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# FC-DFT

*Release 1.0.0*

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# Chapter 1

## Installation

This page explains how to install FC-DFT code.

### 1.1 Requirements

- GeomeTRIC
- PySCF
- PyAMG
- PyAMGCL
- GPU4PySCF (optional)

### 1.2 How to install

Download the latest version of FC-DFT from the repository:

1. Clone the repository:

```
$ git clone https://github.com/Yang-Laboratory/FC-DFT.git
```

2. Change the directory to the cloned repository:

```
$ cd FC-DFT
```

3. Install the package using *pip*:

```
$ pip install .
```

4. Change directory to *\$PYTHONPATH/fcdft/lib* and create build directory.

5. Go into *build* and compile the C shared libraries by *cmake ..* and *make*.



# Chapter 2

## Examples

This chapter introduces brief examples of running FC-DFT and Poisson-Boltzmann solvation calculations

### 2.1 Wide-Band Limit

FC-DFT uses the wide-band limit (WBL) approximation. In particular, the WBL-Molecule approximation is adopted for simple but powerful implementation. WBL-Molecule requires users to provide the imaginary part of the self-energy in WBLMolecule object. Currently, spin-restricted version of FC-DFT (fcdft.wbl.rks) is supported. Below is a sample code, where the self-energy of 0.01 eV is attached to the sulfur atom of methanethiol with 25.95 electrons.

```
>>> from pyscf import gto
>>> from pyscf.dft import RKS
>>> mol = gto.M(atom='''
...     C    -1.718553971   -0.000000250   -0.626147715
...     H    -2.739245971   -0.008907250   -0.227127715
...     H    -1.200493971   -0.879491250   -0.227127715
...     H    -1.215921971    0.888398750   -0.227127715
...     S    -1.718553971   -0.000000250   -2.396147715
...     H    -2.150082583    0.805150681   -2.710667448''',
...     charge=0, basis='6-31g**')
>>> mf = RKS(mol, xc='pbe')
>>> mf.kernel()
>>> from fcdft.wbl.rks import *
>>> wblmf = WBLMolecule(mf, broad=0.01, nelectron=25.95)
>>> wblmf.kernel()
```

### 2.2 Non-Linear Poisson-Boltzmann Solvation Model

We provide the Poisson-Boltzmann solver for general purpose. fcdft.solvent.pbe module supports usual solvation energy calculations as what polarizable continuum model does. To do so, a few attributes of PBE needs to be controlled since it was originally intended to solve the electrostatic potential under the Gouy-Chapman-Stern theory:

```
>>> from pyscf import gto
>>> from pyscf.dft import RKS
>>> mol = gto.M(atom='''
...     O    0.152427064    0.959723218   -2.275350162
```

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```
...      H      0.152427064    1.719060218   -1.679307162
...      H      0.152427064    0.200386218   -1.679307162"',
...      charge=0, basis='6-31g**1')
>>> mf = RKS(mol, xc='b3lyp')
>>> from fcdft.solvent.pbe import *
>>> cm = PBE(mol, cb=1.0, length=15, ngrids=41)
>>> cm.eps = 78.3553
>>> cm.atom_bottom = 'center'
>>> cm.nelectron = mol.nelectron
>>> cm.bias = 0.0e0
>>> cm.surf = 0.0e0
>>> solmf = pbe_for_scf(mf, cm)
>>> solmf.kernel()
```

The following example introduces how to run the Poisson-Boltzmann solver under the Gouy-Chapman-Stern boundary values:

```
>>> from pyscf import gto
>>> from pyscf.dft import RKS
>>> mol = gto.M(atom='''
...      C      -1.718553971   -0.000000250   -0.626147715
...      H      -2.739245971   -0.008907250   -0.227127715
...      H      -1.200493971   -0.879491250   -0.227127715
...      H      -1.215921971    0.888398750   -0.227127715
...      S      -1.718553971   -0.000000250   -2.396147715
...      H      -2.150082583    0.805150681   -2.710667448''',
...      charge=0, basis='6-31g**1')
>>> mf = RKS(mol, xc='pbe')
>>> mf.kernel()
>>> from fcdft.wbl.rks import *
>>> wblmf = WBLMolecule(mf, broad=0.01, nelectron=25.95)
>>> wblmf.kernel()
>>> dm = wblmf.make_rdm1()
>>> from fcdft.solvent.pbe import *
>>> cm = PBE(mol, cb=1.0, length=15, ngrids=41, stern_sam=3.0)
>>> cm.eps = 78.3553
>>> cm.eps_sam = 2.284
>>> cm._dm = dm
>>> solmf = pbe_for_scf(wblmf, cm)
>>> solmf.kernel()
```

## 2.3 Geometry Optimization

Our code supports analytic nuclear gradients of FC-DFT as well as the Poisson-Boltzmann solvation model. We have tested geometry optimization using GeomeTRIC, an external geometry optimizer implemented in PySCF:

```
>>> from pyscf.geomopt.geometric_solver import optimize
>>> moleq = optimize(solmf, maxstep=100)
```

## 2.4 Thermochemistry

We provide a code for numerical Hessian matrix constructed by analytic forces due to the non-Hermitian Hamiltonian resulted by the self-energy. Thermochemical properties can be calculated by utilizing pycf.hessian.thermo module. Hessian offers three-point (default) and five-point finite difference method for calculating the Hessian matrix. The following code introduces how to obtain thermochemical properties using harmonic\_analysis function:

```
>>> from fcdft.hessian numhess import *
>>> hessmf = Hessian(solmf)
>>> hess = hessmf.kernel()
>>> from pycf.hessian.thermo import harmonic_analysis
>>> freq_info = harmonic_analysis(moleq, hess)
```

Once the quantities are obtained, these can be saved into a molden format as implemented in fcdft.tools.molden:

```
>>> from fcdft.tools.molden import dump_freq
>>> dump_freq(moleq, freq_info, 'freq.molden')
```



# Chapter 3

## About

Fractional Charge Density Functional Theory (FC-DFT) is a theory that reformulates open quantum systems in terms of the canonical ensemble. The prototype of FC-DFT is linear interpolation FC-DFT (LI-FC-DFT), which enforces the Perdew-Parr-Levy-Baldur (PPLB) condition to DFT calculations through linear interpolation and bypasses the delocalization error.

### 3.1 Developers

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### 3.2 Citation

Jun-Hyeong Kim, Dongju Kim, Weitao Yang, and Mu-Hyun Baik. Fractional Charge Density Functional Theory and Its Application to the Electro-inductive Effect. *J. Phys. Chem. Lett.* **2023**, *14*, 3329-3334 Jun-Hyeong Kim and Weitao Yang. Fractional Charge Density Functional Theory Elucidates Electro-Inductive and Electric Field Effects at Electrochemical Interfaces. *Submitted*

### 3.3 Bug Report and Feature Request

Please open a thread on the [Issues](#) tab.



## Chapter 4

# Indices and tables

- `genindex`
- `modindex`
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