# **qmpy Documentation**

Release 0.4.9

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qmpy is the backend responsible for creating and running the Open Quantum Materials Database (http://oqmd.org). The OQMD is a project created in Chris Wolverton's group at Northwestern University (http://wolverton.northwestern.edu).

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**CHAPTER** 

ONE

# INSTALLATION

# 1.1 From repo

Install qmpy with pip or easy install:

```
>>> pip install qmpy
or:
>>> easy_install -U qmpy
```

**Note:** Using pip or easy\_install to install scipy or numpy can be unreliable. It is better to install from a proper repository for your linux distribution. However, if that version of SciPy is earlier than 0.12.0 you will need to obtain another installation. If necessary, you can obtain the needed libraries with:

```
$ sudo apt-get install libatlas-dev libatlas-base-dev
$ sudo apt-get install liblapack-dev gfortran
```

# 1.2 From GitHub repo

Obtain the source with:

```
$ git clone https://github.com/wolverton-research-group/qmpy.git
$ cd qmpy
$ python setup.py install
```

Be aware that f you want to install qmpy from source, you will be responsible for ensuring that you have all of the following required packages installed.

# 1.3 Required Packages

- Django (https://www.djangoproject.com/)
- Numpy (http://www.numpy.org/)
- Scipy (http://www.scipy.org/)
- PyYAML (http://pyyaml.org/)
- python-MySQL (https://pypi.python.org/pypi/MySQL-python)

- · python-memcached
- · django-extensions
- PuLP (https://pythonhosted.org/PuLP/) (required for grand canonical linear programming and high-dimensional phase diagram slices)

# 1.4 Recommended Packages

- matplotlib (http://matplotlib.org/) (required for creating figures)
- networx (http://networkx.github.io/) (required for creating spin lattices, and some high-dimensional phase diagram analysis)

**Warning:** In order for pulp to work, you must have a working linear programming package installed. PuLP provides a simple library for this, but it is up to you to make sure it is working.

**CHAPTER** 

**TWO** 

# SETTING UP THE DATABASE

The database can be downloaded from http://oqmd.org/static/downloads/qmdb.sql.gz

Once you have the database file, you need to unzip it and load it into a database MySQL. On a typical linux installation this process will look like:

```
$ wget http://oqmd.org/static/downloads/qmdb.sql.gz
$ gunzip qmdb.sql.gz
$ mysql < qmdb.sql</pre>
```

**Note:** Assuming your install is on linux, and assuming you haven't used MySQL at all, you will need to enter a mysql session as root ("mysql -u root -p"), create a user within MySQL ("CREATE USER 'newuser'@'localhost';"), grant that user permissions ("GRANT ALL PRIVILEGES ON \* . \* TO 'newuser'@'localhost'; FLUSH PRIVILEGES;").

**Note:** The name of the deployed database has changed since previous releases (qmdb\_prod). If your install isn't working, make sure that the database name agrees with what is found in qmpy/db/settings.py.

Once this is done, you need to edit qmpy/db/settings.py. Set the DATABASES variable such that 'USER' is the user with permissions to access the newly installed database.

**Note:** For windows/cygwin users: To use MySQL in Cygwin, you need to install MySQL via the Oracle website for windows. Only after MySQL is install in windows can use mysql in Cygwin. You can find the download for MySQL here: http://dev.mysql.com/downloads/windows/installer.

It is free, but you have to register with Oracle to access. Next, you need to move the database file over to the Windows MySQL data drive. It may vary by version, but you might find it at C:ProgramDataMySQLMySQL Server 5.6data. Copy the downloaded database directory into this folder. Finally, in the db/settings.py file, the HOST has to be set to '127.0.0.1', and set the PORT and PASSWORD variables as well according to your MySQL installation.

To verify that the database is properly installed and has appropriate permissions, run:

```
mysql> select count(*) from entries;
+----+
| count(*) |
+----+
| 173653 |
+-----+
```

The number may not match what is shown above, but as long as you don't recieve any errors, your database should be working properly.

**CHAPTER** 

# THREE

# **TUTORIALS**

qmpy is a package containing many tools for computational materials science.

The qmpy package comes bundled with two executable scripts, qmpy and oqmd. qmpy is a simple bash script that starts an interactive python environment and imports qmpy:

```
$ qmpy >>>
```

To write your own python script that utilizes qmpy functionality, simply start with an import like:

```
from qmpy import *
```

and all of the commands shown below should work.

# 3.1 Database entries

Once the database is installed, you can query it very flexibly and easily. In this section we will explore the data structure of entries in the OQMD and provide several examples of how to make queries. For deeper understanding of how django models work, you should check out the (excellent) django documentation.

First, lets look at how to access an entry from the database. As an example, lets pull up an entry for an Element:

```
>>> fe = Element.objects.get(symbol='Fe')
>>> fe
<Element: Fe>
```

Django models have a number of fields that can be accessed directly once the database entry has been loaded. For example, with an element you can:

```
>>> fe.symbol
u'Fe'
>>> fe.z
26L
```

For a complete list of the model attributes that are stored in the database, refer to the documentation for the model you are interested in, in this case Element.

**Note:** When strings are returned, they are returned as unicode strings, (indicated by the "u" preceding the string) integers are returned as long integers (indicated by the trailing "L"). For most purposes this makes no difference, as these data types will generally behave exactly as expected, i.e.:

```
>>> fe.z == 26
True
```

```
>>> fe.symbol == "Fe"
True
```

In addition to data attributes like these, django models have relationships to other models. In qmpy there are two flavors of relationships: one-to-many and many-to-many. An example of a one-to-many relationship would be the relationship between an Element and an Atom. There are many atoms which are a given element, but each atom is only one element. In the case of Fe:

```
>>> fe.atom_set
<django.db.models.fields.related.RelatedManager object at 0x7f0997fa2690>
>>> fe.atom_set.count()
127585
>>> atom = fe.atom_set.first()
>>> print atom
<Atom: Fe @ 0.000000 0.000000 0.000000>
>>> atom.element
<Element: Fe>
```

A RelatedManager is an object that deals with obtaining other django models that are related to the main object. We can use the objects.count() method to fidn the number of Atom objects that are Fe, and find ~125,000. To obtain one of these atoms, we use the objects.first() method, which simply returns the first Atom which is Fe. Much more functionality of Managers and RelatedManagers will be shown throughout this tutorial and in the examples, but for a proper understanding you should refer to the django docs.

An example of a many-to-many relationship would be the relationship between an Element and a Composition. A composition (e.g. Fe2O3) can contain many elements (Fe and O), and an element can be a part of many compositions (Fe3O4 and FeO as well). This is the nature of a many-to-many relationship. In the case of Fe:

```
>>> fe.composition_set.count()
10882
>>> comp = fe.composition_set.filter(ntypes=2)[0]
>>> comp
<Composition: AcFe>
>>> comp.element_set.all()
[<Element: Ac>, <Element: Fe>]
```

In this example we have taken our base object (the Element) and filtered its composition\_set for Composition objects which meet the condition ntypes=2 (i.e. there are two elements in the composition), and taken the first such Composition (index 0 in the QuerySet that is returned).

# 3.2 Creating a structure

There are several ways to create a structure, but we will start with reading in a POSCAR:

```
>>> s = io.read(INSTALL_PATH+'/io/files/POSCAR_BCC')
```

Once you have the Structure object, the important features of a crystal structure can be accessed readily.:

```
0.500000>]
>>> s.composition
<Composition: Cu>
>>> s.volume
27.0
```

You can also readily construct a Structure from scratch, from the lattice vectors and the atom positions.:

```
>>> s2 = Structure.create([3,3,3], [('Cu', [0,0,0]), ('Cu', [0.5,0.5,0.5])])
>>> s2 == s
True
```

# 3.3 First Principles Calculations

At this time qmpy only supports automation of calculations using the Vienna Ab Initio Simulation Package (VASP). The reading and creation of these calculations are handled by the Calculation model. To read in an existing calculation:

```
>>> path = '/analysis/vasp/files/normal/fine_relax/'
>>> calc = Calculation.read(INSTALL_PATH+path)
```

qmpy will search the directory for an OUTCAR or OUTCAR.gz file. If it is able to find an OUTCAR, it will attempt to read the file. Next, we will demonstrate several of the key attributes you may wish to access:

```
>>> calc.energy # the final total energy
-12.416926999999999
>>> calc.energies # the total energies of each step
array([-12.415236 -12.416596, -12.416927])
>>> calc.volume # the output volume
77.787375068172508
>>> calc.input
<Structure: SrGe2>
>>> calc.output
<Structure: SrGe2>
>>> from pprint import pprint
>>> pprint(calc.settings)
{'algo': 'fast',
 'ediff': 0.0001,
'encut': 373.0,
 'epsilon': 1.0,
 'ibrion': 1,
 'idipol': 0,
 'isif': 3,
 'ismear': 1,
 'ispin': 1,
 'istart': 0,
 'lcharg': True,
'ldipol': False,
'lorbit': 0,
'lreal': False,
 'lvtot': False,
 'lwave': False,
 'nbands': 24,
 'nelm': 60,
 'nelmin': 5,
 'nsw': 40,
```

# 3.4 Searching for models

The documentation for Django for searching for models is ver complete, and should be taken as the ultimate reference for searching for models in qmpy, but a basic overview is provided here.

# 3.4.1 Searching for entries based on stability

Formation energies are stored as FormationEnergy instances, which are associated with an :mod:~qmpy.Entry and a :mod:~qmpy.Calculation. Knowing this, we can search for stable Entries using:

```
>>> stable = Entry.objects.filter(formationenergy__stability__lt=0)
>>> stable.count()
18150
```

The same concept can be applied to searching for other quantities, as long as you can relate them to a FormationEnergy by " constructions:

```
>>> stable_comps = Composition.objects.filter(formationenergy__stability__lt=0)
>>> stable_comps.count()
18150
>>> s = Structure.objects.filter(calculated__formationenergy__stability__lt=0)
>>> s.count()
18150
```

Adding other search criteria lets you explore a little more:

```
>>> stable = FormationEnergy.objects.filter(stability__lt=0)
>>> # Find the number of stable compounds containing 0
>>> stable.filter(composition__element_set='0').count()
4017
>>> # or Fe. Is it surprising that this is smaller?
>>> stable.filter(composition__element_set='Fe').count()
653
>>> # Meta data is also a possiblity. How many stable compounds were found
>>> # in the course of calculations for a particular project?
>>> stable.filter(entry__project_set='prototypes').count()
3119
```

# 3.4.2 Searching for entries based on composition

You can find compositions in a few ways using filters and excludes. If you want a specific region of phase space (including related subspaces):

```
>>> elts = [ 'Fe', 'Li', 'O' ]
>>> others = Element.objects.exclude(symbol__in=elts)
>>> comps = Composition.objects.exclude(element_set=others)
```

This searchs finds every composition that doesn't have any elements that aren't in the region of phase space requested. For binary or ternary phase spaces it can be more efficient to search permutations of sub-spaces:

```
>>> comps = Composition.objects.filter(ntypes=3)
>>> for e in elts:
      comps = comps.filter(element_set=e)
>>>
>>> for e in elts:
>>>
        e_comps = Composition.objects.filter(element_set=e, ntypes=1)
>>>
        comps |= e_comps
>>> for e1, e2 in itertools.combinations(elts, r=2):
       bin_comps = Composition.objects.filter(element_set=el)
>>>
        bin_comps = bin_comps.filter(element_set=e2, ntypes=2)
>>>
>>>
        comps |= bin_comps
>>> comps.distinct().count()
```

However, for larger regions of phase space (4 or 5 or more) the number of subqueries of the second approach rapidly becomes more expensive than the single, more complicated query of the first.

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**CHAPTER** 

# **FOUR**

# **DATA MODELS**

qmpy is a package containing many tools for computational materials science.

# 4.1 Materials models

# 4.1.1 Structure

```
class qmpy.Structure (*args, **kwargs)
    Structure model. Principal attributes are a lattice and basis set.
    Relationships:
```

Entry via entry

Atom via atom\_set: Atoms in the structure. More commonly handled by the managed attributed atoms.

Calculation via calculated. Calculation objects that the structure is an output from.

Calculation via calculation\_set. Returns calculation objects that the structure is an input to.

Composition via composition.

Element via element\_set

Spacegroup via spacegroup

Species via species\_set

Prototype via prototype. If the structure belongs to a prototypical structure, it is referred to here.

Reference Original literature reference.

MetaData via meta\_data

#### **Attributes:**

#### Identification

id

label: key in the Entry.structures dictionary.

natoms: Number of atoms. nsites: Number of sites. ntypes: Number of elements.

measured: Experimentally measured structure?

source: Name for source.

```
Lattice
     x1, x2, x3
     y1, y2, y3
     z1, z2, z3: Lattice vectors of the cell. Accessed via cell.
     volume
     volume_pa
     Calculated properties
     delta_e: Formation energy (eV/atom)
     meta_stability: Distance from the convex hull (eV/atom)
     energy: Total DFT energy (eV/FU)
     energy_pa: Total DFT energy (eV/atom)
     magmom: Total magnetic moment (Μ<sub>b</sub>)
     magmom_pa: Magnetic moment per atom.
     SXX, SXY, SYY
     syx, szx, szz: Stresses on the cell. Accessed via stresses.
Examples:
>>> s = io.read(INSTALL_PATH+'/io/files/POSCAR_FCC')
>>> s.atoms
>>> s.cell
>>> s.magmoms
>>> s.forces
>>> s.stresses
add atom (atom, tol=0.01)
     Adds atom to the structure if it isn't already contained.
atom_types
     List of atomic symbols, length equal to number of atoms.
atomic_numbers
     List of atomic numbers, length equal to number of atoms.
atoms
     List of Atoms in the structure.
cartesian_coords
     Return atomic positions in cartesian coordinates.
cell
     Lattice vectors, 3x3 numpy.ndarray.
comment_objects
     Return list of comments (MetaData objects of type comment)
     Composition dictionary.
compare (other,
                      tol = 0.01,
                                    atom\_tol=10,
                                                      volume=False,
                                                                          allow_distortions=False,
           check_spacegroup=False, wildcard=None)
     Credit to K. Michel for the algorithm.
```

- 2. Convert both structures to primitive form
- 3. Check that the total number of atoms in primitive cells are the same

- 4. Check that the number of atoms of each element are the same in primitive cells
- 4b. Check that the spacegroup is the same.
  - 5.If needed check that the primitive cell volumes are the same
- 6. Convert both primitive cells to reduced form There is one issue here the reduce cell could be type I (all angles acute) or type II (all angles obtuse) and a slight difference in the initial cells could cause two structures to reduce to different types. So at this step, if the angles are not correct, the second cell is transformed as [-1, 0, 0], [0, -1, 0], [0, 0, 1].
- 7. Check that the cell internal angles are the same in both reduced cells.
- 8. Check that the ratios of reduced cell basis lengths are the same. ie a1/b1 = a2/b2, a1/c1 = a2/c2, and b1/c1 = b2/c2 where a1, b1, c1, are the lengths of basis vectors in cell 1 and a2, b2, c2 are the lengths of cell 2.
- 9. Get the lattice symmetry of the reduced cell 2 (this is a list of all rotations that leave the lattice itself unchanged). In turn, apply all rotations to reduced cell 2 and for each search for a vector that overlaps rotated cell positions with positions in reduced cell 1. If a rotation + translation overlaps reduced cells, then they are the same.

MODIFICATIONS: Only need one "base" atom from the first structure Get the distance from the origin for every atom first

Arguments: other: Another Structure.

**Keyword Arguments:** tol: Percent deviation in lattice parameters and angles.

Not Implemented Yet: wildcard: Elements of the specified type can match with any atom.

#### coords

numpy.ndarray of atom coordinates.

#### copy (

Create a complete copy of the structure, with any primary keys removed, so it is not associated with the original.

```
static create (cell, atoms=[], **kwargs)
```

Creates a new Structure.

**Arguments:** cell: 3x3 lattice vector array

**Keyword Arguments:** atoms: List of Atom creation arguments. Can be a list of [element, coord], or a list of [element, coord, kwargs].

Examples:

create vacuum(direction, amount, in place=True)

Add vacuum along a lattice direction.

**Arguments:** direction: direction to add the vacuum along. (0=x, 1=y, 2=z) amount: amount of vacuum in Angstroms.

Keyword Arguments: in\_place: apply change to current structure, or return a new one.

#### Examples:

```
>>> s = io.read(INSTALL_PATH+'/io/files/POSCAR_FCC')
>>> s.create_vacuum(2, 5)
```

#### elements

List of Elements

#### find\_lattice\_points\_by\_transform(transform, tol=1e-06)

Find the lattice points contained within the transformation.

#### find\_lattice\_points\_within\_distance (distance, tol=1e-06)

Find the lattice points contained within radius distance from the origin.

```
find_nearest_neighbors (method='closest', tol=0.05, limit=5.0)
```

Determine the nearest neighbors for all Atoms in Structure.

Calls <code>get\_nearest\_neighbors()</code> and assigns the nearest neighbor dictionary to <code>Structure.\_neighbor\_dict</code>. Each atom is also given a list, <code>nearest\_neighbors</code> that contains the nearest neighbor atoms. For atoms which have the "same" atom as a nearest neighbor across different periodic boundaries, a single atom may appear multiple times on the list.

**Keyword Arguments:** limit: How far to look from each atom for nearest neighbors. Default=5.0.

tol: A tolerance which determines how much further than the closest atom a second atom can be and still be a part of the nearest neighbor shell.

Returns: None

#### forces

numpy.ndarray of forces on atoms.

```
get_distance (atom1, atom2, limit=None, wrap_self=True)
```

Calculate the shortest distance between two atoms.

**Note:** This is not as trivial a problem as it sounds. It is easy to demonstrate that for any non-cubic cell, the normal method of calculating the distance by wrapping the vector in fractional coordinates to the range (-0.5, 0.5) fails for cases near (0.5, 0.5) in Type I cells and near (0.5, -0.5) for Type II.

To get the correct distance, the vector must be wrapped into the Wigner-Seitz cell.

**Arguments:** atom1, atom2: (Atom, Site, int).

#### **Keyword Arguments:**

**limit:** If a limit is provided, returns None if the distance is greater than the limit.

**wrap\_self:** If True, the distance from an atom to itself is 0, otherwise it is the distance to the shortest periodic image of itself.

#### $get_sites(tol=0.1)$

From self.atoms, creates a list of Sites. Atoms which are closer than tol from one another are considered on the same site.

#### get\_spin\_lattice (elements=None, supercell=None)

Constructs a lattice of sites.

# **Keyword Arguments:**

**elements:** If *elements* is supplied, get\_spin\_lattice will return the lattice of those elements only.

**supercell:** Accepts any valid input to Structure.transform to construct a supercell, and return its lattice. Useful for finding AFM orderings for structures which have a smaller periodicity than their magnetic structure.

**Returns:** A SpinLattice, which is a container for a lattice graph, which contains nodes which are atoms and edges which indicate nearest neighbors.

#### Examples:

```
>>> s = io.read(INSTALL_PATH+'/io/files/fe3o4.cif')
>>> sl = s.get_spin_lattice(elements=['Fe'])
>>> sl.set_fraction(0.33333)
>>> sl.fraction
0.3333333333333333333333
>>> sl.run_MC()
```

# get\_volume()

Calculates the volume from the triple product of self.cell

#### group\_atoms\_by\_symmetry()

Sort self.atoms according to the site they occupy.

#### inv

Precalculates the inverse of the lattice, for faster conversion between cartesian and direct coordinates.

# is\_buerger\_cell(tol=1e-05)

Tests whether or not the structure is a Buerger cell.

#### is\_niggli\_cell(tol=1e-05)

Tests whether or not the structure is a Niggli cell.

```
joggle_atoms (distance=0.001, in_place=True)
```

Randomly displace all atoms in each direction by a distance up to +/- the distance keyword argument (in Angstroms).

#### **Optional keyword arguments:**

distance [Range within all internal coordinates are] displaced. Default=1e-3

*in\_place* [If True, returns an ndarray of the applied] translations. If False, returns (Structure, translations).

# Examples:

```
>>> s = io.read('POSCAR')
>>> s2, trans = s.joggle_atoms(in_place=False)
>>> trans = s.joggle_atoms(1e-1)
>>> trans = s.joggle_atoms(distance=1e-1)
```

# keyword\_objects

Return list of keywords (MetaData objects of type keyword)

#### lat param dict

Dictionary of lattice parameters.

# lat\_param\_string (format='screen')

Generates a human friendly representation of the lattice parameters of a structure.

**Keyword Arguments:** format: ('screen'l'html'l'mathtype')

# lat\_params

Tuple of lattice parameters (a, b, c, alpha, beta, gamma).

```
1p
```

Tuple of lattice parameters (a, b, c, alpha, beta, gamma).

#### magmoms

numpy.ndarray of magnetic moments of shape (natoms,).

```
make_conventional (in_place=True, tol=1e-05)
```

Uses spglib to convert to the conventional cell.

#### **Keyword Arguments:**

in\_place: If True, changes the current structure. If false returns a new one

**tol:** Symmetry precision for symmetry analysis

#### Examples:

```
>>> s = io.read(INSTALL_PATH+'io/files/POSCAR_FCC_prim')
>>> print len(s)
1
>>> s.make_conventional()
>>> print len(s)
4
```

#### make\_perfect (in\_place=True, tol=0.1)

Constructs options for a 'perfect' lattice from the structure.

If a site is not fully occupied, but has only one atom type on it, it will be filled the rest of the way. If a site has two or more atom types on it, the higher fraction element will fill the site.

Keyword Arguments: in\_place: If False returns a new Structure, otherwise returns None

tol: maximum defect concentration.

#### Examples:

```
>>> s = io.read(INSTALL_PATH+'/io/files/partial_vac.cif')
>>> s
<Structure: Mn3.356Si4016>
>>> s.make_perfect()
>>> s
<Structure: MnSiO4>
>>> s = io.read(INSTALL_PATH+'/io/files/partial_mix.cif')
>>> s
<Structure: Mn4.264Co3.736Si4O16>
>>> s2 = s.make_perfect(in_place=False)
>>> s2
<Structure: MnCoSiO4>
```

#### make\_primitive (in\_place=True, tol=1e-05)

Uses spglib to convert to the primitive cell.

# **Keyword Arguments:**

in\_place: If True, changes the current structure. If false returns a new one

tol: Symmetry precision for symmetry analysis

#### Examples:

```
>>> s = io.read(INSTALL_PATH+'io/files/POSCAR_FCC')
>>> print len(s)
4
>>> s.make_primitive()
```

```
>>> print len(s)
1
```

# metrical\_matrix

np.dot(self.cell.T, self.cell)

#### name

Unformatted name.

#### nearest\_neighbor\_dict

Dict of Atom:[list of Atom] pairs.

```
pdf_compare (other, tol=0.01)
```

Compute the PDF for each structure and evaluate the overlap integral for all pairs of species.

```
recenter (atom, in_place=True, middle=False)
```

Translate the internal coordinates to center the specified atom. Atom can be an actual Atom from the Structure.atoms list, or can be identified by index.

#### **Keyword Arguments:**

in\_place: If False, return a new Structure with the transformation applied. defaults to True.

**middle:** If False, "centers" the cell by putting the atom at the origin. If True, "centers" the cell by putting the atom at (0.5,0.5,0.5). defaults to False.

# Examples:

```
>>> s = io.read('POSCAR')
>>> s.recenter(s[2])
>>> s2 = s.recenter(s[0], in_place=False)
>>> s2.recenter(2)
>>> s == s2
True
```

#### reciprocal\_lattice

Reciprocal lattice of the structure.

```
reduce (tol=0.001, limit=1000, in_place=True)
```

Get the transformation matrix from unit to reduced cell Acta. Cryst. (1976) A32, 297 Acta. Cryst. (2003) A60, 1

# **Optional keyword arguments:**

*tol* [] eps\_rel in Acta. Cryst. 2003 above. Similar to tolerance for floating point comparisons. Defaults to 1e-5.

limit [] maximum number of loops through the algorithm. Defaults to 1000.

in\_place [] Change the Structure or return a new one. If True, the transformation matrix is returned.If False, a tuple of (Structure, transformation\_matrix) is returned.

#### Examples:

```
>>> s = io.read('POSCAR')
>>> s.reduce()
>>> s.reduce(in_place=False, get_transform=False)
```

#### refine (tol=0.001, recurse=True)

Identify atoms that are close to high symmetry sites (within tol and shift them onto them.

**Note:** "symprec" doesn't appear to do anything with spglib, so I am unable to get "loose" symmetry operations. Without which, this doesn't work.

# Examples:

```
>>> s = io.read('POSCAR')
>>> s.symmetrize()
>>> print s.spacegroup
225L
>>> s.refine()
>>> print s.spacegroup
```

#### set\_magnetism(type, scheme='primitive')

Assigns magnetic moments to all atoms in accordance with the specified magnetism scheme.

#### Schemes:

Key-	Description
word	
None	all magnetic moments = None
"ferro"	atoms with partially filled d and f shells are assigned a magnetic moment of 5 mu_b
"anti"	finds a highly ordererd arrangement arrangement of up and down spins. If only 1 magnetic
	atom is found a ferromagnetic arrangment is used. raises NotImplementedError

#### $set_natoms(n)$

Set self.atoms to n blank Atoms.

#### set nsites(n)

Sets self.sites to n blank Sites.

#### set volume(value)

Rescales the unit cell to the specified volume, keeping the direction and relative magnitudes of all lattice vectors the same.

#### site coords

numpy.ndarray of site coordinates.

#### sites

List of Sites in the structure.

# spec\_comp

Species composition dictionary.

#### species

List of species

#### species\_types

List of species, length equal to number of atoms.

# stresses

Calculated stresses, a numpy.ndarray of shape (6,)

#### sub (replace, rescale=True, in\_place=False, \*\*kwargs)

Replace atoms, as specified in a dict of pairs.

# **Keyword Arguments:**

**rescale:** rescale the volume of the final structure based on the per atom volume of the new composition.

in\_place: change the species of the current Structure or return a new one.

Examples:

```
>>> s = io.read('POSCAR-Fe203')
>>> s2 = s.substitute({'Fe':'Ni', 'O':'F'} rescale=True)
>>> s2.substitute({'Ni':'Co'}, in_place=True, rescale=False)
```

substitute (replace, rescale=True, in\_place=False, \*\*kwargs)

Replace atoms, as specified in a dict of pairs.

#### **Keyword Arguments:**

**rescale:** rescale the volume of the final structure based on the per atom volume of the new composition.

in\_place: change the species of the current Structure or return a new one.

#### Examples:

```
>>> s = io.read('POSCAR-Fe2O3')
>>> s2 = s.substitute({'Fe':'Ni', 'O':'F'} rescale=True)
>>> s2.substitute({'Ni':'Co'}, in_place=True, rescale=False)
```

#### symmetrize (tol=0.001, angle tol=-1)

Analyze the symmetry of the structure. Uses spglib to find the symmetry.

# symmetrize sets:

- spacegroup -> Spacegroup
- uniq\_sites -> list of unique Sites
- orbits -> lists of equivalent Atoms
- rotations -> List of rotation operations
- translations -> List of translation operations
- operatiosn -> List of (rotation, translation) pairs
- for each atom: atom.wyckoff -> WyckoffSite
- for each site: site.multiplicity -> int

#### t (transform, in\_place=True, tol=1e-05)

Apply lattice transform to the structure. Accepts transformations of shape (3,) and (3,3).

# **Optional keyword arguments:**

in place [If False, return a new Structure with the] transformation applied.

#### Examples:

```
>>> s = io.read('POSCAR')
>>> s.transform([2,2,2]) # 2x2x2 supercell
>>> s.transform([[0,1,0],[1,0,0],[0,0,1]]) # swap axis 1 for 2
>>> s2 = s.transform([2,2,2], in_place=False)
```

transform (transform, in\_place=True, tol=1e-05)

Apply lattice transform to the structure. Accepts transformations of shape (3,) and (3,3).

#### **Optional keyword arguments:**

in\_place [If False, return a new Structure with the] transformation applied.

Examples:

```
>>> s = io.read('POSCAR')
          >>> s.transform([2,2,2]) # 2x2x2 supercell
          >>> s.transform([[0,1,0],[1,0,0],[0,0,1]]) # swap axis 1 for 2
          >>> s2 = s.transform([2,2,2], in_place=False)
     translate (cv, cartesian=True, in_place=True)
          Shifts the contents of the structure by a vector.
          Optional keyword arguments:
              cartesian [If True, translation vector is taken to be] cartesian coordinates. If False, translation vector
                  is taken to be in fractional coordinates. Default=True
              in_place [If False, return a new Structure with the] transformation applied.
          Examples:
          >>> s = io.read('POSCAR')
          >>> s.translate([1,2,3])
          >>> s.translate([0.5,0.5, 0.5], cartesian=False)
          >>> s2 = s.translate([-1,2,1], in_place=False)
     unit_comp
          Composition dict, where sum(self.unit_comp.values()) == 1
class qmpy .Prototype (*args, **kwargs)
     Base class for a prototype structure.
     Relationships:
          Composition via composition_set
          Structure via structure set
          Entry via entry_set
     Attributes:
          name: Prototype name.
     classmethod get (name)
          Retrieves a Prototype named name if it exists. If not, creates a new one.
          Examples:
          >>> proto = Prototype.get('Corundum')
4.1.2 Atom
class qmpy . Atom (*args, **kwargs)
     Model for an Atom.
     Relationships:
          Structure via structure
          Element via element
          Site via site
          WyckoffSite via wyckoff
     Attributes:
          id
          x, y, z: Coordinate of the atom
```

```
fx, fy, fz: Forces on the atom
```

magmom: Magnetic moment on the atom (in Μ<sub>b</sub>)

occupancy: Occupation fraction (0-1).

ox: Oxidation state of the atom (can be different from charge)

charge: Charge on the atom

volume: Volume occupied by the atom

#### cart coord

Cartesian coordinates of the Atom.

#### coord

[x,y,z] coordinates.

#### copy()

Creates an exact copy of the atom, only without the matching primary key.

# Examples:

```
>>> a = Atom.get('Fe', [0,0,0])
>>> a.save()
>>> a.id
1
>>> a.copy()
>>> a
<Atom: Fe - 0.000, 0.000, 0.000>
>>> a.id
None
```

#### classmethod create (element, coord, \*\*kwargs)

Creates a new Atom object.

**Arguments:** element (str or Element): Specifies the element of the Atom. coord (iterable of floats): Specifies the coordinate of the Atom.

# **Keyword Arguments:**

**forces:** Specifies the forces on the atom.

magmom: The magnitude of the magnetic moment on the atom.

**charge:** The charge on the Atom.

**volume:** The atomic volume of the atom (Angstroms^3).

#### Examples:

#### forces

Forces on the Atom in [x, y, z] directions.

#### index

None if not in a Structure, otherwise the index of the atom in the structure.

# **is\_on** (*site*, *tol*=0.001)

Tests whether or not the Atom is on the specified Site.

Examples:

```
>>> a = Atom.create('Fe', [0,0,0])
>>> s = a.get_site()
>>> a2 = Atom.create('Ni', [0,0,0])
>>> a2.is_on(s)
True
```

#### species

Formatted Species string. e.g. Fe3+, O2-

# 4.1.3 Site

```
class qmpy.Site(*args, **kwargs)
```

A lattice site.

A site can be occupied by one Atom, many Atoms or no Atoms.

#### **Relationships:**

```
Structure via structure
Atom via atom_set
WyckoffSite via wyckoff
```

#### **Attributes:**

id

x, y, z: Coordinate of the Site

```
add_atom(atom, tol=0.01)
```

Adds Atom to Site.atoms.

**Notes:** If the Site being assigned to doens't have a coordinate, it is assigned the coordinate of *atom*.

**Arguments:** atom (Atom): Atom to add to the structure.

# **Keyword Arguments:**

**tol** (**float**): **Distance between** *atom* **and the Site for the Atom to be** assigned to the Site. Raises a SiteError if the distance is greater than *tol*.

Raises: SiteError: If atom is more than tol from the Site.

# Examples:

```
>>> s = Site.create([0,0,0])
>>> a = Atom.create('Fe', [0,0,0])
>>> s.add_atom(a)
>>> s2 = Site()
>>> s2.add_atom(a)
```

#### atoms

List of Atoms on the Site.

#### cart\_coord

Cartesian coordinates of the Atom.

# comp

Composition dictionary of the Site.

**Returns:** dict: of (element, occupancy) pairs.

Examples:

```
>>> a1 = Atom('Fe', [0,0,0], occupancy=0.2)
>>> a2 = Atom('Ni', [0,0,0], occupancy=0.8)
>>> s = Site.from_atoms([a1,a2])
>>> s.comp
{'Fe':0.2, 'Ni':0.8}
```

#### coord

[Site.x, Site.y, Site.z]

#### static create (coord, comp=None)

Constructs a Site from a coordinate.

**Note:** The Site is created without any Atoms occupying it.

#### **Arguments:**

coord (length 3 iterable): Assigns the x, y, and z coordinates of the Site.

#### **Keyword Arguments:**

**comp** (**dict**, **string**, **or qmpy.Element**): **Composition dictionary.** Flexible about input forms. Options include: <Element: Fe>, 'Fe', {"Fe":0.5, "Co":0.5}, and {<Element: Ni>:0.5, <Element: Co>:0.5}.

**Raises:** TypeError: if *comp* isn't a string, Atom, Element.

#### Examples:

```
>>> s = Site.create([0.5, 0.5, 0.5])
```

# classmethod from\_atoms (atoms, tol=0.0001)

Constructs a Site from an iterable of Atoms.

**Notes:** Site.coord is set as the average coord of all assigned Atoms.

Checks that the Atoms are close together. If the Atoms are further apart than tol, raises a SiteError

**Arguments:** atoms (iterable of *Atom*): List of Atoms to occupy the Site.

**Keyword Arguments:** tol (float): Atoms must be within *tol* of each other to be assigned to the same Site. Defaults to 1e-4.

# Examples:

```
>>> a1 = Atom.create('Fe', [0,0,0])
>>> a2 = Atom.create('Ni', [1e-5, -1e-5, 0])
>>> s = Site.from_atoms([a1,a1])
```

# label

Assigns a human friendly label for the Site, based on its atomic composition. If singly occupied, returns the symbol of the atom on the site. If multiply occupied, returns a comma seperated string

#### Examples:

```
>>> a1 = Atom.create('Fe', [0,0,0], occupancy=0.2)
>>> a2 = Atom.create('Ni', [0,0,0], occupancy=0.8)
>>> s = Site.from_atoms([a1,a2])
```

#### magmom

Calculates the composition weighted average magnetic moment of the atoms on the Site.

**Returns:** float or None

#### occupancy

Calculates the total occupancy of the site.

Returns: float or None

ox

Calculates the composition weighted average oxidation state of the atoms on the Site.

Returns: float or None

#### spec\_comp

Composition dictionary of the Site.

Returns: dict: of (species, occupancy) pairs.

# Examples:

```
>>> a1 = Atom('Fe', [0,0,0], occupancy=0.2)
>>> a2 = Atom('Ni', [0,0,0], occupancy=0.8)
>>> s = Site.from_atoms([a1,a2])
>>> s.comp
{'Fe':0.2, 'Ni':0.8}
```

#### 4.1.4 Element

```
class qmpy.Element (*args, **kwargs)
```

Core model for an element.

# **Relationships:**

```
Atom via atom_set

Species via species_set

Structure via structure_set

Entry via entry_set

Composition via composition_set

Calculation via calculation_set

Potential via potential_set

Hubbard via hubbards

HubbardCorrection via hubbardcorrection_set

ReferenceEnergy via referenceenergy_set
```

#### **Attributes:**

#### Identification

z: atomic number name: full atomic name symbol: atomic symbol group: group in the periodic table period: period in the periodic table

#### Physical properties

mass: Atomic mass, in AMU (float) density: Density at STP, in g/cm^3 (float) volume: Atomic volume at STP, in A^3/atom (float)

atomic\_radii: in A (float)

```
van_der_waals radii: in A (float) covalent_radii: in A (float)
```

scattering\_factors: A dictionary of scattering factor coeffs.

# Thermodynamic properties

melt: melting point in K boil: boiling point in K specific\_heat: C\_p in J/K

# **Electronic properties**

electronegativity: Pauling electronegativity ion\_energy: First ionization energy. (eV)

s\_elec: # of s electrons p\_elec: # of p electrons d\_elec: # of d electrons f elec: # of f electrons

#### **Additional information**

production: Annual tons of element produced. abundance: Amount in earths crust (ppm) radioactive: Are all isotopes unstable?

HHI\_P: Herfindahl-Hirschman Index for production. HHI\_R: Herfindahl-Hirschman Index for reserve

Note: HHI values from Gaultois, M. et al. Chem. Mater. 25, 2911-2920 (2013).

# classmethod get (value)

Return an element object. Accepts symbols and atomic numbers, or a list of symbols/atomic numbers.

#### Examples:

```
>>> Element.get('Fe')
>>> Element.get(26)
>>> Element.get(['Fe', 'O'])
```

# class qmpy . Species (\*args, \*\*kwargs)

Base model for an atomic species. (Element + charge state).

#### **Relationships:**

```
Element via element
Entry via entry_set
Structure via structure_set
```

# **Attributes:**

```
name: Species name. e.g. Fe3+, O2-ox: Oxidation state (float)
```

#### classmethod get (value)

Gets or creates the specified species.

# **Arguments:**

value: Accepts multiple input types. Can be a string, e.g. Fe3+ or a tuple of (symbol, oxidation state) pairs, e.g. (Fe, 3).

**Return:** A Species or list of Species.

#### Examples:

```
>>> Species.get('Fe3+')
>>> Species.get('Fe3')
>>> Species.get(('Fe', 3))
>>> Species.get([ 'Fe3+', 'O2-', 'Li1+'])
```

# 4.1.5 Composition

```
class qmpy.Composition (*args, **kwargs)
```

Base class for a composition.

# **Relationships:**

```
Calculation via calculation_set

Element via element_set

Entry via entry_set

ExptFormationEnergy via exptformationenergy_set

FormationEnergy via formationenergy_set

MetaData via meta_data

Structure via structure_set

Prototype via prototype_set
```

#### **Attributes:**

```
formula: Electronegativity sorted and normalized composition string.
```

```
e.g. Fe2O3, LiFeO2
```

generic: Genericized composition string. e.g. A2B3, ABC2.

mass: Mass per atom in AMUs

meidema: Meidema model energy for the composition

ntypes: Number of elements.

#### comp

Return an element:amount composition dictionary.

### delta\_e

Return the lowest formation energy.

# experiment

Return the lowest experimantally measured formation energy at the compositoin.

#### classmethod get (composition)

Classmethod for getting Composition objects - if the Composition exists the database, it is returned. If not, a new Composition is created.

# Examples:

```
>>> Composition.get('Fe203')
<Composition: Fe203>
```

# ${\bf classmethod\ get\_list\ } (bounds, calculated = False, uncalculated = False)$

Classmethod for finding all compositions within the space bounded by a sequence of compositions.

Examples:

#### ground\_state

Return the most stable entry at the composition.

#### icsd\_delta\_e

Return the lowest formation energy calculated from experimentally measured structures - i.e. excluding prototypes.

#### ndistinct

Return the number of distinct entries.

#### space

Return the set of element symbols

#### unit\_comp

Return an element:amoutn composition dictionary normalized to a unit composition.

# 4.2 Calculation models

# 4.2.1 Calculation

```
class qmpy.Calculation (*args, **kwargs)
Base class for storing a VASP calculation.
```

# **Relationships:**

```
Composition via composition

DOS via dos

Structure via input. Input structure.

Structure via output. Resulting structure.

Element via element_set.

Potential via potential_set.

Hubbard via hubbard_set.

Entry via entry.

Fit via fit. Reference energy sets that have been fit using this calculation.

FormationEnergy via formationenergy_set. Formation energies computed from this calculation, for different choices of fit sets.

MetaData via meta_data
```

# Attributes:

id

```
label: key for entry.calculations dict.
     attempt: # of this attempt at a calculation.
     band_gap: Energy gap occupied by the fermi energy.
     configuration: Type of calculation (module).
     converged: Did the calculation converge electronically and ionically.
     energy: Total energy (eV/UC)
     energy_pa: Energy per atom (eV/atom)
     irreducible_kpoints: # of irreducible k-points.
     magmom: Total magnetic moment (mu_b)
     magmom_pa: Magnetic moment per atom. (mu_b/atom)
     natoms: # of atoms in the input.
     nsteps: # of ionic steps.
     path: Calculation path.
     runtime: Runtime in seconds.
     settings: dictionary of VASP settings.
error_objects
     Return list of errors (MetaData objects of type error)
get_outcar()
     Sets the calculations outcar attribute to a list of lines from the outcar.
     Examples:
     >>> calc = Calculation.read('calculation_path')
     >>> print calc.outcar
     None
     >>> calc.get_outcar()
     >>> len(calc.outcar)
     12345L
static read (path)
     Reads the outcar specified by the objects path. Populates input field values, as well as outputs, in addition
     to finding errors and confirming convergence.
     Examples:
     >>> path = '/analysis/vasp/files/normal/standard/'
     >>> calc = Calculation.read(INSTALL_PATH+path)
read_charges()
     Reads and returns VASP's calculated charges for each atom. Returns the RAW charge, not NET charge.
     Examples:
     >>> calc = Calculation.read('path_to_calculation')
     >>> calc.read_charges()
read_chgcar (filename='CHGCAR.gz', filetype='CHGCAR')
     Reads a VASP CHGCAR or ELFCAR and returns a GridData instance.
read elements()
     Reads the elements of the atoms in the structure. Returned as a list of atoms of shape (natoms,).
```

Examples:

```
>>> calc = Calculation.read('path_to_calculation')
>>> calc.read_elements()
['Fe', 'Fe', 'O', 'O', 'O']
```

#### read\_energies()

Returns a numpy.ndarray of energies over all ionic steps.

Examples:

```
>>> calc = Calculation.read('calculation_path')
>>> calc.read_energies()
array([-12.415236, -12.416596, -12.416927])
```

# read\_lattice\_vectors()

Reads and returns a numpy ndarray of lattice vectors for every ionic step of the calculation.

Examples:

```
>>> path = 'analysis/vasp/files/magnetic/standard'
>>> calc = Calculation.read(INSTALL_PATH+'/'+path)
>>> calc.read_lattice_vectors()
array([[[ 5.707918, 0. , 0.
                  5.707918,
       [ 0.
                            0.
                            7.408951]],
                  0. ,
       [ 0.
      [[ 5.707918, 0.
                        , 0.
                                   1.
      [ 0. , 5.707918, 0.
       [ 0.
                , 0. , 7.408951]])
```

#### read n ionic()

Reads the number of ionic steps, and assigns the value to nsteps.

# read\_natoms()

Reads the number of atoms, and assigns the value to natoms.

#### set chgcar(source)

Copy the CHGCAR specified by *source* to this calculation.

**Arguments:** source: can be another Calculation instance or a string containing a path to a CHGCAR. If it is a path, it should be a absolute, i.e. begin with "/", and can either end with the CHGCAR or simply point to the path that contains it. For example, if you want to take the CHGCAR from a previous calculation you can do any of:

```
>>> c1 # old calculation
>>> c2 # new calculation
>>> c2.set_chgcar(c1)
>>> c2.set_chgcar(c1.path)
>>> c2.set_chgcar(c1.path+'/CHGCAR')
```

#### set\_wavecar (source)

Copy the WAVECAR specified by source to this calculation.

**Arguments:** source: can be another Calculation instance or a string containing a path to a WAVE-CAR. If it is a path, it should be a absolute, i.e. begin with "/", and can either end with the WAVECAR or simply point to the path that contains it. For example, if you want to take the WAVECAR from a previous calculation you can do any of:

```
>>> c1 # old calculation
>>> c2 # new calculation
>>> c2.set_wavecar(c1)
```

```
>>> c2.set_wavecar(c1.path)
>>> c2.set_wavecar(c1.path+'/WAVECAR')
```

static setup (structure, configuration='static', path=None, entry=None, hubbard='wang', potentials='vasp\_rec', settings={}, chgcar=None, wavecar=None, \*\*kwargs)
Method for creating a new VASP calculation.

**Arguments:** structure: Structure instance, or string indicating an input structure file.

#### **Keyword Arguments:**

**configuration:** String indicating the type of calculation to perform. Options can be found with qmpy.VASP\_SETTINGS.keys(). Create your own configuration options by adding a new file to configuration/vasp\_settings/inputs/ using the files already in that directory as a guide. Default="static"

**settings:** Dictionary of VASP settings to be applied to the calculation. Is applied after the settings which are provided by the *configuration* choice.

**path:** Location at which to perform the calculation. If the calculation takes repeated iterations to finish successfully, all steps will be nested in the *path* directory.

**entry:** If the full qmpy data structure is being used, you can specify an entry to associate with the calculation.

**hubbard:** String indicating the hubbard correctionconvention. Options found with qmpy.HUBBARDS.keys(), and can be added to or altered by editing configuration/vasp\_settings/hubbards.yml. Default="wang".

**potentials:** String indicating the vasp potentials to use. Options can be found with qmpy.POTENTIALS.keys(), and can be added to or altered by editing configuration/vasp\_settings/potentials/yml. Default="vasp\_rec".

**chgcar/wavecar:** Calculation, or path, indicating where to obtain an initial CHGCAR/WAVECAR file for the calculation.

#### warning\_objects

Return list of warnings (MetaData objects of type warning)

# 4.2.2 Density of States

```
class qmpy.DOS (*args, **kwargs)
Electronic density of states..
```

#### **Relationships:**

```
Entry via entry
MetaData via meta_data
Calculation via calculation
```

# **Attributes:**

id

data: Numpy array of DOS occupations.

file: Source file. gap: Band gap in eV.

#### energy

Return the array with the energies.

```
read_doscar (fname='DOSCAR')
           Read a VASP DOSCAR file
     site_dos (atom, orbital)
           Return an NDOSx1 array with dos for the chosen atom and orbital.
           atom: int Atom index
           orbital: int or str Which orbital to plot
           If the orbital is given as an integer: If spin-unpolarized calculation, no phase factors: s = 0, p = 1, d = 2
           Spin-polarized, no phase factors: s-up = 0, s-down = 1, p-up = 2, p-down = 3, d-up = 4, d-down = 5 If
           phase factors have been calculated, orbitals are s, py, pz, px, dxy, dyz, dz2, dxz, dx2 double in the above
           fashion if spin polarized.
4.2.3 Potential
class qmpy.Potential(*args, **kwargs)
     Class for storing a VASP potential.
     Relationships:
           calculation
           element
     Attributes:
           name
           date
           electrons: Electrons in potential.
           enmax
           enmin
           gw
           id
           paw
           potcar
```

class qmpy.Hubbard(\*args, \*\*kwargs)

Base class for a hubbard correction parameterization.

#### **Attributes:**

us

calculation convention correction element id 1 ligand ox

u

# 4.3 Thermodynamics models

# 4.3.1 Formation Energies

```
class qmpy.FormationEnergy (*args, **kwargs)

Base class for a formation energy.
```

# **Relationships:**

```
Calculation via calculation
Composition via composition
Entry via entry
FormationEnergy via equilibrium
Fit via fit
```

### **Attributes:**

```
id
delta_e: Formation energy (eV/atom)
description: A label of the source of the formation energy.
stability: Distance from the convex hull (eV/atom)
```

```
class qmpy.ExptFormationEnergy(*args, **kwargs)
```

Experimentally measured formation energy.

Any external formation energy should be entered as an ExptFormationEnergy object, rather than a FormationEnergy. If the external source is also computational, set the "dft" attribute to be True.

# **Relationships:**

```
Composition via composition Fit via fit
```

# **Attributes:**

```
id: integer primary key.
delta_e: measured formation energy.
delta_g: measured free energy of formation.
dft: (bool) True if the formation energy is from a non-OQMD DFT calculation.
source: (str) Identifier for the source.
```

# 4.3.2 Reference energies

```
class qmpy.Fit(*args, **kwargs)
```

The core model for a reference energy fitting scheme.

The Fit model links to the experimental data (ExptFormationEnergy objects) that informed the fit, as well as the DFT calculations (Calculation objects) that were matched to each experimental formation energy. Once the fit is completed, it also stores a list of chemical potentials both as a relationship to ReferenceEnergy and HubbardCorrection objects. These correction energies can also be accessed by dictionaries at Fit.mus and Fit.hubbard\_mus.

## **Relationships:**

Calculation via dft

```
ExptFormationEnergy via experiments
FormationEnergy via formationenergy_set
HubbardCorrection via hubbard_correction_set
ReferenceEnergy via reference_energy_set
```

#### **Attributes:**

name: Name for the fitting

#### Examples:

```
>>> f = Fit.get('standard')
>>> f.experiments.count()
>>> f.dft.count()
>>> f.mus
>>> f.hubbard_mus
```

# class qmpy . ReferenceEnergy (\*args, \*\*kwargs)

Elemental reference energy for evaluating heats of formation.

#### **Relationships:**

```
Fit via fit
Element via element
```

#### **Attributes:**

id

value: Reference energy (eV/atom)

#### class qmpy . HubbardCorrection (\*args, \*\*kwargs)

Energy correction for DFT+U energies.

## **Relationships:**

```
Fit via fit
Element via element
Hubbard via hubbard
```

#### **Attributes:**

id

value: Correction energy (eV/atom)

# 4.3.3 Phase Space

class qmpy.PhaseSpace (bounds, mus=None, data=None, \*\*kwargs)

A PhaseSpace object represents, naturally, a region of phase space.

The most fundamental property of a PhaseSpace is its bounds, which are given as a hyphen-delimited list of compositions. These represent the extent of the phase space, and determine which phases are within the space.

Next, a PhaseSpace has an attribute, data, which is a PhaseData object, and is a container for Phase objects, which are used when performing thermodynamic analysis on this space.

The majority of attributes are lazy, that is, they are only computed when they are requested, and how to get them (of which there are often several ways) is decided based on the size and shape of the phase space.

#### bound\_elements

Alphabetically ordered list of elements with constrained composition.

#### bound space

Set of elements \_of fixed composition in the PhaseSpace.

### Examples:

```
>>> s = PhaseSpace('Fe-Li', '0=-1.4')
>>> s.bound_space
set(['Fe', 'Li'])
```

#### chempot\_dimension

Chemical potential dimension.

#### Examples:

```
>>> s = PhaseSpace('Fe-Li', 'O=-2.5')
>>> s.chempot_dimension
0
>>> s = PhaseSpace('Fe-Li', 'N=0:-5')
>>> s.chempot_dimension
1
>>> s = PhaseSpace('Fe-Li', 'N=0:-5 F=0:-5')
>>> s.chempot_dimension
2
```

### chempot\_scan()

Scan through chemical potentials of *element* from *umin* to *umax* identifing values at which phase transformations occur.

#### clear all()

Clears input data and analyzed results. Same as: >>> PhaseData.clear\_data() >>> PhaseData.clear analysis()

### clear\_analysis()

Clears all calculated results.

## clear\_data()

Clears all phase data.

# cliques

Iterator over maximal cliques in the phase space. To get a list of cliques, use list(PhaseSpace.cliques).

#### comp (coord)

Returns the composition of a coordinate in phase space.

## Examples:

```
>>> space = PhaseSpace('Fe-Li-O')
>>> space.comp([0.2, 0.2, 0.6])
{'Fe': 0.2, 'O': 0.6, 'Li': 0.2}
```

#### comp\_dimension

Compositional dimension of the region of phase space.

```
>>> s = PhaseSpace('Fe-Li-O')
>>> s.comp_dimension
2
>>> s = PhaseSpace('FeO-Ni2O-CoO-Ti3O4')
>>> s.comp_dimension
3
```

#### compute formation energies()

Evaluates the formation energy of every phase with respect to the chemical potentials in the PhaseSpace.

### compute\_stabilities (\*args, \*\*kwargs)

Calculate the stability for every Phase.

#### **Keyword Arguments:**

phases: List of Phases. If None, uses every Phase in PhaseSpace.phases

save: If True, save the value for stability to the database.

**new\_only:** If True, only compute the stability for Phases which did not import a stability from the OQMD. False by default.

#### compute\_stability(p)

Compute the energy difference between the formation energy of a Phase, and the energy of the convex hull in the absence of that phase.

```
coord (composition, tol=0.0001)
```

Returns the barycentric coordinate of a composition, relative to the bounds of the PhaseSpace. If the object isn't within the bounds, raises a PhaseSpaceError.

#### Examples:

```
>>> space = PhaseSpace('Fe-Li-O')
>>> space.coord({'Fe':1, 'Li':1, 'O':2})
array([ 0.25,  0.25,  0.5 ])
>>> space = PhaseSpace('Fe2O3-Li2O')
>>> space.coord('Li5FeO4')
array([ 0.25,  0.75])
```

### dual\_spaces

List of sets of elements, such that any possible tie-line between two phases in phases is contained in at least one set, and no set is a subset of any other.

#### elements

Alphabetically ordered list of elements present in the PhaseSpace.

### find\_reaction\_mus(element=None)

Find the chemical potentials of a specified element at which reactions occur.

### Examples:

```
>>> s = PhaseSpace('Fe-Li-O')
>>> s.find_reaction_mus('O')
```

## $gclp(composition=\{\}, mus=\{\}, phases=[])$

Returns energy, phase composition which is stable at given composition

#### Examples:

```
>>> space = PhaseSpace('Fe-Li-O')
>>> phases, energy = space.gclp('FeLiO2')
>>> print phases
>>> print energy
```

## get\_hull\_points()

Gets out-of PhaseSpace points. i.e. for FeSi2-Li, there are no other phases in the space, but there are combinations of Li-Si phases and Fe-Si phases. This method returns a list of phases including composite phases from out of the space.

```
>>> space = PhaseSpace('FeSi2-Li')
>>> space.get_hull_points()
[<Phase FeSi2 (23408): -0.45110217625>,
<Phase Li (104737): 0>,
<Phase 0.680 Li13Si4 + 0.320 FeSi : -0.3370691816>,
<Phase 0.647 Li8Si3 + 0.353 FeSi : -0.355992801765>,
<Phase 0.133 Fe3Si + 0.867 Li21Si5 : -0.239436904167>,
<Phase 0.278 FeSi + 0.722 Li21Si5 : -0.306877209723>]
```

### get\_minima (phases, bounds)

Given a set of Phases, get\_minima will determine the minimum free energy elemental composition as a weighted sum of these compounds

#### get\_phase\_diagram()

Creates a Renderer attribute with appropriate phase diagram components.

#### **Examples:**

```
>>> space = PhaseSpace('Fe-Li-O')
>>> space.get_renderer()
>>> plt.show()
```

### get\_qhull (phases=None, mus={})

Get the convex hull for a given space.

#### get\_reaction (var, facet=None)

For a given composition, what is the maximum delta\_composition reaction on the given facet. If None, returns the whole reaction for the given PhaseSpace.

#### Examples:

```
>>> space = PhaseSpace('Fe203-Li20')
>>> equilibria = space.hull[0]
>>> space.get_reaction('Li20', facet=equilibria)
```

## get\_reactions (var, electrons=2.0)

Returns a list of Reactions.

#### Examples:

```
>>> space = PhaseSpace('Fe-Li-O')
>>> space.get_reactions('Li', electrons=1)
```

#### get\_tie\_lines\_by\_gclp (iterable=False)

Runs over pairs of Phases and tests for equilibrium by GCLP. Not recommended, it is very slow.

### graph

networkx. Graph representation of the phase space.

#### hull

List of facets of the convex hull.

# $in\_bounds$ (composition)

Returns True, if the composition is within the bounds of the phase space

```
>>> space = PhaseSpace('Fe203-Ni02-Li20')
>>> space.in_bounds('Fe304')
False
>>> space.in_bounds('Li5Fe08')
True
```

#### in\_space (composition)

Returns True, if the composition is in the right elemental-space for this PhaseSpace.

### Examples:

```
>>> space = PhaseSpace('Fe-Li-O')
>>> space.in_space('LiNiO2')
False
>>> space.in_space('Fe2O3')
True
```

### load(\*\*kwargs)

Loads oqmd data into the associated PhaseData object.

# make\_1d\_vs\_chempot (\*\*kwargs)

Plot of phase stability vs chemical potential for a single composition.

### Examples:

```
>>> s = PhaseSpace('Fe', mus={'O':[0,-4]})
>>> r = s.make_vs_chempot()
>>> r.plot_in_matplotlib()
>>> plt.show()
```

### make\_as\_binary(\*\*kwargs)

Construct a binary phase diagram (convex hull) and write it to a Renderer.

#### Examples:

```
>>> s = PhaseSpace('Fe-P')
>>> r = s.make_as_binary()
>>> r.plot_in_matplotlib()
>>> plt.show()
```

# make\_as\_graph(\*\*kwargs)

Construct a graph-style visualization of the phase diagram.

#### make as quaternary(\*\*kwargs)

Construct a quaternary phase diagram and write it to a Renderer.

### Examples:

```
>>> s = PhaseSpace('Fe-Li-O-P')
>>> r = s.make_as_quaternary()
>>> r.plot_in_matplotlib()
>>> plt.show()
```

## make\_as\_ternary(\*\*kwargs)

Construct a ternary phase diagram and write it to a Renderer.

# Examples:

```
>>> s = PhaseSpace('Fe-Li-O-P')
>>> r = s.make_as_quaternary()
>>> r.plot_in_matplotlib()
>>> plt.show()
```

# make\_as\_unary(\*\*kwargs)

Plot of phase volume vs formation energy.

```
>>> s = PhaseSpace('Fe203')
>>> r = s.make_as_unary()
>>> r.plot_in_matplotlib()
>>> plt.show()
```

# make\_vs\_chempot (\*\*kwargs)

Plot of phase stability vs chemical potential for a range of compositions.

### **Examples:**

```
>>> s = PhaseSpace('Fe-Li', mus={'0':[0,-4]})
>>> r = s.make_vs_chempot()
>>> r.plot_in_matplotlib()
>>> plt.show()
```

### phase\_diagram

Renderer of a phase diagram of the PhaseSpace

```
plot_reactions (var, electrons=2.0, save=False)
```

Plot the convex hull along the reaction path, as well as the voltage profile.

#### save tie lines()

Save all tie lines in this PhaseSpace to the OQMD. Stored in Formation.equilibrium

#### shape

(# of compositional dimensions, # of chemical potential dimensions) The shape attribute of the PhaseSpace determines what type of phase diagram will be drawn.

## Examples:

```
>>> s = PhaseSpace('Fe-Li', 'O=-1.2')
>>> s.shape
(1, 0)
>>> s = PhaseSpace('Fe-Li', 'O=0:-5')
>>> s.shape
(1, 1)
>>> s = PhaseSpace('Fe-Li-P', 'O=0:-5')
>>> s.shape
(2,1)
>>> s = PhaseSpace('Fe', 'O=0:-5')
>>> s.shape
(0, 1)
```

#### space

Set of elements present in the PhaseSpace.

#### Examples:

```
>>> s = PhaseSpace('Pb-Te-Se')
>>> s.space
set(['Pb', 'Te', 'Se'])
>>> s = PhaseSpace('PbTe-Na-PbSe')
>>> s.space
set(['Pb', 'Te', 'Na', 'Se'])
```

## spaces

List of lists of elements, such that every phase in self.phases is contained in at least one set, and no set is a subset of any other. This corresponds to the smallest subset of spaces that must be analyzed to determine the stability of every phase in your dataset.

```
>>> pa, pb, pc = Phase('A', 0), Phase('B', 0), Phase('C', 0)
>>> p1 = Phase('AB2', -1)
>>> p2 = Phase('B3C', -4)
>>> s = PhaseSpace('A-B-C', load=None)
>>> s.phases = [ pa, pb, pc, p1, p2 ]
>>> s.spaces
[['C', 'B'], ['A', 'B']]
```

#### stability\_range (p, element=None)

Calculate the range of phase *p* with respect to *element*.

#### stable

List of stable phases

## tie\_lines

List of length 2 tuples of phases with tie lines between them

#### unstable

List of unstable phases.

#### class qmpy.PhaseData

A PhaseData object is a container for storing and organizing phase data. Most importantly used when doing a large number of thermodynamic analyses and it is undesirable to access the database for every space you want to consider.

# add\_phase (phase)

Add a phase to the PhaseData collection. Updates the PhaseData.phase\_dict and PhaseData.phases\_by\_elt dictionaries appropriately to enable quick access.

#### Examples:

```
>>> pd = PhaseData()
>>> pd.add_phase(Phase(composition='Fe203', energy=-3))
>>> pd.add_phase(Phase(composition='Fe203', energy=-4))
>>> pd.add_phase(Phase(composition='Fe203', energy=-5))
>>> pd.phase_dict
{'Fe203': <Phase Fe203 : -5}
>>> pd.phases_by_elt['Fe']
[<Phase Fe203 : -3>, <Phase Fe203 : -4>, <Phase Fe203 : -5>]
```

#### add phases (phases)

Loops over a sequence of phases, and applies *add phase* to each.

#### Equivalent to:

```
>>> pd = PhaseData()
>>> for p in phases:
>>> pd.add_phase(p)
```

# get\_phase\_data(space)

Using an existing PhaseData object return a PhaseData object which is populated by returning a subset which is inside a given region of phase space.

**Arguments:** space: formatted as in qmpy.PhaseSpace.\_\_init\_\_()

```
>>> pd = PhaseData()
>>> pd.read_file('legacy.dat')
>>> new_pd = pd.get_phase_data(['Fe', '0'])
>>> new_pd.phase_dict
```

#### load\_library (library)

Load a library file, containing self-consistent thermochemical data.

### Equivalent to:

```
>>> pd = PhaseData()
>>> pd.read_file(INSTALL_PATH+'/data/thermodata/%s' % library)
```

load\_oqmd (space=None, search={}, stable=False, fit='standard', total=False)
Load data from the OQMD.

# **Keyword Arguments:**

**space:** sequence of elements. If supplied, will return only phases within that region of phase space. i.e. ['Fe', 'O'] will return Fe, O and all iron oxides.

search: dictionary of database search keyword:value pairs.

**stable:** Restrict search to only stable phases (faster, but relies on having current phase stability analyses).

# Examples:

```
>>> pd = PhaseData()
>>> search = {'calculation__path__contains':'icsd'}
>>> pd.load_oqmd(space=['Fe','0'], search=search, stable=True)
```

#### phases

List of all phases.

## read\_file (filename, per\_atom=True)

Read in a thermodata file (named filename).

#### File format:

```
composition energy
Fe 0.0
0 0.0
Li 0.0
Fe304 -0.21331204979
Fe0 -0.589343204057
Fe304 -0.21331204979
FeLiO2 -0.446739168889
FeLi504 -0.198830531099
```

**Keyword Arguments:** per\_atom: If True, the supplied energies are per atom, not per formula unit. Defaults to True.

# 4.4 Database models

#### 4.4.1 Entries

```
class qmpy.Entry (*args, **kwargs)

Base class for a database entry.
```

The core model for typical database entries. An Entry model represents an input structure to the database, and can be created from any input file. The Entry also ties together all of the associated qmpy.Structure, qmpy.Calculation, qmpy.Reference, qmpy.FormationEnergies, and other associated databas entries.

# **Relationships:**

```
Calculation via calculation_set

DOS via dos_set

Entry via duplicate_of

Entry via duplicates

Element via element_set

FormationEnergy via formationenergy_set

Job via job_set

MetaData via meta_data

Project via project_set

Prototype via prototype

Species via species_set

Structure via structure_set

Task via task_set

Reference via reference

Composition via composition
```

#### **Attributes:**

id: Primary key (auto-incrementing int)

natoms: Number of atoms in the primitive input cell ntypes: Number of elements in the input structure

path: Path to input file, and location of subsequent calculations.

label: An identifying name for the structure. e.g. icsd-1001 or A3

#### calculations

Dictionary of label:Calculation pairs.

#### chq

Attempts to load the charge density of the final calculation, if it is done. If not, returns False.

```
\textbf{static create} (source, keywords = [\ ], projects = [\ ], **kwargs)
```

Attempts to create an Entry object from a provided input file.

Processed in the following way:

- 1.If an Entry exists at the specified path, returns that Entry.
- 2. Create an Entry, and assign all fundamental attributes. (natoms, ntypes, input, path, elements, keywords, projects).
- 3.If the input file is a CIF, and because CIF files have additional composition and reference information, if that file format is found, an additional test is performed to check that the reported composition matches the composition of the resulting structure. The reference for the work is also created and assigned to the entry.
- 4. Attempt to identify another entry that this is either exactly equivalent to, or a defect cell of.

**Keywords:** keywords: list of keywords to associate with the entry. projects: list of project names to associate with the entry.

```
do (module, *args, **kwargs)
```

Looks for a computing script matching the first argument, and attempts to run it with itself as the first argument. Sends args and kwargs to the script. Should return a Calculation object, or list of Calculation objects.

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#### Examples:

```
>>> e = Entry.objects.get(id=123)
>>> e.do('relaxation')
<Calculation: 523 @ relaxation settings>
```

#### elements

List of Elements

### energy

If the structure has been relaxed, returns the formation energy of the final relaxed structure. Otherwise, returns None.

#### errors

List of errors encountered in all calculations.

#### hold\_objects

Return list of holds (MetaData objects of type hold)

#### holds

A note indicating a reason the entry should not be calculated

#### html

HTML formatted name

### keyword\_objects

Return list of keywords (MetaData objects of type keyword)

### keywords

Descriptive keyword for looking up entries

#### latex

LaTeX formatted name

#### mass

Return the mass of the entry, normalized to per atom.

```
move (*args, **kwargs)
```

Moves all calculation files to the specified path.

#### name

Unformatted name

# projects

List of Projects

#### red\_comp

Composition dictionary, in reduced form.

#### reset()

Deletes all calculations, removes all associated structures - returns the entry to a pristine state.

```
save (*args, **kwargs)
```

Saves the Entry, as well as all associated objects.

## space

Return the set of elements in the input structure.

```
>>> e = Entry.create("fe2o3/POSCAR") # an input containing Fe2O3
>>> e.space
set(["Fe", "O"])
```

#### spec\_comp

Composition dictionary, using species (element + oxidation state) instead of just the elements.

### species

List of Species

## total\_energy

If the structure has been relaxed, returns the formation energy of the final relaxed structure. Otherwise, returns None.

# unit\_comp

Composition dictionary, normalized to 1 atom.

```
visualize (structure='source')
```

Attempts to open the input structure for visualization using VESTA

#### volume

If the entry has gone through relaxation, returns the relaxed volume. Otherwise, returns the input volume.

```
class qmpy . MetaData (*args, **kwargs)
```

Base class for variable typed model tagging.

Model for arbitrary meta-data descriptors for various qmpy objects. Generally accessed by properties and methods added by the "add\_label" descriptor. See "add\_label" for a more detailed description of its use

#### Relationships

```
Calculation via calculation_set
Composition via composition_set
DOS via dos_set
Entry via entry_set
Structure via structure_set
```

#### **Attributes:**

```
id: Autoincrementing primary key
type: Label for the kind of meta data, e.g. "hold", "keyword"
value: Content of the meta data. e.g. "repeated failure", "known
anti-ferromagnetic"
```

#### Examples:

```
>>> MetaData.get('Keyword', 'ICSD')
<Keyword: ICSD>
```

### 4.4.2 References

```
class qmpy . Reference (*args, **kwargs)
```

Base class for a reference to a publication.

#### **Relatioships:**

```
Author via author_set
Journal via journal
Entry via entry_set
Structure via structure_set
```

#### **Database fields:**

id

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```
title
year
volume
page_first
page_last
```

## 4.4.3 Authors

```
class qmpy . Author (*args, **kwargs)
    Base class for an author.

Relationships:
    Reference via references

Database Fields:
    id
    first
```

# 4.4.4 Journals

last

```
class qmpy.Journal (*args, **kwargs)
Base class for a journal

Relationships:

Reference via references

Database fields:

id

name
code
```

# 4.5 Symmetry models

# 4.5.1 Spacegroup

```
class qmpy.Spacegroup (*args, **kwargs)
    Base class for a space group.

Relationships:
    Structure via structure_set
    Translation via centering_vectors
    Operation via operations
    WyckoffSite via site_set

Attributes:
```

number: Spacegroup #. (primary key)

centrosymmetric: (bool) Is the spacegroup centrosymmetric.

```
hall: Hall symbol.
           hm: Hermann-Mauguin symobl.
          lattice_system: Cubic, Hexagonal, Tetragonal, Orthorhombic,
                Monoclinic or Triclinic.
           pearson: Pearson symbol
           schoenflies: Schoenflies symbol.
     get_site (symbol)
           Gets WyckoffSite by symbol.
     rotations
          List of rotation operations for the spacegroup.
     sym_ops
          List of (rotation, translation) pairs for the spacegroup
     symbol
          Returns the Hermann-Mauguin symbol for the spacegroup
     translations
          List of translation operations for the spacegroup.
     wyckoff_sites
           List of WyckoffSites.
4.5.2 Wyckoff Site
class qmpy . WyckoffSite (*args, **kwargs)
     Base class for a Wyckoff site. (e.g. a "b" site).
     Relationships:
           Spacegroup via spacegroup
           Atom via atom set
           Site via site_set
     Attributes:
          id
           symbol: Site symbol
           multiplicity: Site multiplicity
           x, y, z: Coordinate symbols.
```

# 4.5.3 Symmetry Operations

```
class qmpy.Operation(*args, **kwargs)
     A symmetry operation (rotation + translation).
```

# **Relationships:**

```
Spacegroup via spacegroup
Rotation via rotation_set
Translation via translation_set
```

## **Attributes:**

id

```
Examples:
```

```
>>> op = Operation.get('x+y-1/2,-z-y+1/2,x-z+1/2')
>>> print op
<Operation: +x+y+1/2,-y-z+1/2,+x-z+1/2>
```

### classmethod get (value)

Accepts symmetry operation strings, i.e. "+x, x+1/2, x+y-z" or a tuple of rotation matrix and translation vector.

#### Example:

```
>>> Operation.get("x,y,-y")
>>> Operation.get(( rot, trans ))
```

# class qmpy.Translation(\*args, \*\*kwargs)

A translation operation.

### **Relationships:**

```
Spacegroup via spacegroup Operation via operation
```

#### **Attributes:**

id

x, y, z: Translation vector. Accessed via vector.

## Examples:

```
>>> op = Operation.get('x', 'x+y', 'z-x+1/2')
>>> print op.translsation
<Translation: 0,0,+1/2>
>>> print op.translation.vector
array([ 0. ,  0. ,  0.5])
```

#### class qmpy . Rotation (\*args, \*\*kwargs)

A rotation operation.

# **Relationships:**

```
Spacegroup via spacegroup Operation via operation
```

## **Attributes:**

```
id
a11, a12, a13
a21, a22, a23
a31, a32, a33: Rotation matrix. Accessed via matrix.
```

# 4.6 Resource models

run\_path state

# 4.6.1 Host

```
class qmpy . Host (*args, **kwargs)
      Host model - stores all host information for a cluster.
      Relationships:
           account
           allocation
      Attributes:
           name: Primary key.
           binaries: dict of label:path pairs for vasp binaries.
           check_queue: Path to showq command
           checked time: datetime object for the last time the queue was
                 checked.
           hostname: Full host name.
           ip_address: Full ip address.
           nodes: Total number of nodes.
           ppn: Number of processors per node.
           running: dict of PBS ID:state pairs.
           sub_script: Path to qsub command
           sub_text: Path to queue file template.
           utilization: Number of active cores (based on showq).
           walltime: Maximum walltime on the machine.
           state: State code. 1=Up, 0=Full (auto-resets to 1 when jobs are
                 collected), -1=Down.
      check host()
           Pings the host to see if it is online. Returns False if it is offline.
      check_running()
           Uses the hosts data and one of the associated accounts to check the PBS queue on the Host. If it has been
           checked in the last 2 minutes, it will return the previously returned result.
      static create()
           Classmethod to create a Host model. Script will ask you questions about the host to add, and will return
           the created Host.
4.6.2 Account
class qmpy . Account (*args, **kwargs)
      Base class for a User account on a Host.
      Attributes:
           host
           id
           job
```

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```
user
username
```

# 4.6.3 User

```
class qmpy.User (*args, **kwargs)
    User model - stores an oqmd users information.

Relationships:
    Account via account_set
    Allocation via allocation_set
    Project via project_set

Attributes:
    id
        username
        first_name
```

is\_active

last\_name date\_joined

is\_staff

is\_superuser

last\_login

email

# 4.6.4 Allocation

```
class qmpy . Allocation (*args, **kwargs)
```

Base class for an Allocation on a computing resources.

### **Attributes:**

host

job

key

name

project

state

users

# 4.6.5 Project

```
class qmpy.Project (*args, **kwargs)

Base class for a project within qmpy.
```

#### **Attributes:**

allocations

entry

name

priority state task

users

# 4.7 Queue models

### 4.7.1 Task

```
class qmpy.Task(*args, **kwargs)
    Model for a :Task: to be done.
```

A :Task: consists of a module, which is the name of a computing script, and a set of keyword arguments, specified as a dictionary as the *kwargs* attribute of the task. In order for a Task for be completed, it must also be assigned one or more :Project:s.

# **Relationships:**

```
Entry via entry
Job via job_set
Project via project_set
```

#### **Attributes:**

id

created: datetime object for when the task was created.

finished: datetime object for when the task was completed.

module: The name of a function in scripts

kwargs: dict of keyword:value pairs to pass to the calculation

module.

priority: Priority of the task. Lower values are more urgent.

state: State code, given by the table below.

# Task codes:

Code	Description
-2	being held
-1	encountered error
0	ready to run
1	jobs running
2	completed

### complete()

Sets the Task state to 2 and populates the finished field.

#### errors

List of errors encountered by related calculations.

```
get_jobs (project=None, allocation=None, account=None, host=None)
```

Check the calculation module specified by the *Task*, and returns a list of Job objects accordingly.

Calls the task's entry's "do" method with the *Task.module* as the first argument, and passing *Task.kwargs* as keyword arguments.

**Returns:** List of Job objects. When nothing is left to do for the task, returns empty.

Raises:

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**ResourceUnavailableError:** Raise if for the specified project, allocation, account and/or host there are no available cores.

# jobs

List of jobs related to the task.

# projects

List of related projects.

### 4.7.2 Job

```
class qmpy . Job (*args, **kwargs)
```

Base class for job submitted to a compute cluster.

## **Relationships:**

Task via task

Account via account. The account the calculation is performed on.

Allocation via allocation. The allocation on which the calculation is being performed.

Entry via entry

### **Attributes:**

id

created: datetime object for when the task was created.

finished: datetime object for when the task was completed.

ncpus: # of processors assigned.

path: Origination path of the calculation.

run\_path: Path of the calculation on the compute resource.

qid: PBS queue ID number.

walltime: Max walltime (in seconds).

state: State code, defined as in the table below.

## Job codes

Code	Description
-1	encountered error
0	ready to submit
1	currently running
2	completed

# 4.8 Analysis Tools

```
class qmpy . PDF (structure, limit=6)
```

Container class for a Pair-distribution function.

Attributes: structure: Structure pairs: dict of (atom1, atom2):[distances] limit: maximum distance

```
get_pair_distances()
```

Loops over pairs of atoms that are within radius max\_dist of one another. Returns a dict of (atom1, atom2):[list of distances].

```
class gmpy . XRD (structure=None, measured=False, wavelength=1.5418, min_2th=10, max_2th=60, resolu-
                  tion=0.01)
     Container for an X-ray diffraction pattern.
     Attributes:
           peaks (List): List of Peak instances.
           measured (bool): True if the XRD is a measured pattern, otherwise False.
           min_2th (float): Minimum 2theta angle allowed. Defaults to 60 degrees.
           max_2th (float): Maximum 2theta angle allowed. Defaults to 10 degrees.
           wavelength (float): X-ray wavelength. Defaults to 1.5418 Ang.
           resolution (float): Minimum 2theta angle the XRD will distinguish between.
     get_intensities (bfactors=None, scale=None)
           Loops over all peaks calculating intensity.
           Keyword Arguments:
               bfactors (list) [list of B factors for each atomic site. Care must] taken to ensure that the order of B
                   factors agrees with the order of atomic orbits.
               scale (float) [Scaling factor to multiply the intensities by. If] scale evaluates to False, the intensities
                   will be re-normalized at the end such that the highest peak is 1.
     get peaks()
class qmpy.Peak (angle, multiplicity=None, intensity=None, hkl=None, xrd=None, width=None, mea-
                   sured=False)
      Attributes:
           angle (float): Peak 2*theta angle in radians.
           hkl (list): HKL indices of the peak.
           multiplicity (int): Number of HKL indices which generate the peak.
     lp factor()
           Calculates the Lorentz-polarization factor.
           http://reference.iucr.org/dictionary/Lorentz%E2%80%93polarization correction
     thermal factor (bfactor=1.0)
           Calculates the Debye-Waller factor for a peak.
           http://en.wikipedia.org/wiki/Debye-Waller_factor
class qmpy . Miedema (composition)
```

# H\_form\_ord()

Calculate the enthalpy of formation for an ordered compound of elements A and B with a composition xB of element B.

P

Chooses a value of P based on the transition metal status of the elements A and B.

There are 3 values of P for the cases where: both A and B are TM only one of A and B is a TM neither are TMs.

#### RtoP

Calculate and return the value of RtoP based on the transition metal status of elements A and B, and the elemental values of RtoP for elements A and B.

#### gamma

Calculate and return the value of Gamma\_AB (= Gamma\_BA) for the solvation of element A in element B.

#### $pick_a(elt)$

Choose a value of a based on the valence of element A.

#### class qmpy .GridData (data, lattice=None)

Container for 3d data, e.g. charge density or electron localization function.

#### find min coord (N=1)

Find the *N* lowest valued indices.

#### ind\_to\_cart (ind)

Converts an [i,j,k] index to [X,Y,Z] cartesian coordinate.

#### ind to coord (ind)

Converts an [i,j,k] index to [x,y,z] frational coordinate.

## interpolate (point, cart=False)

Calculates the value at *point* using trilinear interpolation.

**Arguments:** point: point to evaluate the value at.

**Keyword Arguments:** cart: If True, the point is taken as a cartesian coordinate. If not, it is assumed to be in fractional coordinates, default=False.

## local\_min(index)

Starting from *index* find the local value minimum.

Returns: index: shape (3,) index of local minimum. value: Value of grid at the local minimum.

### path (origin, end)

Gets a 1D array of values for a line connecting *origin* and *end*.

#### slice (point, orientation)

Return a 2D array of values for a slice through the GridData passing through *point* with normal vector *orientation*.

## class qmpy . SpinLattice (pairs)

### attempt\_flip()

Randomly selects a lattice point, and attempts to flip it.

dE = 2\*J\*sum(neighboring spins)

## compute\_total\_lattice\_energy()

Compute the total energy of the lattice using the Ising model hamiltonian:

```
H(s) = -J * sum_{\{i, j\}} (s_i * s_j)
```

So, for a positive interaction, J, the energy is minimized when all pairs are alike. Likewise, when J is negative, the energy is minimized when all pairs are unlike.

## run\_GCMC (mu=0)

Run Monte Carlo in the Grand Canonical Ensemble.

Examples: >>> sl = SpinLattice.create\_2d(10) >>> sl.run\_GCMC() >>> sl.run\_GCMC(-1) >>> sl.run\_GCMC(1)

```
run MC (x=None)
```

Run Monte Carlo in the Canonical Ensemble

### Examples:

```
>>> sl = SpinLattice.create_2d(10)
>>> sl.run_MC()
>>> sl.run_MC(0.1)
>>> sl.run_MC(0.25)
```

qmpy.analysis.nearest\_neighbors.find\_nearest\_neighbors (structure, method='closest', limit=5, tol=0.2)

For each atom in the *structure* assign the nearest neighbors.

# **Keyword Arguments:**

```
method ('closest' or 'voronoi'):
```

**closest:** Atoms A and B are neighbors, if and only if there is no atom C such that AC < AB and BC < AB. Once all pairs of this kind have been assigned, the nearest neighbors are those within *tol* of the shortest distance.

**voronoi:** Assign nearest neighbors based on voronoi construction. For each atom which generates a voronoi facet, *tol* specifies the minimum area of the facet before the atom is considered a nearest neighbor.

defaults to 'closest'

**limit:** Range outside of the unit cell that will be searched for nearest neighbor atoms.

tol: Varied depending on the method being used.

**Returns:** dict of Atom:list of Atom pairs. For each atom in the structure, its "neighbors" attribute will be set to the list of its nearest neighbors.

**Note:** Recommended to use the "closest" method unless you are sure you what the "voronoi" method will do. The voronoi neighbors are useful for some purposes, but are often not what are normally considered nearest neighbors. For example in BCC the second nearsest neighbors contribute to facets in the an atoms voronoi cell. The tolerance specification for this method sets the minimum area of such a facet required to add the atom as a nearest neighbor. In the BCC case, this cutoff must be set to at least 2.3 A^2 to exclude the facets due to second nearest neighbors.

# 4.9 Renderer

4.9. Renderer 55

**CHAPTER** 

**FIVE** 

# **ANALYSIS TOOLS**

Basically all thermodynamic analysis in qmpy is done starting from a PhaseSpace instance. If you have the database install and working, these are very easy to construct:

```
>>> ps = PhaseSpace('Li-Si')
>>> ps
<PhaseSpace bound by Li-Si>
```

Since the PhaseSpace was created without any extra arguments, it was assumed that you wanted to pull thermodynamic data from the OQMD, but you can fine tune the data that is included very easily. More on that later.

# 5.1 Convex Hull Construction

To obtain the convex hull for any phase space, simply access the *hull* attribute:

```
>>> ps.hull
set([<Equilibrium: Li13Si4-Li21Si5>, <Equilibrium: Li12Si7-Li7Si3>,
<Equilibrium: Li13Si4-Li7Si3>, <Equilibrium: LiSi-Li12Si7>, <Equilibrium:
Li21Si5-Li>, <Equilibrium: LiSi-Si>])
```

The hull is a set of Equilibrium objects, which have very natural attributes:

```
>>> eq = list(ps.hull)[0]
>>> eq.phases
[<Phase Li13Si4 : -0.240>, <Phase Li21Si5 : -0.212>]
>>> eq.chem_pots
{u'Si': -0.74005684434211016, u'Li': -0.086299552894736051}
```

# 5.2 Phase Stability

Positive for unstable phases, negative for stable phases.

```
>>> p = ps.phase_dict['Li13Si4'] # for just one phase
>>> ps.compute_stability(p)
>>> p.stability
-0.007333029175317474
>>> ps.compute_stabilities()
>>> ps.phase_dict['Li2Si'].stability
0.03116726059829092
```

# 5.3 Grand Canonical Linear Programming

# Examples:

```
>>> energy, phases = ps.gclp('LiSi2')
>>> energy
-0.404968066250002
>>> phases
{<Phase LiSi : -0.202>: 2.0, <Phase Si : 0>: 1.0}
>>> energy, phases = ps.gclp('Si', mus={'Li':-0.4})
>>> phases
{<Phase LiSi : -0.202>: 2.0}
```

# 5.4 Convex Hull Slices

Works by recursively using linear programming to find the lowest point contained within the a specified compositional region.

# Examples:

```
>>> ps = PhaseSpace('Fe2O3-Li2O')
>>> ps.hull
set([<Equilibrium: LiFe5O8-LiFeO2>, <Equilibrium:
LiFeO2-Li5FeO4>, <Equilibrium: LiFe5O8-Fe2O3>, <Equilibrium:
Li5FeO4-Li2O>])
```

# 5.5 Reaction Enumeration

# 5.6 Stability Conditions

**CHAPTER** 

SIX

# **EXAMPLES**

To be filled out in more detail

# 6.1 Identification of FCC decortations

First, we will find all binary entries:

```
>>> binaries = Entry.objects.filter(ntypes=2)
>>> fcc = Composition.get('Cu').ground_state.structure
```

Then we run through every structure, and see if replacing all atoms with Cu results in a structure that is equivalent (on volume scaling) with FCC Cu.:

```
>>> fccs = []
>>> for entry in binaries[:100]:
        struct = entry.structure
>>>
        ## Construct a dictionary of elt:replacement_elt pairs
>>>
        ## where every replacement is Cu
        rdict = dict((k, 'Cu') for k in entry.comp)
>>>
       test = struct.substitute(rdict, rescale=False,
                                        in_place=False)
>>>
        if fcc == test: # simple equality testing will work
>>>
>>>
            fccs.append(entry)
```

**Warning:** If you actually try to run this on the entire database, understand that it will take a pretty long time! Each entry tested takes between 0.1 and 1 second, so it would take most of 24 hours to run through all 80,000+ binary database entries.

# 6.2 Deviation from Vagard's Law

Use the element\_groups dictionary to look get a list of all simple metals:

```
>>> elts = element_groups['simple-metals']
```

Then, for each pair of metals get all of the entries, and their volumes.:

```
>>> vols = {}
>>> for e1, e2 in itertools.combinations(elts, r=2):
>>> entries = Composition.get_list([e1, e2])
>>> for entry in entries:
```

```
>>> vol = entry.structure.volume_pa
>>> vols[entry.name] = vols.get(entry.name, []) + [vol]
```

Then, for every composition get the Vagard's law volume.:

Addendum: \* Calculate an average error for each system \* Make a scatter plot for a few binaries show in volume vs x \* Look for cases where some are above and some are below \* Get relaxed volume of all stable compounds \* What about including the "nearly stable"

# **CHAPTER**

# **SEVEN**

# **CONTRIBUTE**

- Issue Tracker: http://github.com/wolverton-research-group/qmpy/issues
- Source: http://github.com/wolverton-research-group/qmpy

# CHAPTER

# **EIGHT**

# **SUPPORT**

If you are having issues, please let us know. We can be reached at oqmd.contact@gmail.com.

CHAPTER
NINE

# **LICENSE**

The project is licensed under the MIT license.

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# CHAPTER

# **TEN**

# **INDICES AND TABLES**

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