HELAC-Onia 2.0: an upgraded matrix-element and event generator for heavy quarkonium physics

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ABSTRACT

We present an upgraded version (denoted as version 2.0) of the program HELAC-Onia for the automated computation of heavy-quarkonium helicity amplitudes within non-relativistic QCD framework. The new code has been designed to include many new and useful features for practical phenomenological simulations. It is designed for job submissions under cluster environment for parallel computations via Python scripts. We have interfaced HELAC-Onia to the parton shower Monte Carlo programs Pythia 8 and QEDPS to take into account the parton-shower effects. Moreover, the decay module guarantees that the program can perform the spin-entangled (cascade-) decay of heavy quarkonium after its generation. We have also implemented a reweighting method to automatically estimate the uncertainties from renormalization and/or factorization scales as well as parton-distribution functions to weighted or unweighted events. A further update is the possibility to generate one-dimensional or two-dimensional plots encoded in the analysis files on the fly. Some dedicated examples are given at the end of the writeup.
**PROGRAM SUMMARY**

*Program title:* HELAC-Onia 2.0.

*Program obtainable from:* [http://helac-phegas.web.cern.ch/helac-phegas](http://helac-phegas.web.cern.ch/helac-phegas)

*Licensing provisions:* none

*Operating system under which the program has been tested:* Unix-like platform.

*Programming language:* Python, Fortran 77, Fortran 90, C++

*Keywords:* heavy quarkonium, NRQCD, Monte Carlo simulation

*Nature of physical problem:* Heavy quarkonium production processes provide an important way to investigate QCD in its poorly known non-perturbative regime. Its production mechanism has been attracted extensive interests from the high-energy physics community in decades. The qualitative and quantitative description of heavy-quarkonium production requires complex perturbative computations for high-multiplicity processes in the framework of the well established non-relativistic effective theory, NRQCD, and reliable Monte Carlo simulations to reproduce the collider environment.

*Method of solution:* Based on a recursion relation, the program is able to calculate the helicity amplitudes of the high-multiplicity heavy-quarkonium-production processes. Several modules are also designed for dedicated simulations: 1) The code has been interfaced with the parton shower Monte Carlo programs; 2) A decay module to let heavy quarkonia decay with correct spin-correlations has been implemented; 3) The code estimates the theoretical uncertainties and analyzes the generated events on the fly; 4) The code is compilant with multi-threading/multi-core usage or cluster processors.

*CPC classification code:* 4.4 Feynman Diagrams, 11.1 General, High Energy Physics and Computing, 11.2 Phase Space and Event Simulation, 11.5 Quantum Chromodynamics, Lattice Gauge Theory
Typical running time:
It depends on the process to be calculated and the required accuracy.
1 Introduction

Since the breakthrough discovery of the Higgs boson at the LHC, much hope has been put on searching beyond Standard Model (BSM) particles in the next runs of LHC. However, the studies of quantum chromodynamics (QCD) are always playing a crucial role in the LHC objectives mainly because QCD is still a poorly-known theory especially in the non-perturbative region due to its color confinement and because it is crucial to understand QCD background at the hadron colliders. For example, at 14 TeV LHC, each bunch crossing will generate around 50 pile-ups. Such an effect is mainly governed by soft interactions. Hence, all aspects of QCD still deserve to be explored as fully as possible.

For a long time, heavy-quarkonium production and decay at high-energy colliders was thought to provide an ideal opportunity to study both the perturbative and non-perturbative aspects of QCD. Besides, it also shows a rich physics. Fundamental parameters such as the strong coupling constant $\alpha_s$ [1, 2], the heavy-flavor quark mass [1, 2, 3], the elements of the Cabibbo-Kobayashi-Maskawa (CKM) matrix [4], the Yukawa coupling [5] can be measured with heavy-quarkonium-production processes. Non-perturbative parton-distribution functions–either collinear [6, 7] or transverse [8] in the initial hadron–can be constrained from heavy-quarkonium data. B-meson-decay process $B^0 \rightarrow J/\psi + K^0_s$ provides a golden channel to investigate CP violation. Heavy-quarkonium production is also useful in probing the multiple-parton interactions [9, 10]. Other applications include quarkonium in quark-gluon plasma [11], cold nuclear matter effects on quarkonium production [12] and even BSM searching (see e.g. Ref. [13] and references therein) etc.

Despite of its importance, one has very limited choice of Monte Carlo tools for the simulation of the heavy-quarkonium-production processes on the market. From our point of view, this can be attributed to several longstanding puzzles in understanding its mechanism (see e.g Refs. [14, 15]) inspite of the well-established effective theory non-relativistic QCD (NRQCD) [16]. Both MadOnia [17] and HELAC-Onia [18] are such tools dedicated to matrix-element calculations and event generation within the NRQCD framework, which aim at providing general and user-friendly public tools for theorists and experimentalists to study the quarkonium physics. Although there are many similarities in both tools, we wish to emphasize some main differences between MadOnia and HELAC-Onia. HELAC-Onia is based on recursion relations to calculate helicity amplitudes, while MadOnia uses the traditional Feynman diagrams. Moreover, HELAC-Onia is designed to deal with processes containing one or more heavy quarkonium up to P-wave Fock states, while the number of states of heavy quarkonia is restricted to one in MadOnia.

The aim of this writeup is to introduce a 2.0 version of HELAC-Onia, where many new and useful features are included, which are motivated by the practical phenomenological
simulations and the user experience, e.g. interfacing with parton shower Monte Carlo programs. In section 2 we will describe the methodology, the related algorithms and the new features in HELAC-Onia 2.0. Then, we will show how to use the program in section 3. Several examples are given in section 4. In section 5 we draw our conclusions. Some useful information are given in the appendices. The program structure is sketched in appendix A. A summary of the new particle symbols in HELAC-Onia 2.0 is tabulated in appendix B and a few of useful new parameters are introduced in appendix C. Finally, the addon codes in HELAC-Onia 2.0 are introduced in appendix D.

2 Methodology, algorithm and new features

2.1 Heavy-quarkonium-amplitude computation with recursion relations

As it was introduced in our previous document [18], HELAC-Onia is based on the off-shell recursion relation [19]. HELAC-Onia is based on a public package HELAC [20, 21, 22], which is based on the Dyson-Schwinger equations [23, 24, 25] to calculate the helicity amplitude in the SM at parton level. In this section, we will first briefly recall how to calculate a helicity amplitude for a general process with \( n \) external legs. We denote the momenta of these external legs as \( p_1, p_2, \ldots, p_n \), and their quantum numbers, such as color and helicity, are symbolized as \( \alpha_1, \alpha_2, \ldots, \alpha_n \). Any \( k \) external legs can form an off-shell current as \( \mathcal{J}(\{p_{i_1}, \ldots, p_{i_k}\}; \{\alpha_{i_1}, \ldots, \alpha_{i_k}\}) \). We can assign each current \( \mathcal{J} \) a number \( l \), which is called “level”. It is defined as the number of external legs involved in the current \( \mathcal{J} \), i.e. the “level” of \( \mathcal{J}(\{p_{i_1}, \ldots, p_{i_k}\}; \{\alpha_{i_1}, \ldots, \alpha_{i_k}\}) \) is \( k \). The construction of the higher “level” currents is from the lower “level” ones in a recursion relation, where the starting point of the recursion relation is external legs and its end point is to obtain the “level” \( n \) current. The advantages by working in this way is that one is able to avoid computing identical subgraphs contributing to different Feynman diagrams more than once. The summation of all subgraphs contributing to a specific current reduces the total number of objects that should be used in the next recursion procedure.

In HELAC-Onia, we calculate the heavy-quarkonium amplitude in the framework of NRQCD factorization. With this formalism, the cross section for a heavy quarkonium \( Q \) production can be factorized into the perturbative short-distance components and the non-perturbative long-distance matrix elements (LDMEs). For example, at a proton-
proton collider, the cross section can be written as

$$\sigma(pp \rightarrow Q + X) = \sum_{i,j,n} \int dx_1 dx_2 f_{i/p}(x_1) f_{j/p}(x_2) \hat{\sigma}(ij \rightarrow Q\bar{Q}[n] + X) \langle O^Q_n \rangle,$$

(1)

where $f_{i/p}$ and $f_{j/p}$ are the parton distribution functions (PDFs), $\hat{\sigma}(ij \rightarrow Q\bar{Q}[n] + X)$ is the short distance coefficient to produce a heavy quark pair $Q\bar{Q}$ in the specific quantum state $n$. Following the usual notation, the Fock states $n$ can be written in the spectroscopic form $n = 2S + 1L \ [c]$, where $S$, $L$, $J$ identify the spin, orbital momentum, total angular momentum states respectively, and $c = 1,8$ means that the intermediate state $Q\bar{Q}$ can be in a color-singlet or a color-octet state. The LDMEs are denoted as $\langle O^Q_n \rangle$. Their physical interpretation is a probability for a heavy quark pair in the Fock state $n$ to evolve into a quarkonium. The power counting rules in NRQCD yield to the fact that for any quarkonium, there should be only a limited number of Fock states contributing to a specific order of $v$, where $v$ is the relative velocity of the heavy quark pair. The projection method is used to project a heavy-flavor quark pair onto a specific Fock state. The color-singlet projector is $\delta_{ij}$, while the color-octet projector is $\sqrt{2} \lambda a$, where $\lambda$ is the Gell-Mann matrix and it will be projected further onto a color-flow basis \[26, 27, 28\]. The spin projectors \[29\] are in the form of

$$- \frac{1}{2\sqrt{2}(E + m_Q)} \bar{v}(p_2, \lambda_2) \Gamma S \frac{\not{P} + 2E}{2E} u(p_1, \lambda_1),$$

(2)

where $m_Q$ is the mass of the heavy quark, $p_1, p_2$ and $\lambda_1, \lambda_2$ are the momenta and helicity of the heavy quarks respectively. The total momentum of the heavy quark pair is $P^u = p_1^u + p_2^u$ and $E = \sqrt{P^2}$. $\Gamma S$ is $\gamma_5$ for the spin singlet state $S = 0$, and it is $\epsilon^\lambda \gamma^\mu$ for the spin triplet state $S = 1$ with $\lambda_s = \pm,0$. In order to construct the “level” 1 current for the heavy quarkonium, we cut the fermion chain at the place of $\not{P} + 2E$ in the projector shown in Eq.(2). Then, the new “level” $l = 1$ current for $Q$ as $\frac{1}{m_Q} \bar{u}(P, \lambda') (\not{P} + m_Q)$ and for $\bar{Q}$ as $-\frac{1}{2\sqrt{2m_Q}} (\not{p}_2 - m_Q) u(P, \lambda')$. HELAC-Onia is also designed to be able to handle P-wave states. In HELAC-Onia, we introduced numerical stable P-wave currents, which have already been discussed in Ref. \[18\].

### 2.2 Angular distributions of heavy-quarkonium decays

Angular distributions of heavy-quarkonium decays have attracted a lot of attention in the past few years. Theorists are interested in the polarization observables of heavy-quarkonium production because they might provide a “smoking gun” to discriminate

\[1\]Rigorously speaking, the LDMEs are not physical, while only the (differential) cross section is. They are much like the PDFs and fragmentation functions (FFs).
the various heavy-quarkonium-production mechanisms. The understanding of heavy-
quarkonium polarization is also crucial for the simulations in the experimental analy-
yses, e.g. the detector acceptance for lepton pairs from the decay of $J^{PC} = 1^{--}$ heavy
quarkonium strongly depends on its polarization or on the angular distribution of its
decay products. The crue implementation of the polarization in the simulation of heavy-
quarkonium production usually leads to one of the largest systematic uncertainties on
the measurements. However, the current available Monte Carlo programs like PyThIA are
usually very limited in the available decay processes of heavy quarkonium and/or assume
the unpolarized pattern. Hence, it is our motivation to implement some frequently used
decay processes of heavy quarkonium with the polarization pattern.

For the simple decay processes, like $J/\psi \rightarrow \ell^+\ell^-$, we only have to follow the po-
larization of the mother particle and the implementation of the angular distribution
in each spin eigenstate is straightforward. However, for a general decay process, e.g.$\chi_{c,J} \rightarrow J/\psi + \gamma, J = 1,2$, the algorithms for generating the angular distributions of the
decay products is following:

1. Considering the helicity amplitude for the decay process is $A(x)$, where $x$ is the set
   of variables to characterize the kinematics of the decay process.

2. The maximal weight of $|A(x)|^2$ is $W_{\text{max}}$.

3. Randomly generate a phase space point $x$.

4. Uniformly generate a random number $r \in [0,1]$. If $|A(x)|^2 > r \times W_{\text{max}}$, the event
   corresponding to $x$ is retained. Otherwise, go to the former step.

All of the available hard-coded decay processes in HELAC-Onia 2.0 can be found in decay/deca-
list.txt. HELAC-Onia 2.0 also supports the cascade decays. For instance,
a generated $\chi_{c,J}$ meson can be decayed into a $J/\psi$ and a photon $\gamma$ first and then the decay
product $J/\psi$ can be further decayed into a lepton pair. However, the first decay process
may depend on the true masses of $\chi_{c,J}$ and $J/\psi$. The corresponding input values can be
changed by the user in input/decay_param_user.inp.

Such a module is flexible and can be extended to other decay processes. Especially,
we are planning to include EvtGen [30, 31] for B-physics studies in HELAC-Onia.

2.3 Interface to parton shower Monte Carlo programs

Parton shower Monte Carlo programs are widely used in numerical simulations for the
collider environment.
2.3.1 HELAC-Onia+Pythia8

Pythia is a general purpose Monte Carlo program. It provides the QCD and QED parton shower as well as the hadronization. Experiments performed on high-energy colliders rely heavily on it. Hence, the interface between HELAC-Onia and Pythia would surely extend the applications of the program. HELAC-Onia 2.0 has indeed been successfully interfaced with Pythia8\(^2\), which is written in C++. Its usage in HELAC-Onia requires the user to pre-install HepMC\(^3\) and Pythia8\(^3\). Inheriting to its processor Pythia6\(^3\), Pythia8 provides an interface to the external hard matrix element/event generators via Les Houches Event files\(^3\) according to Les Houches accord format\(^3\). In HELAC-Onia 2.0, several files in C++ are written to use the generated Les Houches files and to shower and to hadronize the unweighted events with Pythia8 on the fly. The default output is HepMC event file after passing through Pythia8. However, such a format is usually inefficient to store events since it might result in a huge HepMC file from a relative large Les Houches file (say one million events). Two alternative options are provided. One is to output TopDrawer format plots with Hbook\(^4\). However, such an option requires the user to define all of the observables and the histograms in FORTRAN 90 before calling Pythia. Useful analysis tools, like FastJet\(^3\) (or FJCore) and HEP-TopTagger\(^3\), can also be linked to fill the histograms. Some examples are given in analysis/PYTHIA8. Another option is using the software Root, which however requires the user to pre-install Root. Events after showering and hadronization will be filled into Root tree, and a pre-defined C++ Root script is necessary. We also give some examples in the subdirectory analysis/PYTHIA8. We will described its detailed usage in section 3.

In order to read the events record in a HepMC file, we also provide some useful tools for converting it to a TopDrawer file or a Root tree file in HELAC-Onia 2.0. Their executable files are HepMC2Plot and HepMC2Root in the directory bin.

In principle, such a methodology can be applied to the interfaces to other parton shower Monte Carlo programs as long as it can recognize the Les Houches event files. Although the current version of HELAC-Onia is still not interfaceable with other general-purpose parton shower Monte Carlo programs automatically, it is in our to-do list to write the similar interfaces for Pythia6\(^3\), Herwig6\(^3\), Herwig++\(^4\) and Herwig++\(^5\).

2.3.2 HELAC-Onia+QEDPS

QEDPS is a program for the photon showering from the initial $e^\pm$ in electron-positron collisions. Currently, it only works for Pythia8.1\(^3\) but not for Pythia8.2\(^3\). We used a simplified version of HBOOK written by M. Mangano.
collisions [46, 47, 48, 49], which is based on the fact that the structure function of an electron $D_{e\pm}(x, Q^2)$ should obey the Altarelli-Parisi equation

$$\frac{dD_{e\pm}(x, Q^2)}{d\log Q^2} = \frac{\alpha}{2\pi} \int_x^1 \frac{dy}{y} P_+(\frac{x}{y}) D_{e\pm}(y, Q^2),$$

where $x$ is the longitudinal fraction, $Q^2$ is the virtuality and $P_+(x)$ is the Altarelli-Parisi splitting function. In the leading logarithm approximation, one can solve it to be

$$D_{e\pm}(x, Q^2) = \Pi(Q^2, Q^2_0) D_{e\pm}(x, Q^2_0) + \frac{\alpha}{2\pi} \int_{Q^2_0}^{Q^2} \frac{ds}{s} \Pi(Q^2, s) \int_x^{1-\epsilon} \frac{dy}{y} P(y) D_{e\pm}\left(\frac{x}{y}, s\right),$$

where $Q^2_0 \sim m_e^2$ and $\Pi$ is the Sudakov factor

$$\Pi(Q^2, Q^2_0) = \exp\left(-\frac{\alpha}{2\pi} \int_{Q^2_0}^{Q^2} ds \int_0^{1-\epsilon} dx P(x)\right).$$

As it is well known, the Sudakov factor has the interpretation of the probability of the electron evolving from scale $Q^2_0$ to $Q^2$ without emitting any hard photon. We review the algorithm of the photon shower in QEDPS [46, 47, 48, 49]:

1. Initially, set $x = 1$, where $x$ is the fraction of the light-cone momentum of the electron/positron.

2. Generate a random number $r$. If $r$ is smaller than $\Pi(Q^2, Q^2_0)$, the evolution stops. Otherwise, try to find the next virtuality $Q^2$ with $r = \Pi(Q^2, Q^2_0)$. At $Q^2$, a branching $e^\pm \to e^\pm(Q^2)\gamma$ happens.

3. According to the probability of the splitting function $P(y)$ between $0$ and $1 - \epsilon$, try to determine the value of $y$. Replace the original $x$ to be $xy$. Iterate the step 2 until the evolution stops.

Hence, QEDPS provides a leading logarithm approximation to the initial state radiation in electron-positron collisions.

HELAC-Onia 2.0 provides an interface to QEDPS when calculating an electron-positron annihilation process. Thanks to the release of the source code, QEDPS is self-contained and the user does not need to install it by himself/herself. An input parameter `emep_ISR_shower` is provided in `input/user.inp` or `input/default.inp` to determine whether using QEDPS to perform a photon shower (see its description in appendix C). If one wants to generate TopDrawer, Gnuplot or Root plot files, one should also edit the Fortran subroutine `plot_fill_QEDPS` before compilation. We have performed an application of HELAC-Onia+QEDPS to $J/\psi$ inclusive production at B factories in Ref. [50].
2.4 Estimating the scale and PDF uncertainties

Varying the renormalization scale $\mu_R$ and the factorization scale $\mu_F$ is often thought to be a standard way to estimate the theoretical uncertainty in perturbative computations due to the missing higher-order contributions. Although such an argument can be applied to the scattering or decay processes in general as long as its validation of the perturbative description, there are indeed several cases which we already encountered where it is not applicable, such as where one encounters large coefficients correction (from large logarithms, large $\pi^2$, or large color factors) or new channels (e.g. new initial states, new phase space region, or new fragmentation topology). Unfortunately, the later case frequently happens in heavy-quarkonium-production processes. Because new fragmentation topologies appear only at higher orders in perturbative calculations, it is usually necessary to consider the contributions from the higher-multiplicity processes accompanying with more partons. Some examples indeed already can be seen in the single-quarkonium [51, 52] and double-quarkonium [53] production processes. Nevertheless, after taking into account all of the important topologies, the scale dependence is sufficiently reasonable to estimate the higher-order corrections. At tree level, the renormalization-scale dependence is only in the renormalization running of $\alpha_s(\mu_R)$, while the factorization-scale dependence is in the Dokshitzer-Gribov-Lipatov-Altarelli-Parisi (DGLAP) evolution [54, 55, 56] of the PDFs. Hence, it is straightforward that the estimation of the scale uncertainties are irrelevant to the most time-consuming matrix-element calculations as long as one knows the initial states and the perturbative orders. From the technical point of view, such estimation can be zero CPU cost.\footnote{In a general case, the explicit renormalization and factorization scales dependence is also known at next-to-leading order (NLO) [57].}

In a general case, the explicit renormalization and factorization scales dependence is also known at next-to-leading order (NLO) [57].

In calculating a physical observable or filling a histogram, one just multiplies a weight

$$w_{\text{scale}}(\mu_R, \mu_F; \mu_R^0, \mu_F^0) \equiv \frac{f_1(\mu_F, x_1)f_2(\mu_F, x_2)}{f_1(\mu_R^0, x_1)f_2(\mu_R^0, x_2)} \left(\frac{\alpha_s(\mu_R)}{\alpha_s(\mu_R^0)}\right)^b$$

(6)

to the central value in each phase space point or each event, where $f_i$ is the PDF, $x_i$ is the Bjorken fraction, $\mu_{R,F}^0$ ($\mu_{R,F}$) is the new (central) renormalization and factorization scales, $b$ is the power of $\alpha_s$ in the squared amplitude. Such reweighting procedure has been widely used in other programs, such as MADGRAPH [58, 59, 60].

Another important source of theoretical uncertainty that can be obtained from the reweighting method is the PDF uncertainty, which does not reflect the uncertainty in the hard matrix element but rather the uncertainty in the extraction of the PDF. It is known that the PDF uncertainty stems from at least three sources: the uncertainties in the input (experimental) data, the accuracy of the perturbative calculation, and the method to
extract the PDF. Most of the modern PDF sets provide a way to estimate the impact of such uncertainty to the theoretical calculations. For example, the global-fit PDF MSTW 2008 NLO \[61\] provides 40 error PDFs to quantify its uncertainty. Instead of reevaluating the matrix element with new PDF, one is able to evaluate the PDF uncertainty by multiplying a weight

\[
    w_{\text{pdf}}(f'_1, f'_2; f_1, f_2) \equiv \frac{f'_1(\mu_F, x_1)f'_2(\mu_F, x_2)}{f_1(\mu_F, x_1)f_2(\mu_F, x_2)},
\]

where \( f'_i \) is an error PDF and \( f_i \) is the central PDF. Such a procedure is exact in a parton-level calculation. However, it should be understood as an approximation when incorporating with a parton shower Monte Carlo program, since the backward evolution of the initial state partons in Monte Carlo indeed contains an implicit dependence on the chosen PDF, i.e. the central one in a Les Houches event file.

Although such a procedure is more trivial at tree level than that at NLO \[57\], for completeness, we would like to emphasize that HELAC-Onia 2.0 provides a functionality to estimate scale and PDF uncertainties with such a reweighting method. However, because of the recursion relations, it is non-trivial to separate different coupling orders without degrading its speed advantage. Hence, we want to remind the user that it would be wrong in evaluating renormalization scale dependence if the amplitudes in different orders will contribute to a partonic level process.

### 2.5 A summary of new features

We give a summary before closing this section. HELAC-Onia 2.0 has been improved much compared to the first released version of HELAC-Onia \[18\]. The main changes include:

1. Two completely independent generators based on PHEGAS \[62\] and VEGAS \[63\] are implemented. Both of them can generate unweighted events for \( 2 \to n \) processes when \( n \geq 2 \) at \( pp, p\bar{p} \) and \( e^-e^+ \) collisions. For \( 2 \to 1 \) processes at hadron colliders, only VEGAS is available.

2. Additional internal PDFs are included. The program can also be interfaced with LHAPDF \[64\].

3. Analysis plots are done on the fly. Differential distributions can be plotted at the end of the computation phase.

4. The laboratory frame is not restricted to the center-of-mass frame of the initial collision anymore. The fixed-target mode was also added.
5. An interface from QEDPS to HELAC-Onia is done. One can include the QED photon showering effects from initial $e^\pm$ beams.

6. An interface from Pythia 8 to HELAC-Onia is done. It is able to use the major functionality in Pythia 8.

7. Reweighting method is used to estimate the renormalization/factorization scale and PDF uncertainties.

8. Several spin-entangled decay processes are implemented to take into account the polarization effects.

9. A user-friendly Python script is provided for user to use the program. It also allows us to calculate the cross sections of several subprocesses with multiply CPUs, such as on a multicore computer or on a cluster.

For item 1, in the previous version, the unweighted events can only be generated by PHEGAS [62]. $2 \rightarrow 1$ processes are not handled in this case. This improvement allows us to lift several restrictions. Concerning item 2, beforehand, only CTEQ6 [63] was available. It paves the way to application in nucleus collisions and to estimate PDF uncertainties. With the help of the improvement presented in item 4, we are able to apply HELAC-Onia to more experiments like fixed-target experiments [66]. Item 5 allows us to consider initial radiation effects at $e^-e^+$ collision, which might not be small in several important processes, such as $e^-e^+ \rightarrow J/\psi + gg$ [50], while the improvement of item 6 extends the usage of HELAC-Onia to one of the most widely used multipurpose event generator Pythia 8. For item 7, it will be very useful to estimate the renormalization/factorization scale and PDF uncertainties without the extra cost of recalculating cross sections. The improvement of item 8 is quite useful for practical simulations, and that of item 9 improves the user experience on using the program.

3 How to use the program

In this section, we will first give a brief introduction on how to perform a phenomenological analysis in a basic way. If one is only interested in using the program, one can follow the instruction in this section and ignore the remaining context.

3.1 Standalone

We first introduce how to use HELAC-Onia 2.0 in a standalone way. Before running the code, one should specify the configurations via the configuration file ho_configuration.txt
in the input subdirectory, which is described in section [A]. For example, if one wants to use LHAPDF [64], one should assign the correct path to the parameter lhapdf in the file ho_configuration.txt. Useful comments are also given in the configuration file. If one wants to output plot(s) on the fly, one should also edit the user plot file plot_user.f90 in the subsubdirectory analysis/user. One can follow some example files to write his/her own plot file. After the above preparations, one can set the configuration and make the files via the command line

```
> ./config
```

This procedure should only be done once. Afterward, the program is ready for running.

In HELAC-Onia 2.0, one can use two modes to perform a computation of a cross section. We still keep the initial way to run the program directly via executable file Helac-Onia. To use this way, one should follow the following lines:

1. Specify input parameters in input/user.inp following the format in input/default.inp. Some examples are given in input/demo.

2. Provide the process information in input/process.inp as well as the decay information in input/decay_user.inp.

3. If one wants to define one’s own dynamical renormalization and/or factorization scale, one should edit it in src/setscale.f90 before compiling. Four default scales are defined, i.e. the fixed scale, the transverse mass \( m_{T,1} = \sqrt{m_1^2 + P_{T,1}^2} \) of the first final state, \( \mu_0 = \sqrt{\left( \sum_{f \in \text{final states}} m_f \right)^2 + P_{T,1}^2} \) and \( \mu_0 = \frac{1}{2} \sum_{f \in \text{final states}} \sqrt{m_f^2 + P_{T,f}^2} \).

4. Run the program with the command line

```
> ./Helac-Onia
```

or

```
> ./bin/Helac-Onia
```

The final results will be collected in the output directory.

A second way to run HELAC-Onia 2.0 is by using the Python scripts, which is more user-friendly and hence strongly recommended. It provides the opportunity to avoid mixing the working directory and the HELAC-Onia directory. If one wants to use the majority of the new features in the program, one has to run the program with the Python

```
> python application.py
```
scripts. The basic way of using it after the above first three items is to run the program with the script

```bash
> /PATH/TO/HELAC-Onia/hocluster
```

where /PATH/TO/HELAC-Onia is a path to the HELAC-Onia directory relative to your working directory. Then one will see a prompt starting with “HO >”. There are two phases to compute a cross section for a process, i.e. generate the process and run the program. In the first phase, one should define one process or several subprocesses. For example, if one wants to calculate $J/\psi$ pair production, the syntax should be

```bash
HO> generate g g > cc(3S11) cc(3S11)
```

where the symbol $g$ represents the initial gluon and $cc \sim (3S11)$ means a pair of charm (anti-)quark $c$ and $\bar{c}$ in $\beta$ configuration, which can be found in appendix B. One can also calculate the cross sections for several subprocesses simultaneously. For instance, if one wants to go beyond the leading-order computation of $J/\psi$ pair production, he/she can type the following command lines

```bash
HO> generate g g > cc(3S11) cc(3S11)
HO> generate g g > cc(3S11) cc(3S11) g
HO> generate u g > cc(3S11) cc(3S11) u
HO> generate g u > cc(3S11) cc(3S11) u
```

It will use multiple cores to calculate the cross sections on the cluster or on a multicore machine. One can also run the addon processes, where the available addon processes and the corresponding numbers are listed in addon/addon_process.txt. For instance, if one wants to calculate the double parton scattering (DPS) for $J/\psi$ pair production, one can generate the process via

```bash
HO> generate addon 1
```

where the keyword `addon` should be specified after `generate` and the number for this process is 1 as be seen in addon/addon_process.txt. Before launching the jobs for numerical calculations, one can also change the parameters in input/user.inp via the interactive command syntax

```bash
HO> set <parameter_name> = <value>
```
One example is to take the maximum Monte Carlo integration number to be 10000. Then one just simply types

```
HO> set nmc = 10000
```

If one wants to take VEGAS as the Monte Carlo integration program, please uses

```
HO> set gener = 3
```

Another useful feature is to define the decay process(es) via Python scripts. The syntax is

```
HO> decay <process> @ <branching ratio>
```

The command lines

```
HO> decay cc ~(3 S11) > m+ m- @ 0.06d0
HO> decay w+ > m+ vm @ 1d0
```

means that the $\beta$ charmonium in the final states will decay to a muon pair with the branching ratio 6% and the $W^+$ boson will perform leptonic decay to a muon and a neutrino with 100% probability. After all, one just submits the job via command

```
HO> launch
```

and waits for the final results to be collected in a new created subsubdirectory \texttt{PROC\_HO\_i/results} in the working directory, where \texttt{i} is a number to be assigned uniquely.

In the new way, we take a similar fashion of the widely used program \texttt{MadGraph5\_aMC@NLO}, and we hope that it would become a standard in the future, or at least it will be much easier for the users who are already familiar with \texttt{MadGraph5\_aMC@NLO} to learn to use this program.

### 3.2 HELAC-Onia+Pythia8

Let us start to consider the case of using HELAC-Onia+Pythia8\footnote{It is more or less trivial to use HELAC-Onia+QEDPS by setting the flag \texttt{emep\_ISR\_shower} to be 1 in \texttt{input/user.inp} as described in appendix \cite{appendix}.}. One has to generate the Les Houches file for unweighted events in \texttt{PROC\_HO\_i/P0\_calc\_j/output} first before calling \texttt{Pythia}. This can be achieved by setting the flag \texttt{unvgt} to be T before launching the program.
The program will be able to call **Pythia8** if one sets correct path for Pythia8 and HepMC in the configuration file **ho_configuration.txt** and specifies the shower parameters in the file **shower_card_user.inp** before using the **ho_cluster** script. The corresponding parameter setup for calling Pythia8 in **user.inp** is

```
| HO> set parton_shower = 1
```

All of the above shower related setup can be done before or after generating Les Houches event files. In the former case, the Pythia8 will be called with the command **launch** directly, while in the later case we also provide a new command **shower** with the syntax

```
| HO> shower <working path>
```

where the working path is usually **PROC_HO_i**. The final output from Pythia8 will be collected in the directory **PROC_HO_i/P0_calc_j/shower/HO_PYTHIA8_k**, where i, j and k are integers starting from 0.

We will present some technical details for using Pythia8 in HELAC-Onia 2.0 directly, since the user might have encountered problems in compiling the Pythia8 related code with wrong setup. It will also be useful for extensive usage of Pythia8 in HELAC-Onia. After running the program with **launch** or **shower** commands with the correct shower-related setup, a directory **HO_PYTHIA8_k** will be created in **PROC_HO_i/P0_calc_j/shower**. If the program compiles successfully, an executable file **Pythia8.exe** will be generated. One can also change the Pythia8 setup via the file **Pythia8_lhe.cmnd** in the same directory. A direct using Pythia8 is possible by simply typing

```
| > ./Pythia8.exe
```

If there does not exist the executable file **Pythia8.exe**, it means there is a problem in the compilation. Some useful information can be found in **shower.log** to solve the problems.

---

6Especially, one should be aware of the parameter **ANALYSE** in **shower_card_user.inp**. It determines the output mode as described in section [2.3.1](#). If **ANALYSE** is empty, it will output a HepMC event file from Pythia8. One can also take **ANALYSE** to be a Fortran90 or C++ plot file in the other two output modes in section [2.3.1](#). For example, if **ANALYSE = plot_py8_pp_tj** and there exist a file **plot_py8_pp_tj.f90** in **analysis/PYTHIA8** in the HELAC-Onia root directory, it will output a TopDrawer file from Pythia8. If the extension of the file is .cc in **analysis/PYTHIA8**, it will output a Root file instead.
4 Examples

4.1 NNLO$^*$ level $J/\psi$ and $\psi(2S)$ hadroproduction

$\psi$ and $\Upsilon$ production at hadron colliders have challenged our understanding of heavy-quarkonium mechanism for decades [67, 68, 69]. Since then, the heavy-quarkonium-production data have been removed in the global fit of extracting PDF. Large QCD corrections were found in heavy-quarkonium production due to new $p_T$-enhanced fragmenting Feynman diagrams at higher orders [51]. Hence, it was suggested to look at how partial next-to-next-to-leading order (NNLO) QCD correction impacts the differential cross sections of $\psi$ and $\Upsilon$ production [52], which was called NNLO$^*$. Besides of the complete NLO result [51], it requires to calculate the $\mathcal{O}(\alpha_s^5)$ tree-level $2 \rightarrow 4$ process $pp \rightarrow \psi+3$-jets. MadOnia [17] was able to perform a first numerical computation with such complex. It is a good process to show the robustness of HELAC-Onia and to compare MadOnia.

The calculation of $\mathcal{O}(\alpha_s^5)$ process $pp \rightarrow \psi+3$-jets in the color-singlet mechanism (CSM) consists the following 13 independent subprocesses:

\begin{verbatim}
HO> generate g g > cc"(3S11) g g g
HO> generate g g > cc"(3S11) u u" g
HO> generate u g > cc"(3S11) g g u
HO> generate g u > cc"(3S11) g g u
HO> generate u g > cc"(3S11) u u" u
HO> generate g u > cc"(3S11) u u" u
HO> generate u g > cc"(3S11) d d" u
HO> generate g u > cc"(3S11) d d" u
HO> generate u u" > cc"(3S11) g g g
HO> generate u u" > cc"(3S11) u u" g
HO> generate u u" > cc"(3S11) d d" g
HO> generate u u > cc"(3S11) u u g
HO> generate u d > cc"(3S11) u d g
HO> set quarksumQ = T
HO> set iqnum = 3
\end{verbatim}

In the above command lines, we only include up and down quarks. The other quark-initial-state contribution can be included by

\begin{verbatim}
HO> set quarksumQ = T
HO> set iqnum = 3
\end{verbatim}

The truth of the flag quarksumQ makes sure we will include the initial (anti)quark PDFs
and \texttt{iqnum}=3 means we are working in 3-light-quark-flavor scheme. Moreover, one can type

\begin{verbatim}
HO> set combine_factors = 1. 3. 1. 1. 1. 2. 2. 1. 1. 2. 1. 1.
\end{verbatim}

to explicitly multiply a combination factor for each subprocess. For example, in the subprocess \( gg \rightarrow c\bar{c}\bar{q} + u\bar{u}g \), we take a factor 3 to account for \( gg \rightarrow c\bar{c}\bar{q} + q\bar{q}g \) with \( q = u, d, s \), because they share the same matrix element and PDF. The detailed correspondances are shown in Tab.1. In order to avoid infrared divergence in NNLO\(^*\) calculations, a special cutoff \( s_{ij}^{\text{min}} \geq 2 \) should be applied to the invariant mass squared of any massless parton pair, i.e. \( (p_i + p_j)^2 \geq s_{ij}^{\text{min}} \). In this case, we set

\begin{verbatim}
HO> set minmqqp = 3d0
\end{verbatim}

for any final state massless parton pair \( \sqrt{(p_i + p_j)^2} \geq 3.0 \), and set

\begin{verbatim}
HO> set minmqbeam = 3d0
\end{verbatim}

for one final state massless parton and a initial state parton \( \sqrt{-(k_{1,2} - p_i)^2} \geq 3.0 \).

We have compared the HELAC-Onia result with the MadOnia result, and found they were in perfect agreement. Because HELAC-Onia is based on the recursion relations, HELAC-Onia is faster than MadOnia in the computations. It is much easier for us to extend the MadOnia result to a wider \( p_T \) range. In Ref. \[70\], ATLAS Collaboration already used our result to compare their measurement for \( \psi(2S) \) up to 100 GeV. Here, we present the \( p_T \) distributions of \( J/\psi \) (Fig.1) in the LHCb acceptance at 13 TeV. We should remind the reader the following points. The color-singlet LDME is estimated in potential model \[71\]. The corresponding radial wave function at the origin was derived in the QCD-motivated Buchmuller-Tye potential \[72\]. We used CTEQ6M \[65\] as our PDF set and fixed \( s_{ij}^{\text{min}} = 2m_c \). The error bands in Fig.1 represent the renormalization and factorization scale uncertainties \( \frac{(2m_c)^2 + p_T^2}{2} \leq \mu_R = \mu_F \leq 2\sqrt{(2m_c)^2 + p_T^2} \) and the uncertainty in charm quark mass \( m_c = 1.5 \pm 0.1 \) GeV. In Fig.1 we do not include the NLO contribution, although it should be contained for a real NNLO\(^*\) prediction.

### 4.2 \( pp \rightarrow J/\psi + J/\psi + c\bar{c} \)

As noticed in Ref. \[10\], in double \( J/\psi \) production, \( pp \rightarrow J/\psi + J/\psi + c\bar{c} \) shares the leading-\( p_T \) contribution from charm quark fragmentation diagrams though it is of \( \mathcal{O}(\alpha_s^2) \)

\footnote{We will use the same color-singlet LDME in the following computations.}
| Syntax              | Subprocess                                      |
|---------------------|-------------------------------------------------|
| $gg > cc^{-}(3S_{11}) gg$ | $gg \rightarrow c\bar{c} + gg$                  |
| $gg > cc^{-}(3S_{11}) uu^{-}$ | $gg \rightarrow c\bar{c} + qq$ with $q = u, d, s$ |
| $ug > cc^{-}(3S_{11}) gg u$ | $gg \rightarrow c\bar{c} + gg$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ |
| $ug > cc^{-}(3S_{11}) g g u$ | $gg \rightarrow c\bar{c} + ggq$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ |
| $ug > cc^{-}(3S_{11}) u u^{-} u$ | $gg \rightarrow c\bar{c} + q\bar{q}q$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ and $q, q'$ not in the same flavor |
| $ug > cc^{-}(3S_{11}) uu^{-} u$ | $gg \rightarrow c\bar{c} + q\bar{q}q$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ and $q, q'$ not in the same flavor |
| $ug > cc^{-}(3S_{11}) dd^{-} u$ | $gg \rightarrow c\bar{c} + q'\bar{q}'q$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ and $q, q'$ not in the same flavor |
| $ug > cc^{-}(3S_{11}) dd^{-} u$ | $gg \rightarrow c\bar{c} + q'\bar{q}'q$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ and $q, q'$ not in the same flavor |
| $uu^{-} > cc^{-}(3S_{11}) gg g$ | $q\bar{q} \rightarrow c\bar{c} + gg$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ |
| $uu^{-} > cc^{-}(3S_{11}) uu^{-} g$ | $q\bar{q} \rightarrow c\bar{c} + qqg$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ |
| $uu^{-} > cc^{-}(3S_{11}) dd^{-} g$ | $q\bar{q} \rightarrow c\bar{c} + q'\bar{q}'g$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ and $q, q'$ not in the same flavor |
| $uu > cc^{-}(3S_{11}) uu g$ | $q\bar{q} \rightarrow c\bar{c} + qq$ with $q = u, \bar{u}, d, \bar{d}, s, \bar{s}$ |
| $ud > cc^{-}(3S_{11}) ud g$ | $q\bar{q} \rightarrow c\bar{c} + q'\bar{q}'g$ with $q, q' = u, \bar{u}, d, \bar{d}, s, \bar{s}$ and $q, q'$ not in the same flavor |

Table 1: Subprocesses are calculated with each generation for $pp \rightarrow \psi+3$-jets in CSM.
Figure 1: The transverse momentum distributions of $J/\psi$ from $pp \rightarrow J/\psi + j jj$ in the LHCb acceptance at 13 TeV.

suppressed compared to $pp \rightarrow J/\psi + J/\psi$. Hence, it is necessary to quantify its magnitude by an explicit calculation. For the first time, we performed such a complex calculation with the help of HELAC-Onia 2.0 in Ref. [10], which involves more than 2000 Feynman diagrams. It is a first (and till now the only) $2 \rightarrow 4$ process with at least two quarkonia to be calculated. We take this example to show the uniqueness and the robustness of HELAC-Onia to perform perturbative computations of more than one quarkonium production processes. Because the luminosity of the quark-antiquark initial states is usually much smaller than that of the gluon-gluon initial state at the high-energy colliders, we only include the gluon-gluon initial state here. One can use the following command line to generate the process

```
HO> generate g g > cc^- (3S11) cc^- (3S11) c c^-
```
For illustration, we are working in the CMS acceptance [73]:

\[
\begin{align*}
p_{T}^{J/\psi} & > 6.5 \text{ GeV} \quad \text{if} \quad |y^{J/\psi}| < 1.2, \\
p_{T}^{J/\psi} & > 6.5 \rightarrow 4.5 \text{ GeV} \quad \text{if} \quad 1.2 < |y^{J/\psi}| < 1.43, \\
p_{T}^{J/\psi} & > 4.5 \text{ GeV} \quad \text{if} \quad 1.43 < |y^{J/\psi}| < 2.2,
\end{align*}
\]

where in the rapidity interval \(1.2 < |y^{J/\psi}| < 1.43\), the transverse momentum \(p_{T}^{J/\psi}\) cutoff scales linearly with its absolute rapidity \(|y^{J/\psi}|\). We used the exact setup in Ref. [10].

Some selected differential distributions in \(pp \rightarrow J/\psi + J/\psi + c\bar{c}\) are shown in Fig. 2. The error bands are coming from the variations of the renormalization and factorization scales and the uncertainty of charm quark mass. Following the way in Ref. [10], we have taken into account the feeddown contribution from \(\psi(2S)\) decay. The feeddown contribution enhances the (differential) cross section by a factor of 1.89. The absolute azimuthal difference between the \(J/\psi\) pair \(d\sigma/d\Delta \phi\) is shown in Fig. 2a. It is an observable to distinguish the double-parton scattering (DPS) and the traditional single-parton scattering (SPS) since in the former production mechanism the two \(J/\psi\) are uncorrelated. However, such an observable might be sensitive to the primordial \(k_{T}\) smearing from the beam [53].

Besides the absolute azimuthal difference, the absolute rapidity difference (Fig 2b) and the invariant mass distribution (Fig 2c) are also good kinematical variables to discriminate DPS and SPS. Finally, various transverse momentum spectra are displayed. In Fig. 2d, we presented distribution of the vectorial transverse momentum sum \(P_{T}^{J/\psi J/\psi} = |\vec{p}_{T1}^{J/\psi} + \vec{p}_{T2}^{J/\psi}|\), while Fig. 2e (Fig. 2f) is the yields of the leading \(p_{T} = \max(p_{T1}^{J/\psi}, p_{T2}^{J/\psi})\) (subleading \(p_{T} = \min(p_{T1}^{J/\psi}, p_{T2}^{J/\psi})\)) of the two \(J/\psi\).

### 4.3 \(J/\psi\) hadroproduction with parton shower effect

The inclusive \(J/\psi\) hadroproduction is a first process challenging our understanding of the heavy-quarkonium-production mechanism. For a long time, it was known that the CSM can describe the total cross section of \(J/\psi\) or \(\Upsilon\) (e.g. see Ref. [74]) but not in the transverse momentum \(p_{T}\) distributions. In the recent years, most of the studies were focusing on the interpretation of the yields [51, 52, 75, 76, 77, 79, 80, 81, 82] and the polarization [83, 84, 85, 86, 87, 88, 90, 91, 92] of single-quarkonium hadroproduction at the large \(p_{T}\) regime. Some efforts have also been paid in the small \(p_{T}\) regime [93, 94]. However, none of the consistent matching between large \(p_{T}\) and small \(p_{T}\) results exists.

In Ref. [93], analytical small \(p_{T}\) resummation is performed for the color-octet states only in NRQCD, which lacks the dominant color-singlet contribution and the matching to the fixed-order results. Alternatively, one can perform a resummation with the parton shower (PS) approach, which is formally to be restricted to the leading-log accuracy (although
the partial subleading-log contributions can also be taken into account). It generates the complete events with correct kinematics and can be applied directly on the experimental analysis by including the detector effect.

In this subsection, we will give a simple example to show the importance of parton shower effect for $J/\psi$ hadroproduction in the small $p_T$ region. Let us consider the color-singlet contribution only at leading order (LO) in $\alpha_s$ and in $v$ (the relative velocity between the charm quark pair). Without primordial $k_T$ smearing effect from the beam and the multiple interactions, the LO curve in $p_T$ distribution is indeed significantly smearing by PS as seen in Fig.3, where we have used PYTHIA8.186. In the left panel of Fig.3 one can observe that such a smearing effect is mainly from the initial state radiation (ISR) while the final state radiation (FSR) only distorts the distribution slightly. On the other hand, LO result is good enough to describe the rapidity distribution. We should remind the reader that while the LO+PS color-singlet contribution is expected to describe the small $p_T$ data, the intermediate and the large $p_T$ data will receive substantial higher-order (or real emissions) \cite{51,52} and color-octet contributions. A consistent treatment of $J/\psi$ production in NRQCD is possible with the LO merging of matrix elements and PS in different jet multiplicities \cite{95,96}, where a pioneer work has been done in $\eta_b$ production \cite{97}. Such a detailed analysis is of course interesting but beyond the scope of this paper. We will leave it for a future work.

### 4.4 Validation of decay angular distributions

In subsection 2.2 we have discussed the implementation of the angular distributions in the heavy-quarkonium decay in HELAC-Onia. Here, we will give three examples to validate our implementations: $J/\psi \rightarrow \ell^+ \ell^-$, $\chi_{c1} \rightarrow J/\psi + \gamma$ and $\chi_{c2} \rightarrow J/\psi + \gamma$.

In our first example $J/\psi \rightarrow \ell^+ \ell^-$, we know the angular distribution of one lepton is in the form of

$$\frac{d\sigma}{d\cos \theta} \sim 1 + \lambda_\theta \cos^2 \theta,$$

where $\theta$ is the polar angular respect to the spin quantization axis in the rest frame of $J/\psi$ and $\lambda_\theta$ can be expressed in the longitudinal polarized cross section $\sigma_L$ and the transverse polarized cross section $\sigma_T$ of $J/\psi$

$$\lambda_\theta = \frac{\sigma_T - 2\sigma_L}{\sigma_T + 2\sigma_L}.$$  

\footnote{One should turn on primordial $k_T$ in PYTHIA as well.}
We have compared the numerical result from HELAC-Onia and the analytical result Eq.(9) in Fig.4. The total cross section has been normalized to unity. We selected \( \lambda_\theta = 1, -1, 0, 0.3, 0.5 \) for illustration. Perfect agreement is found.

For \( \chi_{c1} \rightarrow J/\psi + \gamma \), we have the same polar angular distribution Eq.(9) with respect to the decay product \( J/\psi \) or \( \gamma \) [86], while for \( \chi_{c2} \rightarrow J/\psi + \gamma \) the general formula is

\[
\frac{d\sigma}{d\cos\theta} \sim 1 + \lambda_\theta \cos^2 \theta + \lambda_{2\theta} \cos^4 \theta. \tag{11}
\]

In the later case, \( \lambda_{2\theta} \) is suppressed by the higher-order multipole amplitudes [86]. Explicitly, we have for \( \chi_{c1} \)

\[
\lambda_\theta = (1 - 3\delta) \frac{\sigma_{\chi_{c1}} \chi_{c1} - 3\sigma_{\chi_{c1}} \chi_{c1}}{(1 + \delta)\sigma_{\chi_{c1}}^{\chi_{c1}} + (1 - 3\delta)\sigma_{\chi_{c1}}^{\chi_{c1}}}, \tag{12}
\]

and for \( \chi_{c2} \)

\[
\lambda_\theta = 6 \left[ (1 - 3\delta_0 - \delta_1)\sigma_{\chi_{c2}}^{\chi_{c2}} - (1 - 7\delta_0 + \delta_1)(\sigma_{\chi_{c1}}^{\chi_{c1}} + \sigma_{\chi_{c2}}^{\chi_{c2}}) - (3 - \delta_0 - 7\delta_1)\sigma_{\chi_{c2}}^{\chi_{c2}} \right] / R, \\
\lambda_{2\theta} = (1 + 5\delta_0 - 5\delta_1) \left[ \frac{5(\sigma_{\chi_{c2}}^{\chi_{c2}} + \sigma_{\chi_{c2}}^{\chi_{c2}})}{\sigma_{\chi_{c2}}^{\chi_{c2}}} + 5\sigma_{\chi_{c2}}^{\chi_{c2}} \right] / R, \\
R \equiv (1 + 5\delta_0 + 3\delta_1)\sigma_{\chi_{c2}}^{\chi_{c2}} + 3(1 - 3\delta_0 - \delta_1)(\sigma_{\chi_{c2}}^{\chi_{c2}} + \sigma_{\chi_{c2}}^{\chi_{c2}}) + (5 - 7\delta_0 - 9\delta_1)\sigma_{\chi_{c2}}^{\chi_{c2}}. \tag{13}
\]

In the above equation, we denote \( \sigma_{\chi_{c1,2}}^{i,j} \) is the \((i, j)\)-component of the spin density matrix of \( \chi_{c1,2} \) production. In the following, we take \( \sigma_{\chi_{c1,2}}^{i,-j,j} = \sigma_{\chi_{c1,2}}^{i,j} \), which is valid in a CP-conserved process. The spin-summed cross sections can be expressed as

\[
\sigma_{\chi_{c1}}^{\chi_{c1}} = \sigma_{\chi_{c1}}^{\chi_{c1}} + \sigma_{\chi_{c1}}^{\chi_{c1}} + \sigma_{\chi_{c1}}^{\chi_{c1}}, \\
\sigma_{\chi_{c2}}^{\chi_{c2}} = \sigma_{\chi_{c2}}^{\chi_{c2}} + \sigma_{\chi_{c2}}^{\chi_{c2}} + \sigma_{\chi_{c2}}^{\chi_{c2}}. \tag{14}
\]

Parameters \( \delta_0, \delta_1 \) enter into Eq.(12) and Eq.(13). They can be determined by the normalized multipole amplitudes

\[
\delta = \frac{(1 + 2a_1^{j=1}a_2^{j=1})}{2}, \\
\delta_0 = \frac{(1 + 2a_1^{j=2}(\sqrt{5}a_2^{j=2} + 2a_3^{j=2}) + 4a_1^{j=2}(a_2^{j=2} + \sqrt{5}a_3^{j=2}) + 3(a_3^{j=2})^2)}{10}, \\
\delta_1 = \frac{(9 + 6a_1^{j=2}(\sqrt{5}a_2^{j=2} - 4a_3^{j=2}) - 4a_1^{j=2}(a_2^{j=2} + 2\sqrt{5}a_3^{j=2}) + 7(a_3^{j=2})^2)}{30}, \tag{15}
\]

where \( a_1^{j=1}, a_2^{j=1} \) and \( a_3^{j=1} \) are the electric dipole (E1), magnetic quadrupole (M2) and electric octupole (E3) amplitudes for \( \chi_{cj} \). We take the measured values by CLEO collaboration in Ref. [98]. The numerical values are shown in Tab.2. However, the implementation of the cascade decay \( \chi_c \rightarrow J/\psi \gamma \rightarrow \ell^+\ell^-\gamma \) in HELAC-Onia requires its full

\footnote{We have \((a_1^{j=1})^2 + (a_2^{j=1})^2 = 1\) and \((a_1^{j=2})^2 + (a_2^{j=2})^2 + (a_3^{j=2})^2 = 1\).}
Table 2: The normalized multipole amplitudes of $\chi_{cj} \rightarrow J/\psi + \gamma$ from the CELO measurement \cite{98}.

| $j$  | $a_{1}^{J=1,j}$ | $a_{2}^{J=1,j}$ | $a_{3}^{J=1,j}$ |
|------|----------------|----------------|----------------|
| 1    | 0.998          | -0.0626        | -              |
| 2    | 0.996          | -0.093         | 0              |

knowledge of the helicity decay amplitudes in terms of multipole amplitudes. Its complete derivation was performed in Ref. \cite{99} at the amplitude level for the first time. Such amplitude-level expressions will be served as the helicity amplitude $\mathcal{A}(x)$ defined in subsection \ref{subsec:2.2}. The validation of the implementations for $\chi_{c1,2} \rightarrow J/\psi + \gamma$ in HELAC-Onia 2.0 can be found in Fig.5 and Fig.6. The histograms of the decay product $J/\psi$’s angular distributions perfectly agree with the analytical expressions.

5 Conclusions

We have presented a version 2.0 of HELAC-Onia with several important updates for the practical theoretical studies and the Monte Carlo simulations for heavy-quarkonium-production processes. The main improvements are

- a completely new interface for talking between the user and the program written in Python scripts. It is much user-friendly and suitable to submit calculation jobs with multi-threading usage or on a cluster;

- automated interfacing HELAC-Onia to the parton shower Monte Carlo event generators. Two parton shower programs are successfully linked; One is QEDPS for the initial photon showering from the processes in electron-positron collisions, while the other one is the widely used one Pythia 8;

- a decay module for performing the spin-entangled (cascade-) decay of heavy quarkonium. Some dedicated decay processes are implemented such as $J/\psi \rightarrow \ell^{+}\ell^{-}$, $\chi_{c} \rightarrow J/\psi + \gamma$ and the decays of top quark, W-boson, Z boson;

- a reweighting method for estimating the uncertainties from the renormalization/factorization scale and PDF in an automatic manner;

- one-dimensional or two-dimensional histograms generation on the fly. Moreover, we also provide several useful analysis tools.
All of the above improvements are quite useful in the study of the heavy-quarkonium production. It also provides a flexible framework for the future developments like heavy-quarkonium production in heavy ion collisions [100] or in the transverse momentum factorization framework [8].
A Program structure

In this section, we will describe briefly the new program structure of HELAC-Onia 2.0 for the future developments. The main files contained in the program are already described in the README file of the tarball. The files in the program are mainly included in several subdirectories, which are displayed in Fig. 7. There are mainly ten subdirectories under the main directory of HELAC-Onia. Let us explain them in somewhat detail:

- **input.** All of the input files that required by the program are contained in this subdirectory. They are:
  - `user.inp`: a file for user to specify his/her input parameters.
  - `default.inp`: a file that includes all of the default values for the input parameters.
  - `process.inp`: a file for user to tell the program the process information.
  - `ho_configuration.txt`: a configuration file for HELAC-Onia.
  - `seed.input`: a seed for random number generator.
  - `shower_card_user.inp` *(shower_card_default.inp)*: a user (default) card to use parton shower programs.
  - `decay_param_user.inp` *(decay_param_default.inp)*: a list of user-defined (default) parameters for using in the decay module.
  - `decay_user.inp` *(decay_default.inp)*: a file to specify decay chains in this card.

- **output.** All of the output files will be generated here. Initially, it is empty.

- **src.** It contains all of the main source files of the program. They can be mainly divided into two parts. One part is for the matrix elements generator and the other part is for the phase space integration and events generation.

  1. matrix elements generation.
     - `Helac_Global.f90`: It is a file which contains all of the global variables.
     - `Helac_Func_1.f90`: In it, many helper functions and subroutines are defined.
     - `alfas_functions.f90`: Running of $\alpha_S$ which is used in MCFM [101].
     - `Projectors.f90`: It is a file in which the Clebsch-Gordan coefficients are defined.
– **Constants.f90**: Several subroutines are defined for reading input parameters.

– **SM_FeynRule_Helac.f90**: It contains all of the Feynman rules of the Standard Model.

– **Feynman_Helac.f90**: A useful subroutine is written in this file for reconstructing all Feynman diagrams.

– **Helac_wavef.f90**: It is a file to define all of external wave functions.

– **Helac_pan2.f90**: Definition of vertices to be used in Helac_pan1.f90.

– **Helac_pan1.f90**: Off-shell currents generation by using recursion relation.

– **Helac_master.f90**: It is a main file of computing helicity amplitudes.

2. **Phase space integration and events generation**. It is based on several adapted Monte Carlo integration programs.

(a) **PHEGAS**:

– **Phegas.f90**: It is an extensive version of PHEGAS[62] to deal with quarkonium kinematics. It was rewritten in *Fortran* 90.

– **Phegas_Choice.f90**: Some helper functions are defined here that will be used by **Phegas.f90**.

(b) **VEGAS**:

– **MC_VEGAS.f90**: A *Fortran* 90 version of VEGAS[63].

– **Func_PSI.f90**: Some helper functions of phase space integration were written in this file.

– **Colliders_PSI1.f90**: Phase space integration with VEGAS for $2 \rightarrow n(n \geq 1)$ at hadron colliders.

– **Colliders_PSI2.f90**: Phase space integration with VEGAS for $2 \rightarrow n(n \geq 2)$ at electron-positron colliders.

(c) **MINT**:

– **mint-integrator.f90**: It is a *Fortran* 90 version of MINT[102].

(d) **Internal Fortran 90 PDF files**:

– **CTEQ6PDF.f90**: CTEQ6 PDF[65] file in *Fortran* 90 version.

– **Structf_PDFs.f90**: A file for calling PDFs.

(e) **LHAPDF file**:

– **Structf_LHAPDF.f90**: A file for calling PDFs from LHAPDF[64]. User should specify “lhapdf=/path/to/lhapdf-config” in **input/ho_configuration.txt**.
(f) Others:

- **Helac_ranmar.f90**: A random number generation program RANMAR in **Fortran 90**.
- **MC_PARNI_Weight.f90**: PARNI in **Fortran 90**, but it is not used.
- **MC_RAMBO.f90**: Rambo [103] in **Fortran 90**.
- **MC_Helac_GRID.f90**: A grid file.
- **Helac_unwei.f90**: There are some subroutines for dealing with unweighted events in this file.
- **ADAPT.f90**: It is for optimization by using adaption procedure.
- **Phegas_Durham.f90**: Durham in **Fortran 90**. It can only be used to generate phase space points for massless external particles.
- **MC_Func.f90**: There are some helper functions and subroutines for Monte Carlo integrations.
- **Kinetic_Func.f90**: Some kinematical variables are defined in this file.
- **Cuts_Module.f90**: It is a file to provide the user to impose kinematical cutoff.
- **KT_Clustering.f90**: $k_T$ clustering and reweight factor for MLM merging [104, 96].
- **setscale.f90**: It provides the user to specify his/her renormalization and factorization scales.
- **setscale_default.f90**: It is only a default **setscale.f90** file for backup.
- **Helac_histo.f90**: Histogram drawing file in HELAC.
- **SinglePro.f90**: It is the main file for phase space integration and events generation.
- **Summation_Pro.f90**: A file for the summation mode, which is not used yet.
- **unweight_lhe.f90**: A file for writing out Les Houches events files.
- **FO_plot.f90**: A file for plotting fixed-order distributions. In this case, unweight events generation is not necessary.
- **Main_Test.f90**: The **Fortran 90** main program.

- **pdf**: More extensive internal PDFs are located in this subdirectory.
  - **pdf_list.txt**: A summary of internal PDFs in HELAC-ONIA.
- `make_opts.makefile_pdf`: Files of makefile for the PDF related routines. A library `libpdf.a` will be generated in `lib` subdirectory.
- `opendata.f`: A file in FORTRAN 77 for opening PDF data.
- `Partonx5.f`: Standalone FORTRAN 77 Partonx function.
- CTEQ files: They include `cteq3.f,Ctq4Fn.f,Ctq5Par.f,Ctq5Pdf.f,Ctq6Pdf.f`.
- MRST files: They include `mrs98.f,mrs98ht.f,mrs98lo.f,mrs99.f,mrst2001.f,jeppe02.f`.
- `gsdpdf` file: They include GS09 dPDF files [105].

**· shower.** The subdirectory contains files for parton shower.

- `QEDPS`: It contains the files of QEDPS for ISR photon shower form initial $e\pm$ beams.
- `PYTHIA8`: `PYTHIA` 8 subsubdirectory. It includes the main files for interfacing HELAC-Onia to `PYTHIA` 8 for showering.
- `PYTHIA6`: `PYTHIA` 6 [35] subsubdirectory. It will be used for the future development.
- `HERWIG6`: `HERWIG` 6 [42, 43] subdirectory. It will be used for the future development.
- `HERWIGPP`: `HERWIG++` [44, 45] subdirectory. It will be used for the future development.
- `interface`: Some interface files are included in this subdirectory. For example, `QEDPS_interface.f90` is a file to interface HELAC-Onia with QEDPS.

**· analysis.** A subdirectory for performing analysis.

- `hbook`: HBOOK files (a simplified version written by M. Mangano) for plotting.
- `user`: user defined plot files, like `plot_user.f90`. Some examples are also given.
- `PYTHIA8`: the analysis code for generating histograms or `ROOT` trees by using `PYTHIA8` and `FASTJET` [38] (or its core functionality `FJCore`).
- `heptoptagger`: the `HEPTopTagger` [39, 40, 41] source code for top quark tagging in the analysis stage.
- `include`: some including files for example `HEPMC90.INC` for defining `HepMC` [34] common variables.
- `various`: some useful tools at the analysis stage.
- **TMVA**: some examples for using TMVA contained in ROOT for multiply variable analysis.
- **LesHouches**: some useful tools for dealing the Les Houches event files.
- **HepMC**: a code to convert HepMC file to histograms or ROOT trees by using FastJet or FJCore.

• **jets**. A subdirectory containing jet related tools.
  - **fastjet**: code for interfacing FastJet to HELAC-Onia.
  - **fjcore**: the source code of FJCore as well as the interface code to HELAC-Onia.
  - **merge**: the different multiplicity leading-order matrix elements and parton shower merging code.

• **cernlib**. A subdirectory containing cernlib files.
  - **minuit**:MINUIT source files.

• **decay**. A subdirectory for decaying final state particles.
  - **decay_list.txt**: a list of available decay processes.
  - **Decay_interface.f90**: the main decay file.
  - **DecayInfo.f90**: a file to read the decay information from decay_user.inp in input subdirectory.
  - **HOVll.f90**: the angular distribution file for a vector decays into two leptons.
  - **HO_chi2psia.f90**: the angular distribution file for $\chi$ particle decays into a $J^{PC} = 1^{--}$ quarkonium and a photon.
  - **HO_t2bw.f90**: the angular distribution file for top quark decays into a bottom quark and a W boson.

• **cluster**. A subdirectory containing PYTHON scripts.
  - **create_subdir.sh**: a bash shell script for creating subdirectories in the working directory. It is useful for running on the cluster or in the multi-core mode.
  - **bin**: A subsubdirectory that contains executable script file ho_cluster after configuring and make.
  - **pythoncode**: A subsubdirectory that contains the python source codes.
* **cluster.py**: A file includes various cluster classes.
* **misc.py**: Helpful functions defined to perform routine administrative I/O tasks.
* **coloring_logging.py**: A file with logging color.
* **extended_cmd.py**: A file containing different extension of the cmd basic python library.
* **files.py**: A file contains useful classes for dealing with file access.
* **helaconia_run_interface.oy** and **helaconia_interface.py**: A user-friendly command line interface to access HELAC-Onia features.

- **addon**. A subdirectory for some ad hoc codes.
  - **addon_process.dat**: A list of available addon processes.
  - **pp_psiDPS**: An ad hoc code for DPS of \(pp(\bar{p}) \rightarrow Q_1 Q_2 + X\), where \(Q_i = J/\psi, \psi(2S), \Upsilon(1S), \Upsilon(2S), \Upsilon(3S)\).
  - **pp_psiX_CrystalBall**: An ad hoc code for \(pp(\bar{p}) \rightarrow Q + X\) via crystal ball function, where \(Q = J/\psi, \psi(2S), \Upsilon(1S), \Upsilon(2S), \Upsilon(3S), \chi_c0, \chi_c1, \chi_c2\) and \(\chi_{bJ}(nP)\) with \(J = 0, 1, 2, n = 1, 2, 3\).
  - **fit_pp_psiX_CrystalBall**: An ad hoc code for fitting crystal ball function to the experimental data of \(pp(\bar{p}) \rightarrow Q + X\), where \(Q = J/\psi, \psi(2S)\).
  - **fit_pp_upsilonX_CrystalBall**: An ad hoc code for fitting crystal ball function to the experimental data of \(pp(\bar{p}) \rightarrow Q + X\), where \(Q = \Upsilon(1S), \Upsilon(2S), \Upsilon(3S)\).
  - **fit_pp_QQ_CrystalBall**: An ad hoc code for fitting crystal ball function to the experimental data of \(pp(\bar{p}) \rightarrow Q + \bar{Q}\), where \(Q\) is charm or bottom quark.
  - **pp_QQ_CrystalBall**: An ad hoc code for generating events of \(pp(\bar{p}) \rightarrow Q + \bar{Q}\) via crystal ball function.
  - **pp_aajj_DPS**: An event generator for producing \(pp(\bar{p}) \rightarrow \gamma\gamma+dijet\) from DPS.

There are other subdirectories under the main directory. All generated libraries will be put in **lib** subdirectory. All module (object) files will be put in **mod (obj)** subdirectory. Executable files will be generated in the subdirectory **bin**.

## B Particle symbols in HELAC-Onia via Python script

In this appendix, we will introduce the new particle symbols for using HELAC-Onia 2.0 with Python scripts. We list them explicitly in Tabs.3156.
| Particle | Particle ID | Particle Symbol |
|----------|-------------|-----------------|
| $\nu_e, e^-, u, d, \nu_\mu, \mu^-, c, s, \nu_\tau, \tau^-, t, b$ | 1, \ldots, 12 | $ve, e^-, u, d, vm, m^-, c, s, vt, tt^-, t, b$ |
| $\bar{\nu}_e, e^+, \bar{u}, d, \bar{\nu}_\mu, \mu^+, \bar{c}, s, \bar{\nu}_\tau, \tau^+, \bar{t}, \bar{b}$ | $-1, \ldots, -12$ | $ve^-, e^+, u^-, d^-, vm^-, m^+, c^-, s^-, vt^-, tt^+, t^-, b^-$ |
| $\gamma, Z, W^+, W^-, g$ | 31, \ldots, 35 | $a, z, w^+, w^-, g$ |
| $H, \chi, \phi^+, \phi^-$ | 41, \ldots, 44 | $h, g0, g^+, g^-$ |

Table 3: The identity numbers and symbols of the SM “elementary” particles in HELAC-Onia 2.0.

| Fock State | Particle ID | Particle Symbol |
|------------|-------------|-----------------|
| $cc[3P^J_{J=0,1,2}]$ | 4431J1 | $cc^\sim(3P1J)$ |
| $cc[3P^8_{J=0,1,2}]$ | 4431J8 | $cc^\sim(3P8J)$ |

Table 4: The identity numbers and symbols for the charmonia in various Fock states in HELAC-Onia 2.0.
| Fock State | Particle ID | Particle Symbol |
|------------|-------------|-----------------|
| $b\bar{b}[_{0}^{S}]$ | 551001 | $bb\sim(1S01)$ |
| $b\bar{b}[_{0}^{S}]$ | 551008 | $bb\sim(1S08)$ |
| $b\bar{b}[_{1}^{S}]$ | 553011 | $bb\sim(3S11)$ |
| $b\bar{b}[_{1}^{S}]$ | 553018 | $bb\sim(3S18)$ |
| $b\bar{b}[_{1}^{P}]$ | 551111 | $bb\sim(3P11)$ |
| $b\bar{b}[_{1}^{P}]$ | 551118 | $bb\sim(3P18)$ |
| $b\bar{b}[_{3}^{P}]_{J=0,1,2}$ | 5531J1 | $bb\sim(3PJ1)$ |
| $b\bar{b}[_{3}^{P}]_{J=0,1,2}$ | 5531J8 | $bb\sim(3PJ8)$ |

Table 5: The identity numbers and symbols for the bottomonia in various Fock states in HELAC-Onia 2.0.

| Fock State | Particle ID | Particle Symbol |
|------------|-------------|-----------------|
| $c\bar{b}[_{0}^{S}]$ | 451001 | $cb\sim(1S01)$ |
| $c\bar{b}[_{0}^{S}]$ | 451008 | $cb\sim(1S08)$ |
| $c\bar{b}[_{1}^{S}]$ | 453011 | $cb\sim(3S11)$ |
| $c\bar{b}[_{1}^{S}]$ | 453018 | $cb\sim(3S18)$ |
| $c\bar{b}[_{1}^{P}]$ | 451111 | $cb\sim(3P11)$ |
| $c\bar{b}[_{1}^{P}]$ | 451118 | $cb\sim(3P18)$ |
| $c\bar{b}[_{3}^{P}]_{J=0,1,2}$ | 4531J1 | $cb\sim(3PJ1)$ |
| $c\bar{b}[_{3}^{P}]_{J=0,1,2}$ | 4531J8 | $cb\sim(3PJ8)$ |
| $b\bar{c}[_{0}^{S}]$ | -451001 | $bc\sim(1S01)$ |
| $b\bar{c}[_{0}^{S}]$ | -451008 | $bc\sim(1S08)$ |
| $b\bar{c}[_{1}^{S}]$ | -453011 | $bc\sim(3S11)$ |
| $b\bar{c}[_{1}^{S}]$ | -453018 | $bc\sim(3S18)$ |
| $b\bar{c}[_{1}^{P}]$ | -451111 | $bc\sim(3P11)$ |
| $b\bar{c}[_{1}^{P}]$ | -451118 | $bc\sim(3P18)$ |
| $b\bar{c}[_{3}^{P}]_{J=0,1,2}$ | -4531J1 | $bc\sim(3PJ1)$ |
| $b\bar{c}[_{3}^{P}]_{J=0,1,2}$ | -4531J8 | $bc\sim(3PJ8)$ |

Table 6: The identity numbers and symbols for the mixed flavour quarkonium $B_c^\pm$ family in various Fock states in HELAC-Onia 2.0.

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C  New input parameters

Some of the parameters in `input/default.inp` and `input/user.inp` have already been introduced in Ref. [18]. The new parameters we introduced in the new version are:

1. `energy_beam1` and `energy_beam2` are the energies in unit of GeV of the first beam and second beam respectively.

2. `fixtarget` is a flag to compute the cross section in a fixed-target collision environment (T) or not (F).

3. `ranhel` is a parameter to determine whether the program uses the Monte Carlo sampling over the helicity configurations. In HELAC-Onia 2.0, we extend `ranhel` to be 4, which is at the same level of performing Monte Carlo over the helicity configuration with `ranhel`=3. Instead of using $\int_{0}^{2\pi} d\phi \epsilon_{\mu}^{\phi}(\epsilon_{\mu}^{\phi})^{\ast}$ to perform the helicity summation where $\epsilon_{\mu}^{\phi} = \sum_{\lambda=\pm,0}{e^{i\lambda \phi}}{\epsilon_{\mu}},$ we select the helicity eigenstate of external particle when `ranhel`=4 to take a subsequent spin-entangled decay.

4. `pdf` is the PDF set number proposed in LHAPDF [64] or in `pdf/pdflist.txt`. Entering 0 means no PDF is convoluted. If one wants to use LHAPDF, please edit `input/ho_configuration.txt` and set the parameter `lhapdf` to be T.

5. `reweight_pdf` is a flag to use reweighting method to get PDF uncertainty. It only works when using LHAPDF. Correspondingly, one should also specify the first (`pdf_min`) and the last (`pdf_max`) of the error PDF sets.

6. `reweight_Scale` is a flag to use reweighting method to get renormalization and factorization scale dependence, which requires `alphasrun`=T. One can change the lower bound and upper bound for renormalization/factorization scale variations via parameters `rw_RScaledown`, `rw_RScale_up`, `rw_FScaledown` and `rw_FScale_up`.

7. `useMCFMrun` is a flag to perform the strong coupling $\alpha_S$ renormalization group running in the MCFM [101] way.

8. `toodrawer_output`, `gnuplot_output`, `root_output` are flags to ask HELAC-Onia to plot histograms and to output into TopDrawer, Gnuplot and Root files on the fly.

9. `emep_ISR_shower` is a parameter to determine whether use QEDPS to take into account initial state radiation effects in electron-positron collisions (1) or not (0).

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10. **parton shower** is a parameter to determine whether perform parton shower. **parton shower=0** means no shower, i.e. fixed-order calculation. The shower can only be used when the corresponding parton shower program is already installed and the user has already edited properly in `input/ho_configuration.txt`.

All other parameters are listed in `default.inp`. The user can fix his/her values in `user.inp` following the format in `default.inp`.

### D Addon codes

In this section, we will describe some addon codes implemented in HELAC-Onia 2.0 for dedicated studies. All of the addon codes have been listed in `addon/addon_process.dat`.

#### D.1 Single-quarkonium hadroproduction with crystal ball function

In fact, the description of the quarkonium-production mechanisms is still a challenge for theorists, especially current the state-of-the-art computation in NRQCD cannot describe the single-quarkonium-hadroproduction data in the whole kinematical region. It would be quite interesting and might be necessary to use empirical function to describe the single-quarkonium production from $pp$ or $p\bar{p}$ collisions with a data-driven way and use it to test other mechanisms like DPS or $pA$ and $AA$ collisions. Moreover, it also provides an economy way to generate events for single-quarkonium hadroproduction. Therefore, HELAC-Onia 2.0 has already been implemented some dedicated codes to fit the single-quarkonium-hadroproduction data and to generate events of the single-quarkonium hadroproduction.

Let us start with the description of the calculation. The initial-averaged squared amplitude for single-quarkonium $Q$ hadroproduction with the assumption of the dominance of gluon-gluon channel can be expressed in a crystal ball function

$$|A_{gg\rightarrow Q+X}|^2 = \begin{cases} 
K \exp\left(-\kappa \frac{P_T^2}{M_Q^2}\right) & \text{when } P_T \leq \langle P_T \rangle \\
K \exp\left(-\kappa \frac{(P_T)^2}{M_Q^2}\right) \left(1 + \frac{K}{n} \frac{P_T^2 - (P_T)^2}{M_Q^2}\right)^{-n} & \text{when } P_T > \langle P_T \rangle
\end{cases}$$

(16)

where $K = \lambda^2 \kappa \hat{s}/M_Q^2$ and $\hat{s}$ is the partonic center-of-mass energy. Then the cross section of single quarkonium $Q$ production in $pp$ collisions is

$$\sigma(pp \rightarrow Q + X) = \int dx_1 dx_2 f_g(x_1) f_g(x_2) \frac{1}{2\hat{s}} |A_{gg\rightarrow Q+X}|^2 dLIPS,$n$$

(17)
where $f_g$ is the gluon PDF and dLIPS is the Lorentz-invariant phase space measure for $pp \rightarrow Q + X$. The coefficients $\lambda, \kappa, n$ and $\langle P_T \rangle$ can be determined by fitting it to the experimental data.

D.1.1 Fit codes

The codes for fitting $\psi(1S, 2S)$ and $\Upsilon(1S, 2S, 3S)$ are in subdirectories `fit_pp_psiX_CrystalBall` and `fit_pp_upsilonX_CrystalBall` respectively. One can use the command lines

```
HO> generate addon 3
```

and

```
HO> generate addon 4
```

to drive the corresponding codes to perform fitting with Minuit package. The input files dedicated to these ad hoc codes are in `fit_pp_psiX_CrystalBall/input` and `fit_pp_upsilonX_CrystalBall/input`. One can specify the meson in `state.inp` and the fitting parameters in `fit_param_card.inp`. The selected experimental data can be assigned in `data_list_i.inp`, where $i$ is the number in `state.inp`. Some fitted results are contained in `fit_pp_psiX_CrystalBall/fitresults` and `fit_pp_upsilonX_CrystalBall/fitresults`. We have checked the fitted results of Ref. [9] for prompt $J/\psi$ production with the same setup.

For instance, through a combined fit of $d^2\sigma/dP_Tdy$ to the ATLAS [70], CMS [107], LHCb [108] and CDF [109] data, we obtained $\kappa = 0.543$ and $\lambda = 0.118$ for prompt $\psi(2S)$ when $\langle P_T \rangle = 4.5$ GeV and $n = 2$, where $\chi^2 = 242$ for total 90 experimental data. The comparisons are shown in Fig. 8. The result is collected in `fitresults/psi2s/fit1`.

D.1.2 A simple event generator for single-quarkonium hadroproduction

With the fitted parameters, we wrote a simple event generator for $pp(\bar{p}) \rightarrow Q + X$, where $Q = J/\psi, \psi(2S), \Upsilon(1S), \Upsilon(2S), \Upsilon(3S), \chi_{c0}, \chi_{c1}, \chi_{c2}$ and $\chi_{bJ}(nP)$ with $J = 0, 1, 2, n = 1, 2, 3$. The code is put in the subdirectory `addon/pp_psiX_CrystalBall`. One can drive such program with the following command line

```
HO> generate addon 2
```

Some special input parameters can be specified in `pp_psiX_CrystalBall/input`. One can set the type of $Q$ in `state.inp` and its polarization in `polarization.inp`. The file
crystalball.inp is used to input the parameters $\lambda, \kappa, n$ and $\langle P_T \rangle$. We have performed some applications in Ref. [110];

D.2 Double parton scattering for double-quarkonium production

We also implemented the code for calculating DPS for double-quarkonium production in a $pp$ or $p\bar{p}$ collider. One of its application can be seen in Refs. [10, 111]. We used a simple but widely-used “pocket formula” to describe DPS for double-quarkonium production $pp \rightarrow Q_1 Q_2 + X$

$$
\sigma_{Q_1 Q_2}^{DPS} = \frac{1}{1 + \delta_{Q_1 Q_2}} \frac{\sigma_{Q_1} \sigma_{Q_2}}{\sigma_{\text{eff}}} , \tag{18}
$$

where $\sigma_{Q_i}$ is the cross section for single quarkonium $Q_i$ production and $\sigma_{\text{eff}}$ is a parameter to characterise an effective spatial area of the parton-parton interactions. $\sigma_{\text{eff}}$ can be related to the parton spatial density $F(b)$ inside the proton as

$$
\sigma_{\text{eff}} = \left[ \int d^2b \left( F(b) \right)^2 \right]^{-1} . \tag{19}
$$

Within the factorization, $\sigma_{\text{eff}}$ should be independent of final states but it might change with different species of initial partons and its Bjorken fraction $x$. A first order assumptions of $\sigma_{\text{eff}}$ is independent of process and energy, which however should be checked case by case.

For single-quarkonium production, we used the crystal ball function described in appendix D.1 to estimate the squared amplitude. This special code can be found in addon/pp_psipsi_DPS. The command line to generate this process is

```
HO > generate addon 1
```

Similar to other addon codes, the input parameters dedicated to this code is in pp_psipsi_DPS/input. The parameter $\sigma_{\text{eff}}$ can be specified in sigma_eff.inp. The user can change the type of the quarkonium pair $Q_1$ and $Q_2$ in the file states.inp. The polarizations of $Q_i$ can be fixed in polarization_<name>.inp, where <name> is the name of $Q_i$, i.e. $J/\psi = jpsi, \psi(2S) = psi2S, \Upsilon(1S) = Y1S, \Upsilon(2S) = Y2S, \Upsilon(3S) = Y3S$. The parameters $\lambda, \kappa, n$ and $\langle P_T \rangle$ in the crystal ball function should be told in the files crystalball_<name>.inp.
D.3 Heavy-flavor quark pair hadroproduction with crystal ball function

Using the crystal ball function for heavy-flavor quark pair or open heavy-flavor meson pair production, one can also perform a fit to experimental data at the hadronic colliders in the $P_T$ spectrum of the quark/meson production. Hence, it also provides an opportunity to use a data-driven way to analyze the corresponding heavy-flavor quark/meson pair production, which usually suffers large theoretical uncertainties in a perturbative computation. Such a method indeed has been applied in the open charm production at a proposed fixed-target experiment at the LHC (AFTER@LHC) in Ref. [110]. The fit can be performed using the following commands

\begin{verbatim}
HO> generate addon 5
\end{verbatim}

Some input parameters for fitting are needed to be specified in fit_pp_QQ_CrystalBall/input. Moreover, with the fitted parameters, one can use the

\begin{verbatim}
HO> generate addon 6
\end{verbatim}

to generate the unweighted events for the heavy quark pair production in proton-proton or proton-antiproton collisions.

D.4 Double parton scattering for associated production of diphoton and dijet

Similar to Eq.(18), we have a “pocket” formula for diphoton and dijet production via DPS mechanism in $pp$ collisions

\begin{equation}
\sigma_{\gamma\gamma+jj}^{DPS} = \frac{\sigma_{\gamma\gamma}\sigma_{jj}}{\sigma_{\text{eff}}} + \frac{\sigma_{\gamma+j}\sigma_{\gamma+j}}{2\sigma_{\text{eff}}}.
\end{equation}

The LO matrix elements of $ab \to \gamma + \gamma, \gamma + j, j + j$ have been implemented in HELAC-Onia with the correct color flow. To the diphoton production, we also implemented the gluon-gluon initial state process, which is a loop-induced process for diphoton production. However, due to the high luminosity of the gluon-gluon initial state at a high energy collider, such a contribution might be substantial. Unweighted events for the DPS contributions to diphoton and dijet production can be generated by the command

\begin{verbatim}
HO> generate addon 7
\end{verbatim}
One can change the input $\sigma_{\text{eff}}$ in the input file `pp_aajj_DPS/input/sigma_eff.inp`. There is also a file `pp_aajj_DPS/input/subprocess.inp` for user to select/drop some partonic subprocesses and to choose to generate the unweighted events of $pp \to \gamma + \gamma$, $pp \to \gamma + j$, $pp \to j + j$ instead of the DPS process. Such a functionality is very useful to cross check and to specify a global K-factor from the missing higher-order quantum corrections. Some studies on the inclusive DPS production rates of this process at the Tevatron \[112\] and the LHC \[113\] have been explored in the literature.

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Figure 2: The differential distributions for $pp \rightarrow J/\psi + J/\psi + c\bar{c}$ in the CMS fiducial region [73]: (a) absolute azimuthal difference; (b) absolute rapidity difference; (c) invariant mass distribution; (d) the vectorial transverse momentum sum; (e) leading $p_T$; (f) sub-leading $p_T$. 
Figure 3: Illustrative plots for $J/\psi$ production at 13 TeV LHC with parton shower from Pythia8.186. We presented the fixed-order LO calculation (solid curve), LO+PS (long-dashed curve), LO+PS but turning off ISR (short-dashed curve) and LO+PS but turning off FSR (dotted curve).
Figure 4: Validation of lepton angular distributions in $J/\psi \rightarrow \ell^+\ell^-$. 
Figure 5: Validation of $J/\psi$ angular distributions in $\chi_{c1} \rightarrow J/\psi + \gamma$. 
Figure 6: Validation of $J/\psi$ angular distributions in $\chi_{c2} \rightarrow J/\psi + \gamma$.

Figure 7: Program structure of HELAC-Onia with version 2.0.
Figure 8: Combined fit of $d^2\sigma/dP_Tdy$ to ATLAS [70] (1st-3rd plots), CDF [109] (4th plot), CMS [107] (5th-7th plots), LHCb [108] (8th plot) for prompt $\psi(2S)$ production. The plots are generated automatically by HELAC-Onia 2.0.