadGraph, which provides Fortran code for the necessary scattering amplitudes. Checks of the numerical stability are discussed.

Program summary

Program title: AutoDipole
Catalogue identifier: AEGO_v1_0
Program summary URL: http://cpc.cs.qub.ac.uk/summaries/AEGO_v1_0.html
Program obtainable from: CPC Program Library, Queen’s University, Belfast, N. Ireland
No. of lines in distributed program, including test data, etc.: 138042
No. of bytes in distributed program, including test data, etc.: 1117665
Distribution format: tar.gz
Programming language: Mathematica and Fortran
Computer: Computers running Mathematica (version 7.0)
Operating system: The package should work on every Linux system supported by Mathematica. Detailed tests have been performed on Scientific Linux as supported by DESY and CERN and on openSUSE and Debian.
RAM: Depending on the complexity of the problem, recommended at least 128 MB RAM
Classification: 11.5
External routines: MadGraph (including HELAS library) available under http://madgraph.hep.uiuc.edu/ or http://madgraph.phys.ucl.ac.be/ or http://madgraph.roma2.infn.it/. A copy of the tar file, MG_ME_SA_V4.4.30, is included in the AutoDipole distribution package.
Nature of problem: Computation of next-to-leading order QCD corrections to scattering cross sections, regularization of real emission contributions.
Solution method: Catani–Seymour subtraction method for massless and massive partons [1,2]; Numerical evaluation of subtracted matrix elements interfaced to MadGraph [3–5] (stand-alone version) using helicity amplitudes and the HELAS library [6,7] (contained in MadGraph).
Restrictions: Limitations of MadGraph are inherited.
Running time: Dependent on the complexity of the problem with typical run times of the order of minutes.

References:

✩ This paper and its associated computer program are available via the Computer Physics Communications homepage on ScienceDirect (http://www.sciencedirect.com/science/journal/00104655).
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1. Introduction

The Large Hadron Collider (LHC) allows us to explore an energy regime far beyond what has been accessible in direct measurements up to now. Operating at the TeV scale it will shed light on the mechanism of electroweak symmetry breaking. The large center-of-mass energy allows the production of new heavy particles — if they exist. It is expected that the results obtained from the LHC will significantly influence our future understanding of nature. The high potential of the LHC in making an important step forward towards a deeper understanding of elementary particle physics comes not for free. Experimentally as well theoretically the LHC is a very challenging experiment. The large multiplicity in the individual event together with pile-up and the complications due to the underlying event make the experimental analysis at the LHC highly non-trivial. The separation of known Standard Model (SM) physics from new physics is very demanding, in particular since in many cases the new physics signals are overwhelmed by the large SM backgrounds. Sophisticated methods have been devised in the past to cope with this situation within the experiments with respect to theoretical predictions.

Colliding protons at LHC interact primarily through strong interactions and Quantum Chromodynamics (QCD) plays an important role in this context. It is well known that perturbative QCD predictions are in general plagued by a large (residual) renormalization and factorization scale dependence. It is not rare that predictions for cross sections change by 100% when the scales are varied in a reasonable range. To reduce the large scale uncertainty next-to-leading order (NLO) calculations are required, which generally consist of two ingredients: one contribution from the virtual corrections and a second contribution from real emission of one additional parton. With respect to the virtual contributions much progress has been achieved recently, although the evaluation of one-loop amplitudes for “large” multiplicities (2 → 3, 2 → 4) is still a highly non-trivial enterprise. Fortunately, as far as the real corrections are concerned the situation is much better and efficient methods exist for the numerical evaluation of the required matrix elements. However, the integral over the phase space leads to soft and collinear singularities, which we denote here generically as infrared (IR) divergencies. In combination with the virtual corrections, all IR cancel between the two contributions for the physical observables of interest [8–10].

General algorithms, typically classified either as slicing or subtraction methods, are available for the extraction of soft and collinear singularities encountered in the real corrections. In both cases one makes use of the universal behavior of QCD amplitudes for soft and collinear configurations. IR singularities are nowadays usually regulated within dimensional regularization, so the same regulator has to be applied to the real corrections in a way that numerical integration over the phase space in 4 dimensions is possible in the end.

In slicing methods the idea is to separate the phase space into resolved and unresolved contributions [11–14]. Unresolved regions at NLO are those where one parton becomes soft or two become collinear. In these regions the matrix elements are approximated using the QCD factorization theorems. After this simplification the unresolved regions can be integrated analytically in $d = 4 - 2\epsilon$ dimensions and the emerging singularities cancel analytically against the corresponding ones in the virtual corrections. Resolved regions on the other hand exclude all IR singularities by definition and can be integrated numerically in 4 dimensions. A one-dimensional illustration of the slicing approach is shown below,

$$\int_0^1 \frac{f(x)}{x^{1-\epsilon}} \, dx \approx f(0) \int_0^\delta \frac{1}{x^{1-\epsilon}} \, dx + \int_\delta^1 \frac{f(x)}{x} \, dx + O(\epsilon)$$

$$= \frac{1}{\epsilon} f(0) + \ln(\delta) f(0) + \int_\delta^1 \frac{f(x)}{x} \, dx + O(\epsilon),$$

where $f(x)$ is an arbitrary function, which is regular at $x = 0$. It is a drawback of the slicing method that partial results exhibit a logarithmic dependence on the cut $\delta$ which separates resolved from unresolved configurations. This logarithm — which cancels when the two parts (resolved and unresolved) are combined — is manifest in the analytic integration of the unresolved terms. However, for the resolved terms the logarithmic dependence arises from the numerical phase space integration. Since the matrix elements in the unresolved phase space regions are only approximate one tends to make the respective region around singular configurations as small as possible. This procedure, though, would result in large numerical cancellations and a loss of accuracy in the sum of the resolved and unresolved parts compared to the accuracy reached in the numerical integration. In practice this requires a compromise between the quality of the approximation and the numerical effort to achieve a certain precision in the sum of both, resolved and unresolved contributions.

Subtraction methods make use of our knowledge about QCD factorization in the soft and collinear limits to construct suitable “counter-terms”. These have to match pointwise all singularities in the real-emission matrix elements and, at the same time, should be simple enough to be integrated analytically in $d = 4 - 2\epsilon$ dimensions over the entire phase space [1,5,2]. A one-dimensional example is shown in Eq. (1.2).

$$\int_0^1 \frac{f(x)}{x^{1-\epsilon}} \, dx = \int_0^1 \frac{f(x) - f(0)}{x} \, dx + f(0) \int_0^\epsilon \frac{1}{x^{1-\epsilon}} \, dx + O(\epsilon)$$

$$= \int_0^1 \frac{f(x) - f(0)}{x} \, dx + \frac{1}{\epsilon} f(0) + O(\epsilon).$$

Here the cancellation of the singularity takes place at the integrand level — i.e. the divergencies cancel pointwise — and the integrals are easier to evaluate numerically. However, care must be taken with respect to the numerical accuracy. Deep in the singular regions the individual contributions become arbitrary large and, due to limited numerical precision of floating point arithmetic, their cancellation might be incomplete leading to potentially wrong results. While our one-dimensional example in Eq. (1.2) admits the construction of a suitable subtraction term in a straightforward manner this no longer true when considering complicated scattering amplitudes. Fortunately, this formidable problem has been solved with the Catani–Seymour dipole formalism [1,2]. Similar algorithms are presented in Refs. [15,16]. Based on the factorization of soft and collinear singularities in an SU(N) gauge theory all subtraction terms are constructed from universal functions and process specific amplitudes. It turns out, though, that for complicated processes the complete subtraction term is a sum over many...
different contributions — tedious to derive by hand. On the other hand, the method is completely algorithmic, thus an automation is feasible. This is a timely problem and its solution is the aim of the present work [17,18]. We note that it has recently also been addressed by other groups [19–22], see also [23].

In passing let us briefly mention that the attractive features of the subtraction approach (universality of counter-terms and numerical stability) have led to extensions of the formalism to next-to-next-to-leading order (NNLO), see e.g. [24–27]. Presently, the proposed schemes at NNLO apply to processes without colored partons in the initial state and an arbitrary number of massless particles (colored or colorless) in the final state. The necessary counter-terms are either derived from so-called antenna functions [28] or alternatively defined as universal counter-terms based on QCD factorization in the various soft and collinear limits of singly- and double-unresolved parton configurations [29–32]. Ongoing work at NNLO is concerned with extensions to colored partons in the initial state as needed for the LHC.

The outline of the paper is as follows. In Section 2 we review the general features of the Catani–Seymour algorithm [1,2] and in Section 3 we describe the details of its implementation in the AutoDipole package. Particular emphasis is put on the details of the numerical evaluation of subtraction terms via an interface to MadGraph [3–5], which uses the HELAS library [6,7] for the computation of helicity amplitudes. Section 4 illustrates with a few examples of how to use AutoDipole in practice. The examples serve also as a non-trivial cross check of the implementation when compared with existing results from the literature. Finally, we conclude in Section 5 and list some technical details of the implementation and the comparison with the literature in Appendices A–C.

2. Review of the Catani–Seymour subtraction formalism

We consider a generic scattering process for the production of an n-parton final state. At NLO accuracy in QCD the corresponding cross section may be written as:

\[
\sigma_{\text{NLO}} = \sigma_{\text{LO}} + \delta \sigma_{\text{NLO}},
\]

where \(\sigma_{\text{LO}}\) denotes the Born contribution at leading order (LO) and the genuine NLO correction \(\delta \sigma_{\text{NLO}}\) receives contributions from three different sources,

\[
\delta \sigma_{\text{NLO}} = \int \frac{d\sigma_{\text{virt}}}{n} + \int \frac{d\sigma_{\text{real}}}{n+1} + \int dx \int \frac{d\sigma_{\text{fact}}}{n}. \tag{2.2}
\]

By \(d\sigma_{\text{virt}}\) we denote here the contributions from the virtual corrections and by \(d\sigma_{\text{real}}\) the ones from the real emission of one additional parton. For hadrons in the initial state there is a third contribution \(d\sigma_{\text{fact}}\) due to the factorization of initial state singularities. The subscript on the integral signs in Eq. (2.2) indicates the dimensionality of the phase space. The virtual corrections are integrated over an n-parton phase space while the real emissions are integrated over an (n + 1)-parton phase space. All three contributions are individually divergent and thus require regularization in intermediate steps until the divergencies are canceled. In the Catani–Seymour formalism the expression for \(\delta \sigma_{\text{NLO}}\) in Eq. (2.2) is rewritten schematically:

\[
\delta \sigma_{\text{NLO}} = \int \frac{(d\sigma_{\text{real}} + dA)}{n+1} + \int \frac{(d\sigma_{\text{virt}} + dA')}{1} + \int dx \int \frac{(d\sigma_{\text{fact}} + dA'')}{n}. \tag{2.3}
\]

with

\[
0 = \int \frac{dA}{n+1} + \int \frac{dA'}{1} + \int dx \int \frac{dA''}{n}. \tag{2.4}
\]

The expressions \(dA, dA'\) and \(dA''\) are defined such that they render the individual pieces in Eq. (2.3) finite. In the Catani–Seymour formalism, they correspond to the sum of all dipoles \((dA)\), the integrated dipoles \((\int dA')\), I-term, and the terms arising from mass factorization \((dA'')\) including the so-called P-, K- and H-terms.

The explicit form of \(dA\) is constructed from the knowledge about the soft and collinear factorization of QCD amplitudes, which exhibit a simple factorization for collinear configurations. The expression for \(dA\) in Eq. (2.3) is obtained as a sum over potentially collinear partons since the factorization for soft configurations can be derived from the collinear behavior of the amplitudes. For soft singularities however only color-ordered amplitudes exhibit a simple factorization, which implies in general non-trivial color correlations at the level of the squared matrix elements. These color correlations are reflected by the reference to an additional (so-called) spectator parton. The subtraction \(dA\) is thus written as a sum of individual dipoles in the following form:

\[
dA = \sum \mathcal{D}(i, j; k), \tag{2.5}
\]

where the sum runs over all colored partons in scattering process and the possible configurations for \((i; j; k)\) are determined from the real corrections for which the subtraction term is constructed. The generic form of the dipoles is given by

\[
\mathcal{D}(i, j; k) = d_{ij} \times \{ 1, \ldots, \bar{i}, \ldots, \bar{k}, \ldots, n | \mathbf{V}_{ij, k} | 1, \ldots, \bar{i}, \ldots, \bar{k}, \ldots, n \}, \tag{2.6}
\]

As a concrete example we display the expression for \(\mathcal{D}(g_i, g_j, k)\) with all three partons in the final state, i.e.

\[
\mathcal{D}(g_i, g_j, k) = \frac{1}{2 p_i \cdot p_j} \{ 1, \ldots, \bar{i}, \ldots, \bar{j}, \ldots, n | T_k \cdot T_{ij} V_{g_i g_j, k} | 1, \ldots, \bar{i}, \ldots, \bar{j}, \ldots, n \}. \tag{2.7}
\]

\(\mathcal{D}(g_i, g_j, k)\) describes the situation of a collinear splitting of the gluons \(i\) and \(j\) (emitter pair) in the presence of a spectator parton \(k\). Here, the \(T_i\) are the color charge operators depending on the partons being in the fundamental or adjoint representation of the color SU(3) (for details we refer to [1,2]). The function \(V_{g_i g_j, k}\) is given by

\[
V_{g_i g_j, k}^{\mu \nu} = \langle \mu | V_{g_i g_j, k} | \nu \rangle
\]

\[
= 16 \pi a_s \mu R^2 C_A \left[ -g_{\mu \nu} \frac{1}{1 - z_i (1 - y_{ij, k})} + \frac{1}{1 - z_j (1 - y_{ij, k})} - 2 \right] + (1 - e) \frac{1}{p_i \cdot p_j} (z_i p_i^{\mu} - z_j p_j^{\mu})(z_i p_i^{\nu} - z_j p_j^{\nu}). \tag{2.8}
\]

where \(z_i\) and \(y_{ij, k}\) are some functions of the Lorentz scalars \(s_{ij}, s_{ik}\), and \(s_{jk}\) and \(a_s\) is the strong coupling. The quantity

\[
\{ 1, \ldots, \bar{i}, \ldots, \bar{k}, \ldots, n | T_k \cdot T_{ij} | 1, \ldots, \bar{i}, \ldots, \bar{k}, \ldots, n \}. \tag{2.9}
\]
is called the color linked Born amplitude squared (CLBS). The amplitude \((\ldots)\) is deduced from the real emission by the factorization of one splitting. The color factor is extended by two color operator insertion, \(T_i \cdot T_j\). The CLBS has the reduced kinematics as input. In the example the original momenta \((p_i, p_j, p_k)\) are reduced to \((\tilde{p}_i, \tilde{p}_j)\) where for example, the \(\tilde{p}_j\) is defined as

\[
\tilde{p}^\mu_j = p^\mu_j + p^\mu_k - \frac{y_{ij,k}}{1 - y_{ij,k}} p_k^\mu.
\]  

(2.10)

Generally, \((n+3)\) original momenta are reduced to the \((n+2)\) momenta in each dipole term. One distinguished feature of the dipole subtraction is that the reduced kinematics satisfy the on-shell conditions and momentum conservation, which makes it possible to evaluate the reduced Born amplitude by existing codes for LO calculations. In the gluon emitter case like the example above, the second term in the square bracket in Eq. (2.8) introduces spin correlations of the gluon. In that case we need also to evaluate the CLBS where the helicity of the gluon in the amplitudes is different from the one in the corresponding Born amplitudes. In the next section we explain how the three main ingredients, the generation of the dipole terms, the CLBS, and the dipoles with a gluon emitter, are implemented in our package.

The other two functions \(f_i dA'\) and \(dA''\) in Eq. (2.3) are easily explained. The first one \((\tilde{f}_i dA')\) denotes the integrated dipoles which cancel the corresponding IR divergences in the virtual contributions at NLO. They can be constructed from universal insertion operators \(I\), to be sandwiched between the bra \((\ldots)\) and ket \((\ldots)\) amplitudes of the corresponding Born process. This includes all algorithms for the generation of the dipole terms in the final state. These remarks conclude our brief review of the Catani–Seymour formalism. For details the reader is referred to the original literature [1,2].

3. The AutoDipole package

The AutoDipole package constructs all necessary subtraction terms of the Catani–Seymour formalism [1,2] for a given scattering process with \(n\) colored partons in a fully automatic manner. It can handle both massless and massive partons and it also allows for additional (non-colored) SM particles in the scattering process, e.g. couplings of quarks to \(γ, Z, W^-\)-bosons and so on.

Let us briefly sketch the details of our implementation of the dipole subtraction formalism, because there is a large freedom in this particular implementation. For instance, in Refs. [35,36] the implementation was realized in form of two independent C/C++ libraries providing all the necessary functions to evaluate the dipole terms. As a slight disadvantage of this approach the produced code is non-local and there is some redundancy in the calculation. In the present work we follow a different strategy. The main idea here is to have a code generator which will produce an optimized flat code. To realize this we have constructed a Mathematica program which acts as such a generator and its output is interfaced with MadGraph [3–5]. To that end we found the stand-alone version of MadGraph suitable and for the numerical evaluation of all subtraction terms, we use helicity amplitudes provided by MadGraph (which are based on the HELAS library [6,7]).

3.1. Structure of the code

The complete layout of the code of the AutoDipole package is displayed in Fig. 1. The major part of the source code in the package (all contained in the directory lib) is written in Mathematica. This includes all algorithms for the generation of the dipole terms as well as the evaluation of the \(I\)-, \(P\)- and \(K\)-operator insertions sandwiched between the Born amplitudes in the appropriate kinematics. The use of a computer algebra program like Mathematica has the clear advantage here that all terms are accessible to symbolic manipulation. This feature is very useful for general studies of the IR behavior of scattering amplitudes.

In using AutoDipole we are primarily interested in the generation of Fortran code for all subtraction terms of Eq. (2.3). The essential parts of the flowchart are displayed in Fig. 2. From a given input process, the Mathematica code generates all dipole terms. The important files here are dipole.f, reducedm.f, and the shell script for the interface with MadGraph, where dipole.f contains all dipole terms except for the reduced kinematics and the CLBS. The routine reducedm.f calculates the reduced kinematics of each dipole term as defined in Eq. (2.10). The subsequent run of a patched version of MadGraph (contained in the directory patch) via the interface produces the files, cmatrix.f and allcolormat.inc. The routine cmatrix.f evaluates all CLBS as in Eq. (2.9). It includes the file allcolormat.inc, which contains all extended color matrices by the two-color operator insertions. We use the latest version of MadGraph.
In short, the evaluation of the generated Fortran code for the dipole terms proceeds along the following chain, see Fig. 2. The routine reducedm.f receives a given phase space point as input and calculates the reduced kinematics of each dipole term. Next, the routine cmatrix.f (with the reduced kinematics as input) computes the CLBS of each dipole term together with the extended color matrices in allcolormat.inc. Finally the routine dipole.f returns the values of all dipole terms having received the reduced kinematics and the CLBS as input. All this is done automatically when the user executes the command ./runD (see Sections 4.2 and 4.3) to calculate the subtraction term for a specific phase space point.

Then, the first step in the construction of the dipoles is the choice of the emitter pair, that is the root of the splitting of the quarks and gluons. This also specifies the kind of splitting. At NLO in QCD, there exist seven possible splittings, which we group into four classes and we enumerate these types of dipoles accordingly, Dipole 1, …, and Dipole 4. The ordering is shown in Fig. 3. Due to the factorization of the splitting each dipole is associated to a certain reduced Born process and the following equations show schematically the operations performed on the sets of partons in Eq. (3.1) in order to construct the corresponding reduced Born process,

\[
\text{Dipole 1: } \{\text{initial}\} \rightarrow \{\text{final}\} - g, \\
\text{Dipole 2: } \{\text{initial}\} \rightarrow \{\text{final}\} - f\bar{f} + g, \\
\text{Dipole 3: } \{\text{initial}\} - f + g \rightarrow \{\text{final}\} - f, \\
\text{Dipole 4: } \{\text{initial}\} - g + \bar{f} \rightarrow \{\text{final}\} - f,
\]

where \(g\) is a gluon and \(f(\bar{f})\) a quark (anti-quark). The notation in Eqs. (3.2)-(3.5) indicates which partons are removed from or added to the respective sets \{initial\} and \{final\}. The cases Dipole 3 and 4 can also occur with a reduced Born cross section where \(f\) and \(\bar{f}\) are exchanged. The groups Dipole 2, 3 and 4 also need to account for the existence of various quark flavors, i.e. they exhibit manifest flavor dependence, see Table 2 for details.

3.2. Algorithm of dipole generation

Let us start with a short description of the Mathematica code generating all dipole terms. This procedure makes use of the following three steps:

1. Choose all possible emitter pairs from the external legs.
   For a given real emission \(2 \rightarrow (n+1)\)-particle scattering process we abbreviate the set of initial (final) state partons collectively as \{initial\} \{final\}, i.e. the scattering reaction reads generically

\[
\{\text{initial}\} \rightarrow \{\text{final}\}.
\]

2. Choose all possible spectators for each emitter pair.
   The spectator is one external field which is different from both fields of the emitter pair and its choice is a purely combinatorial
procedure. We use indices, $i$, $j$, and $k$, for fields in a final state and, respectively, indices, $a$ and $b$, for an initial state field. For a spectator in the final (initial) state denoted by $k$ ($b$), this condition means $k \neq i, j$ ($b \neq a$). It emerges from a special feature of the subtraction formalism namely that the squared matrix elements with the Casimir operator, $(T^2_i)$, is expressed, due to color conservation, through CLBS $(T_i, T_k)$ ($i \neq k$) which always includes one color dipole configuration in the part.

3. Construct the dipole terms from the chosen combinations of emitter and spectator.

The previous steps provide all such combinations as pairings of the type (emitter, spectator) $=$ ($ij$, $k$), ($jj$, $b$), ($ai$, $k$), and ($ai$, $b$). Each case corresponds to one dipole term, for which we use the shorthand $D_{ij,k}$, $D_{ij}$, $D_{ai,k}$, and $D_{ai,b}$, respectively. Detailed information about all pairings of emitter and spectator as well as the potential presence of a mass parameter is given in Table 2 in Appendix A. For explicit expressions we refer to [1,2].

Here it is worth to stress a few points. First of all, the chosen order for the generation has the advantage of grouping together the reduced Born matrix elements. The CLBS in the same category exhibit the same Lorentz structure, only the extended color matrix is different. This ordering leads to a block-diagonal structure in the matrix with the color operator, $\mathcal{C}$, in the denominator always cancels against the same one in the dipole splitting function. This cancellation leads to manifest simplifications. Finally, in our set-up we have access to the symbolic expression and can easily identify or extract partial subset of dipole terms, if needed. For example we may want to discard the $t - \bar{t}$ (heavy-quark) splitting in the category Dipole 2, because this splitting does not give rise to poles in $\bar{t}$. Our way of generating the dipoles in the package easily allows this selection. The user can discard dipoles of a specific category in the input file parameter.m (see Section 4).

At this stage it remains to discuss the generation of all terms involving the $I$-, $P$-, and $K$-operators. This is sketched in Fig. 4. Given, that these operators originate from the phase space integral over unresolved parton of the dipole terms, it is clear that we only need a subset of all previously generated terms. From all reduced Born amplitudes generated for the regularization of the real emission contributions (i.e. Fig. 3) we need only the CLBS in the category Dipole 1 for all $I$-, $P$-, and $K$-operator insertions. Hence, no new information is required and the Mathematica package assembles the respective expressions (along with the color correlations, see Section 3.3) in the Fortran files $I$term.f and $PK$term.f.

3.3. The color linked Born amplitude squared

The numerical evaluation of Born amplitudes (or their squares) is a routine task for many publicly available packages designed for automated LO calculations. As announced above, we choose the stand-alone version of MadGraph [3–5] for this purpose and the added feature needed is the evaluation of the color link operators, see e.g. Eq. (2.7). This can be done by a patch and allows us to obtain all CLBS in an automatic way, see Fig. 2.

The generation of the amplitudes in MadGraph proceeds via Feynman diagrams and the color factors are separated from each diagram. During the evaluation everything is expressed in terms of generators of the fundamental representation of the color SU(3). A typical example is that the factor $f^{abc}$ of the gluon three-point vertex is rewritten in terms of the fundamental generator $t^a$ with the help of the identity,

$$f^{abc} = -2i(\text{Tr}[t^a t^b t^c] - \text{Tr}[t^c t^a t^b]).$$

(3.6)

The color factors of each diagram are sorted in a unique order and they are expressed in a sum. When a specific term of a diagram is identical to one of the other diagrams, both are combined as

$$M = \sum_a C_a \bar{J}_a.$$  

(3.7)

where $C_a$ denotes the independent color factors. Each $C_a$ has fundamental and adjoint color indices corresponding to the external quarks and the gluons, respectively, $\bar{J}_a$ is the joint amplitude, e.g. $\bar{J}_1 = +A_1 - A_2 + \cdots$ where $A_i$ is the partial amplitude of $i$-th diagram (with the color factor stripped off). The invariant matrix element squared is finally expressed in the form,

$$|M|^2 = \langle \bar{J} | \mathcal{C} | \bar{J} \rangle.$$  

(3.8)

where the color matrix $\mathcal{C}$ is defined as

$$(\mathcal{C})_{ab} = \sum_{\text{color}} C_a C_b.$$  

(3.9)

For the CLBS we need to evaluate Eq. (3.8) with an insertion of two additional color operators to the emitter and spectator legs. This is precisely what our patch of MadGraph does (see Fig. 1 and the directory lib/patch). The subroutines of MadGraph for the color factor calculations are well structured and the original routines to add the color factors $t^a_i$ and $f^{abc}$ can be applied to the additional color insertions for the CLBS. The color algebra of the SU(3) is performed numerically and the resulting extended color matrix $\mathcal{C}$ is written to the file allcolormat.inc, see Fig. 2.
We have realized the two color operator insertions for all CLBS in an automatic way and we have also checked that MadGraph with our interface works for a set of rather involved processes. As a simple example let us discuss the color insertions required for the process $g(a)g(b) \to u(1)\bar{u}(2)g(3)$. The reduced Born process $g(a)g(b) \to u(1)\bar{u}(2)$ has three diagrams and the color factors are combined into two independent ones, $(C_1, C_2) = (t^a t^b)^{1/2}, (t^b t^a)^{1/2})$. The components of the color matrix are given by the traces, $(C_{F11} = (CF)_{12} = Tr(t^a t^b t^c t^d)$ and $(CF)_{12} = (CF)_{21} = Tr(t^b t^a t^d t^c)$. Then the color matrix is calculated as

$$CF = \begin{pmatrix} 16/3 & -2/3 \\ -2/3 & 16/3 \end{pmatrix}.$$  \hfill (3.10)

In the CLBS we need for instance the fundamental operator insertions into the legs 1 and 2. The components of the color matrix $CF$ are modified to $(CF)_{11} = Tr(t^b t^d t^a t^c)$ and $(CF)_{12} = Tr(t^d t^a t^b t^c)$. Then the modified color matrix is obtained as

$$CF' = \begin{pmatrix} 1/9 & 10/9 \\ 10/9 & 1/9 \end{pmatrix}. \hfill (3.11)$$

One of the more complicated examples consists of two color operator insertions into the process $g(a)g(b) \to t(1)t(2)g(3)g(4)$. In MadGraph the normal SU(3) color matrix for the process is a $24 \times 24$ matrix. The first 15 components in the first row read (we refrain from spelling out the rest),

$$CF = \frac{1}{54} \begin{pmatrix} 512, 8, -64, 80, 8, -10, -1, -64, -64, 8, \\ -1, -10, -1, 62, -10, \ldots. \end{pmatrix}. \hfill (3.12)$$

Upon insertions of two adjoint color operators for a gluon into the legs 3 and 4 the extended routines calculate the modified color matrix as

$$CF' = \frac{1}{4} \begin{pmatrix} 8, 0, 8, 16, 0, -2, 0, 8, -1, -1, 1, 2, -8, -7, 1, \ldots. \end{pmatrix}, \hfill (3.13)$$

and, of course, the result in Eq. (3.13) agrees with independent checks.

3.4. Spin correlations for gluon emitters

As we have seen above the dipoles for the splittings $g \to gg$ and $g \to q\bar{q}$ (which involve a gluon emitter) introduce spin correlations (see e.g. Eq. (2.7)). Since our numerical evaluation of the reduced Born amplitudes uses helicity amplitudes, we have to derive the components of the tensor for the spin correlations in the helicity formalism as well. To that end, we choose the definitions in [37] which we here call the XZC gluon polarization vector.

Let us illustrate the necessary steps (formulated in [38]) with the case of the massless dipole term $D(g_i, g_j, k)$ in Eq. (2.7),

$$D(g_i, g_j, k) = -\frac{1}{2p_i \cdot p_j} V^{\mu \nu} \frac{1}{g_{i\delta j,k}} (A_+^\mu T_k \cdot T_j A_+), \hfill (3.14)$$

where $V^{\mu \nu}$ is written schematically as

$$V^{\mu \nu} = 16\pi \alpha_s \frac{1}{2} \epsilon^\mu \epsilon^\nu \epsilon^\rho \epsilon^\sigma,$$

with $L^\mu = z_i p_i^\mu - z_j p_j^\mu$ and some Lorentz scalars $C_{1,2}$ (e.g. given in Eq. (2.8)). Moreover, in Eq. (3.14) we have expressed the CLBS in Eq. (2.9) through amplitudes $A_\mu$ and $A_{\nu}$, where the polarization vector $\epsilon^\mu_{\rho}(p_i)$ of the emitter gluon with momentum $p_i$ has been amputated. Then we can transform both the dipole splitting function $V^{\mu \nu}$ and the amplitude $A_\mu$ to a helicity basis by inserting the polarization sum $\sum_a \epsilon_{\rho a}^\mu \epsilon_{\sigma a}^\nu$ as

$$A^\mu_{\nu} V^{\mu \nu} A_\nu = \sum_{\lambda, \lambda'} A^{\mu}_{\nu} V^{\lambda \lambda'} A_{\lambda'}, \hfill (3.16)$$

$$V^{\lambda \lambda'} = \epsilon^{\lambda \lambda'}_{\rho a} V^{\mu \nu} \epsilon^\rho_{\sigma a}, \hfill (3.17)$$

and we obtain the dipole $D(g_i, g_j, k)$ in a $\pm$-helicity basis as

$$D(g_i, g_j, k) = \frac{1}{2p_i \cdot p_j} \epsilon^{\mu} \delta^{\nu}_{\rho a} \epsilon^\rho_{\sigma a} \epsilon_{\rho a}^\mu (C_1 + C_2 |E_+|^2) (A_+^\mu T_k \cdot T_j A_+ + A_+^\mu T_k \cdot T_j A_- + 2C_2 Re(\bar{E}_+^\mu E_-^\nu A_+^\mu T_k \cdot T_j A_-)). \hfill (3.18)$$

Here we have used gauge invariance, the on-shell condition for the gluon and orthogonality of the polarization vectors, which gives rise to the relations $\epsilon^{\mu \nu} (p_{ij}) = A \cdot \bar{p}_{ij} = L \cdot \bar{p}_{ij} = \bar{p}_{ij}^2 = 0$. The quantity $E_+$ needed to express the dipoles in a helicity basis is defined as (see e.g. [38])

$$E_+ = \epsilon_{\rho a} \cdot L. \hfill (3.19)$$

As a subtlety, we would like to point out the following. In the HELAS library [7,6] the gluon polarization vector is calculated by the subroutine $VXXXXX$ and taken to be in a helicity basis but the phase conventions are different from the ones in [37]. Thus we have to relate these conventions to our choice [37]. Although it may be natural to use the same definition for the gluon polarization vectors as in HELAS for the dipole splitting functions (e.g. $V^{\mu \nu}_{g\bar{g},k}$ in Eq. (3.15)) we have the freedom not to do so because of gauge invariance and the on-shell condition. In our implementation we have chosen the latter option with the advantage that the calculation of the dipole splitting functions can be completely separated from the part for evaluation of the CLBS. With the XZC definitions $E_+$ in Eq. (3.19) is expressible in terms of spinor products which can be easily implemented in Fortran code.

The polarization vectors of HELAS [7,6] are written in a helicity basis as

$$\epsilon^{\mu \nu}_{H E L A S}(k, q) = \frac{1}{\sqrt{2}} (\bar{\epsilon}_{\mu(1)} \epsilon_{\nu(2)} - i \bar{\epsilon}_{\mu(2)} \epsilon_{\nu(1)}),$$

with the explicit expression for the vectors in a linear basis, $\epsilon_{\mu(1,2)}$ given in [7,6]. Likewise, for XZC [37] we have

$$\epsilon^{\mu \nu}_{X Z C}(k, q) = \frac{\langle q - |y_{\mu}| k^- \rangle}{\sqrt{2} (q^0 k^0)^*}, \hfill (3.21)$$

where the relation $\epsilon^{\mu \nu}_{X Z C} = \epsilon^{\mu \nu}_{H E L A S}$ holds and $q$ is an arbitrary reference momentum. The difference between the conventions Eq. (3.20) and Eq. (3.21) amounts to a complex phase,

$$\epsilon^{\mu \nu}_{X Z C}(k, q) \cdot \epsilon^{\mu \nu}_{H E L A S}(k, q) \propto e^{i\phi(q)}.$$  \hfill (3.22)

The phase difference contributes to only the second term in the square bracket in Eq. (3.18) and it is then rewritten as

$$+ 2C_2 \Re(\bar{E}_+^\mu E_- A_+^\mu T_k \cdot T_j A_-)_{X Z C} \hfill (3.23)$$

where the CLBS is also taken in the helicity basis for the gluon emitter as in HELAS. The quantity $E_+ = \epsilon^{\mu \nu}_{X Z C} \cdot L$ in Eq. (3.23) can be computed according to Eq. (3.19) in the conventions of [37] in terms of spinor products as (see e.g. [38])

$$e^{i\phi(q)} E_+ = e^{i\phi(p)} \frac{Z(p_{ij}^\mu p_j^\nu)}{\sqrt{2(p_i p_j)}}.$$  \hfill (3.24)
in Eq. (3.19) for all required momentum configurations (massless and massive) are given in Appendix B. For the dipole terms including the massive partons, the result for $E_+$ in Eq. (3.19) contains terms like $(p_i - |p_k|\tilde{p}_{ai})$ with on-shell momenta $p_k^2 = m_k^2$. In order to express these terms through spinor products we have used the flat momenta,

$$p_i^\flat = p_i - \frac{m_k^2}{2p_k \cdot \tilde{p}_{ai}} \tilde{p}_{ai},$$

(3.25)

which are massless because $(p_i^\flat)^2 = 0$. Due to the equations of motion $\tilde{p}_{ai} |\tilde{p}_{ai}\rangle = 0$, we can rewrite the quantities under consideration, e.g.,

$$\langle p_i - |p_k|\tilde{p}_{ai}\rangle = \langle p_i - |p_k^\flat|\tilde{p}_{ai}\rangle,$$

(3.26)

so that they are accessible to standard spinor calculus, see also [39] for the spinor helicity formalism including massive fermions. Our treatment of the spin correlations is contained in the Mathematica sources where the expressions for $E_+$ (as given in Appendix B) are implemented. During the automated generation of all subtraction terms for numerical evaluation the results are written to the output in the Fortran file, dipole.f, see Fig. 2.

3.5. Checking the soft and collinear limits

Let us discuss the checks for the AutoDipole package. First, there is the standard quality check on the automatic generation of the dipole terms because the result according to Eq. (2.3) has to be finite when approaching the singular regions. Second, since the Catani–Seymour formalism is well established, we can compare the output of AutoDipole with results of independent implementations in the literature.

The AutoDipole package performs automatically for all generated subtraction terms a numerical check of all IR limits. For that purpose the subtracted squared matrix element for the process under consideration are constructed and sampled over all limits to test the cancellation of the IR singularity. For a given soft/collinear limit $L_i$, we pick up the set $S(i)$ of the corresponding dipoles to test whether the quantity

$$\lim_{L_i} \left[ |M|^2 - \sum_{j \in S(i)} D(j) \right],$$

(3.27)

is soft/collinear safe. The code lists all limits $L_i$, and the corresponding sets $S(i)$. A concrete example will be presented in Section 4.2. The leading soft/collinear singularity of the squared matrix element $|M|^2$ behaves in the soft limit as $1/k^2$ for a gluon of momentum $k$ and in the collinear one as $1/(2p_i \cdot p_j) = 1/s_{ij}$ for two collinear momenta $p_i$ and $p_j$. The set of the corresponding dipoles cancels this leading singularity pointwise as

$$|M|^2 - \sum_{j \in S(i)} D(j) = \frac{1}{k^2} (a_0 + a_1 x + a_2 x^2 + \cdots) - \frac{1}{k^2} a_0,$$

(3.28)

where $x = k(\sqrt{s_{ij}})$ is the respective Lorentz scalar depending on the momenta in the soft (collinear) limit. The difference in Eq. (3.28) is integrable over the real emission phase space. The numerical accuracy, however, has to be controlled, because deep in the singular regions the individual contributions in Eq. (3.28) become very large and with the limited numerical precision of floating point arithmetic the cancellation might be imperfect. Fortunately, the stability of the numerical cancellation can be tested automatically. Normalizing Eq. (3.28) one can check the slope of the quantity

$$\frac{|M|^2 - \sum D}{|M|^2} = \frac{1}{a_0} (a_1 x + a_2 x^2 + \cdots).$$

(3.29)

Based on Eq. (3.29) AutoDipole tests the cancellation and returns values for the fiducial regions for the Lorentz invariants $s_{ij}$.

We have extensively tested the soft and collinear finiteness of the generated subtraction terms for numerous scattering processes in $e^+ e^-$, $ep$ and $pp$-collisions including massive quarks and weak gauge bosons. Among those are a large class of $2 \to 4$ scattering processes as needed in the NLO QCD correction to $2 \to 3$ reactions. We have also tested scattering processes with $2 \to 5$ and $2 \to 6$ partons, like e.g.,

$$u\bar{u} \to d\bar{d}ggg,$$

$$gg \to W^+W^-,$$

which are currently under investigation in view of phenomenological applications for the LHC.

In addition, we have been able to obtain also perfect agreement with published results [36,40]. For the NLO QCD corrections to $pp \to t\bar{t} + 1$ jet production results for the real emission contributions have been presented at individual phase space points in [36]. We agree at least to 14 digits for the matrix elements squared and at least to 12 digits for the sum of the subtraction terms. All details of our comparison are shown in Table 5 in Appendix C. We have also checked against the results for the NLO QCD corrections to $pp \to W^+W^- + 1$ jet [40], in particular the real emissions in the channel $u\bar{u} \to W^+W^- gg$. Also here we have obtained very good agreement as documented in the last entry of Table 5. The AutoDipole package provides a shell script to reproduce some of the numbers in that Table 5 (see Section 4.4).

Let us briefly comment on the integrated dipoles in Eq. (2.3) originating from the $I$-operators. Here, a complete check is more involved because for a given process scheme dependence enters, i.e. whether the singularities of the NLO virtual contributions are factorized with respect to the $4$-dimensional or $3$-dimensional Born amplitude.

Nevertheless, partial checks of the singularity structure are straightforward. The matrix element squared with the sum of all $I$-insertions is written schematically as

$$\langle 1, \ldots, n|1, \ldots, n\rangle = C_{-2} \frac{1}{\epsilon^2} + C_{-1} \frac{1}{\epsilon} + C_0,$$

(3.30)

where $C_{-1}$ and $C_{-2}$ are process (and kinematics) dependent coefficients. For the leading pole $\sim 1/\epsilon^2$, the coefficient $C_{-2}$ obeys the following simple relation

$$C_{-2} = \frac{\alpha_s}{2\pi} |M_{\text{Born}}|^2 \left( n_q + n_{\bar{q}} \right) C_F + n_q C_A,$$

(3.31)

where $M_{\text{Born}}$ is the corresponding Born amplitude. The number of external massless (anti-)quarks is $n_q$ ($n_{\bar{q}}$) and $n_q$ the number of external gluons. In QCD the standard SU(N) color factors take the numerical values $C_F = 4/3$ and $C_A = 3$.

Apart from simple relations like Eq. (3.31) the coefficients for color and spin averaged results for the $I$-operator could again be compared with the published literature for $pp \to t\bar{t} + 1$ jet production at NLO in QCD [36]. We have obtained agreement to at least 14 digits as shown in Table 6. The numbers in Table 6 can again be reproduced with a simple command explained in Section 4.4.

4. Running AutoDipole

This section is meant to be a short manual to the AutoDipole package. We explain the installation procedure, the use and discuss a few processes as examples.
4.1. Install

The package, AutoDipole_V1.2.3.tar, is available for download from [41] (or else from the authors upon the request). In addition, one must also obtain the stand-alone version of MadGraph (MG_ME_SA_V4.4.30) e.g. by download from [42] and put it (in the form of .tar.gz or .tar) in the AutoDipole package directory. Then execute the installation procedure by

```
./install.sh
```

and the directory structure as displayed in Fig. 1 emerges. Repetition of this command always allows the user to recover the initial settings described in this section.

4.2. A short example

Let us next illustrate the use of AutoDipole. First of all, one has to specify all partonic real emission processes which appear at NLO in the observable under consideration. To that end, as a concrete example let us choose a simple process, $\bar{u}u \to d\bar{d}g$ which contributes to hadronic di-jet production, $pp \to 2 \text{jets}$ at NLO. This example exhibits all features of the Catani–Seymour formalism, i.e. it needs dipoles and I-operators as well as the P- and K-terms. We start with the Mathematica part of AutoDipole. The package can simply be included through the driver file as

```
<<driver_user.m
```

Next, we can run the package for the process $\bar{u}u \to d\bar{d}g$ with the command

```
GenerateAll[{u,ubar},{d,dbar,g}]
```

Upon running the AutoDipole package successfully, all generated Fortran files are stored in a (newly created) directory under `process`, for the example at hand `/process/Proc_uux_ddxg`. AutoDipole returns the message:

```
******************************************************************************************
Run has been succeeded.
In order to run the generated code, please enter in the newly created directory under
./process
make runD
******************************************************************************************
```

The real matrix element squared $|M|^2$ and the sum of all dipole terms $\sum_i D(i)$ are evaluated at 10 phase space points with this command. Also the check of cancellations in all infrared limits for the subtracted matrix element can performed. The respective command is:

```
make checkIR
./checkIR
```

Likewise, the integrated dipoles (I-operators) as well as the contribution of the P- and K-terms at 10 points in phase space can be evaluated in the subdirectories `Proc_uux_ddxg/Virtual` and `Proc_uux_ddxg/PK` with the commands:

```
make runI
./runI
make runPK
./runPK.
```

The present example $\bar{u}u \to d\bar{d}g$ runs with the default settings of AutoDipole. For further illustration the package also comes along with a prepared list of examples so that explicit numbers can be obtained by executing the shell scripts explained in the next subsection.

If the user wants to include the generated Fortran code with the subtraction terms into his/her own project for further numerical evaluations, he/she needs (i.e. has to copy) a number of files for that purpose. The complete list of generated files is given in Table 1. Note that most of the functions are only needed internally and are not meant to be called by the user directly. We assume here that all input momenta $(p_i)$ are generated from phase space routines supplied by the user and that the HELAS library libdhelas3.a is linked.

**Dipole terms:** directory `process/Proc_uux_ddxg`

From the file `check.f` the call of the relevant subroutines for the evaluation of the dipole terms is obvious. The user needs to copy the following files:

- `matrix.f` (generated by AutoDipole)
- `dipole.f`
- `reducedm.f`
- `cmatrix.f`
- `allcolormat#.inc`
- `emitspecinfo.inc`
- `nexternal.inc`
- `nexternal2.inc` (generated by AutoDipole)
- `coup1.inc` (process independent file)
- `couplings.f` (process independent file)
- `param_card.dat` (parameter input file)

To use the generated code, one first has to initialize MadGraph. The parameters are determined in `param_card.dat` and are assigned to the definitions in `coup1.inc` with the help of the subroutine `setpara` in `couplings.f` as

```
include ‘coup1.inc’
call setpara(‘param_card.dat’, .true.)
```

to be used subsequently in the subroutines `cmatrix#`, `cmatrix#dh`, and `smatrix` through the common blocks. The symbol # here and above denotes an integer number (e.g. # = 1). For the dipoles only the strong coupling constant $\alpha_s$ as well as the top and bottom masses are needed in the evaluation. These three parameters appear as

```
AL=g**2*(4.0*pi)**(-1)
mt=tmass
mb=bmass
```

and are passed to the subroutine `dipole` through the common blocks,
Table 1

<table>
<thead>
<tr>
<th>File (subroutine)</th>
<th>Input</th>
<th>Output</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Dipole terms]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>matrix.f (smatrix)</td>
<td>{p_1}</td>
<td>[</td>
</tr>
<tr>
<td>dipole.f (dipole)</td>
<td>{p_1}</td>
<td>D(i)</td>
</tr>
<tr>
<td>reducedm.f (reduced)</td>
<td>{p_1}</td>
<td>reduced {p_1}</td>
</tr>
<tr>
<td>cmatrix.f (cmatrix#)</td>
<td>{p_1}</td>
<td>CLBS (like helicity)</td>
</tr>
<tr>
<td>cmatrix.f (cmatrix#d)</td>
<td>{p_1}</td>
<td>CLBS (unlike helicity)</td>
</tr>
<tr>
<td>check.f (program checkdipole)</td>
<td>{p_1} in inputm.h</td>
<td>check of D(i), [</td>
</tr>
<tr>
<td>[P-terms]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>matrixLO1.f (smatrix)</td>
<td>{p_1}</td>
<td>[</td>
</tr>
<tr>
<td>Iterm.f (Iterm)</td>
<td>{p_1}</td>
<td>Ci of I-terms</td>
</tr>
<tr>
<td>cmatrix.f (cmatrix#)</td>
<td>{p_1}</td>
<td>CLBS</td>
</tr>
<tr>
<td>checkVirt.f (program checkVirt)</td>
<td>{p_1} in inputmLO.h</td>
<td>check of Ci of I-terms</td>
</tr>
<tr>
<td>[Check of soft(collinear limits)]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>collinear.f</td>
<td>{p_1}</td>
<td>collinear limits of {p_1}</td>
</tr>
<tr>
<td>checkIR.f (program checkIR)</td>
<td>{p_1}</td>
<td>((</td>
</tr>
<tr>
<td>Common input files</td>
<td></td>
<td></td>
</tr>
<tr>
<td>allcoloromat#.inc</td>
<td>extended color matrices (used in cmatrix#)</td>
<td></td>
</tr>
<tr>
<td>emitspecinfo.inc</td>
<td>information of emitter for dipoles (used in cmatrix#)</td>
<td></td>
</tr>
<tr>
<td>coupl.inc</td>
<td>common block of parameters</td>
<td></td>
</tr>
<tr>
<td>couplings.f</td>
<td>reads parameters in param_card.dat and assigns them in coupl.inc</td>
<td></td>
</tr>
</tbody>
</table>

Next, the evaluation of the squared matrix element for the real emission process \([|M|^2]\) proceeds in exactly the same way as in MadGraph with the call

```
call smatrix(p,msq)
```

Finally, the dipole terms are evaluated by calling the subroutine

```
call dipole(p,dip,SumD)
```

which returns an array with the values for each dipole term \((dip)\) and the sum of the all dipoles \((\text{SumD})\) as output. For our example process \(uu \rightarrow ddg\) we have 15 individual dipoles. The reduced kinematics of each dipole (as determined by the subroutine \(\text{reducedm}\)) can be accessed by the common block

```
double precision ptil(0:3,1:n1,n2)
common /OUTPUTM/ ptil
```

where \(n1 = 4\) and \(n2 = 15\) in the definition of the momenta for the example at hand.

**I-terms:** directory process/Proc_uux_ddxg/Virtual

Our program \(\text{checkVirt.f}\) for checks of the I-terms again provides an example for the use of the subroutines to evaluate the I-terms. In addition to the eight files from \(\text{cmatrix.f}\) to \(\text{param_card.dat}\) listed above for the evaluation of the dipoles, the user has to copy one file:

```
Iterm.f
```

Then, the initialization phase is exactly the same as before and for the evaluation of the I-terms, one has to call the subroutine

```
call Iterm(p,coef,SumI)
```

which provides the value of the coefficients \((i.e.\ \text{the array coef})\) as well as the sum \((\text{SumI})\) of all I-terms as output. If the user also wants to compute the LO matrix element \([M_{Born}]^2\) he needs the file \(\text{matrixLO1.f}\). The execution of the subroutine with the command

```
call smatrix(p,msqLO)
```

is performed again exactly as in MadGraph.

**P- and K-terms:** directory process/Proc_uux_ddxg/PK

The user needs the following file to evaluate the P- and K-terms:

```
PKterm.f
```

as well as the eight files from \(\text{cmatrix.f}\) to \(\text{param_card.dat}\) already discussed for the dipoles. The initialization phase is unchanged but the user has to provide the parton momentum fraction \(x\) from the mass factorization as additional input. The P- and K-terms for the first leg are computed by the call

```
call PKterm1(p,x,SumP,SumK)
```
which returns separately the sum of all P- and K-terms (SumP and SumK). For the second parton in initial state (as in hadron-hadron collisions) the corresponding P- and K-terms are evaluated by calling PKterm2.

4.3. General usage

For a specific application of AutoDipole, the user needs to perform the following three steps:

1. Setup of parameters : parameter.m
2. Run of package : GenerateAll[{initial},{final}]
3. Run and checks of the generated code at :
   /process/Proc_xx_xx/.

Let us explain each of these steps in more detail.

1. Setup of parameters

   Generally, the user has to supply all process parameters needed for the dipole subtraction procedure by editing the file,

   parameter.m

   where (following the MadGraph conventions) all values with the dimension of a mass are in units of [GeV]. Let us explain all variables in the file together with the chosen default values.

Parameters for dipole terms

   ep=0:
   The parameter of dimensional regularization of space–time, $D = 4 - 2\epsilon$. The default choice eliminates higher orders (positive powers) of $\epsilon$ in the dipole terms.

   kap=2/3:
   $\kappa$ is free parameter in some expressions for massive dipoles [2]. The value $\kappa = 0$ leads to the simplest results for the dipole terms, while $\kappa = 2/3$ produces the simplest expressions of the I-terms.

   skipdipole={};
   This set specifies the kind of dipoles to be skipped during the creation. For example, the set skipdipole={2t} omits the creation of the Dipole 2-{5} with the $t$-$\bar{t}$ splitting (see Table 2 in Appendix A).

   murt=174.30:
   The value of the renormalization scale. The default value is the same as the default value of the top-quark mass in MadGraph.

   acccut=10^{-3}:
   This value is used for the checking the quality of the infrared cancellations in the generated code upon scanning over all possible collinear configurations.

Parameters for I-terms

   mFlist={};
   This set lists all heavy quarks which possibly contribute to the I-terms with a gluon emitter in final state, e.g. as mFlist={t,b}.

   lightflavors=1:
   This is the number of the light flavors which contributes to the I-terms with a gluon emitter in initial and final states.

   replistVirtual={};
   This set enables user defined symbolic replacements to simplify the virtual contribution, e.g. of the type, replistVirtual={Gamma[1-eps]->(4^P)^(eps)*some_symbol^(-1)} which results in the replacement, $\frac{d\Gamma}{d(1-\epsilon)} = \text{somesymbol}$.

   sijexpansion=0:
   This parameter determines whether the typical factors of dimensional regularization like $\left(\frac{4\pi\epsilon}{2P_iP_j}\right)^{\epsilon}$ in the I-terms are expanded (0) or not (1).

Parameters for P- and K-terms

   KFSff=0, KFSgg=0, KFSfg=0, and KFSgf=0:
   Theses parameters specify the factorization scheme. In the MS scheme (default choice) they are all vanishing.

   lightquarksPK3={u}:
   This set specifies all light flavors to be added to the initial gluon of the reduced Born process in the P- and K-terms (see group (6) in Table 4 in Appendix A).

   muFfort=174.30:
   This is the value of the factorization scale $\mu_F$, which has the same default value as the renormalization scale $\mu_R$.

   The other parameters which are used by MadGraph are determined in the file patch/Models/sm/param_card.dat.

   The default values for the most important ones are the following (in units of [GeV] for masses),

   $\alpha_s = 0.1180$
   $\alpha_{em} = 0.007547169811320755$

   Top mass = 174.3
   Bottom mass = 4.7
   W mass = 80.419
   Z mass = 91.188
   Higgs mass = 120.0

   These parameters are used consistently throughout the run of the package. Any other changes of SM parameters like e.g. $G_F$, the CKM-parameters, the decay widths of heavy fields, and so on, are done in the same MadGraph file, param_card.dat. Similarly, also control over all interactions in the SM proceeds entirely through MadGraph, i.e. through the file, patch/Models/sm/interactions.dat in the exactly same way as in the normal use of MadGraph. All MadGraph parameters are explained in [42].

2. Run of package

   For general QCD processes (including leptons and SM bosons) we run AutoDipole by executing the Mathematica command:

   GenerateAll[{initial},{final}]

   where the sets labeled initial and final contain all external particles of the respective scattering process. The following fields are available in the present version of the package:

   Fermions:
   $(u, d, b, t, e^-, \mu^-, \bar{u}, \bar{d}, \bar{b}, \bar{t}, e^+, \mu^+)$ = $(u, d, b, t, e, \muon, \bar{u}, \bar{d}, \bar{b}, \bar{t}, \bar{e}, \muonbar)$

   Bosons:
   $(\text{gluon}, \text{photon}, W^+, W^-, Z, \text{Higgs})$ = $(g, \gamma, \Wp, \Wm, Z, h)$

   where the right hand side denotes the input format of AutoDipole. The available quark fields include the first family $(u$ and $d$-quarks) which serves as a template for light quarks and, respectively, the third family $(b$ and $t$-quarks) as one for heavy quarks. In practice this suffices as far as the subtraction formalism is concerned, because generally QCD scattering processes at NLO will be insensitive to the individual flavor of the quarks, and the sum over all light quark flavors can be trivially taken afterwards. If one intends to
study particular channels, e.g. in $W^\pm$ production, flavor separation is required, though. As a result of the run, AutoDipole produces the following information about the created dipoles and the I-, P- and K-terms:

- All analytical expressions for the dipole terms are written in the order given in Table 2 along with the reduced Born process (emitter,spectator) and the reduced kinematics for each dipole term. Also, the number of created dipole terms is shown group by group, namely Dipole 1, 2, 3, and 4 and their subgroups (see Fig. 3). The number of massive dipole terms is given separately.

- All soft and collinear limits together with the corresponding dipole terms are returned. For example, for the process $g(1)g(2)\rightarrow t(3)f(4)g(5)g(6)$ (see next Subsection) the fifth collinear limit is denoted $Lc(5) = (5,6)$: $Sc(5) = (17,18,19,20)$. This means that the collinear divergence of the pair $(5g), (6g)$ is canceled by the set of the dipoles, $Sc(5)$ and the set consists of the dipoles number 17, 18, 19, and 20. In the same way, the soft limit and the corresponding dipole terms are displayed. E.g. for the soft limit of gluon $5g$, $Ls(1) = (5)$: $Ss(1) = (Sc(1),Sc(2),Sc(5),Sc(6),Sc(7))$, which means that the soft divergence of $5g$ is canceled by the set of the dipoles, $Ss(1)$. This set in turn consists of the sets of collinear limits $Sc(1),Sc(2)$, and so on.

- The analytical expressions of all I-terms are shown in the order as given in Table 3 prior to the expansion in $\epsilon$ as well as the individual coefficients of the poles $C_{-2}$, $C_{-1}$, and $C_0$ (see Eq. (3.30)) after the $\epsilon$ expansion. Also the total number of all created I-terms is displayed by group by group (see Table 3).

- The analytical expressions of all created P- and K-terms are shown in the order given in Table 2 together with the total number of P- and K-terms.

- Finally, AutoDipole outputs a summary with the numbers of all created dipoles, I-, P- and K-terms and the reduced Born processes.

This information, being rather detailed, helps with the identification of individual terms according to the order of creation. The Mathematica part of AutoDipole provides analytical and symbolic expressions, which are easily accessible, if needed. These features are primarily for an experienced user.

Upon running the AutoDipole package it occurs (on purely algorithmic grounds) in some cases though, that a particular reduced Born cross section does not exist. For example, the process $e^-e^+\rightarrow u\bar{u}g$ does give rise through the dipole $2-(5)-(u,\bar{u})$ to the Born process $e^-e^+\rightarrow gg$, which does not exist at tree level. In such cases, an error message is displayed:

Reduced Born amplitude $B_{2u}$ does not exist.
Dipole $2u$ must be switched off.
Please set in the file ./parameter.m skipdipole=$2u$

Following the message the user has to modify the command skipdipole and repeat the Mathematica run. In this way only the relevant partial subsets of dipole terms is extracted and the offending Born processes are skipped (thanks to skipdipole) before calling the MadGraph part.

3. Run and checks of the generated code

Finally, as briefly explained above, the user can run the generated Fortran code for the regularized real emission process as well as the I-, P- and K-terms. The checks proceed as follows:

Dipole terms

As mentioned above, the evaluation of the real matrix element squared $|M|^2$ and the sum of all dipole terms $\sum D(i)$ at 10 randomly chosen phase space points as well as the check of cancellations in all infrared limits for the subtracted matrix element are automatically executed:

```
make
./runD
make checkIR
./checkIR
```

The latter command performs a scan over all possible collinear configurations. For the expression

$$\frac{|M|^2 - \sum D}{|M|^2},$$

of Eq. (3.29), it is checked whether the cancellation works at least to the accuracy defined by the value of the parameter acccut in the file parameter.m. If the absolute value of the normalized subtracted squared matrix element in Eq. (3.29) drops below the pre-defined value of acccut sufficiently deep in the collinear limit, the message,

Infrared safety of all collinear limits is confirmed for $S_{ij}/S > ...$

is printed.

I-terms

The Fortran code to evaluate all I-terms is collected in the subdirectory ./Virtual. There, the coefficients of the double and single poles in the $\epsilon$ expansion $C_{-2}$, $C_{-1}$, $C_0$ in Eq. (3.30) can be evaluated at 10 randomly chosen phase space points by:

```
make runI
./runI
```

This includes in particular the calculation of the double poles according to

$$C_{-2} = \frac{C_2}{(\alpha_s/2\pi)|M_{Born}|^2},$$

and the user can easily confirm that Eq. (3.31) holds.

P- and K-terms

Likewise, the mass factorization terms originating from the P and K-terms is stored in ./PK and also evaluated at 10 randomly chosen phase space points by the command:

```
make runPK
./runPK
```

This concludes our brief description of the user specific features of the AutoDipole package.

4.4. Examples

The package includes a command to demonstrate the run for several example processes. This helps to explain the package and at the same time confirms that the installation was done successfully and that the package works properly. These examples include the computation of some of the numbers given in Tables 5 and 6 in Appendix C.
generates five processes in different kinematics,

1. $e^- e^+ \rightarrow u\bar{u} g$
2. $e^- e^+ \rightarrow e^- u g$
3. $g g \rightarrow t\bar{t} g$
4. $u g \rightarrow t u g$
5. $u u \rightarrow W^+ W^- g$

provides suitable settings for the parameters of each process and runs the package. For illustration, we only explain in the following those settings which are changed with respect to the default ones.

The processes from 1 to 4 use a value of $\alpha_s = 0.1075205492734706$ from [36] for the strong coupling constant. For the processes 1 and 2 the contribution of the Z-boson exchange is switched off in the file interactions.dat for simplicity. Moreover, in parameter.m the splitting $2\mu$ in process 1 (and $3\mu$ in 2) is skipped by the command skipdipole = (2u) (and skipdipole = (3u) in 2, respectively) to exclude an inexistent reduced Born process.

For the processes 3 and 4 the QED interaction is switched off in the file interactions.dat in order to focus on the QCD corrections only. Furthermore, a value of 174 [GeV] for the top mass is used following [36] and the decay width of the top-quark must be switched off in the file param_card.dat. For the comparison with [36] the $t$–$\bar{t}$ splitting is skipped in parameter.m, i.e. skipdipole = (2t), and the renormalization and factorization scales are set equal to the top mass value. For the $I$-terms five light flavors are taken into account, lightflavors = 5.

Finally, process 5 uses the following settings [40] in the file param_card.dat: $W$-mass = 80.425, $\alpha_s = 0.1202629003906369$, $\alpha_{em} = 0.00754359570869335$ and Fermi’s constant $G_F = 1.16637 \times 10^{-5}$, and the decay width of $W$-boson is switched off. The results of processes 3, 4, and 5 should reproduce the corresponding numbers in Tables 5 and 6.

The test run of all examples is finally checked by comparing all computed values of the dipoles, $I$-terms, and the averaged matrix element squared $|M|^2$ for the processes 1 to 5 the with ones in the data files in ./lib/check/. At the end of the run, e.g. for example 3, the following message appears:

```
3.gg_ttxgg
|M|^2(Real) :Confirmed
Dipole :Confirmed
|M|^2(LO) :Confirmed
I-term :Confirmed
```

This confirms the successful installation and proper functioning of the package. The generated code for all examples is stored in the directory ./process as in a usual run and the directory of each process includes in the file res_std the result of the comparison with the data files from ./lib/check/. All parameters used for each process are accessible in param_card.dat, so the user can also run the respective files individually with usual commands of the previous subsection.

5. Conclusion

We have presented the package AutoDipole which provides an implementation of the Catani-Seymour dipole formalism [1, 2] to compute subtracted matrix elements squared in an automatic way. The package (partial aspects of which have been discussed before [17,18]) consists of essentially three ingredients: the automatic generation of all dipole terms via Mathematica routines as well as the calculation of the color-correlated matrix elements and the evaluation of different helicity amplitudes with the help of MadGraph [3–5] (i.e. a patched stand-alone version).

We have presented a complete discussion of the set-up of the program and the details of our implementation as well as the numerical evaluation of the squared matrix elements. Particular emphasis has been put on explaining the use of AutoDipole which has been illustrated with sufficiently non-trivial examples. The finiteness in the soft and collinear limits has been demonstrated for a large number of processes in $e^- e^-$, $ep$ and $pp$-scattering. Also, checks for the complete set of dipoles have been performed by comparing to various processes in the recent literature.

In comparison with other available software to generate subtraction terms for the real emission contributions AutoDipole has a number of advantages. First of all, the algorithm is implemented in a way so that analytic expressions are generated which are accessible to symbolic manipulation. Then, the chosen order for the dipole generation produces flat Fortran code and allows to handle scattering processes with $2 \rightarrow 5$ and $2 \rightarrow 6$ partons of relevance for LHC phenomenology including the computation of the CLBS. Finally, the structure of the package is highly modular and flexible for further extensions.

Future improvements of the AutoDipole package will address restricting the dipoles to small regions of phase space, as realized e.g. through $\alpha$-parameters [43]. This is advantageous when performing the Monte Carlo integration over the phase space of the real emission process. The implementation of such cuts affects also the integrated dipoles and the case of massless partons is available in the literature whereas for the case of massive partons most cases can be found in [44,45]. The remaining integrals for exhaustive coverage can be done e.g. with the methods of [32]. Next, the current version of AutoDipole does not cover processes with identified hadrons in the final state where one needs fragmentation. Here, the so-called $H$-terms have to be taken into account, which will be implemented in a later version of AutoDipole. Finally, the structure of our code is particularly suited for extensions to physics beyond the Standard Model (BSM). The main modification needed is the so-called model file of MadGraph, which is currently already available for a large class of BSM scenarios including e.g. the minimal supersymmetric extension of Standard Model for which the necessary modifications of the dipole formalism are known [2]. All the extensions mentioned above can be easily realized within the existing structure of our code.

The AutoDipole package is available for download from [41] or from the authors upon request. The MadGraph package (stand-alone version) which includes the HELAS library may be obtained from [42].

Acknowledgements

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Appendix A. Creation order of subtraction terms

Table 2
The emitter pair is denoted \((i, j)\) or \((a, i)\) and, respectively, the spectator \(k(\neq i, j)\) or \(b(\neq a, d)\). D is a massless dipole and DM is a massive dipole. The equation numbers refer to [1] for massless dipoles (D) and to [2] for massive ones (DM). Dipole 2 has the other flavors, massless down and massive bottom quarks. Dipole 3 and 4 have also the other light flavors, \(f = \bar{u}, \bar{d}, \bar{d}\).

<table>
<thead>
<tr>
<th>Emitter</th>
<th>Spectator</th>
<th>Dipole</th>
<th>Condition</th>
<th>Eq.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dipole 1</td>
<td>(1) ((i, j) = (f, g))</td>
<td>1. (k)</td>
<td>(D_{ij,k})</td>
<td>(m_i = m_j = 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. (b)</td>
<td>(D_{ij,b})</td>
<td>(m_i \neq 0) or (m_j \neq 0)</td>
</tr>
<tr>
<td></td>
<td>(2) ((i, j) = (g, g))</td>
<td>3. (k)</td>
<td>(D_{ij,k})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4. (b)</td>
<td>(D_{ij,b})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td></td>
<td>(3) ((a, i) = (f, g))</td>
<td>5. (k)</td>
<td>(D_{ip,k})</td>
<td>(m_i = 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>6. (b)</td>
<td>(D_{ip,b})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td></td>
<td>(4) ((a, i) = (g, g))</td>
<td>7. (k)</td>
<td>(D_{ip,k})</td>
<td>(m_i = 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>8. (b)</td>
<td>(D_{ip,b})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td>Dipole 2</td>
<td>((5) (i, j) = (f, \bar{f}))</td>
<td>9u. (k)</td>
<td>(D_{ij,k})</td>
<td>(m_i = 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10u. (b)</td>
<td>(D_{ij,b})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td></td>
<td>((i, j) = (t, \bar{t}))</td>
<td>9t. (k)</td>
<td>(D_{ij,k})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10t. (b)</td>
<td>(D_{ij,b})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td>Dipole 3</td>
<td>(6) ((a, a') = (f, f))</td>
<td>11u. (k)</td>
<td>(D_{pi,k})</td>
<td>(m_i = 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>12u. (b)</td>
<td>(D_{pi,b})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td>Dipole 4</td>
<td>(7) ((a, a') = (g, f))</td>
<td>13u. (k)</td>
<td>(D_{pi,k})</td>
<td>(m_i = 0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>14u. (b)</td>
<td>(D_{pi,b})</td>
<td>(m_i \neq 0)</td>
</tr>
</tbody>
</table>

Table 3
The emitter is denoted \(i\) or \(a\) and, respectively, the spectator \(k(\neq i, a)\) or \(b(\neq a, d)\). I is the massless insertion operator and IM is a massive one. The equation numbers refer to [1] for massless insertions (I) and to [2] for massive ones (IM). The set of the heavy-quark masses running in the quark loop are abbreviated \([m_j]\).

<table>
<thead>
<tr>
<th>Emitter</th>
<th>Spectator</th>
<th>Condition</th>
<th>Eq.</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) (i = f)</td>
<td>1. (k)</td>
<td>(I_{f,i,k})</td>
<td>(m_i = m_j = 0)</td>
</tr>
<tr>
<td></td>
<td>2. (b)</td>
<td>(I_{f,i,b})</td>
<td>(m_i \neq 0) or (m_j \neq 0)</td>
</tr>
<tr>
<td>(2) (i = g)</td>
<td>3. (k)</td>
<td>(I_{g,i,k})</td>
<td>(m_i = [m_j] = 0)</td>
</tr>
<tr>
<td></td>
<td>4. (b)</td>
<td>(I_{g,i,b})</td>
<td>(m_i \neq 0) or ([m_j] \neq 0)</td>
</tr>
<tr>
<td>(3) (a = f)</td>
<td>5. (k)</td>
<td>(I_{f,a,k})</td>
<td>(m_i = 0)</td>
</tr>
<tr>
<td></td>
<td>6. (b)</td>
<td>(I_{f,a,b})</td>
<td>(m_i \neq 0)</td>
</tr>
<tr>
<td>(4) (a = g)</td>
<td>7. (k)</td>
<td>(I_{g,a,k})</td>
<td>(m_i = 0)</td>
</tr>
<tr>
<td></td>
<td>8. (b)</td>
<td>(I_{g,a,b})</td>
<td>(m_i \neq 0)</td>
</tr>
</tbody>
</table>

Appendix B. Correlation functions of gluon emitter

We use the following definitions of spinor products [37]:

\[
\langle k_1 k_2 \rangle = \sqrt{k_1^+ k_2^-} e^{i\phi(k_1)} - \sqrt{k_1^- k_2^+} e^{-i\phi(k_2)} \tag{B.1}
\]

\[
k^\pm \equiv |k| \pm k_z \tag{B.2}
\]

\[
e^{i\phi(k)} = \frac{k_i}{|k|} \tag{B.3}
\]

\[
k_{\perp} = k_x + ik_y \tag{B.4}
\]

If the reduced momentum of the initial gluon emitter lies in z-axis with \(l = E_l(1,0,0,\pm 1)\) the spinor product are defined as \((p_{\perp} p_{\perp}^L)|_L\) to avoid an ambiguity in the intermediate steps of the calculation,

\[
(k_L)|_L = \sqrt{2E_l k^- e^{i\phi(k)}} (E_l = l_z) \tag{B.5}
\]

\[
z (l_L l_L^L) = \left[ \frac{2 E_l E_2}{2 E_l E_2} \right] \left( E_1 = l_{1z} \text{ and } E_2 = -l_{2z} \right) \tag{B.6}
\]

The explicit forms of \(e^{i\phi(k)}|_{L}\) in Eq. (3.23) are shown for all massless and massive dipoles with a gluon emitter. Here we write the quantity including the phase difference as \(E_{\perp} = e^{i\phi(k)}|_{L}\). The reduced momenta are ones used in the corresponding equations.
in [1,2], which are shown in Table 2. The \( \tilde{p}_{ij} \) and \( \bar{p}_{ai} \) are massless and the \( p^2 M_{ij} \) are massive ones.

### B.1. Massless momenta

[Dipole 1-(2)-3\((m_k = 0)\) and Dipole 2-(5)-9\((m_k = 0)\)]

\[
E_{1+} = e^{+i\phi(\tilde{p}_{ij})} z_i^*(p_i^j p_j^i) \sqrt{2/(p_i^j p_j^i)} \tag{B.7}
\]

[Dipole 1-(2)-4 and Dipole 2-(5)-10\((m_k = 0)\)]

\[
E_{2+} = e^{+i\phi(\tilde{p}_{ij})} z_i^*(p_i^j p_j^i) \sqrt{2/(p_i^j p_j^i)} \tag{B.8}
\]

[Dipole 1-(4)-7\((m_k = 0)\) and Dipole 3-(6)-11\((m_k = 0)\)]

\[
E_{3+} = \frac{(p_i^j p_j^i)}{\sqrt{2(1-u_i)(p_i^j p_j^i)}} \tag{B.9}
\]

[Dipole 1-(4)-8 and Dipole 3-(6)-12u]

\[
E_{4+} = \frac{-(s_{ai} p_i^a p_j^i)}{\sqrt{2s_{ab}(p_i^a p_j^i)}} \tag{B.10}
\]

### B.2. Massive momenta

[Dipole 1-(2)-3\((m_k \neq 0)\) and Dipole 2-(5)-9\((m_k \neq 0)\)]

\[
EM_{1+} = e^{+i\phi(p^2 M_{ij})} z_i^m(p_i^j p_j^i) \sqrt{2/(p_i^j p_j^i)} \tag{B.11}
\]

[Dipole 2-(5)-9t]

\[
EM_{1k+} = e^{+i\phi(p^2 M_{ij})} z_i^m(p_i^j p_j^i) \sqrt{2/(p_i^j p_j^i)} \tag{B.12}
\]

with flat momentum

\[
P_{i ij} = p_{ij} - \frac{m^2_{ij}}{2p_{ij} \cdot p^2 M_{ij}} p^2 M_{ij} \tag{B.13}
\]

[Dipole 2-(5)-10t]

\[
EM_{2+} = e^{+i\phi(p^2 M_{ij})} z_i^m(p_i^j p_j^i) \sqrt{2/(p_i^j p_j^i)} \tag{B.14}
\]

with

\[
P_{i ij} = p_{ij} - \frac{m^2_{ij}}{2p_{ij} \cdot p^2 M_{ij}} p^2 M_{ij} \tag{B.15}
\]

[Dipole 1-(4)-7\((m_k \neq 0)\) and Dipole 3-(6)-11\((m_k \neq 0)\)]

\[
EM_{3+} = \frac{(p_i^j p_j^i)}{\sqrt{2(1-u_i)(p_i^j p_j^i)}} \tag{B.16}
\]

with

\[
P_{k} = p_{k} - \frac{m^2_{k}}{2p_{k} \cdot p_{ai}} \bar{p}_{ai} \tag{B.17}
\]

### Appendix C. Comparison with the \( t\bar{t} + 1 \) jet

#### Table 5

The results of the \( |M|^2 \) and the sum of all dipole terms are shown. The two implementations agree at least to 14 digits for the matrix elements squared and at least to 12 digits for the sum of the subtraction terms. The phase space point used in the last entry corresponds to the first one in ./lib/check/check5_uuc_wv-wq/input_m_uuc_wv-wq_gg.

<table>
<thead>
<tr>
<th>( b_0 ) [GeV(^{-4})]</th>
<th>( d_0 ) [GeV(^{-4})]</th>
</tr>
</thead>
<tbody>
<tr>
<td>( g g \rightarrow t\bar{t} g )</td>
<td>( 1.2353289076588 \times 10^{-11} )</td>
</tr>
<tr>
<td>AutoDipole</td>
<td>( 1.2353289076588 \times 10^{-11} )</td>
</tr>
<tr>
<td>Ref. [36]</td>
<td>( 1.2353289076588 \times 10^{-11} )</td>
</tr>
<tr>
<td>( u \bar{u} \rightarrow t\bar{u} g )</td>
<td>( 9.06309024880356 \times 10^{-12} )</td>
</tr>
<tr>
<td>AutoDipole</td>
<td>( 9.06309024880356 \times 10^{-12} )</td>
</tr>
<tr>
<td>Ref. [36]</td>
<td>( 9.06309024880356 \times 10^{-12} )</td>
</tr>
<tr>
<td>( u g \rightarrow t\bar{u} )</td>
<td>( 6.8153074525503 \times 10^{-5} )</td>
</tr>
<tr>
<td>AutoDipole</td>
<td>( 6.8153074525503 \times 10^{-5} )</td>
</tr>
<tr>
<td>Ref. [36]</td>
<td>( 6.8153074525503 \times 10^{-5} )</td>
</tr>
</tbody>
</table>

#### Table 6

The coefficients for the color and spin averaged results for the 1-operator. At least 14 digits agreements are obtained.

<table>
<thead>
<tr>
<th>( c_{-2} )</th>
<th>( c_{-1} )</th>
<th>( c_0 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( g )</td>
<td>( 2.49467966948004 \times 10^{-4} )</td>
<td>( 6.3898976683705 \times 10^{-4} )</td>
</tr>
<tr>
<td>AutoDipole</td>
<td>( 2.49467966948004(1) \times 10^{-4} )</td>
<td>( 6.38989766837061(1) \times 10^{-4} )</td>
</tr>
<tr>
<td>Ref. [36]</td>
<td>( 2.49467966948004(1) \times 10^{-4} )</td>
<td>( 6.38989766837061(1) \times 10^{-4} )</td>
</tr>
<tr>
<td>( u )</td>
<td>( 1.3849987972387 \times 10^{-5} )</td>
<td>( 2.88738914839178 \times 10^{-5} )</td>
</tr>
<tr>
<td>AutoDipole</td>
<td>( 1.3849987972387(0) \times 10^{-5} )</td>
<td>( 2.887389148391791(0) \times 10^{-5} )</td>
</tr>
<tr>
<td>Ref. [36]</td>
<td>( 1.3849987972387(0) \times 10^{-5} )</td>
<td>( 2.887389148391791(0) \times 10^{-5} )</td>
</tr>
<tr>
<td>( g )</td>
<td>( 3.8458076607470 \times 10^{-6} )</td>
<td>( 7.73777480040817 \times 10^{-6} )</td>
</tr>
<tr>
<td>AutoDipole</td>
<td>( 3.8458076607470(0) \times 10^{-6} )</td>
<td>( 7.73777480040817(1) \times 10^{-6} )</td>
</tr>
<tr>
<td>Ref. [36]</td>
<td>( 3.8458076607470(0) \times 10^{-6} )</td>
<td>( 7.73777480040817(1) \times 10^{-6} )</td>
</tr>
<tr>
<td>( g g \rightarrow t\bar{t} g )</td>
<td>( 6.22738241305372 \times 10^{-5} )</td>
<td>( 6.8153074525503 \times 10^{-5} )</td>
</tr>
<tr>
<td>AutoDipole</td>
<td>( 6.22738241305372(0) \times 10^{-5} )</td>
<td>( 6.81530745255037(1) \times 10^{-5} )</td>
</tr>
</tbody>
</table>
References