

# CIPPRES User Manual

Nicolas SISOURAT

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# Contents

<b>1</b>	<b>Introduction</b>	<b>2</b>
1.1	What is CIPPRES?	2
1.2	How to install CIPPRES?	2
1.3	List of tools	2
<b>2</b>	<b>Quickstart tutorial</b>	<b>4</b>
2.1	Computing CI (ORMAS-like) states	4
2.2	Computing Dyson norms	4
2.3	Computing Fano resonance widths	5
2.3.1	Cases where the P-space is obtained using only one CI run	5
2.3.2	Cases where the P-space is obtained using several CI runs	5
2.4	Computing dipole transition moments and Photoionization cross sections	6
2.5	Computing electronic processes cross sections in ion-molecule collisions	6
2.5.1	Total and partial cross sections	6
2.5.2	Cross sections for a given ionized state using the ad hoc projection operator method	6
2.6	Computing Time Dependent Density Matrices, Entropies	6
2.6.1	Time Dependent Dyson Norm	7
<b>3</b>	<b>The theory behind CIPPRES</b>	<b>8</b>
3.1	Fano theory of resonances and Fano-CI-Stieltjes method	8
3.1.1	Fano theory of resonances	8
3.1.2	Fano-CI method	8
3.1.3	Stieltjes procedure	9
3.1.4	Choice of basis sets and Hartree-Fock reference	10
3.1.5	Choice of the PQ partitioning	10
3.2	Impact parameter method and Target-states expansion	10
3.2.1	The Impact parameter method	10
3.2.2	Close-coupling approach	12
3.3	Ad hoc projection operator method	12
	<b>Acknowledgements</b>	<b>13</b>
	<b>Bibliography</b>	<b>14</b>

# Chapter 1

## Introduction

### 1.1 What is CIPPRES?

CIPPRES stands for Configuration Interaction Plugin for Photoionized and Resonant Electronic States. It is a set of tools to compute resonant widths, dipole transitions moments, and photoionization cross sections using ORMAS-like CI states. CIPPRES contains also tools to compute the electron-loss and excitation cross sections in ion-molecule collisions.

CIPPRES is a plugin for Quantum Package 2.0 (QP2). It uses as much as possible the QP2 machinery (e.g. HF orbitals, integrals,...)

### 1.2 How to install CIPPRES?

CIPPRES can be installed as any QP2 plugin. Look at <https://quantum-package.readthedocs.io/en/master/> for the QP2 documentation (installation, plugins, scf calculations,...). CIPPRES plugin is here: [https://github.com/sisourat/qp2\\_plugins\\_nsourat](https://github.com/sisourat/qp2_plugins_nsourat)

### 1.3 List of tools

**cippres\_gencsf** generate lists of CSF to built the CI matrices

**cippres\_runci** built and diagonalize the CI matrices defined by **cippres\_gencsf**

**cistate\_analysis** provides some infos on the CI states

**mymulliken\_analysis** Mulliken analysis

**cippres\_dip** computes the dipole transition moments between CI states (e.g. to compute photoionization cross sections)

**cippres\_fano** computes 2e matrix elements between CI states (to compute decay widths)

**cippres\_stieltjes** computes decay widths or photoion cs using Stieltjes imaging

**cippres\_setup\_collision** read input collision xml file and set up the calc.

**cippres\_prop\_collision** and computes  $V_{\text{proj}}$  (defined in **cippres\_setup\_collision**) matrix elements between CI states solves the TDSE (i.e. TDCI) using the matrix elements from **cippres\_collision**

**cippres\_timedep\_entropy** computes the von Neumann entropy of a time dependent wavefunction.

**Compute\_Dyson.py** Python script that computes the norm of the Dyson orbitals  $= \langle \psi^{(N-1)} | \psi^{(N)} \rangle$

**Partial\_cross\_section.py** Python script that using the Projector method computes the ionization cross sections for  $\psi^{(N-1)}$  states.

**merge\_fano\_files.py** Python script that merges Fano output files in the case the P-space is built on several CI runs.

**generate\_xml.py** Python script to generate XML files for **cippres\_gencsf**

**Compute\_density\_matrix\_fast.py** Python script to compute the density matrix and natural energies using CI states.

**Compute\_Dyson.py** Python script to compute the Dyson norms (i.e. projecting N-electron states on (N-1) electron states).

**CorrelationIntegralDyson.py** Python script to compute the Correlation Integral using the Dyson norms.

**Dyson\_Partial\_cross\_section.py** Python script to compute the partial ionization cross sections in ion-molecule collisions using the Dyson norms.

**Even\_temp\_basis\_set.py** Python script to generate even tempered basis sets.

**TimeDependent\_ComputeDyson.py** Python script to compute the Time Dependent Dyson norms in ion-molecule collisions.

**cippres\_timedep\_entropy** computes the von Neumann entropy of a time dependent wavefunction.

# Chapter 2

## Quickstart tutorial

### 2.1 Computing CI (ORMAS-like) states

In the following example, one can compute CI-states for helium dimer. HF orbitals are optimized for  $\text{He}_2^+$ . Two kind of CI calculations are done (defined in hehe.xml): in the first CRun, CSFs with 2e in the two lowest orbitals and 2e in orbitals from 3 to 10 are included. In the second CRun, CSFs with 3e in the two lowest orbitals and 1e in orbitals from 23 to 134 are included.

```
qp create_ezfio -b "he-test" HeHe.xyz -o HeHe.ezfio
qp set electrons elec_alpha_num 1
qp set electrons elec_beta_num 1
qp run scf
qp set electrons elec_alpha_num 2
qp set electrons elec_beta_num 2
qp set cippres finput_cippres hehe.xml
qp run cippres_gencsf
qp run four_idx_transform
qp run cippres_runci
```

hehe.xml file:

```
<input>
<cirun>
<general>
<spin> 1 </spin>
<electron na=' 2 ' nb=' 2 ' />
</general>
<block>
<space ne='2' imo='1' fmo='2' />
<space ne='2' imo='3' fmo='10' />
</block>
</cirun>
<cirun>
<general>
<spin> 1 </spin>
<electron na='2' nb='2' />
</general>
<block>
<space ne='3' imo='1' fmo='2' />
<space ne='1' imo='23' fmo='134' />
</block>
</cirun>
</input>
```

### 2.2 Computing Dyson norms

One computes the N-electrons and (N-1)-electrons CI states in two separate EZFIO directories. Then using cistate\_analysis, the CI states as linear combination of Slater determinants are obtained. The Dyson norms are computed using the Python script ComputeDyson.py.

```

qp create_ezfio -b "aug-cc-pvdz-kbj-4spd" He.xyz -o He.ezfio
qp run scf
qp set cippres finput_cippres he.xml
qp run cippres_gencsf
qp run cippres_runci
qp run cistate_analysis
mv cistates_det.txt cistates_det_he.txt
qp create_ezfio -b "aug-cc-pvdz-kbj-4spd" He.xyz -o Hep.ezfio
qp run scf
qp set cippres finput_cippres hep.xml
qp set electrons elec_beta_num 0
qp run cippres_gencsf
qp run cippres_runci
qp run cistate_analysis
mv cistates_det.txt cistates_det_hep.txt
python /home/common/quantum_package/plugins/qp2_plugins_nsisourat/nicotools/Compute_Dyson.py cistates_det_he.txt
cistates_det_hep.txt ncistates_he — tee dyson_log

```

## 2.3 Computing Fano resonance widths

### 2.3.1 Cases where the P-space is obtained using only one CI run

Once the CI states are computed, they can be used in the Fano calculations: `ici1` is the Q-space and `ici2` is the P-space. "`i_stj_job 1`" == Stieltjes should perform a Fano calculations ("`i_stj_job 2`" is used for photoionization cross sections). `ifanosta` is the index of the CI state to be used as the resonance (from `ici1` CRun).

```

qp set cippres ici1 1
qp set cippres ici2 2
qp run cippres_fano
qp set cippres ifanosta 1
qp run cippres_stieltjes

```

### 2.3.2 Cases where the P-space is obtained using several CI runs

```

qp_create_ezfio -b "he-cs-nico" -p bfd HeCs.xyz -m 3
qp run scf
qp set cippres finput_cippres cshe.xml
qp set electrons elec_alpha_num 10
qp set electrons elec_beta_num 9
qp run cippres_gencsf
qp run four_idx_transform
qp run cippres_runci
qp set cippres ici1 1
j=1
for i in 2..189
do
rm HeCs.ezfio/cippres/cfano_cippres.gz
rm HeCs.ezfio/cippres/efano_cippres.gz
qp set cippres ici2 $i
qp get cippres ici2
qp run cippres_fano
mv HeCs.ezfio/cippres/cfano_cippres.gz HeCs.ezfio/cippres/cfano_cippres$j.gz
mv HeCs.ezfio/cippres/efano_cippres.gz HeCs.ezfio/cippres/efano_cippres$j.gz
((j++))
done
python /home/common/quantum_package/plugins/qp2_plugins_nsisourat/cippres/nicotools/merge_fano_files.py
HeCs.ezfio 188
/home/common/quantum_package/plugins/qp2_plugins_nsisourat/cippres/libs/stieltjes/stieltjes < HeCs.ezfio/cippres/fanot

```

## 2.4 Computing dipole transition moments and Photoionization cross sections

More or less as computing Fano resonance widths:

```
qp set cippres ici1 1
qp set cippres ici2 2
qp run cippres_dip
qp set cippres ifanosta 1
qp run cippres_stieltjes
```

## 2.5 Computing electronic processes cross sections in ion-molecule collisions

### 2.5.1 Total and partial cross sections

ici1 is the CRun to be used in the collision simulations

```
qp set cippres finput_coll coll.input.xml
qp set cippres ici1 1
qp run cippres_setup_collision
qp run cippres_prop_collision
```

coll.input.xml:

```
<CollInput>
<ImpactParam type='linear' bmin='1.05' bmax='2.2' nb='1' />
<ImpactVel vx='0.1' vy='0.0' vz='2.0' />
<Zgrid type='exp' zmin='-100' zmax='100' nzgrid='200' />

  <States stamin='1' stamax='11' />

  <InitState type='target' state='1' />
<Potential charge='-1.0' exponent='0.0' />
</CollInput>
```

### 2.5.2 Cross sections for a given ionized state using the ad hoc projection operator method

One first needs to obtain the norm of the Dyson orbitals (see above) and the cross sections. The latter is obtained with

```
/home/common/quantum_package/plugins/qp2_plugins_nsisourat/nicotools/coll_cross_section Prop_collision.out
| tee cross_section.txt
```

(Do not forget to delete the first and two list lines of cross\_section.txt). Then

```
python /home/common/quantum_package/plugins/qp2_plugins_nsisourat/nicotools/Partial_cross_section.py cross_section.txt
Dyson_norms.txt nbound_hep
```

## 2.6 Computing Time Dependent Density Matrices, Entropies

### **2.6.1 Time Dependent Dyson Norm**

python2.7 TimeDependent\_ComputeDyson.py cistates\_N.txt cistates\_Nm1.txt nstate\_dyson Psit\_collision.out

### **2.6.2 Density Matrix**

python2.7 Compute\_density\_matrix\_fast.py cistates\_N.txt

### **2.6.3 Time Dependent Entropy**

qp run cippres\_timedep\_entropy (needs first Density\_matrices.txt and Psit\_collision.out)



# Chapter 3

## The theory behind CIPPRES

### 3.1 Fano theory of resonances and Fano-CI-Stieltjes method

We briefly outline the Fano theory of resonances following the derivation of Howat et al. [HrG78]. Next, we present and discuss the Fano-CI method. We then sketch the Stieltjes procedure needed to extract a continuous approximation to the decay width from a discrete representation of the continuum. Finally, we discuss the choice of basis set and Hartree-Fock reference in the calculation of Fano-CI decay widths.

#### 3.1.1 Fano theory of resonances

Similarly to the Fano-ADC method [AC05], the Fano-CI approach is devised for the treatment of Feshbach or Fano resonances. In the Fano formalism [Fan61, HrG78], the wave function at energy  $E$  in the vicinity of the resonance is represented as a superposition of a discrete component,  $\Phi$ , and continuum components,  $\chi_{\beta,\epsilon}$ , corresponding to the  $N_c$  decay channels

$$\Psi_{\alpha,E} = a_{\alpha}(E)\Phi + \sum_{\beta=1}^{N_c} \int d\epsilon C_{\alpha,\beta}(E,\epsilon)\chi_{\beta,\epsilon}, \alpha = 1, \dots, N_c. \quad (3.1)$$

Here the index  $\beta$  runs over the  $N_c$  decay channels and  $\epsilon$  is the kinetic energy of the outgoing electron. The bound part of the wave function,  $\Phi$ , is assumed to be an isolated resonance, not interacting with other resonances. It has a mean energy

$$E_{\Phi} = \langle \Phi | \hat{H} | \Phi \rangle \quad (3.2)$$

where  $\hat{H}$  is the full electronic Hamiltonian of the system. The continuum states  $\chi_{\beta,\epsilon}$ , in their turn, are assumed to diagonalise the Hamiltonian

$$\langle \chi_{\beta,\epsilon} | \hat{H} - E | \chi_{\beta',\epsilon'} \rangle \approx \delta_{\beta,\beta'} \delta(E_{\beta} - E_{\beta'} + \epsilon - \epsilon') (E_{\beta} + \epsilon - E) \quad (3.3)$$

and are thus also non-interacting.

Solving the Schrödinger equation  $(\hat{H} - E)\Psi_{\alpha,E} = 0$ , one can determine the bound and continuum amplitudes,  $a_{\alpha}(E)$  and  $C_{\alpha,\beta}(E,\epsilon)$ , respectively [Fan61, HrG78]. From the expression for  $|a_{\alpha}(E)|^2$  [HrG78], one obtains the relationship between partial  $\Gamma_{\beta}$  and total widths

$$\Gamma = \sum_{\beta} \Gamma_{\beta} = 2\pi \sum_{\beta} |\langle \Phi | \hat{H} - E_r | \chi_{\beta,\epsilon_{\beta}} \rangle|^2 \quad (3.4)$$

where  $\epsilon_{\beta}$  is the asymptotic kinetic energy of the emitted electron and  $E_r$  is the energy of the resonance. In practice, the energy of the resonance is approximated with the expectation value of the Hamiltonian with respect to the bound component of the resonance, i.e.  $E_r \approx E_{\Phi}$  [AC05]. This approximation is justified provided that the shift of the resonance, resulting from the interaction with the continuum, is smaller than the error in the calculation of the energy of the bound state [AC05].

As can be seen from Eq. (3.4), the total decay width is a sum of the partial decay widths to different decay channels. The expression for the partial decay widths given in (3.4) is known as “single channel” approximation, where no interaction between the decay channels is assumed.

#### 3.1.2 Fano-CI method

The evaluation of expression (3.4) requires suitable approximations for the bound ( $\Phi$ ) and continuum ( $\chi_{\beta,\epsilon_{\beta}}$ ) parts of the resonance. In the following, we discuss the particular approximations made in the Fano-CI method. The discussion is focused on a singly ionised resonance of a closed shell atom or molecule. However, the method

can be generalised to  $n$ -tuply ionised states, as well as ionisation satellites and excited states of both closed and open shell species<sup>1</sup>

Let us consider the specific case of an electronic decay process following single ionisation of an  $N$ -electron atom or molecule. In this case, the bound part of the resonance state is the initial, singly ionised state of the system, whereas the continuum components comprise all doubly ionised states with an electron in the continuum. Normal Auger decay, ICD and ETMD are examples of such electronic decay processes. Within the *original version of the* (see footnote) Fano-CI method, the initial singly ionised state is approximated as a one-hole (1h) configuration, i.e. an  $(N - 1)$ -electron Slater determinant, where an electron is removed from orbital  $\varphi_i$

$$|\tilde{\Phi}\rangle = |\Psi_i\rangle = c_i|\Psi_0\rangle \quad (3.5)$$

Here,  $c_i$  indicates the annihilation operator and  $|\Psi_0\rangle$  denotes the Hartree-Fock ground state of the system. The continuum states  $|\chi_{\beta,\epsilon}\rangle$  are approximated as discrete square integrable states  $|\tilde{\chi}_q^a\rangle$ , which are linear combinations of two-hole-one-particle (2h1p) configurations  $|\Psi_{akl}\rangle$

$$|\chi_{\beta,\epsilon}\rangle \approx |\tilde{\chi}_q^a\rangle = \sum_{k,l: \epsilon_{k,l} > \epsilon_i}^{n_{occ}} C_{akl}^q |\Psi_{akl}\rangle \quad (3.6)$$

Here, the indices  $k, l$  and  $a$  stand for occupied and virtual Hartree-Fock orbitals, respectively. The 2h1p configurations  $|\Psi_{akl}\rangle$  are characterised with a particle in  $\varphi_a$  and two holes in the restricted space of only those Hartree-Fock states  $\{\varphi_k\}$ , whose energy  $\epsilon_k$  obeys the relation  $\epsilon_k > \epsilon_i$ . Note that in our implementation all 2h1p configurations entering the expansion of a given final state  $|\tilde{\chi}_q^a\rangle$ , have a fixed virtual orbital,  $\varphi_a$ .

The coefficients  $C_{akl}^q$  are determined by solving the eigenvalue problem

$$\hat{H}|\tilde{\chi}_q^a\rangle = E_q^a |\tilde{\chi}_q^a\rangle \quad (3.7)$$

in the basis of the following doublet spin adapted configurations (or configuration state functions)

$$|\Psi_{akl}^{S_1}\rangle = \frac{1}{\sqrt{2}}(\bar{c}_a^\dagger c_k \bar{c}_l - \bar{c}_a^\dagger \bar{c}_k c_l) |\Psi_0\rangle \quad (3.8)$$

$$|\Psi_{akl}^{S_2}\rangle = \bar{c}_a^\dagger c_k \bar{c}_k |\Psi_0\rangle \quad (3.9)$$

$$|\Psi_{akl}^T\rangle = \frac{1}{\sqrt{6}}(\bar{c}_a^\dagger c_k \bar{c}_l + \bar{c}_a^\dagger \bar{c}_k c_l + 2c_a^\dagger c_k c_l) |\Psi_0\rangle \quad (3.10)$$

Here, the superscripts denote the singlet ( $S_{1,2}$ ) and triplet ( $T$ ) spin states of the two holes. A bar over the creation ( $\bar{c}^\dagger$ ) or annihilation ( $c$ ) operators stands for creation or annihilation of a spin orbital with spin  $\beta$ . The size of the resulting CI matrix is of the order of  $n_{occ}^2$ , where  $n_{occ}$  is the number of occupied Hartree-Fock orbitals. In order to obtain all final states  $|\tilde{\chi}_q^a\rangle$ , one needs to solve the eigenvalue problem (3.7)  $n_{virt}$  times, where  $n_{virt}$  is the number of virtual orbitals. The multiple diagonalisation of a small matrix is a substantial reduction in the computational effort compared to, for example, the effort needed to diagonalise a single ADC(2)x matrix, where the size of the 2h1p block is of the order of  $n_{occ}^2 n_{virt}$ . The cost for the substantial reduction in the size of the Hamiltonian matrix in the Fano-CI method is that the coupling between the pseudocontinuum final states  $|\tilde{\chi}_q^a\rangle$  is completely neglected. The 2h1p configurations are however partially coupled through Eq. (3.6).

Finally, the states  $|\tilde{\chi}_q^a\rangle$  are used to compute the coupling matrix elements

$$\gamma_q^a = 2\pi |\langle \tilde{\Phi} | \hat{H} - E | \tilde{\chi}_q^a \rangle|^2 \quad (3.11)$$

These coupling matrix elements cannot be directly used in Eq. (3.4) to compute the decay width. However, the total decay width can be recovered from them employing the Stieltjes procedure described below.

### 3.1.3 Stieltjes procedure

The states  $|\tilde{\chi}_q^a\rangle$  computed using the Fano-CI method (Eq. (3.6)) cannot be directly associated with the true continuum wave functions. First of all, these states are normalised to unity rather than energy normalised, and as such they do not satisfy the appropriate scattering boundary conditions. Moreover, owing to the finite basis sets employed in practical calculations, these final states do not satisfy the energy conservation condition for a non-radiative decay process

$$E_\beta + \epsilon_\beta = E_r \quad (3.12)$$

except by a coincidence.

These difficulties can be resolved by applying a mathematical approach known as the Stieltjes imaging technique [Haz79, Lan79, MPD89]. It relies on the fact that even though the square integrable ( $\mathcal{L}^2$ ) pseudocontinuum states cannot be used to compute the decay width directly, the spectral moments obtained from them are good approximations to the true spectral moments of the decay width.

$$S(k) = \sum_{\beta=1}^{N_c} \int dE E^k |\langle \tilde{\Phi} | \hat{H} - E_r | \chi_{\beta,\epsilon_\beta} \rangle|^2 \approx \sum_q \sum_a E_q^k |\langle \tilde{\Phi} | \hat{H} - E_r | \tilde{\chi}_q^a \rangle|^2 \quad (3.13)$$

<sup>1</sup>There is no specific PQ partitioning implemented in CIPPRES, any can be done manually which makes it flexible but also up to the users to do it well.

The moment theoretical approach is based on the observation that the pseudocontinuum  $\mathcal{L}^2$  wave functions approach the behaviour of the true continuum wave functions in the molecular interaction region provided that the basis set is sufficiently large to ensure a high density of pseudocontinuum states around the resonance energy. Usually, a series of calculations with an increasing number of spectral moments  $S(k)$  is performed until a consistent result is obtained [Haz79, Lan79, MPD89, AC05].

In practice it is difficult to obtain converged results without requiring a prohibitively large basis set. A question thus arises in the choice of the maximal Stieltjes order to be employed. In principle, higher orders provide more accurate results. However, owing to the finite basis sets, high order moments become inaccurate [MPD89]. By computing the standard deviation for each averaged decay width, one can give an estimate of the error resulting from the Stieltjes imaging procedure. This error is generally small even for moderate basis set size.

### 3.1.4 Choice of basis sets and Hartree-Fock reference

The choice of basis set is crucial for the accurate calculation of decay widths as the basis set has to provide a reliable description of both the discrete state and the continuum region of interest. This is usually achieved by employing large Gaussian basis sets (see e.g. [AC05, KAGC08, MCK<sup>+</sup>14b, MCK<sup>+</sup>14a, KS15, SGC16, JSCG16]), which are a combination of the standard basis sets and diffuse or compact basis functions centred around the atom or molecule. In the case of ICD due to the low energy of the emitted electrons, the most commonly used basis sets include the standard Gaussian basis sets augmented with a large number of diffuse basis functions specifically designed for the description of Rydberg and continuum states [KBJ89]. In contrast, the Auger electrons are much faster and in order to describe them the standard Gaussian basis sets are often uncontracted or augmented with compact basis functions [AC05, SGC16], which ensures a non-zero density of states in the high energy region of the continuum. For ICD widths we augment the standard Gaussian basis sets with diffuse functions of the Kaufmann-Baumeister-Jungen (KBJ) type [KBJ89] on the atomic centres. In the case of Auger decay, we uncontract the basis set and we augment it with sets of compact even tempered basis functions.

To construct the bound and continuum parts of the resonance state in the Fano-CI method, one needs to provide Hartree-Fock orbital energies and two-electron integrals as input. In the case of a singly ionised decaying state considered here, the ionisation step is accompanied by orbital relaxation effects, which are not accounted for in the Hartree-Fock calculation on the neutral system. These effects can be included by performing a restricted open-shell Hartree-Fock (ROHF) calculation, in which one can force an electron to be removed from the Hartree-Fock orbital of interest. Subsequently, the orbital energies and two-electron integrals generated in the ROHF step can be used in the Fano-CI procedure.

### 3.1.5 Choice of the PQ partitioning

As mentioned above, there is no specific PQ partitioning implemented in CIPPRES, any can be done manually (using the several CI runs). It makes CIPPRES flexible but also it is up to the users to do PQ partitioning well. For singly ionized states of closed-shell systems, the partitioning discussed above has proven to be rather accurate for many systems. There are also several schemes proposed in the Fano-ADC papers. For other kind of states and open-shell systems, more investigations are needed.

## 3.2 Impact parameter method and Target-states expansion

### 3.2.1 The Impact parameter method

An exact theoretical treatment of the electron dynamics requires to solve the full time-independent Schrödinger equation:

$$\hat{H}^{lab}\Psi_{sys} = E^{lab}\Psi_{sys} \quad (3.14)$$

where  $\hat{H}^{lab}$  and  $E^{lab}$  are the Hamiltonian operator and the total energy of the system in the laboratory frame, respectively. For a system with  $n_N$  nuclei and  $n_e$  electrons, the Hamiltonian  $\hat{H}^{lab}$  reads

$$\hat{H}^{lab} = \hat{T}^{lab} + \hat{V} \quad (3.15)$$

$$\hat{T}^{lab} = \sum_{I=1}^{n_N} -\frac{1}{2M_I} \nabla_{R_I}^2 - \frac{1}{2m_e} \sum_{i=1}^{n_e} \nabla_{r_i}^2 \quad (3.16)$$

$$\hat{V} = \sum_{I=1}^{n_N} \sum_{i=1}^{n_e} V_{Ii} + \sum_{I=1}^{n_N} \sum_{J=I}^{n_N} V_{IJ} + \sum_{i=1}^{n_e} \sum_{j=i}^{n_e} V_{ij} \quad (3.17)$$

where  $\vec{R}_I$  and  $\vec{r}_i$  are the nuclei and electron positions, respectively. The potential energy terms  $V_{IJ}$ ,  $V_{ij}$  and  $V_{Ii}$  represent the Coulombic potential between the nuclei, the electrons and between the nuclei and the electrons, respectively. Using that the nuclei masses are much larger than the electron mass (i.e.  $M_I \gg m_e$ ), the

Hamiltonian can be written as

$$\hat{H}^{lab} = -\frac{1}{2\mu} \sum_{i=1}^{n_e} \nabla_{r_i}^2 - \frac{1}{2\mu_{TP}} \nabla_R^2 - \frac{1}{2M_{tot}} \nabla_{R_G}^2 + \hat{V} \quad (3.18)$$

where the masses are defined as

$$\mu = m_e, \quad \mu_{TP} = \frac{M_T M_P}{M_T + M_P} \quad \text{and} \quad M_{tot} = M_T + M_P \quad (3.19)$$

The indexes T and P are used to label the target and the projectile, respectively. The vector  $\vec{R}_G$  represents the position of the center of mass of all nuclei in the laboratory frame. The kinetic energy  $\hat{T}^{lab}$  can be written as

$$\hat{T}^{lab} = \hat{T} - \frac{1}{2M_{tot}} \nabla_{R_G}^2 \quad (3.20)$$

where  $\hat{T}$  is the internal kinetic energy. Since the potential energy terms depend only of the internal coordinates of the system, one can split the full Hamiltonian into two contributions

$$\hat{H}^{lab} = \hat{H} - \frac{1}{2M_{tot}} \nabla_{R_G}^2 \quad (3.21)$$

Furthermore,  $\Psi_{sys}$  and  $E^{lab}$  can be written as

$$\Psi_{sys} = \Psi_{int} e^{i\vec{k}_G \cdot \vec{R}_G} \quad (3.22)$$

$$E^{lab} = E + \frac{k_G^2}{2M_{tot}} \quad (3.23)$$

Substituting Eqs. (3.21) to (3.23) into Eq. (3.14), one is left with

$$\hat{H} \Psi_{int} = E \Psi_{int}. \quad (3.24)$$

The internal Hamiltonian  $\hat{H}$  reads

$$\hat{H} = \hat{H}_{el} - \frac{1}{2\mu_{TP}} \nabla_R^2 \quad (3.25)$$

where  $\hat{H}_{el}$  is the electronic Hamiltonian:

$$\hat{H}_{el} = -\frac{1}{2m_e} \sum_{i=1}^{n_e} \nabla_{r_i}^2 + \hat{V}. \quad (3.26)$$

To go further,  $\Psi_{int}$  is written as a product of an electronic wavefunction  $\Psi$  and a nuclear one  $\Xi$ :

$$\Psi_{int} = \Psi(\vec{R}, \{\vec{r}\}) \Xi(\vec{R}) \quad (3.27)$$

In this approximation, which is widely known as the Born-Oppenheimer approximation, most dependence on  $\vec{R}$  is included in  $\Xi(\vec{R})$ . In that case,  $\nabla_R^2 \Psi(\vec{R}, \{\vec{r}\})$  can be neglected. Furthermore, in the intermediate energy range the wavefunction  $\Xi(\vec{R})$  can be written as a plane-wave [BM92]:

$$\Xi(\vec{R}) = e^{i\vec{k} \cdot \vec{R}} \quad (3.28)$$

where  $\vec{k}$  with a wave vector defined by

$$|\vec{k}| = \sqrt{2\mu_{TP} E} \quad (3.29)$$

Using Eqs. (3.25) to (3.29), Eq.(3.24) reads

$$[\hat{H}_{el} - \frac{i}{\mu_{TP}} \vec{k} \cdot \nabla_R] \Psi = 0 \quad (3.30)$$

From there one can introduce the semiclassical approximation where it is assumed that the relative motion of the nuclei  $\vec{R}$  is described by a classical trajectory  $\vec{R}(t)$ . Moreover, for collision energies above 1 keV/amu the projectile scattered dominantly in the forward direction and at very small angles. It is therefore reasonable to further assume that  $\vec{R}(t)$  describes a straight-line constant velocity trajectory:

$$\vec{R}(t) = \vec{b} + \vec{v}_P t \quad (3.31)$$

where  $\vec{b}$  is the impact parameter and  $\vec{v}_P$  is the velocity vector. The latter relates to  $\vec{k}$  as follows

$$\vec{v}_P = \frac{1}{\mu_{TP}} \vec{k} \quad (3.32)$$

Using this relationship, Eq. (3.30) reads

$$i\hbar \frac{\partial \Psi}{\partial t} = H_{el} \Psi \quad (3.33)$$

which is known as the eikonal equation, or equivalently the time-dependent Schrödinger equation describing the electron dynamics in the moving field of the nuclei.

This equation cannot be solved analytically even for the simplest collision systems (e.g. proton-Hydrogen). A full numerical solution for systems having one active electron is now possible given the tremendous progress in computational technologies. However, in the following we present a more general approach in which Eq. 3.33 is solved by assuming that the electronic wavefunction can be expanded using a finite set of basis functions.

### 3.2.2 Close-coupling approach

In a close-coupling description, the electron dynamics is constrained to a configuration space defined by a finite set of basis functions  $\chi_k(\{\vec{r}\})^2$ . The time-dependent electronic wavefunction is approximated by the following expansion:

$$\Psi(\{\vec{r}\}, t) = \sum_{k=1}^N a_k(t) \chi_k(\{\vec{r}\}) \quad (3.34)$$

If the set of basis functions  $\chi_k(\{\vec{r}\})$  is complete, an exact solution is then obtained. For obvious numerical reasons, the expansion must be truncated and the configuration space is thus not complete. It is therefore important to choose wisely the basis functions used in the calculations.

Substituting the wavefunction ansatz (Eq. 3.34) into Eq. 3.33 and projecting into each of the basis functions leads to a set of coupled first-order differential equations for the expansion coefficients. This set of equations can be written in a matrix form as

$$i\dot{\mathbf{S}}\mathbf{a} = \mathbf{M}\mathbf{a} \quad (3.35)$$

where  $\mathbf{a}$  is a vector containing the time-dependent coefficients and  $\mathbf{S}$  and  $\mathbf{M}$  are the overlap and coupling matrices, respectively. Various algorithms have been developed and implemented to solve numerically Eq. 3.35<sup>3</sup>.

We recall here that the electronic Hamiltonian depends on time due to the classical relative motion of the nuclei  $\vec{R}(t)$ . The latter depends explicitly on the impact parameter  $\vec{b}$ . For a given collision energy, Eq. 3.35 must therefore be solved for a set of impact parameter that reproduces a given experiment. Assuming an uniform and cylindrical ion beam, the total cross section for an inelastic process leading to a final state  $\phi_f(\{r\})$  is given by

$$\sigma_f = 2\pi \int_0^\infty b P_f(b) db \quad (3.36)$$

where the transition probability  $P_f(b)$  reads

$$P_f(b) = \lim_{t \rightarrow \infty} | \langle \phi_f | \Psi(t) \rangle |^2. \quad (3.37)$$

In the latter equation, we have used the Dirac *bra-ket* notation for simplicity.

Within the impact parameter method, the above derivation is general. In CIPPRES,  $\chi_k(\{\vec{r}\})$  are the (target) CI-states obtained with **cippres\_runci**.

### 3.3 Ad hoc projection operator method

Since cippres employs gaussian functions, it does not allow one to formulate a rigorous procedure for the calculation of the partial widths/cross sections. Such a rigorous calculation must involve the true degenerate continuum functions corresponding to the various channels. Despite the fact that these functions are not available within the framework of the method used in cippres, it is still possible to estimate partial quantities by an ad hoc procedure.

Let us assume the true quantity of interest is

$$| \langle \psi_f^{(N-1)} k | O | \psi^{((N))} \rangle |^2 \quad (3.38)$$

where  $\psi^{((N))}$  is a bound state with  $N$  electrons,  $\psi_f^{(N-1)}$  the final *ionic* state with  $(N-1)$  electrons and  $k$  is a true continuum state. Furthermore, we define the CI states of the  $P$ -space as  $\psi_p^{(N)}$ . A projection operator can be defined as

$$P = \sum_p | \psi_p \rangle \langle \psi_p |. \quad (3.39)$$

Inserting this operator into Eq. 3.38 one obtains

$$| \langle \psi_f^{(N-1)} k | O | \psi^{((N))} \rangle |^2 = \left| \sum_p \langle \psi_f^{(N-1)} k | \psi_p^{(N)} \rangle \langle \psi_p^{(N)} | O | \psi^{(N)} \rangle \right|^2 \quad (3.40)$$

where  $\langle \psi_f^{(N-1)} | \psi_p^{(N)} \rangle$  is a Dyson orbital. The norm of Dyson orbitals are usually smaller to one. Assuming  $k$  and the Dyson orbitals form a orthogonal basis set, one obtains

$$| \langle \psi_f^{(N-1)} k | O | \psi^{((N))} \rangle |^2 = \sum_p N_{fp} | \langle \psi_p^{(N)} | O | \psi^{(N)} \rangle |^2. \quad (3.41)$$

In the latter equation,  $N_{fp}$  is the norm of the Dyson orbital obtained when projecting  $\psi_f^{(N-1)}$  and  $\psi_p^{(N)}$ . These states are computed with independent cippres\_runci runs and  $| \langle \psi_p^{(N)} | O | \psi^{(N)} \rangle |^2$  are obtained from cippres\_fano for example.

<sup>2</sup>It should be mentioned that it is also possible, and sometimes numerically advantageous, to employ time-dependent basis functions. However, in the following we restrict the presentation to the case where the time-dependence of the electronic wavefunction is solely included in the expansion coefficients.

<sup>3</sup>In CIPPRES the predictor-corrector Adams-Moulton-Bashford (as implemented in the Heidelberg MCTDH package) is used.

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