# **Frazier-PipeLine Documentation**

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**CHAPTER** 

ONE

### FRAZIER PIPELINE

Pipeline for submitting solubility simulations. Bonus output may include Unsaturated Mayer Bond Order amongst other things.

# 1.1 Installing

Currently installation involves first installing clancelot:

```
cd ~; git clone git@github.com:clancylab/clancelot2.0.git
```

And then cloning this project:

```
cd ~; git clone git@github.com:hherbol/frazier-pipeline.git
```

Note, you'll also have to append the frazier-pipeline/pys folder to your PYTHONPATH variable.

echo 'nexport PYTHONPATH="'/PATH/TO/FRAZIER/PIPELINE/pys:\$PYTHONPATH"" >> ~/.zshrc

### 1.2 Documentation

Documentation is necessary, and the following steps MUST be followed during contribution of new code:

### **Setup**

- 1. Download Sphinx. This can be done simply if you have pip installed via pip install -U Sphinx
- 2. Wherever you have *frazier-pipeline* installed, you want another folder called *frazier-pipeline-docs* (NOT as a subfolder of frazier-pipeline).

```
cd ~; mkdir frazier-pipeline-docs; cd frazier-pipeline-docs; git clone -b gh-pages git@github.com:hherbol/frazier-pipeline.git html
```

3. Forever more just ignore that directory (don't delete it though)

### **Adding Documentation**

Documentation is done using ReStructuredText format docstrings, the Sphinx python package, and indices with autodoc extensions. To add more documentation, first add the file to be included in *docs/source/conf.py* under *os.path.abspath('example/dir/to/script.py')*. Secondly, ensure that you have proper docstrings in the python file, and finally run *make full* to re-generate the documentation and commit it to your local branch, as well as the git *gh-pages* branch.

For anymore information on documentation, the tutorial follwed can be found here\_.

# 1.3 Using FPL

Automated calculations for the Frazier Pipeline are as follows:

- get\_UMBO()
- get\_MBO()

The following is still prone to bugs. It should work for num\_solvents=1; however, any more and it is prone to blowing up (ie, irrationally large enthalpy of solvations) - get\_enthalpy\_solvation()

```
fpl_auto.get_MBO (cation, halide, solvent)
```

Get the mayer bond order. The Mayer Bond Order (see get\_UMBO) for more details.

### **Parameters**

cation: str The cation within the perovskite.halide: str The halide within the perovskite.solvent: str The solvent of the system.

```
fpl_auto.get_UMBO (cation, halide, solvent, offset=2.0)
```

Get the unsaturation (average?) mayer bond order. The Mayer Bond Order (MBO) is well described here. In short, it is a numerical representation of the probability of how many electrons partake in a bond. For instance, a single bond would have a theoretical bond order of 1.0; however, in practice it may have more or less depending on how electrons distribute across the molecule. The MBO helps describe this, and the Unsaturated MBO (UMBO) helps represent this in a more understandable fashion. That is, if the UMBO is larger than zero, the bond is weaker than theory. If the UMBO is less than zero, then the bond is stronger than theory.

### **Parameters**

cation: str The cation within the perovskite.halide: str The halide within the perovskite.

**solvent:** *str* The solvent of the system.

**offset:** *float, optional* The offset supplied to get the UMBO. In most cases we consider, this is 2.0 as that is the theoretical bond order of a double bonded oxygen to sulfur.

### Return

**UMBO** [float] The Unsaturation Mayer Bond Order

# **CHAPTER**

# TWO

# **INDICES AND TABLES**

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