

Running RIXS calculations with eSPec: practical guide

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I. GENERAL IDEA

The eSPec package, is in a wave packet propagation program which has been applied to a wide variety of dynamical problems, from absorption spectrum to reaction dynamics. In order to apply it to RIXS cross section calculations we ought to perform two different wavepacket propagations. The procedure goes as follows:

I) first we propagate the nuclear wavefunction from the initial state of the system on the intermediate state potential surface

II) we apply the eSPec-Raman module to obtain the new starting conditions, which depend on the incoming photon frequency.

II) finally we perform a second propagation of the obtained wave packets on the final state potential surface.

The eSPec already computes the auto-correlation function and the corresponding cross-section in the .SPECTRUM mode.

II. THE eSPec-RAMAN MODULE

The eSPec-RAMAN module was coded to provide the appropriate wave packets for the final state propagation, in other words it solves the following integral

$$|\Phi(0)\rangle = \int_0^\infty e^{-\Gamma t} e^{-i(\hat{H}_c - E_0 - \omega)t} |0\rangle dt, \quad (1)$$

now, eSPec does the propagation shifting the potentials to zero, so we should make $\hat{H}'_c = \hat{H}_c - \omega_{c0}$, then we get

$$|\Phi(0)\rangle = \int_0^\infty e^{-\Gamma t} e^{-i(\hat{H}'_c + \omega_{c0} - E_0 - \omega)t} |0\rangle dt, \quad (2)$$

defining

$$\phi_c(t) = e^{-i\hat{H}'_c t} |0\rangle \quad (3)$$

we can identify eq. (2) as a fourier transform (acutally here we have a half fourier transform, but it is possible to rewrite it as an ordinary transform, check Faris's file for detailed derivation)

$$|\Phi(0)\rangle = \int_0^\infty e^{-\Gamma t} e^{-i(\omega_{c0} - E_0)t} \phi_c(t) e^{i\omega t} dt, \quad (4)$$

since $|\Phi(0)\rangle$ is function of the detuning, we can write this dependence explicitly as $|\Phi(0)\rangle \equiv \Phi(0, \omega)$ now, if we consider the following shift property of the fourier transform:

$$\int_{-\infty}^\infty f(t) e^{i\omega t} dt = g(\omega), \quad (5)$$

$$\int_{-\infty}^\infty e^{\pm iat} f(t) e^{i\omega t} dt = g(\omega \pm a) \quad (6)$$

we can readily see that eq. (4) can be written

$$\Phi(0, \omega - \omega_{c0} + E_0) = \int_0^\infty e^{-\Gamma t} e^{-i(\omega_{c0} - E_0)t} \phi_c(t) e^{i\omega t} dt \quad (7)$$

where we can define $\bar{\omega} = \omega - \omega_{c0} + E_0$, which is the shifted frequency used in the calculation of the fourier transform. Now, we define the shifted resonance frequency, which is the one associated with $\bar{\omega}$ rather than ω . Since the original vertical resonance frequency is defined as $\omega_{res} = V_d(R_0^g) - V_g(R_0^g) - E_0$, where V_d and V_g are the core-excited and ground state potentials respectively, and R_0^g is the minimum position of the ground state potential, we may write $\bar{\omega}_{res} = \omega_{res} - \omega_{c0} + E_0$, so, summarizing, we have

$$\bar{\omega} = \omega - \omega_{c0} + E_0 \quad (8)$$

$$\omega_{res} = V_d(R_0^g) - V_g(R_0^g) - E_0 \quad (9)$$

$$\bar{\omega}_{res} = \omega_{res} - \omega_{c0} + E_0 \quad (10)$$

$$\Omega = \omega - \omega_{res} = \bar{\omega} - \bar{\omega}_{res} \quad (11)$$

this is the way the program computes different photon frequencies, by using the shift property of the Fourier transform. And it requires as input the shifted resonance frequency $\bar{\omega}_{res}$ and the desired values of detuning Ω .

Then we solve it using FFT techniques (such as the FFTW library). We note, however, that to take full advantage of this methodology, one should request all desired wavepackets in a single run, since the computational cost of obtaining the intermediate wavepacket $|\Phi(0)\rangle$ for a single photon frequency, and for several ones is virtually the same. For this reason everytime you run the raman code, it will save to disk a file with the spline coefficients (fft_spline.bcoef) of the fourier transformed wavepacket, so that on a second run, the computational time of obtaining a different $|\Phi(0)\rangle$ will be just the one of reading the spline file and printing the desired function.

III. THE run_raman.sh SCRIPT

The run_raman.sh script, was written following the ideas outline in the previous section. Before running, you should edit the path to the three required programs in the header of the scripts, aside from that, no further editing is necessary. It runs in several modes, which are given as flags when running the script. the -all flag, runs all three steps of the calculation. -init only runs the propagation of the initial state on the intermediate potential. -cond only generates the new initial conditions (you must have run -init previously) from a given propagation

So you should run it like this

```
$ ./run_raman.sh -flag input
```

-init in this mode, the script solves the time independent problem to obtain the ground state wavefunction $|0\rangle$, then proceeds to propagate it on the core-excited potential. The wavefunction evolution is saved on disk to a series of files called ReIm_, which are kept in the folder wf_data for the next step

-cond in this mode, we run the eSPec-raman module to obtain the $|\Phi(0)\rangle$ for the desired values of detuning (and bending states). The wavefunctions generated are saved to files, for the next step.

-fin in this mode, we run eSPec to propagate the generated $|\Phi(0)\rangle$ on the final state potential, the program automatically computes the auto-correlation function, and subsequently the spectrum. The spectra for each detuning required is saved to files *.spec and have to be shifted like $\omega' = \omega - \bar{\omega}' - \omega_{0f} + E_0$, here $\bar{\omega}'$ is the abscissa in the eSPec output, ω' is the emitted photon frequency (desired abscissa), ω is the incoming photon frequency, ω_{0f} is the energy difference between the ground and final state, finally E_0 is the ground state vibrational energy. This happens because eSPec shifts all potentials to zero.

-all intuitively, runs all the option above on a single shot (This mode can be a bit messy on first runs, due to input errors and such, so I recomend running step-by-step at the beginning)

IV. INPUT KEYWORDS

Now we'll go into details of the input file the script reads.

here is an example of some essential keywords required

```
# RIXS input
#
jobid test
```

```

dimension .2D
npoints 100 100
mass 12.00 12.00

initial_pot test2d.init
decaying_pot test2d.decay
final_pot test2d.final

gamma 0.05
step 5D-4
detuning -2.0 -1.0 -0.1 0.0 0.1 1.0 2.0 4.0

Vg_min 0.0000
Vd_min 90.7708
Vf_min 0.367493
Vd_vert 90.7758

fin_time 140.0

absorb_cond 0.0001
absorb_range 0.00 10.00 0.000 10.00

```

Now, we detail each one of them

jobid reads a string to serve as name base for all output files (necessary)

dimension options .1D, .2D and .2DCT, which specifies the dimension of the calculation

npoints number of discretization points on each dimension

mass mass in AMU, associated with each dimension (remember to input the three masses in .2DCT calculations)

initial_pot expects the name of the file containing the ground state potential.

initial_wf if you have already computed the initial wavefunction, then you should use this keyword followed by the file containing the initial wavefunction. In this case the keyword **initial_pot** is not used. It is desirable that the wavefunction is normalized with unitary euclidian norm.

E0 if you are using a wavefunction computed previously (**initial_wf**) then you might need to include this keyword, followed by the energy of said wavefunction. The exception being if you already have a file named \$jobid_init.out, from a previous -init run.

decayin_pot expects the name of the file containing the intermediate state potential.

final_pot expects the name of the file containing the final state potential.

gamma Core-excited lifetime broadening in eV.

step Time step to be used in the propagations (10^{-4} recommended, for test runs larger steps may be used).

fin_time Propagation time (femtoseconds) on final state potential.

Vg_min Potential at the minimum of the ground state potential $V_g(R_0^g)$, in a.u.

Vd_min Potential at the minimum of the decaying state potential $V_d(R_0^d)$, in a.u.

Vf_min Potential at the minimum of the final state potential $V_f(R_0^f)$, in a.u.

Vd_vert Potential at the vertical transition point in the decaying potential $V_d(R_0^g)$

All potential files must be given in a.u., and must contain the first line as a coment line starting with #. Also there must be only one empty line at the end of the file (empty in the sense that you just broke the line above, but no empty spaces). This is due to the way eSPec expects the potential files to be formatted.

absorb_cond reads the strenght of the damping applyied to the wavefunction. (there is no default, if the keyword is not given, then no absorbing conditions will be applied)

absorb_cond reads the range beyond which the absorbing conditions will be applied (a.u.). For instance, absorb_cond 0.000 10.000 on a .1D calculation, applies the damping function from 10.000 a.u. until the end of the grid. For .2D/.2DCT calculations four values are expected, naturally.

When choosing absorbing conditions, some trials might be required, since large values for **absorb_cond**

and short ranges might still result in unphysical reflections. Usually a value around 10^{-4} and a range of 1-2 a.u. may give good results, note however that the kinetic energy of the wavefunction also plays a role, since for large energies we may use a stronger damping, whereas for small energies such strong damping would result in reflections. In sum, some tweaking is advised, but usually it is fairly easy to find working conditions for parameters.

shift if this keyword is present, the script automatically shifts the final spectrum to be a function of the emitted photon energy ω' in eV.

print_level This keyword accepts four modes **minimal**, **essential**, **intermediate**, **full**. In **minimal** level, the script saves only the very important info to files, in each section, then after the -cross option is run, it erases everything, leaving only the final result. This erases the initial state propagation files, and also erases the `fft_spl.bcoef` file, so be cautious, as running this mode does not save any computational time if you wish to run a similar calculation a second time. In **essential**, which is the default, we save to disk only important output files, the initial propagation, and the `fft_spl.bcoef` file, the correlation functions are also saved. All input files are erased, as well as any temporary calculation files (which are relevant for debug). In **intermediate** everything from essential is saved, plus all output files. Finally **full** everything is saved to files in the disk, this causes the directory to be very messy, but can be very useful if you are facing some sort of error.

window this expects the parameter for the supergaussian window applied to the final spectrum (used to avoid aliasing). The default value is 0.00001. Small values result in broader peaks in the spectrum, naturally, large values of this parameters will lead to sharper peaks.

cross_term This keyword reads the value of $\cos\theta$ which multiplies the kinetic cross term of the hamiltonian in valence coordinates, according to Eq. (39) from Faris's file. We also remember that running this type of calculations you should input three masses in the **mass** keyword (μ_1 , μ_3 and m_2 in Eq. (39))

V. SPECIFIC CASE OF OUR 2D+1D MODEL

Here we take into account the bending modes, which, as we shall see, also can be obtained using the shift of the Fourier transform. In the same fashion as in the second section of this file, we will have

$$\Phi_{vc}(0, \omega - \omega_{c0} + E_0 - \epsilon_{vc} + \epsilon_0) = \int_0^\infty e^{-\Gamma t} e^{-i(\omega_{c0} - E_0 + \epsilon_{vc} - \epsilon_0)t} \phi_c(t) e^{i\omega t} dt, \quad (12)$$

so in this case we will have the shifted frequency and properties being

$$\bar{\omega} = \omega - \omega_{c0} + E_0 - \epsilon_{vc} + \epsilon_0 \quad (13)$$

$$\omega_{res} = V_d(R_0^g) - V_g(R_0^g) - E_0 \quad (14)$$

$$\bar{\omega}_{res} = \omega_{res} - \omega_{c0} + E_0 \quad (15)$$

$$\Omega = \omega - \omega_{res} = \bar{\omega} - \bar{\omega}_{res} + \epsilon_{vc} - \epsilon_0 \quad (16)$$

$$\bar{\omega} = \Omega + \bar{\omega}_{res} - \epsilon_{vc} + \epsilon_0 \quad (17)$$

so in the program we use eq (17) to obtain the functions associated with each bending state, using the shift keyword with the value $-\epsilon_{vc} + \epsilon_0$.

VI. run_raman_water.sh SCRIPT

in addition to the flags contained in the general raman script, we now have the following additional ones

-fc in this mode, we run the eSPec to obtain the bending mode franck-condon factors, which are saved into files `fc_0vc.dat` and `fc_vcvf.dat` for later use.

-cross in this mode, the final program `fcorrel` is run, which puts together all the previous steps into the final cross section (see eq. (51) from `WP_water.pdf` file).

VII. SPECIFIC INPUT KEYWORDS

In addition to the previous general keywords, we need a few extra ones in the case of our 2D+1D model

```
nvc 3
nvf 5
```

```
bend_npoints 256
bend_mass 12.00
```

```
bend_init_pot bend.init
bend_decay_pot bend.decay
bend_fin_pot bend.final
```

nvc number of core-excited bending states to be included.
nvf number of final bending states to be included.
bend_npoints number of discretization points in the bending potential.
bend_mass mass in AMU, associated with the bending mode.
bend_init_pot name of the file with the initial bending potential.
bend_decay_pot name of the file with the intermediate bending potential.
bend_fin_pot name of the file with the final bending potential.

VIII. COMPILING THE PROGRAMS

The raman and fcorrel programs can be downloaded from <https://github.com/LTCC-UFG/eSPec-RAMAN>.

Compiling is very simple, all three programs have their own makefile, so for eSPec, you should just run

```
$ make espec
```

eSPec does not require any additional libraries, as they are included in the code.

For raman and fcorrel, you should just run make, with no specific targets.

```
$ make
```

For these programs, however you should have installed the fftw library (available at <http://www.fftw.org/>).

On a final note, we suggest that you compile the programs with ifort (intel compiler), specially eSPec, since we've had problems compiling it with the latest gfortran.