Contents:
PyChemia is a python library for automatize atomistic simulations. PyChemia is build around a core module with two classes and a set of some other modules that offers a variety of operations in order to perform more complex operations.

The core is made of two classes ‘Structure’ and ‘Composition’. Other modules are:

- **analysis**: Uses pure structure information for changing structures, matching atoms between two structures, create slabs, surfaces and some other geometrical operations on structures.
- **code**: This module deals with creating inputs and reading outputs from several atomistic simulation codes. Right now, we support ABINIT, DFTB+, Fireball, an internal calculator for LennardJones Clusters, Octopus and VASP.
- **core**: The core of PyChemia are two classes that are imported at the root level of the library: Structure and Composition. For PyChemia, Structure is a set of sites with one or more atoms located on each site with a probability associated to them. The Structure could be finite or periodic in one or more directions. In the case of a crystal the Structure will have also a lattice. Composition is a set of atoms of a define species. For crystals is the set of atoms on each unit cell. No geometry is store on a Composition object, and the order of atoms is irrelevant.
- **crystal**: For the case of periodic structures in three directions, a set of modules is created for three basic properties of a crystal. The class ‘KPoints’ store the description a a k-point mesh, path, or direct list of points on the reciprocal lattice. The class ‘Lattice’ store and manipulate the cell vectors of a crystal. The third class is ‘CrystalSymmetry’ for computing the Space Group, getting structures on the Bravais cell and finding the primitive cell.
- **db**: PyChemia uses MongoDB as Database for storing Structures and the properties computed by atomistic simulation codes. There are two classes defined, PyChemiaDB to store structures and properties and PyChemiaQueue to store a massive set of calculations and the status of those calculations.
- **dm**: This is a module in development, it contains classes for DataMining PyChemia Databases and global searches. Right now it contains a class for Network analysis, but in the future will provide interfaces with some other libraries for machine learning.
- **evaluator**: There are two circumstances where a atomistic simulation can be perform. If you have a machine without a queue system PyChemia will provide a very simplified queue for computing concurrent calcu-
lations under the constrains of your number of cores. If you use a queue system, the evaluator will setup the batch scripts, submit the jobs and monitor the status of those jobs, finally when the job is done, it will collect the final data and update the databases.

- **external**: There are some other packages for which PyChemia worth interact, ASE (Atomistic Simulation Environment) is a python package supporting a remarkable number of calculators. Findsym is a close software to compute spacegroups and computing the CIF file of a give structure. ‘pymatgen’ is python code behind ‘Materials Project’ and a outstanding piece of very well written code. Originally implemented for VASP only now includes support for ABINIT.

- **io**: There are a pletora of formats for describing atomistic structures. We support three basic file-formats, ascii, CIF and XYZ. This module provides the classes for reading and writing them.

- **md**: Molecular Dynamics is an important operation for atomistic simulations. This module provide a ‘in-house’ calculator for MD simulations, using the forces computed from static calculations from an external atomistic code.

- **population**: A population is basically a collection of candidates collected somehow by a global searcher or from a DB query. PyChemia includes classes for populations of Lennard-Jones clusters, populations of vectors on a N-dimensional space.

- **runner**: Controls the executions in cluster by creating batch scripts and checking the status of the queue. Right now only supports Torque

- **searcher**: Global search operations, the prototypical case is structural search but any pychemia population could be use for searching. Several metaheuristic algorithms have been implemented

- **utils**: Several small routines, like a periodic table, mathematical operations, conversions and serializers.

- **visual**: PyChemia includes a set of classes and routines for interfacing with some other libraries and external software for data visualization, 3D plotting and graphic representation. pyprocar plots band-structures, LatticePlot and StructurePlot uses mayavi for visualizing atomic configurations and lattices. We have also interfaces for Povray and a developing interface to D3.js.

- **web**: On development, creation of a web interface for looking into the pychemia databases and controlling executions. The web interface is build on top of Python-Bottle and CherryPy to access the database.

PyChemia is a open-source Python library for High-throughput first-principles materials discovery. The focus of this library is on structural search and data analysis. The ultimate purpose of the code is to optimize the search of new materials using a variety of methods such as Minima hopping method (MHM), soft-computing-based methods and statistical methods.

The main objectives of the code are:

1. Provide flexible classes for atomic structures such as molecules, clusters, thin films and crystals.
2. Manipulate both input and output for DFT and Tight-binding codes such as VASP, ABINIT, Fireball and DFTB+
3. Offer a robust architecture for storing a large collection of structures. Structural search methods generate many structures that are stored in repositories. Calculations done on those structures are also store in repositories.
4. Similarity analysis based on fingerprints, pair correlation and comparators.
5. Stability analysis for crystals. Including thermal analysis (Enthalpy) and dynamic stability (Phonons)
6. Tools for producing comprehensive reports, convex hulls, band structures, density of states, etc
7. Datamining tools to extract knowledge from the structures found, identify patterns in the data and identify suitable candidates for technological applications
8. A web interface

This is a new project and many classes and methods are refactored frequently. This is and will be a work in progress. We hope to stabilize the most critical classes for the release 1.
This code is open-source. We also welcome extra hands to improve this library with your own contributions. At present only one developer has been in charge of the project. More hands and eyes are very welcomed.
CHAPTER 2

Installation

2.1 Quick and Simple

From a terminal install the latest stable version with pip:

```
pip install pychemia --user
```

It will install pychemia and all its dependencies in your $HOME folder. If you want a central installation you can install directly with:

```
sudo pip install pychemia
```

2.2 Installing PyChemia with pip

This is probably the easiest way, pip will download the code, check and eventually install dependencies and installing the package on a system-wide place or the home directory. All you have to do is execute this command:

```
sudo pip install pychemia
```

or for python 3.x:

```
sudo pip3 install pychemia
```

If you are on a machine where you do not have superuser privileges you can install pychemia on your home directory by adding the command ‘--user’:

```
pip install pychemia --user
```

or for python 3.x:

```
pip3 install pychemia --user
```
Uninstall or upgrade pychemia is also very simple with pip:

```
pip uninstall pychemia
pip install pychemia --upgrade
```

### 2.3 Installing PyChemia with virtualenv

PyChemia requires a relatively recent version of Python (Python 2.7 appeared on July 3, 2010). However, it is common for some HPC infrastructures to use Linux installations such as RHEL 6 that only provides python 2.6 dating back to 2008. You can contour this situation by using Software Collections:

[https://www.softwarecollections.org/en/](https://www.softwarecollections.org/en/)

Install a more recent version of python and virtualenv. After that you can install PyChemia and its dependencies contained on a virtual environment following this commands:

1. Create a new virtual environment, let’s call it “venv_pcm”
   ```
   virtualenv venv_pcm
   ```
2. Activate the environment by sourcing the activate script
   ```
   source venv_pcm/bin/activate
   ```
3. You will notice that the prompt changes and you get your path pointing to a containerized python environment where you can install PyChemia and its dependencies
   ```
   pip install pychemia
   ```
4. Shortly after that, you will get all the dependencies installed and the software ready for use without touching the libraries and modules installed by your Linux distribution. You can leave the virtual environment with the command:
   ```
   deactivate
   ```

To re-enable the environment all you have to do is execute step 2 and you get the environment ready to work.

### 2.4 Installing PyChemia from github

The current stable repository for PyChemia is on Github, you can download the master branch with the command:

```
git clone https://github.com/MaterialsDiscovery/PyChemia.git
```

If you get a message such as:

```
$ git clone https://github.com/MaterialsDiscovery/PyChemia.git
-bash: git: command not found
```

You need to install `git` first. On machines from the Debian ‘lineage’ (Ubuntu, Mint, and many others) you can use the command:

```
sudo apt-get install git
```

On systems with `yum` you can use:
Now that you have ‘cloned’ the PyChemia repository you have two options. Install the package by using the set of commands:

```
    cd PyChemia
    python setup build
    python setup install --user
```

Use `python3` for the commands above, if you want to use python 3.x instead.

Another alternative is add path where you downloaded the repository to the variable $PYTHONPATH. You can do that by editing your `.bashrc` file. Supposing that you execute the `git clone` command directly on your home directory you can add the path for PyChemia adding this line to your `.bashrc`:

```
    export PYTHONPATH=$HOME/PyChemia:$PYTHONPATH
```

If you want the changes on `.bashrc` take immediate effect execute:

```
    source $HOME/.bashrc
```

## 2.5 Dependencies

PyChemia relies on the presence of at least this libraries:

```
    numpy >= 1.12.0
    scipy >= 0.18.0
    future >= 0.16.0
    spglib >= 1.9.9
    pymongo >= 3.4.0
```

Most Linux distributions will probably include older versions of those dependencies. HPC Linux clusters usually have a very conservative approach related to packages and even older versions of those dependencies will be installed the system or completely absent from it.

The packages `numpy` and `scipy` are well known python packages for Linear Algebra and general scientific computing. PyChemia is able to run on Python 2 and 3 using the same sources. The python package `future` is a small package that helps keeping compatibility with python 2 and python 3 on the same source code. The package `spglib` is a C-library with python wrappers for computing space groups and related functionality. The package `pymongo` offers connectivity with a MongoDB database. MongoDB is a document-based NoSQL database engine. PyChemia uses MongoDB databases for storing collections of structures or properties and parameters associated to them.

We try to keep the dependencies of PyChemia to a very minimum. Some other libraries provides extra functionality that could be necessary for some tasks. Consider install nose and matplotlib. You can do that using pip with the command:

```
    sudo pip install nose matplotlib
```

Remember that you can use `--user` if you want to install on your home directory without special privileges. Matplotlib is the ‘standard de-facto’ for 2D plots on python. Many of the functionalities on ‘pychemia.visual’ subpackage depends on it. Nose is a python package for executing automatize tests for PyChemia. If you want to use it more information is below.
2.6 Importing the library

No matter how you installed PyChemia, you should be able to load the library. You can use the traditional python terminal, for example:

```
$ python3.5
Python 3.5.1 (default, Mar 2 2016, 03:38:02)
[GCC 4.2.1 Compatible Apple LLVM 7.0.2 (clang-700.1.81)] on darwin
Type "help", "copyright", "credits" or "license" for more information.
>>> import pychemia
>>> pychemia.info()
PyChemia
--------
Version: 0.1.2
Path: /Users/guilleaf/PyChemia/pychemia
Date: May 13, 2016

Python version=3.5.1 (default, Mar 2 2016, 03:38:02)
[GCC 4.2.1 Compatible Apple LLVM 7.0.2 (clang-700.1.81)]

pymongo Not Found
numpy 1.11.0 /opt/local/Library/Frameworks/Python.framework/Versions/3.5/
   → lib/python3.5/site-packages/numpy
scipy 0.17.1 /opt/local/Library/Frameworks/Python.framework/Versions/3.5/
   → lib/python3.5/site-packages/scipy
mayavi Not Found
Scientific Not Found
matplotlib 1.5.1 /opt/local/Library/Frameworks/Python.framework/Versions/3.5/
   → lib/python3.5/site-packages/matplotlib
future 0.15.2 /opt/local/Library/Frameworks/Python.framework/Versions/3.5/
   → lib/python3.5/site-packages/future
coverage 4.0.3 /opt/local/Library/Frameworks/Python.framework/Versions/3.5/
   → lib/python3.5/site-packages/coverage

spglib 1.9.4 /Users/guilleaf/Library/Python/3.5/lib/python/site-packages/
   → spglib
pyhull Not Found
pymatgen Not Found
qmpy Not Found
ase Not Found
```

The method `pychemia.info()` will inform about the several libraries that PyChemia uses, both mandatory and optional, their versions and path. That could be informative in case of something not working as expected.

2.7 Testing with nose

It is always important to test a library, not only from the developer point of view, but also for an user. Nose is a python package that offers a simple command to execute predefined test for a python package and report any errors or inconsistencies from the expected results.

Direct the terminal to the place where PyChemia is located. Lets suppose that you have `pychemia` on `/Users/guilleaf/PyChemia`, test PyChemia using the command:
The name of the command could have small variations according to your distribution. On a MacOS using Macports the name could be for example nosetests-2.7 or nosetests-3.5 for python 2 and 3 respectively.

nosetests will search for tests on the entire package and subpackages. If everything is fine (and you use \texttt{--v} for verbose output) you will get something like:

\begin{verbatim}
... 
Example of a simple calc : ... ok
Example of a multiple calc : ... ok

Ran 38 tests in 5.469s
OK
\end{verbatim}

That is an indication that all tests were successful and eventually you are ready to use the library.
PyChemia is built from several layers of code, the foundations being the ability to manipulate conglomerates of atoms, either molecules or crystals. Before doing more complex operations such execute tasks on various electronic structure codes or running structural searches we need to understand how the fundamental blocks work and that is the purpose of this quick start section.

For the examples presented here you can use the official python terminal, however, the IPython terminal or a Jupyter notebook offers you more power to go across the examples and explore what PyChemia can do for you. At least, installing IPython is very simple on many operating systems.

For example on machines running a debian derivative (Ubuntu, Mint and several others) you can use the following command to install the IPython terminal:

```
sudo apt-get install python-ipython
```

or for python 3.x:

```
sudo apt-get install python3-ipython
```

On MacOS using macports:

```
sudo port install py27-ipython
```

or for python 3.x:

```
sudo port install py35-ipython
```

### 3.1 Element

PyChemia is about materials, molecules, cluster and crystals. A class for elements can become handy to serve as a periodic table and inquire for the properties of a given specie. Alchemy was all about transmuting materials into gold. Lets explore what PyChemia can tell us about that element:
In [1]: `import pychemia`

In [2]: `ele=pychemia.Element('Au')`

In [3]: `ele.atomic_number`
Out[3]: 79

In [4]: `ele.name`
Out[4]: 'Gold'

In [5]: `ele.electronegativity`
Out[5]: 2.54

In [6]: `ele.group`
Out[6]: 11

In [7]: `ele.period`
Out[7]: 6

In [8]: `ele.block`
Out[8]: 'd'

In [9]: `ele.mass`
Out[9]: 196.96654

In [10]: `ele.electronic_configuration`
Out[10]: ['Xe', '6s1', '4f14', '5d10']

In [11]: `ele.is_madelung_exception`
Out[11]: True

In [12]: `ele.oxidation_states`
Out[12]: (-3, -2, -1, 1, 2, 3, 5)

In [13]: `ele.oxidation_states_common`
Out[13]: (3,)

In [14]: `ele.covalent_radius`
Out[14]: 1.36

The Element class provide a set of useful properties about all elements in the periodic table. The chapter about Element will explain the several properties that can be accessed on the Element class.

### 3.2 Composition

Now, atoms group themselves into molecules, clusters or crystals. Without entering into the geometry, we can think only about the composition, the list of atoms contained in the molecule or cluster or the list of atoms in the cell that builds the crystal. The Composition class offers a set of routines to manipulate lists of atoms. Lets consider for example Gold(III) oxide (Au2O3):

In [1]: `import pychemia`

In [2]: `comp=pychemia.Composition('Au2O3')`

In [3]: `comp`

(continues on next page)
Compositions can be created from the formula and also from a python dictionary and after you can use the composition object to ask for the number of atoms of each specie.

### 3.3 Structure

In most cases, pychemia uses Structures. A PyChemia Structure stores atomic positions and cell parameters for periodic structures. Let's start with a very simple structure, the FCC structure of gold. One way of creating and structure is by directly specifying atomic positions and cell parameters, for example:

```python
In [1]: import pychemia
In [2]: a=4.05
In [3]: b=a/2
In [4]: fcc = pychemia.Structure(symbols=['Au'], cell=[[0, b, b], [b, 0, b], [b, b, 0]], periodicity=True)

In [5]: fcc
Out[5]: Structure(symbols=['Au'], cell=[[0.0, 2.025, 2.025], [2.025, 0.0, 2.025], [2.025, 2.025, 0.0]], reduced=[[0.0, 0.0, 0.0]], periodicity=True)
```

You have created your first structure on PyChemia. The structure contains one atom of Gold and the cell is the primitive cell of a FCC structure with one atom. You can 'see' the structure using the 'print' function:

```python
In [6]: print(fcc)
```

<table>
<thead>
<tr>
<th>Symb ( Positions )</th>
<th>Cell-reduced coordinates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Au ( 0.0000 0.0000 0.0000 )</td>
<td>0.0000 0.0000 0.0000</td>
</tr>
</tbody>
</table>

Periodicity: X Y Z

Lattice vectors:

| 0.0000 2.0250 2.0250 |
| 2.0250 0.0000 2.0250 |
| 2.0250 2.0250 0.0000 |

We can now be interested in see the spacegroup of the structure just created. A pychemia.crystal.CrystalSymmetry object allow us to get access to symmetry calculations provided by the spglib library:
Both the space group number and the International symbol are accessible from the CrystalSymmetry object. Now we can use the CrystalSymmetry object to recreate the conventional FCC cell:

```python
In [10]: fcc_conv=cs.refine_cell()

In [11]: fcc
fcc    fcc_conv

In [12]: fcc_conv
Out[12]: Structure(symbols=['Au', 'Au', 'Au', 'Au'], cell=4.05, reduced=[[0.0, 0.0, 0.0], [0.0, 0.5, 0.5], [0.5, 0.0, 0.5], [0.5, 0.5, 0.0]], periodicity=True)

In [11]: print(fcc_conv)
4
Symb  (Positions)  [Cell-reduced coordinates]
Au   (0.0000 0.0000 0.0000) [0.0000 0.0000 0.0000]
Au   (0.0000 2.0250 2.0250) [0.0000 0.5000 0.5000]
Au   (2.0250 0.0000 2.0250) [0.5000 0.0000 0.5000]
Au   (2.0250 2.0250 0.0000) [0.5000 0.5000 0.0000]

Periodicity: X Y Z
Lattice vectors:
4.0500 0.0000 0.0000
0.0000 4.0500 0.0000
0.0000 0.0000 4.0500

The lattice vectors are now making a cube, and the structure is made of 4 gold atoms. Now we can want to create a supercell, 2x2x1 using the supercell method:

```python
In [13]: fcc221=fcc_conv.supercell(size=(2,2,1))

In [14]: print(fcc221)
16
Symb  (Positions)  [Cell-reduced coordinates]
Au   (0.0000 0.0000 0.0000) [0.0000 0.0000 0.0000]
Au   (0.0000 2.0250 2.0250) [0.0000 0.2500 0.5000]
Au   (2.0250 0.0000 2.0250) [0.2500 0.0000 0.5000]
Au   (2.0250 2.0250 0.0000) [0.2500 0.2500 0.0000]
Au   (0.0000 4.0500 0.0000) [0.0000 0.5000 0.0000]
Au   (0.0000 6.0750 2.0250) [0.0000 0.7500 0.5000]
Au   (2.0250 4.0500 2.0250) [0.2500 0.5000 0.5000]
Au   (2.0250 6.0750 0.0000) [0.2500 0.7500 0.0000]
Au   (4.0500 0.0000 0.0000) [0.5000 0.0000 0.0000]
Au   (4.0500 2.0250 2.0250) [0.5000 0.2500 0.5000]
Au   (6.0750 0.0000 2.0250) [0.7500 0.0000 0.5000]

(continues on next page)
PyChemia is much more than just elements, compositions and structures. For the purpose of this introduction, those are the most basic building blocks for most of the of higher level operations such as populations and global searches.

### 3.4 Lattice

For periodic structures, the lattice is fundamental to describe how the geometry of the atoms is repeated. PyChemia offers a class Lattice for manipulate cell parameters both for the real space lattice and the reciprocal one. For example from the example before:

```python
In [1]: import pychemia

In [2]: a=4.05

In [3]: b=a/2

In [4]: fcc = pychemia.Structure(symbols=['Au'], cell=[[0, b, b], [b, 0, b], [b, b, 0]], periodicity=True)

In [5]: lat=fcc.lattice

In [6]: print(lat)
Cell= 0.000 2.025 2.025
    2.025 0.000 2.025
    2.025 2.025 0.000
Angles: alpha = 60.000
        beta = 60.000
        gamma = 60.000
Lengths:  a = 2.864
          b = 2.864
          c = 2.864

In [7]: rlat = lat.reciprocal()

In [8]: print(rlat)
Cell= -0.247 0.247 0.247
     0.247 -0.247 0.247
     0.247 0.247 -0.247
```

(continues on next page)
As you can see from the example above, the reciprocal lattice can be obtained from the real space lattice. The class is useful to obtain the Brillouin zone via computing the Wigner Seitz from the reciprocal lattice. Those and several other functionality can be accessed from the class as described on the corresponding chapter.

| Angles: alpha =     | 109.471 |
| beta =             | 109.471 |
| gamma =            | 109.471 |

| Lengths: a =       | 0.428   |
| b =                | 0.428   |
| c =                | 0.428   |
4.1 Global Minimization of Lennard-Jones Clusters

This tutorial will guide you on how to search for global minima using the methods implemented on PyChemia.

4.1.1 Quick version

The shortest version of a global search using the FireFly method will look like this:

```python
>>> from pychemia.searcher import FireFly
>>> from pychemia.population import LJCluster
>>> popu = LJCluster('LJ13', composition='Xe13', refine=True, direct_evaluation=True)
>>> searcher = FireFly(popu, generation_size=16, stabilization_limit=10)
>>> searcher.run()
```

For this case, you should have a mongo server running on your local machine, no SSL encryption and no authorization with username and password. The population will be created with Lennard-Jones clusters with 13 particles each. Each new candidate is locally relaxed when created. The searcher will use 16 candidates on each generation and will stop when the best candidate survives for 10 generations.

4.2 PyChemia Software Framework

PyChemia is a framework for materials discovery, more than just compute DFT calculations for structures already present in databases such as ICSD, PyChemia searches for new structures that could never have been synthesized or
To achieve its goal, PyChemia relies on a number of methods of structural search such as minima hoping method, genetic algorithms and other population based algorithms.

As a software framework, PyChemia is structurated around five axis. They are:

1. Structural Search Methods
2. Storage and Databases
3. Data Mining or Knowledge Discovery in Data
4. Execution and Analysis
5. Reporting and Visualization

We will develop those five axis and the relations between them.

### 4.2.1 Structural Search Methods

One of the distinctive characteristics of PyChemia is its ability to search for new structures. Ab-initio calculations of structures present of databases such as ICSD becomes limited to the extension of the original databases. Even if such effort is indeed a big challenge with large databases, those databases represent a small portion of the structures that could be created. The ability to predict new structures and target those findings to specific applications is a task of technological relevance.

PyChemia was created with an strong focus on structural prediction. PyChemia implements several methods, from minima hoping method to metaheuristics.

#### 4.2.1.1 Minima Hoping Method

#### 4.2.1.2 Metaheuristics

### 4.2.2 Storage and Databases

If PyChemia were a software to compute ab-initio calculations from structures taken from another database or from very predictive set of prototypes only one database will be enough. However, as we mention before PyChemia is focused on structural search.

The kind of algorithms that we describe above typically explore thousands of different structures before selecting a reduced subset of thermodynamically stable or metastable ones.

Also, the search for new structures must be guided by specific applications of interest, batteries, thermoelectrics, superconductors, etc. Different applications are associated to different properties in the electronic structure.

Those two elements, the dynamic nature of structural search methods and the flexibility in the data that we would like to store is the reason why we are not using a single database and why we departed from traditional SQL schemas.

PyChemia was designed to create new databases for each structural search. We still keep one large database with a curated selection of structures but the ability to create small and flexible databases is central for the success of PyChemia.

PyChemia relies on MongoDB. MongoDB is an open-source document database, and the leading NoSQL database. We take advantage of dynamic schemas that offer simplicity and power and are in harmony with the own principles of scientific research. Different structures are intended for different applications and different applications requires the calculation of different physical properties.
4.2.3 Data Mining or Knowledge Discovery in Data

Data Mining or more correctly Knowledge Discovery in Data is the computational process of discovering patterns in large data sets involving methods at the intersection of artificial intelligence, machine learning, statistics, and database systems.

Structural Search algorithms has the ability to generate in the long term more structures that those that could be synthesized in reasonable time. As the database becomes large traditional techniques based on estimate good candidates based on just a few properties will be replaced by automatize algorithms that search patterns in structures and predict were new stochiometries deserve exploration.

4.2.3.1 Bayesian networks and Gaussian Processes

4.2.4 Reporting and Visualization

Doing electronic structure calculations for thousands of materials is challenging but there is not point in doing that without the ability to communicate to experimentalist structures that worth to be synthesized. PyChemia uses Django as a web-frontend to display and explore the computations not only on the global database but also on the specific database product of a structural search run.

4.2.5 Execution and Analysis

There are two levels of execution in structural search runs. Ab-initio calculations and structural analysis. PyChemia provides concurrent execution and basic job management for running multiple ab-initio calculations on clusters or non-queue systems

PyChemia relies on state-of-the-art ab-initio software packages to compute electronic structure and their properties. In particular PyChemia has support for VASP, ABINIT, Octopus, Fireball and DFTB+

Different levels of theory allows for a better compromise between accuracy and computational cost.

Structural analysis is defined as all kinds of procedures that relies only on structural data, for example structural fingerprints, topological analysis, hardness calculations, comparators and symmetry calculations. PyChemia provides multi-threaded execution of structural analysis routines

4.3 Optimization of Magnetic Moments (VASP)

This tutorial shows how to perform global search for optimal magnetic moments on VASP.

In VASP there are two variables that control the magnetic moments imposed on atoms and for contrain magnetic moments along predefined directions.

Lets start with one INCAR file that looks like this:

```
$SYSTEM  =  CaMnO3 Pnma
PREC     =  Accurate
NELMIN   =  6
NELM     =  100
EDIFF    =  1E-09
EDIFFG   =  -5E-6
IBRION   =  1
ISIF     =  2
LREAL    =  .FALSE.
ADDGRID  =  .TRUE.
```

(continues on next page)
NSW = 0
ISMEAR = -5
ENCUT = 500
ISPIN = 2
LORBIT = 11
LMAXMIX = 4
ISYM = 0
LSORBIT = .TRUE.
RWIGS = 2.00 1.87 1.78 1.24 # d-d/2
SAXIS = 0 0 1
LPLANE = .TRUE.
NPAR = 2
LSCALU = .FALSE.
NSIM = 4
LWAVE = .FALSE.
AMIX = 0.8
BMIX = 0.9
AMIX_MAG = 0.4
BMIX_MAG = 0.9

MAGMOM = 12x0
0.0000000 0.0416472 2.3859677
0.0000000 0.0416472 -2.3859677
0.0000000 0.0416472 -2.3859677
0.0000000 0.0416472 2.3859677
60x0
M_CONSTR = 12x0
0.0000000 0.0416472 2.3859677
0.0000000 0.0416472 -2.3859677
0.0000000 0.0416472 -2.3859677
0.0000000 0.0416472 2.3859677
60x0
I_CONSTRAINED_M = 1
LAMBDA = 10

We have commented the variables related with LDA+U but the procedure works enabling those variables too. The vari-
ables MAGMOM and M_CONSTR controls the initial direction of Magnetic Moments and the constrained direction
using LAMBDA as a parameter to control the intensity of the contrain.

Now, for different values of MAGMOM you can get variations on the total energy and the optimal magnetization can
only be obtained by covering all possible directions.

The population ‘NonCollinearMagMoms’ defined the procedures to create a pool of candidates with random direc-
tions for Magnetic Moments and modify their directions in several ways suitable for being used by the global search
algorithms implemented on PyChemia.

On this tutorial we will explore step by step how the methods on ‘NonCollinearMagMoms’ were implemented and
how use them for efficiently
4.4 Global optimization of correlation matrices for DFT+U (Abinit)

4.4.1 The variable dmatpawu

The objective of this global search is finding the optimal values for the density matrices in DFT+U. ABINIT allows to locally optimize the density matrices from a given initial value. The initial density matrices used in LDA+U are kept fixed during the first usedmatpu SCF iterations. For SCF iterations beyond that, the density matrices change. The challenge here is that from a given initial set of density matrices the system gets easily trapped into a local minimum, the usual procedure then is to start from several initial options hoping to reach the global minimum at some point.

A global minimizer takes the responsibility of efficiently explore the configuration space of the problem. PyChemia implements several global searchers as we saw on previous tutorials. Those global searchers joined by an efficient evaluation infrastructure allows many evaluations being perform without human assistance and an effective chance of reaching a global minimum if the search is long enough.

For this tutorial consider the following problem:

'pychemia/test/data/abinit_dmatpaw/abinit.in'.

First, we can read the abinit input and access the contents of the variable ‘dmatpawu’:

```python
import pychemia
import numpy as np
pychemia_path = pychemia.__path__[0]
abiinput = pychemia.code.abinit.InputVariables(pychemia_path + '/test/data/abinit_dmatpaw/abinit.in')
dmatpawu = np.array(abiinput['dmatpawu']).reshape(-1,5,5)
```

The variable ‘dmatpawu’ stores the contents of 4 5x5 matrices, the correlation matrices for the corresponding 4 Co atoms in the crystal. The matrices can be converted into a numpy array with shape (4, 5, 5) and they look like this:

```
array([[[ 0.06256, 0. , 0.01218, 0. , 0. ],
        [ 0. , 0.481 , 0. , 0.45877, 0. ],
        [ 0.01218, 0. , 0.07148, 0. , 0. ],
        [ 0. , 0.45877, 0. , 0.481 , 0. ],
        [ 0. , 0. , 0. , 0. , 0.94038]],
       [[ 0.97852, 0. , 0.00305, 0. , 0. ],
        [ 0. , 0.95493, 0. , -0.00449, 0. ],
        [ 0.00305, 0. , 0.98115, 0. , 0. ],
        [ 0. , -0.00449, 0. , 0.95493, 0. ],
        [ 0. , 0. , 0. , 0. , 0.95168]],
       [[ 0.97852, 0. , -0.00305, 0. , 0. ],
        [ 0. , 0.95493, 0. , 0.00449, 0. ],
        [-0.00305, 0. , 0.98115, 0. , 0. ],
        [ 0. , 0.00449, 0. , 0.95493, 0. ],
        [ 0. , 0. , 0. , 0. , 0.95168]],
       [[ 0.06256, 0. , -0.01218, 0. , 0. ],
        [ 0. , 0.481 , 0. , -0.45877, 0. ],
        [-0.01218, 0. , 0.07148, 0. , 0. ],
        [ 0. , -0.45877, 0. , 0.481 , 0. ],
        [ 0. , 0. , 0. , 0. , 0.94038]]])
```

The objective is to find the set of correlation matrices that minimize the energy. Those are density matrices so even if we have 100 numbers, any set of numbers is a valid set of correlation matrices. We will now convert this set of matrices into a reduced set of variables that can be treated independently.
A correlation matrix can be expressed as the following product:

\[ R \cdot O \cdot R^{-1} \]

Where \( R \) is a rotation matrix, \( O \) is a diagonal matrix with a trace that is the total number of electrons correlated. We need to find a set of independent variables to re-create any correlation matrix. We know that not any arbitrary set of 25 numbers is a good rotation matrix. However, a 5x5 rotation matrix can be effectively decomposed into 10 independent numbers, the so-called “Generalized Euler angles”, this set of angles reduces the 25 values from a 5x5 rotation matrix into 10 independent variables. We should also be aware that the occupations on the diagonal of the matrix ‘\( O \)’ are not exactly integers, we will account for the small differences into a separate set of values. With those premises a 5x5 correlation matrix is converted into a set with 10 euler angles, 5 occupations and 5 deltas. This is done by the routines ‘dmatpawu2params’ and ‘params2dmatpawu’ that allow us to go back and forward from the set of correlation matrices into the set of ‘euler_angles’, integer ‘occupations’ and the small ‘deltas’:

```python
params = pychemia.population.orbitaldftu.dmatpawu2params(dmatpawu, 5)
```

The variable ‘params’ is a dictionary with values for ‘occupations’, ‘deltas’ and ‘euler_angles’:

```python
{'deltas': array([[ 0.02223 , 0.054049, 0.079991, 0.06023 , 0.05962 ],
                  [ 0.04956 , 0.04832 , 0.04058 , 0.023486, 0.016844],
                  [ 0.04956 , 0.04832 , 0.04058 , 0.023486, 0.016844],
                  [ 0.02223 , 0.054049, 0.079991, 0.06023 , 0.05962 ]]),
 'euler_angles': array([[ 0. , 0. , 0. , 0. , 0. , 0.785398, 0. , 0. , 0. , 0.785398, 0. , 0. , 0. , 0. , 0. , 0. , 0. , 0. , 0. , -0.609893, 0. , -1.570796],
                        [ 1.570796 , 1.570796 , 3.141593],
                        [-0.581865, 0. , -1.570796, 0. , 0.785398, 0. , 0.785398, 0. , 0. , 0. , -0.609893, 0. , 1.570796]],
 'ndim': 5,
 'num_matrices': 4,
 'occupations': array([[0, 0, 0, 1, 1],
                       [1, 1, 1, 1, 1],
                       [1, 1, 1, 1, 1],
                       [0, 0, 0, 1, 1]]))
```

This is in fact the set of independent variables that we can use to optimize the correlation using a global searcher. We can also go back and recover the correlation matrices using the inverse procedure, ie, from the dictionary params recover the correlation matrices:

```python
dmatpawu_new = pychemia.population.orbitaldftu.params2dmatpawu(params)
```

The dmatpawu is recovered from the values stored in ‘params’:

```python
array([[ 0.06256, -0. , -0.01218, -0. , 0. ],
        [-0. , 0.481, -0. , 0.45877, 0. ],
        [-0.01218, -0. , 0.07148, -0. , 0. ],
        [-0. , 0.45877, -0. , 0.481 , 0. ],
        [ 0. , 0. , 0. , 0. , 0.94038]],
       [[ 0.97852, -0. , 0.00305, -0. , 0. ],
        [-0. , 0.95493, -0. , -0.00449, -0. ],
        [ 0.00305, -0. , 0.98115, -0. , -0. ],
        [-0. , -0.00449, -0. , 0.95493 , 0. ],
        [-0. , -0. , -0. , 0. , 0.95168]])
```

(continues on next page)
Each correlation matrix contains 25 values, using the procedure above, we reduce this number to 20: 10 euler angles, 5 integer occupations and 5 deltas. The values of deltas can be ignored for the purpose of the global searcher and the occupations are contrained by the condition that their sum must be equal to the number of electrons in the correlated orbital. We have now the ingredients to move into the next step, create a population of correlation matrices.

### 4.4.2 The population

The most simple way of creating the population requires just the name of the mongo database to be created and one abinit input file. The relevant information to setup the search will be inferred from the contents of the abinit input file:

```python
popu=pychemia.population.orbitaldftu.OrbitalDFTU('test', abinit_input=pychemia_path + '/test/data/abinit_dmatpawu/abinit.in')
```

Orbital population:
- Species [znucl]: [19, 27, 9]
- Orbitals corrected:
  - 19: False
  - 27: True (l=2)
  - 9: False
- Number of atoms where DFT+U is applied: 4
- Correlation of 'd' orbitals
- Variables controlling the total number of matrices:
  - nsppol: 1
  - nspden: 2
- Total number of matrices expected on dmatpawu: 4
- Number of electrons for each correlation matrix: [2 5 5 2]
- Number of independent matrices: 4

Create random correlation matrices can be done with:

```python
popu.add_random()
```

The return is the Identifier of the new entry on the database. Also a set of new random correlation matrices can be created with:

```python
popu.random_population(16)
```

We have the basic ingredients for creating the first population for the global searcher. How the correlation matrices are evaluated is out of scope of the population and depends on the particularities of the machines where Abinit is used.
to evaluate them. We will move our focus to the methods needed to produced new correlation matrices based on the results of a given set of correlation matrices.
CHAPTER 5

Indices and tables

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- modindex
- search