# **PYQCTools Documentation**

Release 1.0

**Enrico Ronca** 

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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  $\omega$ -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 timedependent 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 *omega*-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.