GR@PPA 2.8: initial-state jet matching for weak boson production processes at hadron collisions

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Abstract

The initial-state jet matching method introduced in our previous studies has been applied to the event generation of single $W$ and $Z$ production processes and diboson ($W^+W^-$, $WZ$ and $ZZ$) production processes at hadron collisions in the framework of the GR@PPA event generator. The generated events reproduce the transverse momentum spectra of weak bosons continuously in the entire kinematical region. The matrix elements (ME) for hard interactions are still at the tree level. As in previous versions, the decays of weak bosons are included in the matrix elements. Therefore, spin correlations and phase-space effects in the decay of weak bosons are exact at the tree level. The program package includes custom-made parton shower programs as well as ME-based hard interaction generators in order to achieve self-consistent jet matching. The generated events can be passed to general-purpose event generators to make the simulation proceed down to the hadron level.

Keywords: GRACE; Hadron collision; Event generator; Jet matching; Parton shower; Weak boson production

PROGRAM SUMMARY

Manuscript Title: GR@PPA 2.8: initial-state jet matching for weak boson production processes at hadron collisions
Authors: S. Odaka, Y. Kurihara
Program Title: GR@PPA 2.8
Journal Reference:
Catalogue identifier: none
Licensing provisions: none
Programming language: Fortran; with some included libraries coded in C and C++
Computer: all
Operating system: any UNIX-like system
RAM: 1.6 Miga bytes at minimum
Supplementary material: none
Keywords: GRACE; Hadron collision; Event generator; Jet matching; Parton shower; Weak boson production
Classification: 11.2
External routines/libraries: bash and Perl for the setup, and CERNLIB, ROOT, LHAPDF, PYTHIA according to the choice of users
Does the new version supersede the previous version?: No, this version supports only a part of the processes included in previous versions.
Nature of problem: We need to combine those processes including 0 jet and 1 jet in the matrix elements using an appropriate matching method, in order to simulate weak-boson production processes in the entire kinematical region.
Solution method: The leading logarithmic components to be included in parton distribution functions and parton showers are subtracted from 1-jet matrix elements. Custom-made parton shower programs are provided to ensure satisfactory performance of the matching method.
Reasons for the new version: An initial-state jet matching method has been implemented.

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1. Introduction

The inclusive production cross section of a final state, \( A \), from collisions of two hadrons, \( h_1 \) and \( h_2 \), at a squared center-of-mass (cm) energy of \( s \) is usually evaluated as

\[
\sigma_{h_1h_2 \rightarrow A + X} = \sum_{a,b,i} \int_0^1 dx_1 \int_0^1 dx_2 \, d\hat{\Phi}_{A,f_{h_1 \rightarrow a}(x_1, \mu_F^2)} f_{h_2 \rightarrow b}(x_2, \mu_F^2) \frac{d\hat{\sigma}_{ab \rightarrow A_i}(\hat{s})}{d\hat{\Phi}_{A_i}} \delta(\hat{s} - x_1 x_2 s),
\]

where \( f_{h_1 \rightarrow a}(x_1, \mu_F^2) \) is the parton distribution function (PDF) representing the existence probability of the parton \( a \) (a light quark or a gluon) inside the hadron \( h_k \) with a momentum fraction of \( x_k \) at a certain energy scale of \( \mu_F \) (factorization scale). The factor \( d\hat{\sigma}_{ab \rightarrow A_i}(\hat{s})/d\hat{\Phi}_{A_i} \) represents the differential cross section of the hard interaction that produces the final state \( A_i \) from the collision of two partons, \( a \) and \( b \), perturbatively calculated according to fixed-order matrix elements (ME) at a squared cm energy of \( \hat{s} \). The final state \( A \) may consist of several sub-states \( A_i \) at the parton level; for instance, \(^1\!\!J\!e\!t\) production includes a variety of light quark and gluon productions. Such a generalization is necessary in hadron collisions because it is difficult to separate the sub-states experimentally.

Monte Carlo (MC) event generators are, in principle, an MC integration of Eq. (1), in which the distribution of the sampling points is controlled so that the frequency is proportional to the differential cross section. Thus, each sampling point (event) can be treated just like an event produced by actual interactions. MC event generators are mainly used for evaluating the detection efficiency and acceptance of experiments; therefore, they are desired to exclusively reproduce the actual phenomena as precisely as possible. The initial-state QCD activities reproduced by PDFs in Eq. (1) not only alter the longitudinal momentum distribution of constituent partons, but also produce transverse activities which result in a finite transverse recoil of the hard interaction system and additional hadronic activities visible in detectors. Experimentalists want MC event generators to reproduce such transverse activities as well, since all these effects may alter the performance of detectors.

Parton showers have been developed to fulfill the above requirement; they reproduce the transverse activities of QCD radiations as well as the resultant longitudinal evolution evaluated in PDFs. Parton showers are a recursive solution of the DGLAP equation [1, 2, 3] from which the perturbative part of PDFs is derived. Thus, in principle, the PDFs in Eq. (1) can be replaced with parton showers, except for the non-perturbative components of the PDFs. However, in most of the MC event generators currently available, PDFs are directly employed for evaluating the initial-state conditions of the hard interactions, and parton showers are adopted only as models for simulating the initial-state QCD activities by using a technique of the "backward evolution" [4]. The identity between the PDF and PS is not seriously considered in such simulations.

PDFs sum up the collinear QCD corrections, which result in large logarithmic terms, to all orders of the coupling constant \( \alpha_s \). In order to improve the convergence of the perturbative calculations for hard interactions, the summation is limited by an arbitrary energy scale (factorization scale), and the resultant cross section depends on the choice of this parameter. The factorization scale, \( \mu_F^2 \), is defined as the upper limit on the momentum transfer, \( Q^2 = -t \), of the radiations considered in the summation. This scale is usually taken to be equal to a typical energy scale, such as \( \hat{s} \) or \( |\hat{t}| \), of the considered hard interaction because non-collinear components missing in PDFs become significant around this scale. Parton showers should reproduce the perturbative part of PDFs; hence, if we consider the identity between PDF and PS seriously, \( Q^2 \) of PS branches must also be limited by \( \mu_F^2 \).

As a result, the transverse recoil of the system \( A \) in Eq. (1) is limited. Since there is no such limitation
In actual phenomena, the inclusion of harder radiations is necessary to reproduce the phenomena in the entire phase space.

In order to add the hard radiation effect to Eq. (1), we have to anyhow include the effects of interactions that produce at least one additional parton in association with the production of $A (A + 1$-jet process). A straightforward approach is to replace the hard interaction part in Eq. (1) with that for the $A + 1$-jet process, and then, to add the resultant cross section or an event sample to the results for the production of $A (A + 0$-jet process). Equation (1) does not have any problem if the final state $A$ is a color-singlet state, for instance, a single weak boson. However, we encounter serious problems if we replace it with $A + 1$-jet processes.

Partons in the final state (jets) are in most cases produced as a result of QCD interactions. Double counting may occur because PDFs and parton showers are also consequences of QCD interactions. We encounter this problem even when we simply try to generate those events including partons in the final state of the hard interaction. The same problem also occurs when we try to combine processes having different jet multiplicities, for instance, the $A + 0$-jet process and the $A + 1$-jet process. The simplest solution to this problem is to clearly separate the phase space for the radiations using $\mu_F$. Since radiation effects at $Q^2 < \mu_F^2$ are taken into account in PDFs and parton showers, the hard interaction part should include only those radiations having $Q^2 > \mu_F^2$. Although double counting can be avoided with this solution, another serious problem (mismatch) may occur owing to the existence of non-collinear contributions disregarded in PDFs and parton showers, which may become significant around $\mu_F^2$.

Several solutions to the double counting problem have been proposed and implemented in MC event generators, such as the ME correction in PYTHIA [5] and HERWIG [6], and the CKKW method [7] implemented in Sherpa [8]. The MLM prescription in AlpGen [9] can be considered as an alternative to the CKKW method. These are solutions for leading-order (LO) event generators. For complete next-to-leading order (NLO) event generation, a subtraction method is applied in MC@NLO [10, 11] and a suppression method is used in POWHEG [12, 13].

GR@PPA (GRace At Proton-Proton/Antiproton)\(^1\) is a Monte Carlo event generator package for simulating interactions at proton-proton and proton-antiproton collider experiments. It is an extension of the GRACE system [14, 15] to hadron collision interactions. GRACE is a powerful tool for deriving the differential cross section of hard interactions at the parton level, $d\hat{\sigma}_{ab\rightarrow A_i}(\hat{s})/d\hat{\Phi}_{A_i}$ in Eq. (1), and for generating events according to it with the help of BASES/SPRING [16, 17]. GR@PPA provides a mechanism for adding the effects of the initial-state variation in the flavor and momentum according to PDF and for achieving the generalization of the final state, as described symbolically in Eq. (1). The previous releases of GR@PPA [18, 19] include many multi-body (multi-jet) production processes, such as $W$ and $Z +$ jets, diboson ($W^+W^-$, $WZ$, $ZZ$) + jets, top-pair + jet, and QCD multi-jets. However, its application is restricted because the jet matching discussed above is not taken into consideration.

As described in previous reports, we have proposed a solution (matching method) [20] to the double counting problem, and we have shown its feasibility in $W$ boson production [21]. We have also shown that if we apply the method to $Z$ boson production in the GR@PPA event generator, the generated events reproduce the $p_T$ spectrum of $Z$ bosons measured at Fermilab Tevatron with surprisingly high precision over the entire measurement range [22]. In this report, we describe a new version of the GR@PPA event generator package, GR@PPA 2.8, in which our matching method is applied not only to the single $W$ and $Z$ boson production processes but also to the diboson ($W^+W^-$, $WZ$ and $ZZ$) production processes in proton-proton and proton-antiproton collisions.

Although our matching method is designed with the objective of developing NLO event generators, where the inclusion of one additional parton in the final state is necessary, the event generation in GR@PPA 2.8 is currently at the tree level because virtual corrections are yet to be included. Jet matching is accomplished by subtraction, as in the case of MC@NLO. However, subtraction is carried out in a limited phase space in our method, whereas there is no such limitation in MC@NLO. The limited application of subtraction can potentially enable us to easily extend the matching method to the final state. If the extension is realized successfully, it can be used for the matching between multi-jet production processes, similar to the CKKW method. Because our method is based on the concept

\(^1\)http://atlas.kek.jp/physics/nlo-ug/grappa.html.
that higher jet multiplicity generators are used to supplement the deficits in lower jet multiplicity generators, the necessity for higher jet multiplicity processes will not be as critical as that in the CKKW method.

Although the process-independent framework and process-dependent packages were provided separately in the previous version [19], the current version provides an all-in-one package because our matching method can be presently applied only to a subset of the processes implemented in the previous version. Though other process packages in the previous version can be imported in principle, such a modification is not recommended because careful treatments must be required for the execution, and the performance has not been tested. The features supported in previous versions [18, 19], such as inclusion of weak-boson decays in matrix elements, finite decay widths of weak bosons, branching ratios of weak bosons tuned to measurement data, inclusion of CKM non-diagonal couplings, and Z-photon mixing, are also supported in the current version.

The remainder of this report is organized as follows. The matching method applied in GR@PPA 2.8 is described in Section 2. In general, the detailed implementation of the solution to the double counting problem depends on the parton shower used in the event generation. We provide our own parton showers in this package in order to ensure satisfactory performance of our method. Parton showers are provided for the initial state and the final state. The former is crucial for our matching method, whereas the latter remains experimental and is implemented for completeness. These parton showers are described in Section 3. Section 4 contains instructions for the installation of libraries and execution of sample programs. Some results that have not been presented in previous reports are presented in Section 5. Practical performance parameters such as the program size and CPU time are presented in Section 6. Finally, a summary is provided in Section 7.

2. Initial-state jet matching

The concept of our matching method has been described in a previous report [20]. Hereafter, we assume that A represents a single weak boson or a diboson system. Our concept is as follows. We preserve the event generation according to Eq. (1), with the factorization scale \( \mu_F \) chosen as usual. Then, we separately generate events simulating the radiation contributions that are missing in Eq. (1), non-collinear contributions and larger \( Q^2 (> \mu_R^2) \) contributions, using the matrix elements for the \( A + 1 \)-jet process. If these two event samples are combined, we should obtain an event sample that covers all the radiation effects that may affect the production kinematics of the system A. Since the \( A + 1 \)-jet process is of the first order in QCD, it contains only the leading-order contribution of parton showers. This contribution can be factorized and can be evaluated using the \( A + 0 \)-jet matrix element and a radiation factor. Therefore, we can derive the desired \( A + 1 \)-jet process by subtraction. We call this method the limited leading-log (LLL) subtraction.

The LLL subtraction is carried out at the matrix-element (ME) level as

\[
\left| M_{A+1}^{(\text{sub})}(\hat{s}_{A+1}, \hat{\Phi}_{A+1}; \mu_R) \right|^2 = \left| M_{A+1}(\hat{s}_{A+1}, \hat{\Phi}_{A+1}; \mu_R) \right|^2 - \sum_i \left| M_A(\hat{s}_A, \hat{\Phi}_{A,i}; \mu_R) \right|^2 f_{LL,i}(Q_i^2, z) \theta(\mu_F^2 - Q_i^2),
\]

where the first term on the right-hand side represents the exact ME for \( A + 1 \)-jet production at a squared cm energy of \( \hat{s}_{A+1} \). The factor \( f_{LL,i}(Q_i^2, z) \) is the radiation factor in the leading-log approximation, and \( |M_A(\hat{s}_A, \hat{\Phi}_{A,i}; \mu_R)|^2 \) represents the ME for the non-radiative (\( A + 0 \)-jet) subsystem having a squared cm energy of \( \hat{s}_A = z\hat{s}_{A+1} \). The Lorentz boost and angular rotation of the final state \( A \), owing to the jet radiation, is taken into account in the calculation of the \( A + 0 \)-jet ME. The matrix elements are evaluated using the first-order strong coupling, expressed as

\[
\alpha_s(Q^2) = \frac{4\pi}{\beta_0 \ln(Q^2/\Lambda^2)}
\]

with \( \beta_0 = 11 - 2n_f/3 \), where the energy scale \( Q^2 \) is fixed to a given renormalization scale \( \mu_R^2 \). The \( \theta \) function "limits" the subtraction at the factorization scale \( \mu_F^2 \) since parton showers are limited by this scale.

4
We define the radiation factor as
\[
f_{\text{LL},i}(Q_i^2, z) = \frac{\alpha_s(\mu_R^2)}{2\pi} \frac{P_i(z) 16\pi^2}{z Q_i^2},
\]
where \(P_i(z)\) represents the leading-order Altarelli-Parisi splitting functions, given by
\[
P_{q\to qg}(z) = C_F \frac{1 + z^2}{1 - z},
\]
\[
P_{g\to gg}(z) = N_C \frac{(1 - z(1 - z))^2}{z(1 - z)},
\]
\[
P_{g\to q\bar{q}}(z) = T_R \{z^2 + (1 - z)^2\},
\]
for the parton branches \(q \to qg, g \to gg,\) and \(g \to q\bar{q}\), respectively. The parameters are given as \(C_F = 4/3, N_C = 3,\) and \(T_R = n_f/2\) with \(n_f = 5\). The parameter \(Q^2\) is the squared momentum transfer of the radiation, defined as \(Q^2 = -t\).

The sum in Eq. (2) is taken over all possible sources of radiation in the picture of parton showers. We can consider a unique source, \(g \to q\bar{q}\) for the final-state quark in the process \(gg \to A + q\), whereas there are two possible sources, \(q \to qg\) and \(g \to q\bar{q}\), that may produce the gluon in \(q\bar{q} \to A + g\). The \(Q^2\) of the branch is different for the two possible branches in the latter case. The orientation of the subsystem \(A\) in its cm frame, for which the non-radiative ME is evaluated, may also be different since the boost and rotation due to the branch may be different. There are two solutions of \(Q^2\) for a given \(p_T\) of the radiation:
\[
Q_i^2 = \left\{ \begin{array}{ll}
\frac{1 - z}{2} \pm \sqrt{\left(\frac{1 - z}{2}\right)^2 - \frac{p_T^2}{s_{A+1}}}
\end{array} \right\} \hat{s}_{A+1}.
\]
These two solutions correspond to the two possible branches that we consider, and the sum of the inverses of the two solutions is
\[
\frac{1}{Q_+^2} + \frac{1}{Q_-^2} = \frac{1 - z}{p_T^2}.
\]

Therefore, since the \(A + 0\)-jet MEs for the two possible branches become identical at the collinear limit, \(p_T^2/\hat{s}_{A+1} \to 0\), the definition in Eq. (2) with Eq. (4) agrees with the definition in the previous paper [20] at the collinear limit in this case.

The LLL subtraction works well for all processes supported in GR@PPA 2.8. The divergences in the \(A + 1\)-jet MEs are properly subtracted, and the remaining MEs are all finite. Though we apply a small \(p_T\) cut, \(p_T > 1\) GeV/c, to the additional parton in the \(A + 1\)-jet processes for numerical stability, the cut effects are negligible because the differential cross sections converge to zero as \(p_T \to 0\) in all the processes.

In order to achieve appropriate matching, we have to be careful about high \(p_T\) behavior, too. Since the radiation effect is separated by the energy scale \(\mu_F\), the scale has to be consistently defined in the 0-jet and 1-jet processes. Namely, we have to assign the same \(\mu_F\) value to an \(A + 1\)-jet event produced by the \(A + 1\)-jet ME and to an \(A + 0\)-jet event having the same topology after the application of a parton shower. The problem is not serious in the case of single weak boson production processes. We can choose a fixed \(\mu_F\), typically, equal to the weak boson mass. On the other hand, there may be many possible choices for diboson production processes. For the 0-jet processes, the scale is frequently defined as
\[
\mu_F^2 = m_V^2 + p_T^2,
\]
where \(m_V^2\) is the average of the squared masses of the produced weak bosons, and \(p_T\) is the transverse momentum of the hard interaction, \(qq' \to VV'\). The definition in Eq. (10) cannot be directly applied to 1-jet processes since the diboson system has a transverse momentum, \(q_T\), due to the radiation of the jet. If we assume that the jet is radiated from one of the initial-state partons, we can define the 0-jet subsystem and define \(\mu_F\) using Eq. (10) in its cm frame. However, this assumption is ambiguous and may be incorrect especially for those events having high \(p_T\) radiation, for which the consistency has
to be most seriously considered. There should not be a "correct" answer to this question. Therefore, we adopt a simple definition in the sample program as one of the possible options, that is

$$\mu_F^2 = m_T^2 + \left| \vec{p}_{T,1} - \vec{p}_{T,2} \right|^2,$$

where $\vec{p}_{T,i}$ represents the transverse momentum vector of a weak boson. Equation (11) can be calculated only from the properties of weak bosons, and is in agreement with Eq. (10) at the limit of $q_T \to 0$.

3. Parton showers

We provide three parton shower (PS) routines in GR@PPA 2.8: a forward-evolution initial-state PS (QCDPS), a backward-evolution initial-state PS (QCDPSb), and a final-state PS (QCDPSf). All of them are based on the Sudakov form factor at the leading order, which is expressed as

$$S(Q_1^2, Q_2^2) = \exp \left[ -\int_{Q_1^2}^{Q_2^2} dQ^2 \int_\epsilon^{1-\epsilon} dz \frac{\alpha_s(Q^2)}{2\pi} \sum_i P_i(z) \right],$$

where $P_i(z)$ are the leading-order splitting functions in Eqs. (5-7), and they are summed over all possible branches. The parameter $\epsilon$ cuts off the divergences in the splitting functions. We set the cutoff to be very small so that physical quantities should not be affected by the choice of this arbitrary parameter; $\epsilon = 10^{-6}$ for the initial state and $10^{-3}$ for the final state as the default. A very small cutoff is applied to the initial state in order to make the total cross section stable at the level of one percent, though kinematical distributions of weak bosons are already stable at $\epsilon = 10^{-3}$.

The PS branches are therefore ordered in $Q^2$. For instance, in QCDPS, when we have a branch at $Q^2 = Q_1^2$, the $Q^2$ of the next branch $Q_2^2$ is determined by solving the equation $S(Q_1^2, Q_2^2) = \eta$, using a random number $\eta$ that is uniformly generated in the range of 0 to 1. $Q_1^2$ is derived from $Q_2^2$ in the same way in QCDPSf. The splitting parameter $z$ is randomly determined in proportion to the relevant splitting functions $P_i(z)$. We set the lower limit of $Q^2$ to be $Q_0^2 = (4.6 \text{ GeV})^2$, and the upper limit at the factorization scale $\mu_F^2$ if there is no other limitation.

3.1. Initial state

The initial-state PS plays a crucial role in the matching method. We use a forward-evolution PS, named QCDPS, as the primary tool for this purpose. The forward evolution in the initial state is in general very inefficient, especially for those interactions which require tight constraints on the parton momenta after the evolution, such as narrow resonance productions, because the final momentum is usually unpredictable in the forward evolution. This problem has been solved by introducing the "x-deterministic" forward-evolution technique [20]. As discussed previously, parton showers can in principle replace the perturbative part of PDFs. QCDPS realizes this concept in practical event generators. When we use QCDPS, a PDF is employed only for setting the initial condition at the lower limit, $Q^2 = Q_0^2$.

There is no ambiguity in the procedure for determining PS branches characterized by two parameters, $Q^2$ and $z$, on the basis of Eq. (12). However, since the parameters are defined in the infinite-momentum frame, the correspondence of the determined parameters to the kinematical variables in a finite-momentum frame is not trivial. We need to introduce a certain model to construct a practical PS. In a previous paper [22], we have pointed out that the definition of the $p_T$ of each PS branch is important in such models, and shown that the parton shower we have developed reproduces experimental data with surprisingly high precision. We apply the same kinematics model in GR@PPA 2.8, where the $p_T$ of each branch is "prefixed" according to the definition

$$p_T^2 = (1 - z)Q^2.$$

\[ \text{The lower limit is increased to 5.0 GeV since the GR@PPA 2.8.1 update.} \]
Since the branch kinematics do not affect the production cross section, this branch model is applied to events that are already unweighted, i.e., they already have an event weight of ±1. Though there is an option to include the kinematical smearing by PS in the cross section integration, this option remains experimental and it is not recommended for use in the present version.

Although the "x-deterministic" forward evolution technique significantly improves the generation efficiency, the application of QCDPS requires a long CPU time, compared to widely used backward-evolution parton showers. Besides, the use of QCDPS requires another constraint. Since the leading-order (LO) Sudakov form factor is used, QCDPS cannot reproduce the evolution in next-to-leading order (NLO) or next-to-next-to-leading order (NNLO) PDFs, nor in the recently proposed modified leading-order (LO*) PDFs [23, 24]. If we use one of such PDFs for the initial condition at $Q^2 = Q_0^2$, we will obtain an incorrect result on the parton distribution at larger energy scales relevant to the interactions of interest. In order to overcome these problems in QCDPS, we also provide a backward-evolution PS named QCDPSb in GR@PPA 2.8. The algorithm for generating PS branches is the same as PYSHOW in PYTHIA [25], the so-called "old model" of the PYTHIA PS. Though PYSHOW is known to give an unsatisfactory softer $p_T$ spectrum in weak boson productions, we can expect QCDPSb to have a performance similar to that of QCDPS because we apply the same kinematics model. However, QCDPSb does not strictly reproduce the evolution in PDFs, even when we compare it with LO PDFs. We suggest that users should consider QCDPSb as a model for simulating initial-state hadronic activities, and check whether the desired properties are effectively reproduced by QCDPSb by comparing the results with those obtained using QCDPS.

3.2. Final state

The final-state PS in GR@PPA 2.8, named QCDPSf, remains experimental. It is implemented mainly for consistency because the LHA user-process interface [26], which is used to pass the event information to other programs, has only one energy scale in each event. When the QCD evolution is simulated down to $Q_0$ in the initial state by QCDPS or QCDPSb, the evolution has to be simulated to the same level also in the final state. We describe the implemented final-state PS in some detail in this section because it is based on a new concept that we have learned in the development of the initial-state PS.

We have learned that consequences from discussions in the infinite-momentum frame, which may affect observable quantities, must be followed as strictly as possible in parton showers in a finite-momentum frame. The effective definition of $p_T$ in each PS branch is most important because it determines the visible $p_T$ of the hard interaction system. Another related consequence is that the evolution should not depend on the ordering of the branches. However, these consequences are not consistent with energy-momentum conservation. PS branches produce non-zero virtuality in at least one of the partons participating in each branch. If the exact energy-momentum conservation is required, this virtuality necessarily affects the kinematics of preceding or subsequent branches; thus, the evolution becomes dependent on the ordering of branches and the originally applied $p_T$ definition is altered. The "$p_T$-prefixed" kinematics model in QCDPS is designed to minimize the effect of this difficulty. However, there may be a kinematics model that can more strictly preserve the consequences in the infinite-momentum frame.

We have to violate energy-momentum conservation in order to realize the independence of PS branches. If we disregard energy conservation, we can consistently determine the momenta of partons participating in a branch on the basis of arguments in the infinite-momentum frame. Energy conservation can be restored by adjusting the overall scale of the momenta after completing the momentum determination for all branches. This adjustment alters the momenta of the color-singlet products, such as weak bosons, as well. However, we expect the alternation to be small and to have an insignificant effect on observable quantities. We have developed QCDPSf on the basis of this concept.

We consider a final-state (time-like) PS branch of a parton having momentum $p$. The branch is characterized by two parameters, $Q^2$ and $z$, as shown in Fig. 1. The two branch products are assumed to be massless. The product that we are now going to study has momentum $k$, with a longitudinal component $k_L$ and a transverse component $k_T$ with respect to the direction of the mother; i.e., $k^2 = k_L^2 + k_T^2$. This branch product is assumed to have a "momentum fraction" $z_k$, which is either $z_k = z$ or $z_k = 1 - z$. 
We define the momentum fraction in an infinite-momentum frame. We consider a boost having a large Lorentz factor of $\tilde{\beta} \approx 1$ along the direction of the mother parton. In order to carry out the Lorentz transformation consistently, we assume that the mother has an invariant mass $Q$, i.e., it has the energy expressed as

$$E^2 = p^2 + Q^2.$$  \hfill (14)

The kinematical variables in the infinite-momentum frame are denoted by tildes, i.e., $\tilde{E} = \tilde{\gamma}(E + \tilde{\beta}p)$ and $\tilde{p} = \tilde{\gamma}(p + \beta E)$ with $\tilde{\gamma}^2 = 1/(1 - \tilde{\beta}^2)$. The transverse momentum of the branch product is given by

$$k_T^2 = z_k (1 - z_k) Q^2 = z(1 - z) Q^2.$$  \hfill (15)

This is invariant against the Lorentz boost. On the other hand, using quantities in the infinite-momentum frame, the longitudinal momentum of the branch product can be expressed as

$$k_L = \tilde{\gamma}(k_L - \tilde{\beta} \tilde{k}) = \tilde{\gamma} \tilde{k}_L \left(1 - \tilde{\beta} \sqrt{1 + k_T^2 / k_L^2}\right).$$  \hfill (16)

Since $k_T^2 / k_L^2 \ll 1$, it must be sufficient to take a first-order expansion of the square root as

$$k_L = \tilde{\gamma} \tilde{k}_L \left\{1 - \tilde{\beta} \left(1 + \frac{k_T^2}{2 k_L^2}\right)\right\}. \hfill (17)$$

Here, we define the momentum fraction $z_k$ as

$$\tilde{k}_L = z_k \tilde{p}$$  \hfill (18)

in the infinite-momentum frame. Since $\tilde{p} = \tilde{\gamma}(p + \tilde{\beta}E)$, $\tilde{k}_L$ can be expressed as $\tilde{k}_L = z_k \tilde{\gamma}(p + \tilde{\beta}E)$. Substituting this relation in Eq. (17) and taking $\tilde{\beta} = 1$, we obtain the definition of the longitudinal momentum as

$$k_L = \frac{z_k(p + E)}{2} - \frac{k_T^2}{2 z_k(p + E)}. \hfill (19)$$

This definition has an undesirable feature that it may produce backward branches ($k_L < 0$) which may form a jet-like structure in the backward direction.

The presence of these backward branches has already been discussed by Kato and Munehisa [27] for the $e^+e^- \rightarrow q\bar{q}$ process. In principle, the QCD evolution for the final-state $q\bar{q}$ system can be completed only by applying a PS to either $q$ or $\bar{q}$ in an infinite-momentum frame ("single cascade"). Soft but not very collinear radiations are boosted backwards in the cm frame, and they reproduce radiations to be generated by the other quark. Though theoretically the radiations become symmetric in the cm frame, the actual implementation of PS may produce a substantial asymmetry. In order to overcome this technical difficulty, Kato and Munehisa proposed a "double cascade" scheme in which, from a study of the $e^+e^- \rightarrow q\bar{q}g$ differential cross section, they derived a condition to consistently separate the radiations. The condition gives a constraint on $z$ in each PS branch and effectively selects forward-going branches in the cm frame. An independent application of a PS to $q$ and $\bar{q}$ with this condition is theoretically equivalent to the "single cascade" in the infinite-momentum frame, and technically it produces symmetric radiations in the cm frame.

From the discussions on the "double cascade" scheme, we learn that the backward branches in Eq. (19) can be interpreted as branches associated with a system compensating the color flow of the
parton of interest, and that it is reasonable to reject these backward branches in an actual implementation of parton showers. Hence, the question is: what is the compensating system in our case? We cannot select any other parton or parton system in the PS since the branches must be independent of the ordering. It would be natural to consider the rest of the whole colliding system as the compensating system because the color flow may be connected even with the remnants of beam collision. By the way, we need not be very sensitive to the detailed definition of the "forward" and "backward" branches because most of the branches are concentrated in very forward and backward regions.

In the present version of QCDPSf, we apply the final-state PS in the cm frame of the hard interaction. We take the momentum in this frame as $p$ in Eq. (19), and retain only those branches having $k_L > 0$. This selection yields the constraint

$$z_k^2 (p + E)^2 > k_T^2.$$  \hfill (20)

Since $k_T$ is defined in Eq. (15), Eq. (20) can be written as

$$z_k > \frac{Q^2}{(p + E)^2 + Q^2}.$$  \hfill (21)

The condition in Eq. (21) has to be satisfied by the two branch products. Therefore, it is effectively a constraint on the lower-momentum product having $z_k < 1/2$. The right-hand side of Eq. (21) is always smaller than 1/2. Thus, Eq. (21) does not give any clear constraint on $Q^2$. However, it strongly limits the branch of low momentum partons. The condition in Eq. (21) gives a constraint of $z_k > 1/2$ at the limit $p/Q \to 0$; thus, no branch is allowed at this limit. The constraint is rapidly relaxed as $p/Q$ increases. The right-hand side of Eq. (21) is smaller than $10^{-2}$ for $p/Q > 3$.

In the actual implementation, the condition in Eq. (21) is required each time a set of $Q^2$ and $z$ is generated, using the momentum $k = \sqrt{k_L^2 + k_T^2}$ tentatively evaluated in the previous branch. If the condition is not satisfied, the branch is discarded and the evolution proceeds by taking the discarded $Q^2$ as the maximum value of the next branch. The evolution starts from the factorization scale $\mu_F$, in principle. If the $p_T$ of the parton with respect to the initial-state direction is smaller than $\mu_F$, the $p_T$ is chosen as the initial value. The evolution is terminated when the newly determined $Q^2$ becomes smaller than $Q_0^2$. The full kinematics of the branches are determined after completing the generation of branches. The branch direction is randomly chosen in azimuth in each branch in order to determine the three-momenta of the products. The momenta of the final-state particles are then adjusted in order to restore the energy conservation. We adjust (decrease) the overall scale of the momenta of all particles that constitute the final state of the hard interaction so that the total invariant mass should match the value before the parton shower is applied. Weak bosons are treated as particles in order to preserve their invariant masses, and the momentum adjustment is carried out such that the decay directions in the cm frame of weak bosons remain unchanged.

The final-state PS can also be applied to the radiations produced by the initial-state PS. If it is turned on, QCDPSf is applied when the kinematics of each initial-state branch is determined, with the maximum $Q^2$ set to the $p_T^2$ of the branch. We assume that the radiation has a momentum equal to $p_T$. The application of QCDPSf produces a non-zero mass of the radiation if it results in additional branches. If any branch is added, the kinematics of the initial-state PS branch is adjusted by taking the produced mass into consideration, keeping the $p_T$ definition unchanged as far as possible. The momenta of branch products are then rotated and boosted so that the total momentum matches the adjusted radiation momentum.

4. How to use

4.1. Distribution package

The program package is distributed as a gzipped tar file named GR@PPA-2.8.tgz, which can be obtained from the GR@PPA Web page\(^3\). The compressed file can be expanded, for instance, by typing

\(^3\)http://atlas.kek.jp/physics/nlo-ug/grappa.html#GRAPPA2.8
on UNIX systems. When the file is expanded, users have a directory named `GR@PPA-2.8` containing the following files and directories:

(miscellaneous files)
- **READMe**: readme file describing how to use this package,
- **VERSION-2.8**: file to show the version number,

(files for setup)
- **config.input**: file to specify the configuration for the setup,
- **Install**: shell script for the installation,
- **config**: shell script to configure the setup,
- **config.perl**: Perl script called by **config**,
- **proc.list**: process list, which is referred to in **config.perl**,

(GR@PPA framework)
- **grckinem**: source files of the framework,
- **basesv5.1**: BASES 5.1 source package with some customization,
- **chanel**: CHANEL source package to define the interaction model,
- **inc**: directory containing common include files,
- **example**: directory to be used for the setup of sample programs,
- **diagrams**: directory containing PS files illustrating typical Feynman diagrams,
- **lib**: directory to store object libraries; initially empty,

(process directories)
- **wjets**: W production processes,
- **zjets**: Z production processes,
- **diboson**: diboson (W+W−, WZ, ZZ) production processes.

### 4.2. Installation

The installation of GR@PPA is easy on Unix/Linux systems on which the `bash` shell and the Perl interpreter are available. Users have to specify the user-dependent configuration for the installation in the file `config.input`. The items to be specified are the choice of a PDF library, compile and archive commands and their options, and the paths to external libraries and include directories. Concerning the PDF library, it is most convenient to choose the built-in CTEQ6 PDF [28] for users who just want to test the included event generators, while it must be better to choose LHAPDF4 [29] if they want to do intensive studies. Users can also choose the classical PDFLIB library5 in CERNLIB, if they want. Fortran and C compilers are necessary for building GR@PPA libraries, while a C++ compiler is required only if users want to use RBOOK6 in sample programs. The external libraries are referred to for building sample programs. Though they can be given later by editing the *Makefile* for sample programs, it is better to specify them in `config.input` in order to avoid problems arising from inconsistency in the compile conditions. Some directory paths are also referred to in the configuration of sample programs.

A PYTHIA 6.4 library7 is necessary if users want to perform the simulations down to the hadron level. When LHAPDF is chosen as the PDF library, it is required to specify the paths to its `bin`, `lib`, and PDF data directories separately because their location depends on the installation of LHAPDF. The CERNLIB libraries are necessary to use HBOOK for histogramming in sample programs, while a ROOT library8 is necessary to use RBOOK. Sample programs require at least one of the two libraries. Unnecessary items can be left blank in `config.input`.

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5[http://projects.hepforge.org/lhapdf/](http://projects.hepforge.org/lhapdf/)
6[http://wwwasd.web.cern.ch/wwwasd/cernlib/](http://wwwasd.web.cern.ch/wwwasd/cernlib/)
7A simple Fortran interface to the ROOT histogramming coded by W. Verkerke (NIKHEF). The source code was copied from the MC@NLO 3.4 package.
8[http://root.cern.ch/](http://root.cern.ch/)
Here, users can delete any process directories which they are not interested in. The installation is done only for the retained processes. If the configuration is given properly in `config.input`, users can complete the installation of the libraries by invoking a shell script as

```
./Install
```

at the top directory of GR@PPA. The script `Install` creates a Makefile by executing the script, `config`, and then compiles the source files to install object libraries in the `lib` directory by invoking the `make` command. It may be better to invoke `./config` and `make` separately, instead of `./Install`, when users encounter a problem. The script `config` invokes `config.perl` to create the Makefile and to make other miscellaneous setups according to the configuration described in `config.input`. Parameters defined in `config.input` are exported as environmental variables to be referred to in a Perl script `config.perl`. This export does not affect the shell environment of users because `config` is executed on a separate shell. The process directories are searched in `config.perl` to add necessary descriptions to the Makefile and some source files in `grckinem` by referring to the process list in `proc.list`. No description is added for those processes for which the corresponding process directories are not found. The installation is confined in the GR@PPA-2.8 directory and nothing is created outside of it. Therefore, users can completely clean up the installation by deleting the GR@PPA-2.8 directory, though some clean-up utilities are provided in the Makefile created by the `Install` script.

### 4.3. Sample programs

Sample programs are provided separately for each process. For instance, those for the Z production processes are placed in the `zjets/example` directory. There are two examples for the matched generation of the Z + 0-jet and the Z + 1-jet processes: `z1j_matched.hbook` and `z1j_matched.root`. The former uses HBOOK for histogramming and the latter uses RBOOK. The produced histograms can be viewed and processed by PAW in CERNLIB in the former and by ROOT in the latter. Moving to one of these directories, one can build and execute the program by invoking the commands

```
./Config
make
./run
```

The sample programs produce an event file using the LhaExt utility. The event data in the produced file can be further processed by PYTHIA [31]. A sample program is provided under the `pythia` directory in each sample. The parameters in PYTHIA are left unchanged from the default in the sample programs, except for \texttt{PARP(67)}= 1.0 and \texttt{PARP(71)}= 1.0 explicitly written in the source program `Pythia.f`. This setting should be applied when one uses the default "old" PS in PYTHIA. One can build and execute the sample program by invoking the above three commands again in the `pythia` directory.

The sample programs are fully relocatable once they are configured by executing the `Config` scripts. Users can copy the sample directory to anywhere within the same file system. The configuration has to be done also in the `pythia` directory before copying, if one wants to apply PYTHIA to the generated events.

Each sample program contains four source files: `grappa.f`, `grcpar.F`, `upevnt.f`, and `upinit.F`. The file `grappa.f` contains the main program of each sample, in which an analysis example is provided together with the necessary steering procedure. Users can modify the execution of GR@PPA by customizing the three files, `grappa.f`, `grcpar.F` and `upinit.F`. Users will not need to edit `upevnt.f` because it contains only one subroutine, `UPEVNT`, which simply calls the GR@PPA steering routine `GRCPYGEN` with `mode` = 0 in the event generation stage. The `usr` directory in each sample program directory is used to store common utility routines and component files to be used to compose some sample program routines.

The file `upinit.F` contains the subroutine `UPINIT`. This routine is called once before starting the event generation. The optimization of the random number generation and the cross section integration

\textsuperscript{9}http://atlas.kek.jp/physics/nlo-ug/grappa.html#LhaExt
Table 1: Parameters to control the matching method and parton showers, which can be set in \texttt{UPINIT}. "D" denotes the default choice in \texttt{GRCPYGEN}.

| Parameter | Description |
|-----------|-------------|
| matching  | switch for the matching method. (D = 0)  
|           | = 0 : no matching method is applied.  
|           | = 1 : the LLL subtraction is applied to 1-jet processes. |
| ishower   | switch for the initial-state PS. (D = 0)  
|           | = 0 : no PS is applied.  
|           | = 1 : QCDPS is applied.  
|           | = 2 : QCDPSb is applied.  
|           | = 3 : QCDPS with a final-state PS for the radiations.  
|           | = 4 : QCDPSb with a final-state PS for the radiations. |
| ishwfin   | switch for the final-state PS. (D = 0)  
|           | = 0 : no PS is applied.  
|           | = 1 : QCDPSf is applied. |

are done by calling \texttt{GRCPYGEN} with \texttt{mode} = 1 at the end of \texttt{UPINIT}. Parameters controlling the matching method and parton showers have to be set in this routine. The allowed choices are listed in Table 1. The recommended combination is \texttt{matching} = 1, \texttt{ishower} = 3 or 4, and \texttt{ishwfin} = 1.

Other parameters which can be set in \texttt{UPINIT} are listed in Table 2. Users have to set basic parameters concerning the beam conditions and the processes to be generated. \texttt{GR@PPA} supports multiple process generation, in which generated events are randomly mixed according to the production cross sections. The processes to generate have to be specified using a parameter \texttt{NPRUP} and an array \texttt{LPRUP} defined in the LHA user-process interface [26]. In order to apply the matching method, we have to set \texttt{NPRUP} = 2 and set the process numbers for the 0-jet and 1-jet processes in \texttt{LPRUP(1)} and \texttt{LPRUP(2)}. Though the appropriate numbers are already set in the sample programs, users can find the process numbers defined in \texttt{GR@PPA in grcpar.F} and \texttt{proc.list}.

The PDF to be used for evaluating the cross sections and/or for setting the initial condition of QCDPS also has to be specified in \texttt{UPINIT}. Users have to modify appropriate lines according to the choice of the library in \texttt{config.input}. Users do not need to care about those parts irrelevant to the choice because they are discarded in the preprocessing before compilation. The setting is simplified for LHAPDF because a symbolic link, \texttt{PDFsets}, pointing to the PDF data directory given in \texttt{config.input} is created by \texttt{Config} in the sample program directory when LHAPDF is chosen. Refer to the manual of the chosen PDF library for more details.

In addition, users can choose decay modes of weak bosons and customize kinematical cuts in \texttt{UPINIT}. However, it should be noted that the cuts are applied to the quantities before applying the parton showers, and cuts on the weak-boson decay products never dramatically improve the generation efficiency. Moreover, any change concerning the jets may affect the performance of the matching method. Therefore, we recommend that those parameters relevant to the kinematical cuts should be kept unchanged from the preset values in \texttt{UPINIT} of the sample programs. The default setting corresponds to the no-cut condition. If users want to apply any cuts relevant to a restricted detection condition, it would be better to apply them in the analysis part in \texttt{grappa.f} and \texttt{Pythia.f}. Refer to the previous reports [18, 19] for more details of the parameters in Table 2.

The file \texttt{grcpar.F} contains routines which are frequently called during the execution of \texttt{GR@PPA}. This file is created by \texttt{config.perl} and copied to a sample program directory by \texttt{Config} in each sample. The subroutine \texttt{GRCPAR} in this file defines parameters depending on the process. The parameters that users are allowed to change are listed in Table 3. These parameters have to be given for every process. For single weak-boson production processes, it must be best to choose fixed values for the renormalization scale and the factorization scale by setting \texttt{ICOUP} = 5 and \texttt{IFACT} = 5. The parameters \texttt{GRCQ} and \texttt{GRCFAQ} have to be given explicitly in \texttt{GRCPAR} in this case. On the other hand, there may be many possible definitions for diboson production processes. We have chosen \texttt{ICOUP} = 6 and \texttt{IFACT} = 6, and as an example we have defined the scales according to Eq. (11) in the subroutine \texttt{GRCUSRSETQ}, in which we take an identical definition for the two scales.
Table 2: Other GR@PPA parameters which users can set in UPINIT. "D" denotes the default choice.

| Parameter          | Description                                                                                                                                 |
|--------------------|---------------------------------------------------------------------------------------------------------------------------------------------|
| CBEAM              | "PP" for pp collisions and "PAP" for pp collisions. (D = "PP")                                                                |
| GRCECM             | CM energy of the beam collision in GeV. (D = 14000.0)                                                                                      |
| GPTCUT             | Minimum $p_T$ in GeV for jets, except for those from weak-boson decays. (D = 20.0)                                                          |
| GRCPTCUT(8)        | Minimum $p_T$ in GeV for each final state particle. (D = 0.0)                                                                               |
| GETACUT            | Largest pseudorapidity in the absolute value for jets, except for those from weak boson decays. (D = 3.0)                                    |
| GRCETACUT(8)       | Largest pseudorapidity cut for each final state particle. (D = 10.0)                                                                      |
| GRCONCUT           | Minimum separation cut in $\Delta R = \sqrt{\Delta \phi^2 + \Delta \eta^2}$ for jets except for those from weak boson decays. (D = 0.4)       |
| GRCRCONCUT(8)      | Minimum separation cut in $\Delta R = \sqrt{\Delta \phi^2 + \Delta \eta^2}$ for each final state particle. (D = 0.0)                      |
| IGWMOD(20)         | Switch for the decay modes of W: 1 to activate and 0 to deactivate.                                                                       |
| IGZMOD(16)         | Switch for the decay modes of Z: 1 to activate and 0 to deactivate.                                                                       |
| IWIDCOR            | Switch for the decay widths correction for W and Z. (D = 1)                                                                               |
|                    | = 0: no correction; i.e., the lowest-order values are used.                                                                             |
|                    | = 1: corrected to match the values given in SETMAS and branching ratios in GRCWBR and GRCZBR.                                           |
| GRCWBR(20)         | Branching ratios of W; valid only when IWIDCOR = 2.                                                                                    |
| GRCZBR(16)         | Branching ratios of Z; valid only when IWIDCOR = 2.                                                                                     |
| IGJFLV(7)          | Switch for the jet flavors to be produced; 1 to activate and 0 to deactivate. These are irrelevant to those from weak boson decays.        |
| IGRCGEF            | Switch for the $Z-\gamma$ interference effect in $Z$ production processes; turned on if set to 1 and ignored if 0. (D=1)                    |
| GRCKKM(3,3)        | Squared CKM matrix.                                                                                                                         |
| IGAUGE             | Choice of the scheme to determine the electroweak parameters. The $G_\mu$ scheme is used as the default. (D=1)                             |

The parameter NCALL defines the number of sampling points in each step of the BASES iteration. The numbers preset in the sample programs are optimized for the generation in the LHC condition. Users need to optimize them for other conditions. The guiding principle is to achieve a statistical accuracy of 0.2% in the first stage of BASES and to eliminate any unexpected jump (increase) in accuracy in each iteration.

In order to implement QCDPS, a small modification has been applied to the BASES library included in the package. Because the random numbers in QCDPS are not handled by BASES, the application of QCDPS appears as a random distribution of the event weight for each sampling point. Therefore, the application of QCDPS degrades the statistical accuracy of BASES. In order to compensate for this degradation, the number of sampling points that is originally determined by NCALL is multiplied by a factor, $mfbps$, when QCDPS is applied. In the first stage of BASES, the distribution of random numbers in BASES is optimized by adjusting a multi-dimensional grid for the sampling. QCDPS is not applied there even if its use is required. Instead, the given PDF is directly referred to since QCDPS has nothing to do with this optimization. QCDPS is applied and the multiplication to the number of sampling points is activated in the second stage, where the maximum event weight...
Table 3: Parameters which can be set in GRCPAR. “D” denotes the default choice.

| Parameter | Description |
|-----------|-------------|
| ICOUP     | Choice of the renormalization scale. |
|           | = 1: $\sqrt{s}$ of the hard interaction. |
|           | = 2: average of squared transverse mass ($\langle m_T^2 \rangle$). |
|           | = 3: total squared transverse mass ($\sum m_T^2$). |
|           | = 4: maximum squared transverse mass ($\max m_T^2$). |
|           | = 5: fixed value. Set GRCQ in GeV. |
|           | = 6: user defined scale. Set GRCQ in the subroutine GRCUSRSETQ. |
| IFACT    | Choice of the factorization scale. (D=0) |
|           | The definition is the same as ICOUP. |
|           | If IFACT = 0, the same value as the renormalization scale is used. |
|           | In case IFACT = 5 or 6, set GRCFAQ in GeV. |
| GRCFILE  | Output file name for the BASES integration. |
| IBSWRT   | Mode selection for BASES integration: = 0 for calling the BASES integration, and 1 for skipping. (D=0) |
|           | If IBSWRT = 1, the file defined in GRCFILE is used. |
| NCALL    | Number of sampling points in each step of the iterative grid optimization in BASES. |
| mfabsps  | Multiplication factor to the number of sampling points in BASES, to be applied in the 2nd stage when QCDPS is used. |
| INPFL    | Number of flavors used in the coupling calculation and PDF. (D=5) |

is determined in each hypercube by increasing the statistics. Therefore, when the use of QCDPS is required, the second stage of BASES consumes longer CPU time than the first stage. Users can check the performance of QCDPS for the longitudinal QCD evolution by comparing the cross section results from the two stages of BASES. A statistical accuracy of about 0.5% would be enough in the second stage of BASES when QCDPS is applied. The factor mfabsps should be increased if an unnatural structure is observed in the rapidity distribution of the hard interaction system.

The subroutine SETMAS in grcpar.F defines basic properties of particles and interactions, such as masses, decay widths, and coupling constants. Users may change these parameters if they want. In addition, users can apply their own cuts to the events to be generated by customizing the subroutine GRCUSRSET in grcpar.F. They can define any cuts which cannot be accomplished by using the parameters in Table 2. However, as we have already mentioned, it is not recommended to apply any cut relevant to jets in this version because they may deteriorate the performance of the jet matching. It is safe to apply those cuts relevant to Lorentz-invariant quantities of weak bosons. For instance, it is necessary to define, at least, the lower limit of the $Z$-boson mass when the interference with the photon exchange is turned on ($\text{IGRCGEF} = 1$). The subroutine GRCUSRSETQ defines the energy scales, GRCQ and GRCFAQ, when ICOUP = 6 and/or IFACT = 6. In GRCUSRSET and GRCUSRSETQ, users can access the internal event information through the arrays, PGR and PLGRC. Some utility functions are also available there. Refer to the comments in grcpar.F for details. The internal particle numbering can be found in Feynman diagrams illustrated in the figures in the diagram directory under the top directory. Please ignore the subroutine grclabcut because its implementation is still in an experimental phase.

The simulation can proceed to the hadron level by applying PYTHIA to the generated events. The sample programs that we provide in the pythia directory employ the so-called “old” PS as the default. If we change it to the ”new” PS by setting, for instance, MSTP(81) = 21, PYTHIA issues many warnings and sometimes the execution hangs. This would not be a problem caused by the combined use of PYTHIA and GR@PPA because similar warnings frequently appear even in stand-alone event generations. Though the reason is yet to be intensively investigated, the problem becomes less severe if we turn off parton showers in GR@PPA. As we have shown in a previous report [21], the PYTHIA ”new” PS shows transverse activities very similar to QCDPS. If users are eager to use
the "new" PS, it may be better to turn off all parton showers in GR@PPA by setting matching = 1, ishower = 0, and ishwin = 0, and pass the generated hard interaction events directly to PYTHIA. In this case, the given factorization scale value (GRCFAQ) is passed to PYTHIA as the energy scale in the LHA user-process interface. When the "new" PS is applied, users have to comment out the line setting PARP(67) in the main program Pythia.f.

Users may encounter another problem in the PYTHIA simulation when they turn on the parton showers in GR@PPA. The parton showers sometimes generate very large number of particles, and PYTHIA issues a warning and sometimes generate severe errors when the total number of particles passed through the LHA interface exceeds 80, even though up to 500 particles are allowed in the LHA interface. Because it is very rare for the total number of particles to exceed 80 even in the LHC condition with the full implementation of parton showers in GR@PPA, it may be better to skip such events in the subroutine UPEVNT included in Pythia.f.

The HERWIG PS [32, 33] seems to be incompatible with the parton showers in the present version of GR@PPA. The execution is immediately terminated with an error. Contrary to the case of the PYTHIA "new" PS, it would not be a good choice to fully replace the GR@PPA parton showers with the HERWIG PS because there is a marked difference between them [22]. Therefore, we do not officially support the combination with HERWIG in the present version.

5. Simulation results

In this section, we show some results to verify the performance of GR@PPA. The presented results are all obtained with the default setting in the sample programs, except for the number of events to generate. Concerning the decay modes of the weak bosons, only the modes $W \to e\nu$ and $Z \to \mu^+\mu^-$ are activated. The presented quantities are extracted after applying the PYTHIA simulation to the generated events.

5.1. W and Z productions

We have shown in a previous report [22] that the simulation employing our matching method and QCDPS reproduces the $p_T$ spectrum of the $Z$ bosons measured at Tevatron experiments [34, 35, 36] with a very good precision. Figure 2 shows the result of a simulation on the same quantity, in which, together with QCDPS, QCDPSf is applied to partons from the hard interaction and those radiated in QCDPS, as described in the previous section. The factorization scale ($\mu_F$) and the renormalization scale ($\mu_R$) are taken to be equal to the $Z$-boson mass (91.17 GeV/$c^2$), and the built-in CTEQ6L1 is used for PDF. PYTHIA 6.421 is applied to the generated events in order to add simulations at lower energy scales, such as the primordial $k_T$ effect, hadronization, and decays. Though this PYTHIA version is slightly newer than that used in the previous study, no significant change is observed in the quantities that we are currently interested in. Together with the summed spectrum, the contributions from the $Z + 0$-jet process and the LLL-subtracted $Z + 1$-jet process are separately shown in the figure. We can see that the two components are smoothly combined. The $Z + 0$-jet events are strongly suppressed at $p_T > \mu_F$ and the spectrum is totally determined by the $Z + 1$-jet events at high $p_T$. The LLL subtraction does not sharply suppress the $Z + 1$-jet events at $p_T \lesssim \mu_F$ because non-logarithmic contributions are comparable around $Q^2 = \mu^2$. We can see that the $Z + 0$-jet component overwhelms the remaining non-logarithmic $Z + 1$-jet component at low $p_T$, and the non-logarithmic component converges to zero as $p_T \to 0$. Though the summed spectrum is a little bit softer compared to the previous result because of the application of QCDPSf, the simulation is still in good agreement with the measurements.

Figure 3 shows the $Z$-boson $p_T$ spectrum expected for the LHC condition, proton-proton collisions at a cm energy of 14 TeV, with the same choice of $\mu_F$, $\mu_R$ and PDF. The GR@PPA simulation is compared with those from PYTHIA 6.421 [31] and HERWIG 6.510 [32, 33]. The compared PYTHIA simulation employs its built-in event generator with the "new" PS model, and a primordial $k_T$ effect is added to the HERWIG simulation, as described in our previous report [22]. The $Z$-boson invariant mass is required to be greater than 60 GeV/$c^2$ in all simulations. The overall tendencies are the same as those observed for the simulations in the Tevatron condition [22]. The GR@PPA simulation and the PYTHIA new-PS simulation are nearly identical in the whole $p_T$ range.
Figure 2: $p_T$ spectrum of $Z$ bosons at Tevatron Run 1, $p\bar{p}$ collisions at a cm energy of 1.8 TeV. The GR@PPA simulation (histograms) is compared with the measurements by CDF [34] (circles) and D0 [35] (triangles). Together with a result covering the $p_T$ range up to 200 GeV/c (a), a result to cover the range up to 20 GeV/c (b) is presented to show the low-$p_T$ behavior. In addition to the summed spectrum (solid), the spectra of events from the $Z + 0$-jet (dashed) and $Z + 1$-jet (dotted) processes are separately shown for the GR@PPA simulation in (a).
Figure 3: $p_T$ spectrum of $Z$ bosons at LHC, $pp$ collisions at a cm energy of 14 TeV. The GR@PPA simulation (solid) is compared with the predictions from PYTHIA 6.421 (dashed) and HERWIG 6.510 (dotted).
The rapidity distribution of $Z$ bosons is predominantly determined by the parton momentum distribution inside protons. The distributions from the simulation using QCDPS and that using QCDPSb are compared in Fig. 4. The longitudinal momentum evolution is simulated by the parton shower in the former, while the evolution in the PDF is directly referred to in the latter. The built-in CTEQ6L1 is used for the PDF, and the $Z$ bosons are tagged by $Z \rightarrow e^+e^-$ decays with $m_{e^+e^-} \geq 60$ GeV/$c^2$ in both simulations. The two simulations are in good agreement. Note that the absolute values of the differential cross section are compared in Fig. 4. The agreement shows that the evolution by the parton shower in QCDPS satisfactorily reproduces the analytical evolution in CTEQ6L1.

Hereafter, we present some results concerning $W$ boson production in the LHC condition, where we set the renormalization and factorization scales to the $W$-boson mass (80.42 GeV), and we use the built-in CTEQ6L1 for the PDF. The simulation results are compared with the predictions from PYTHIA. The PYTHIA simulation employs the ”new” PS model and uses CTEQ6L1 in LHAPDF for the PDF with the help of LHAGLUE\textsuperscript{10}. We expect that this PYTHIA simulation is at the same level as GR@PPA, except for the overall normalization. PYTHIA does not add the $W + 1$-jet contribution to the total cross section. Hence, the results are compared in the form of the relative shape of distributions.

The rapidity distribution is separately shown in Fig. 5 for $W^+$ and $W^-$ together with the summed distribution. Figure 6 shows the pseudo-rapidity ($\eta$) distributions of the $e^+$ and $e^-$ from $W^+$ and $W^-$ decays, respectively. The summed $p_T$ distribution is shown in Fig. 7. The contributions from the $W + 0$-jet and $W + 1$-jet processes are also separately shown in Fig. 7. We can see that the tail of the distribution extends to a very high $p_T$ region owing to the existence of hard radiations in the $W + 1$-jet process. The distribution is slightly smeared due to photon radiations because we have taken final stable electrons in the PYTHIA event record. Figure 8 shows the transverse mass distribution.

\textsuperscript{10}See the LHAPDF online manual at \url{http://projects.hepforge.org/lhapdf/manual#tth_sEc3.2}.
Figure 5: Rapidity distribution of $W$ bosons at LHC. The GR@PPA simulation (plots) is compared with the PYTHIA simulation (histograms). CTEQ6L1 is used for PDF in both simulations.

Figure 6: Pseudo-rapidity distribution of electrons from $W$-boson decays at LHC. The GR@PPA simulation (plots) is compared with the PYTHIA simulation (histograms). CTEQ6L1 is used for PDF in both simulations.
Figure 7: $p_T$ distribution of electrons from $W$-boson decays at LHC. The GR@PPA simulation (circles) is compared with the PYTHIA simulation employing the "new" PS model (histogram). The contributions from the $W + 0$-jet and the LLL-subtracted $W + 1$-jet processes are separately shown as well for the GR@PPA simulation.

Figure 8: Transverse mass ($m_T$) distribution of $W$ bosons at LHC, where $m_T$ is calculated from the momenta of the products from electronic decays, $W \rightarrow e\nu$. The GR@PPA simulation (circles) is compared with the PYTHIA simulation (histogram).
Table 4: Benchmark cross sections in pb for $VV + N$ jets processes, correcting Table 4 in the previous report [19].

|          | Tevatron Run II |          |          | LHC       |          |          |
|----------|----------------|----------|----------|-----------|----------|----------|
| $N$ jets | $W^+W^-$       | $WZ$     | $ZZ$     | $W^+W^-$  | $WZ$     | $ZZ$     |
| 0        | 7.06(1)×$10^{-2}$ | 6.10(2)×$10^{-3}$ | 7.71(1)×$10^{-4}$ | 4.91(1)×$10^{-1}$ | 4.66(2)×$10^{-2}$ | 5.40(1)×$10^{-3}$ |
| 1        | 1.769(3)×$10^{-2}$ | 1.608(4)×$10^{-3}$ | 1.860(3)×$10^{-4}$ | 3.275(7)×$10^{-1}$ | 4.57(1)×$10^{-2}$ | 3.031(6)×$10^{-3}$ |

Here, we define the transverse mass ($m_T$) as

$$m_T^2 = 2 \left( p_T^{(e)} p_T^{(\nu)} - \vec{p}_T^{(e)} \cdot \vec{p}_T^{(\nu)} \right), \quad (22)$$

where $\vec{p}_T^{(e)}$ and $\vec{p}_T^{(\nu)}$ denote the transverse momentum vector of the electron and neutrino, respectively, with $p_T^{(e)}$ and $p_T^{(\nu)}$ representing their absolute values. The tail to large $m_T$ values is an off-shell effect of $W$ bosons. The measurement of this tail provides us with an opportunity to directly measure the $W$-boson decay width.

The results from GR@PPA and PYTHIA are in good agreements in all distributions shown in Figs. 5-8. The agreement is not trivial because there are many differences in the simulation technique, not only in the matching method but also in the treatment of $W$ bosons. PYTHIA generates $W$ bosons based on the on-shell approximation. The finite decay width and asymmetry in the decay angle are attached afterwards. On the other hand, the off-shell effects and decay dynamics are included in the matrix elements in the GR@PPA simulation. It should also be noted that GR@PPA employs only the $u\bar{d} \rightarrow W^+$ interaction as the base process. The $W^-$ production and other flavor contributions are derived by applying the charge conjugation and flavor exchanges to the base process. The agreement that we can see in Figs. 5-8 implies that these techniques are properly implemented in both simulations.

5.2. Diboson productions

Some bugs have been identified in the diboson production processes included in the previous version of GR@PPA, and have been fixed for the $VV + 0$-jet and $VV + 1$-jet production processes included in the present version. As a result, some numbers presented in Table 4 in the previous report [19] were incorrect. The corrected numbers are presented in Table 4. Hereafter, we show some results on diboson production processes. Figure 9 shows the $p_T$ distribution of the diboson system in the $W^+W^-$ production process at LHC, $pp$ collisions at 14 TeV. Together with the summed distribution (solid circles), the contribution from the 0-jet (open circles) and the LLL-subtracted 1-jet (open squares) processes are shown separately. The two simulations are smoothly combined also in this process. The GR@PPA result is compared with the results of MC@NLO 3.4.2 [11] and PYTHIA 6.421 [31]. The MC@NLO result has been obtained with the mode $\text{IL1 = IL2 = 1}$. Thus, we can expect the decay width and spin correlations to be simulated properly. The PYTHIA result has been obtained using its built-in generator and the "new" PS model, as in the case of the previous results for single weak-boson productions. The effects of the decay width and spin correlations are simulated also in PYTHIA.

We find a reasonable agreement between the GR@PPA and MC@NLO results. Note that the absolute values of the differential cross section are compared, and not the relative shapes. The overall difference that we can see in the figure can be attributed to the lack of non-divergent corrections at the next-to-leading order (NLO) in GR@PPA. A substantial difference at low $p_T$ must be due to the difference between PYTHIA and HERWIG, employed for small-$Q^2$ simulations in GR@PPA and MC@NLO, respectively. The PYTHIA built-in generator shows a significantly small value of the total cross section because it does not include the 1-jet cross section. In addition, PYTHIA shows a high-$p_T$ behavior apparently different from that of GR@PPA and MC@NLO because hard radiations are simulated with an extrapolation of the collinear approximation.
Figure 9: $p_T$ distribution of the diboson system in the $W^+W^-$ production process at LHC. The results are shown in the absolute values of the cross section. The GR@PPA result is separately shown for the 0-jet (open circles) and LLL-subtracted 1-jet (open squares) contributions together with the summed result (filled circles). The result is compared with the predictions from MC@NLO (solid histogram) and PYTHIA (dashed histogram).

Figure 10: Invariant mass distribution of the diboson system in the $W^+W^-$ production process at LHC. The predictions from GR@PPA (filled circles), MC@NLO (solid histogram), and PYTHIA (dashed histogram) are compared. The results are normalized to the total yield.
Figure 11: Relative shape of the azimuthal opening angle (\(\Delta \phi\)) distribution between the two decay electrons from the \(W^+W^-\) pair to be produced at LHC via known weak interactions. The GR@PPA prediction (plot) is compared with the predictions from MC@NLO (solid histogram) and PYTHIA (dashed histogram).

Figure 10 shows the relative shape of the invariant mass spectrum of the \(W^+W^-\) system. The results of GR@PPA, MC@NLO, and PYTHIA are in very good agreement with each other, except for a small peak below the threshold. This peak corresponds to the \(Z\) resonance in which \(Z\) decays to a highly virtual \(W\) boson pair; it is visible in the GR@PPA and PYTHIA results, whereas it is absent in the MC@NLO result. The peak vanishes if we require a substantial \(p_T\) value for the decay electrons, e.g., \(p_T > 20\) GeV/c.

The azimuthal correlation between the two decay leptons is frequently used as a quantity to distinguish Higgs boson decays to \(W^+W^-\) from non-resonant weak interactions that produce them. The predictions on the opening angle (\(\Delta \phi\)) distribution from GR@PPA, MC@NLO, and PYTHIA are compared in Fig. 11. We can see a substantial difference between the three predictions, even though the overall tendencies are identical. This difference reflects the difference in the \(p_T\) spectrum of the diboson system. Therefore, a detailed understanding/tuning of the \(p_T\) spectrum will be crucial to identify the possible small signal originating from the Higgs boson production.

The \(p_T\) spectrum and the invariant mass spectrum of the diboson system are presented for the \(W^+W^-\), \(ZW\), and \(ZZ\) production processes at LHC in Figs. 12 and 13, respectively. The GR@PPA predictions are compared with those from MC@NLO 3.4.2 and PYTHIA 6.421 in the figures. Here, we present the MC@NLO results in the mode of \(IL1 = IL2 = 7\), in which MC@NLO produces \(W\) and \(Z\) bosons as on-shell particles having decay widths equal to zero. The mode \(IL1 = IL2 = 1\) used for the \(W^+W^-\) production is not available for the \(ZW\) and \(ZZ\) productions\(^{11}\). As a result of the zero decay widths, MC@NLO cannot produce diboson pairs having invariant masses below the threshold. The behaviors of the three simulations are similar to those previously observed for the \(W^+W^-\) production in the \(p_T\) spectrum. At present, we do not identify the reason of a substantial difference between GR@PPA and PYTHIA in the \(ZW\) mass spectrum below the threshold.

As a result of the on-shell approximation, the spin information is not propagated to the decay kinematics in these MC@NLO simulations. Although this effect is small in observable quantities, it may affect detailed studies on the production mechanism. As an example, in Fig. 14, we show the

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\(^{11}\)It has been announced that the spin correlation is included in \(ZW\) production since the version 4.0 of MC@NLO.
Figure 12: $p_T$ spectrum of the diboson system for the $W^+W^-$, $ZW$, and $ZZ$ production processes at LHC. The GR@PPA results (plots) are compared with those of MC@NLO (solid histograms) and PYTHIA (dashed histograms). The MC@NLO simulations were carried out in the mode IL1 = IL2 = 7.

Figure 13: Relative shape of the diboson invariant mass spectrum for the $W^+W^-$, $ZW$, and $ZZ$ production processes at LHC. The GR@PPA results (plots) are compared with those of MC@NLO (solid histograms) and PYTHIA (dashed histograms). The MC@NLO simulations were carried out in the mode IL1 = IL2 = 7.
distribution of the difference between the pseudorapidity of the electron and the rapidity of its parent $W$ boson in the $ZW$ production process at Tevatron. We can see an apparent difference between the distributions for $e^+$ and $e^-$ in the GR@PPA simulation result, whereas there is no such difference in the MC@NLO simulation. Though it is not shown here, the asymmetry observed in the GR@PPA simulation is in good agreement with the result of the PYTHIA simulation.

6. Practical performance

The portability of the program has been tested on some recent Linux systems: Scientific Linux CERN (SLC) 4 and 5, Fedora 10 and 13, and Ubuntu 9. We have successfully executed the sample programs on these systems. Hence, the compilers that we have tested are gcc version 3 and 4 up to 4.4. The Fortran compilers are g77 in gcc version 3 and gfortran in gcc version 4. The difficulties that we have encountered are mostly related to the external libraries. Although there is no difficulty in the use of g77, we need to add the `-fsecond-underscore` option to the Fortran compilation when we use RBOOK/ROOT together with gfortran. In addition, some codes in PYTHIA are incompatible with the simultaneous use of RBOOK/ROOT and gfortran. Necessary modifications for the use of gfortran are described in the file named `config.input-gcc4` in the GR@PPA 2.8 package. In general, we have to be careful about the incompatibility between minor versions of gcc4. It is better for users to compile external libraries by themselves in the same environment as GR@PPA, unless the libraries are provided as standard packages of the distribution. Though this is independent of GR@PPA, we need to replace `/bin/sh` in the setup scripts of LHAPDF with `/bin/bash` when we install it on Ubuntu because `/bin/sh` is a symbolic link to `/bin/bash`. In addition, when we install LHAPDF on a computer having relatively small memory, it is better to add the `--enable-low-memory` option in the configuration because LHAPDF requires a memory larger than 1 GB if it is installed without this option.

The practical performance of GR@PPA 2.8 was intensively tested on the SLC4 operating system installed on an Intel Xeon 5160 CPU operated at a clock speed of 3.0 GHz. The compiler was gcc version 3.4, i.e., g77 for Fortran. Though the operating system was 64 bit, all the tests were carried out with the 32-bit mode, i.e., the `-m32` option was used in the compilation. The used external
Table 5: CPU time consumed by GR@PPA 2.8 on a 3.0-GHz Intel Xeon processor with the SLC4 operating system. The time required for the cross-section integration by BASES and that for the event generation by SPRING are separately presented. The former is in the unit of minutes and the latter is in seconds for generating every 1k events. Results are presented for two choices of parton showers (PS). If QCDPS is used for simulating the initial-state parton shower, it is applied in the second stage of BASES for the integration, while no PS is applied during the integration if QCDPSb is chosen.

|                | QCDPS             | QCDPSb            |
|----------------|-------------------|-------------------|
|                | integ (min) gen (sec/1k) | integ (min) gen (sec/1k) |
| W              | 9.9 23            | 0.72 4.2          |
| Z              | 9.3 18            | 0.73 4.2          |
| W+W−           | 350 260          | 120 27            |
| ZW             | 200 190          | 75 16             |
| ZZ             | 220 96           | 89 12             |

Libraries are LHAPDF 5.7.0, CERNLIB 2006, and ROOT 5.22.00. CERNLIB was installed from a standard package repository, and the binary package of ROOT was downloaded from the official Web page. LHAPDF was locally compiled from the source distribution.

The size of the executable module, grappa, is 1.6 MB for the sample programs using HBOOK, and 1.2 MB for those using RBOOK, when all the processes are retained. The memory size required for the execution is about 6.6 MB with HBOOK and 30 MB with RBOOK when the built-in CTEQ6 PDF is used. If LHAPDF is used instead, the memory size is increased to 1.5 GB in both cases. However, if we use LHAPDF built with the \(--\text{enable-low-memory}\) option, the size is reduced to 150 MB and 170 MB, respectively. The program size can be reduced by removing unnecessary processes. Actually, if we retain only the \wjets\ directory, the total size of the package is reduced from the original size of 5.1 MB to 3.0 MB. The size of grappa is also reduced to 0.96 MB using HBOOK and to 0.57 MB using RBOOK. However, the reduction in the memory size is small, at most 10\%, even when the built-in CTEQ6 is used for the PDF.

The CPU time consumed for the execution of GR@PPA 2.8 is summarized in Table 5. The program was executed with the default settings in the sample programs for the LHC condition in the results denoted by QCDPS. The results are separately presented for the integration by BASES and the event generation by SPRING. In general, the BASES integration time is dominated by that for the 1-jet processes, and by the second stage where QCDPS is applied. The time consumption when QCDPSb (ishower = 4) is used instead of QCDPS (ishower = 3) is also presented in Table 5, in which parton showers are not applied in the integration; instead, PDF is used to derive the parton distribution at large \(Q^2\). The integration time is significantly reduced by this change, as expected. The event generation time is predominantly determined by that for the 0-jet processes because the 1-jet cross section is far smaller. The generation time is presented in seconds consumed for the generation of 1k events. The actual running time increases linearly according to the number of events that users require.

7. Summary

We have described a new release of the GR@PPA event generator package, GR@PPA 2.8, for proton-proton and proton-antiproton collisions. This release supports an initial-state jet matching method, the limited leading-log (LLL) subtraction, that we have proposed. The matching method can be applied to single \(W\) and \(Z\) production processes and diboson (\(W^+W^-, ZW\) and \(ZZ\)) production processes. Custom-made parton shower (PS) programs are included in the package in order to ensure satisfactory performance of the matching method. Though the used matrix elements remain at the tree level, we can reproduce the recoil effects of QCD radiations in the entire phase space of the weak-boson system by combining the "0-jet" and "1-jet" processes with the help of the matching method. The decay widths of weak bosons and the spin effects in the decay products are exactly simulated at the tree level because the decays are included in the matrix elements, as in the previous versions.

The event generators support the use of the LHAPDF library for evaluating the parton distribution function (PDF) inside the proton and antiproton. In addition, a built-in PDF is provided for tests. The classical PDFLIB library can also be used. Though we can only use leading-order PDFs when a
forward-evolution PS for the initial state (QCDPS) is chosen, no such restriction exists if we choose a backward-evolution PS (QCDPSb). A PS for the final state (QCDPSf) based on a new concept is also provided for completeness. The generated events are stored in a file using the LhaExt utility, a set of simple I/O routines for the LHA user-process interface. The event records written in the file can be fed to PYTHIA in order to perform simulations down to the hadron level. The parton showers in GR@PPA can be turned off if users want to apply them in external libraries. The "new" PS model in PYTHIA exhibits a performance nearly identical to QCDPS and QCDPSb.

The program is distributed as an all-in-one package, including process-dependent routines together with the event generator framework. Sample programs are provided for each process. The installation and execution of sample programs are easy in UNIX/Linux systems. The installation of the GR@PPA libraries can be completed in one action. Users are allowed to remove processes in which they are not interested before the installation. The procedure for building sample programs consists of two steps: a configuration and the compile/link. Once it is configured, the sample programs become relocatable. Users can copy the program directory to any place in order to customize it for their own studies.

We provide two sample programs for each process. The difference between the two programs is in the histogramming tool only. HBOOK in CERNLIB is used in one of the two programs, while RBOOK, a simple Fortran interface to ROOT, is used in the other. Therefore, in order to test the sample programs, users have to prepare the CERNLIB library or the ROOT library in addition to PYTHIA and LHAPDF libraries according to their preference. The produced histogram files can be manipulated by PAW in the former and by ROOT in the latter.

The performance of the event generators in GR@PPA 2.8 has been tested by comparing the results with those of PYTHIA and MC@NLO. In general, the results are in excellent agreement. Most of the observed differences can be understood as the effects of different implementations and approximations in these programs. We can, at least, conclude that there is no apparent mistake in GR@PPA 2.8 and the other programs that we have compared.

The portability of GR@PPA 2.8 has been tested on several recent Linux systems. Problems which users may encounter are mostly related to compatibility with external libraries. Users need to be careful about the consistency in the compiler version and compile options. The program size and execution time have also been studied extensively. The program size is predominantly determined by external libraries which users select, while the execution time is nearly independent of them. The application of QCDPS consumes much CPU time, not only in integration but also in event generation. The use of QCDPSb reduces them, as expected.

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