
PYQCTools Documentation

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PYQCTools is a collection of python scripts useful to dump quantum chemistry integrals and to perform data post-processing for different quantum chemistry methods.

PYQCTools requires the following prerequisites to work:

- Python 2.6, 2.7, 3.2, 3.3, 3.4
- Numpy 1.6.2 or higher
- Scipy 0.10 or higher (0.12.0 or higher for python 3.3, 3.4)
- [PySCF](#) for Integrals Calculation in Integrals Dumpings scripts.

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1.1 Tools for Omega Space Green's Functions

gf_trace.py: Calculate the Density of States (DOS) value from an ω -dependent Green's Function. It makes the trace of the Green's Function associated with a certain frequency value.

Example :

```
>>> python gf_trace.py /PATH/green.txt omega_value
```

green.txt: formatted text file containing the Green's function.

omega_value: double frequency value.

1.2 Tools for Real-Time Green's Functions

rtgf.py: Calculate the Density of States (DOS) values during a time propagation. It makes the trace of time-dependent Green's Functions calculated along a time propagation and return the DOS values as a function of time both for the real and for the imaginary part of the Green's Function.

Example :

```
>>> python rtgf.py prop_time time_step /PATH/scratch
```

prop_time: double value of the full propagation time (period).

time_step: double value of the time-step.

scratch: directory containing text files of the real (green.\$t.\$t.txt) and imaginary (green.30000+\$t.30000+\$t.txt) Green's Functions, where \$t indicate the specific time-step.

The script save two output files, `rt_real.txt` and `rt_imag.txt`, containing the real and imaginary part of the time-dependent DOS respectively.

fft.py: Perform the fourier transform of the time-dependent Density of States (DOS). It reads the `rt_real.txt` and `rt_imag.txt` generated by the `rtgf.py` script and produces the `ldos.out` and `real_part.txt` files containing the imaginary and real parts of the ω -dependent DOS respectively.

Example :

```
>>> python fft.py broad rem_add
```

broad: double value of the imaginary broadening.

rem_add: string specifying if we are working with the addition or removal part of the Green's Function. It can assume only the values 'add' or 'rem'.

1.3 Tools for Integrals Dumping

You can also download the [PDF version](#) of this manual.