# MOF-Adsorbate-Initializer Documentation

Release v1.0

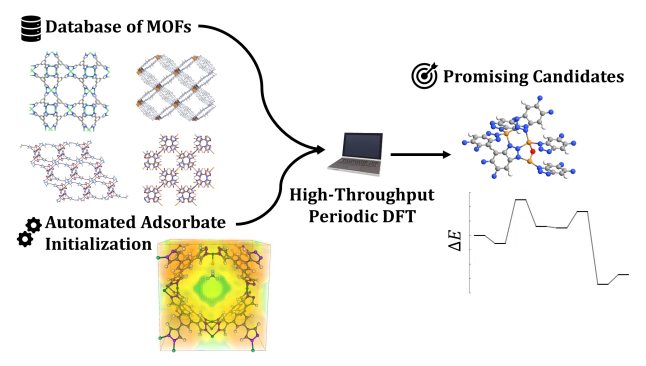
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The MOF Adsorbate Initializer (MAI) is a set of Python tools to initialize the positions of small molecule adsorbates at coordinatively unsaturated sites in metal-organic frameworks (MOFs) in a high-throughput, fully automated manner. The main benefit of MAI is the ability to systematically construct initial structures for large-scale density functional theory (DFT) calculations on many hundreds of materials. With a dataset of initial structures, workflow tools like PyMOFScreen or Atomate can be used to run the structural optimizations with your favorite optimizer of choice.



# CHAPTER 1

### Contents

### **1.1 Installation**

The MAI code can be found on GitHub and can be downloaded from the webpage or cloned via git clone git@github.com:arosen93/mof-adsorbate-initializer.git.

### 1.1.1 Dependencies

- 1. Python 3.6 or newer
- 2. Pymatgen 2018.11.30 or newer
- 3. ASE 3.16.0 or newer
- 4. OpenMetalDetector (recommended)

### 1.1.2 Installation Instructions

- 1. If you don't already have Python installed on your machine, you'll need to install Python 3.6 or newer. I recommend using the Anaconda distribution of Python 3 if you don't already have it installed.
- 2. You will then need to install MAI and all of the required dependencies. The easiest way to do this is to find the requirements.txt file in the MAI base directory and use the command pip install -r requirements.txt; pip install ..

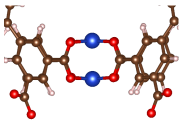
### **1.2 Tutorials**

It is probably best to lead by example, so in this tutorial I will walk you through a few examples meant to highlight MAI's capabilities and get you up to speed. Each one progressively builds on the one prior, so if you're new around here, I recommend you start at the top. Let's get started! In addition to the tutorials shown here, please refer to the examples folder in the main MAI directory for example scripts you can run and play around with.

### 1.2.1 Contents

### **Monatomics**

In this example, we'll work through how to add a single atom adsorbate to open metal site in MOFs. The CIF for the MOF we'll use in this example can be found here. This MOF is known as Cu-BTC and has the structure shown below:



The metal (Cu) sites here are shown in blue. There are multiple Cu sites per unit cell, and each Cu site is in a paddlewheel-like structure. For this example, we will consider the initialization of an O atom adsorbate to a single coordinatively unsaturated Cu site.

We'll start with the code that can do the job. Then we'll walk through what it all means.

```
import os
from mai.adsorbate_constructor import adsorbate_constructor
mof_path = os.path.join('example_MOFs','Cu-BTC.cif') #path to CIF of MOF
#add 0 adsorbate to index 0
ads = adsorbate_constructor(ads='0',d_MX1=1.75)
new_mof_atoms = ads.get_adsorbate(atoms_path=mof_path,site_idx=0)
```

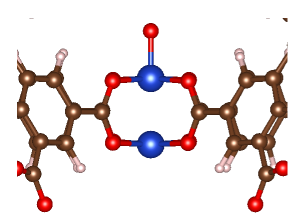
Okay, let's dive right in! MAI requires the calling of an object known as the *adsorbate\_constructor*, which tells MAI what kind of adsorbate you'd like to make. For simple monatomic species, there are only a few arguments you need to worry about.

- 1. The ads keyword argument is a string of the element that you want to add to the structure.
- 2. The d\_MX1 keyword argument is the desired distance between the adsorption site (i.e. the Cu species) and the adsorbate (in Å).

That takes care of initializing the *adsorbate\_constructor* object. Now we can use this object to call a function to initialize the adsorbate. This is done via *get\_adsorbate()*. The output of calling *get\_adsorbate()* is a new ASE Atoms object with the adsorbate initialized. The commonly used keywords for monatomic speies are as follows:

- 1. The atoms\_path keyword argument is the filepath to the starting CIF file of the MOF.
- 2. The site\_idx keyword argument is an integer representing the ASE Atoms index of the adsorption site (i.e. the Cu species). Later in this guide we'll show how this parameter can be determined automatically, but for now we have manually set it to the 0-th Atoms index via site\_idx=0, which corresponds to one of the Cu atoms. To find out the ASE indices for a given structure, you can inspect or visualize the Atoms object associated with the CIF file. Generally, it is the same indexing order as you'd find in your favorite CIF viewer (e.g. VESTA).

Now let's see what happens as a result of running this code! The initialized structure is shown below:



Exactly what we'd expect! Generally, MAI aims to satisfy two major conditions. The first condition is that it tries to maximize the symmetry of the first coordination sphere when the adsorbate is added. In this case, the geometry is square planar prior to adsorption, so MAI makes a square pyramidal structure when the monatomic species is added. The second condition is that MAI tries to minimize steric interactions when possible. In the case of a paddlewheel structure like Cu-BTC, the monatomic adsorbate could have been initialize in one of two directions normal to the square planar first coordination sphere. However, only one of those directions is geometrically accessible (the other is pointed inward between the Cu paddlewheel, which would not be reasonable).

That concludes our tutorial with monatomic adsorbates. Join me as we move onto more complicated systems! Up next is diatomics!

### **Diatomics**

In this example, we'll work through how to add two-atom adsorbates to open metal sites in MOFs. The CIF for the MOF we'll use for this example can be found here. This MOF is known Ni3(BTP)2, or Ni-BTP for short, and has a sodalite-like structure with square planar Ni cations.

### Homoatomic

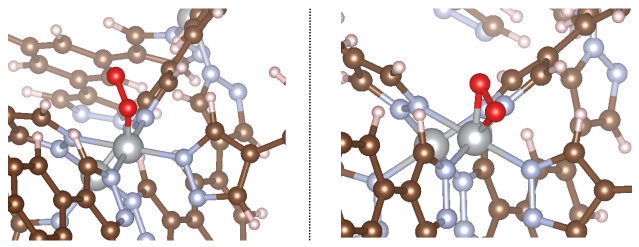
For this example, we will consider the initialization of an O2 molecule to a single coordinatively unsaturated Ni site. O2 can bind in an end-on ( $\eta$ 1-O) or side-on ( $\eta$ 2-O) mode depending on the structure. We'll consider both for this example. The code to handle this is shown below.

Like with the monatomic example, we need to initialize an *adsorbate\_constructor* object and then provide it the MOF of interest. In the case of diatomics, we have a few new keywords to introduce. In addition to the arguments described in the monatomic tutorial, we now need to be able to tell MAI what kind of denticity we would like (i.e.

end-on or side-on adsorption) and what we want the X1-X2 bond length and M-X1-X2 bond angle to be (if X1-X2 is our diatomic of interest and M is our metal adsorption site). The arguments used here are described below:

- 1. The ads argument is a string of the molecule that you want to add to the structure. Note that MAI will internally strip any characters following (and including) an underscore, so ads='02\_end' and ads='02\_side' both get stripped to '02'. That being said, the full string for the ads argument will be used when writing the filenames of the new CIFs, so using an underscore can be helpful for organizational purposes.
- 2. The d\_MX1 argument is the desired distance between the adsorption site (i.e. the Ni species) and the adsorbate (in Å). If the adsorbate is bound in an end-on fashion, this represents the M-X1 distance. If the adsorbate is bound in a side-on fashion, this represents the distance between M and the midpoint between X1 and X2. Here, we set d\_MX1=1.5.
- 3. The eta keyword argument is an integer representing the denticity. In other words, eta=1 would be an end-on adsorption mode, whereas eta=2 would be a side-on adsorption mode. By default, eta=1 if unspecified. For this example, we decided to explore both options.
- 4. The d\_X1X2 keyword argument is the desired distance between X1 and X2 (in Å). If not specified, it will default to the value for d\_MX1. Here, we decided to set d\_X1X2=1.2, which is a reasonable O-O bond distance.
- 5. The ang\_MX1X2 keyword argument is the angle between the adsorption site and the adsorbate (in degrees). If the adsorbate is bound in an end-on fashion, this represents the M-X1-X2 bond angle. If the adsorbate is bound in a side-on fashion, this represents the angle between M, the midpoint between X1 and X2, and X2. By default, it assumes ang\_MX1X2=180 if eta=1 and ang\_MX1X2=90 if eta=2. For this example, we use ang\_MX1X2=120 and ang\_MX1X2=90, respectively, which is representative of common O2 binding geometries.

That takes care of initializing the *adsorbate\_constructor* object. With this, we provide the object with the path to the MOF and the site index, and it will initialize the adsorbate for us. Now let's see what happens as a result of running this code! The initialized structures are shown below:



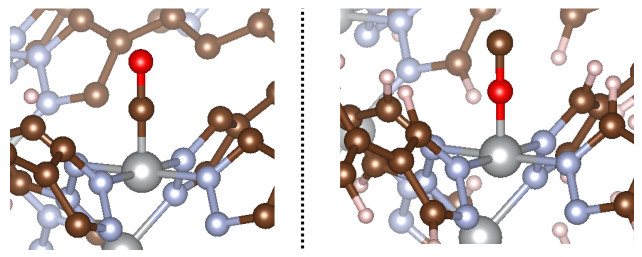
Exactly what we'd expect yet again! You can see that in the first example, O2 is bound end-on, whereas in the second it is bound side-on, as specified in the example script. The bond angles and distances are the same as those specified in the input file.

### Heteroatomic

MAI also supports heteratomic adsorbates. In this example, we'll consider the adsorption of a single CO molecule with the same MOF. The only thing that changes for heteroatomic adsorbates is that you need to tell MAI which atom is the "connecting atom" (i.e. the atom of the adsorbate bound to the metal adsorption site) if bound in an end-on fashion. By default, MAI will assume that the first atom in ads is the connecting atom. Therefore, setting ads='CO' or ads='OC' would yield M-C-O or M-O-C binding modes, respectively.



#### The result of running this code is shown below:



That concludes our tutorial for diatomic adsorbates. Now onto triatomics!

#### **Triatomics**

In this example, we'll work through how to add three-atom adsorbates to open metal sites in MOFs. The CIF for the MOF we'll use for this example can be found here. This MOF is known as MIL-8BB and has trimetallic nodes with each metal cation in a square pyramidal geometry.

### **Contiguous Adsorbate**

For this example, we will consider the initialization of a "contiguous" triatomic adsorbate. What I mean by this is that if we treat the adsorbate as an arbitrary molecule X1-X2-X3, then the adsorption process is described by M-X1-X2-X3. In this example, we'll consider the adsorption of N2O to an Fe site, in both an  $\eta$ 1-N and  $\eta$ 1-O binding mode. The code to handle this is shown below.

```
import os
from mai.adsorbate_constructor import adsorbate_constructor
mof_path = os.path.join('example_MOFs','Fe-MIL-88B.cif') #path to CIF of MOF
#add N20 adsorbate in η1-N mode
ads = adsorbate_constructor(ads='N20',d_MX1=2.0,d_X1X2=1.13,ang_MX1X2=180,d_X2X3=1.19)
new_mof_atoms1 = ads.get_adsorbate(atoms_path=mof_path,site_idx=0)
```

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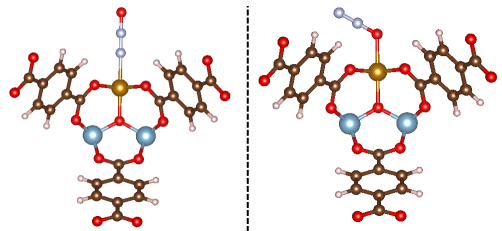
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```
#add N2O adsorbate in η1-O mode
ads = adsorbate_constructor(ads='ON2',d_MX1=2.0,d_X1X2=1.19,ang_MX1X2=120,d_X2X3=1.13)
new_mof_atoms2 = ads.get_adsorbate(atoms_path=mof_path,site_idx=0)
```

Like with the previous examples, we need to initialize an *adsorbate\_constructor* object and then provide it the MOF of interest. In the case of triatomics, we have a few new keywords to introduce. In addition to the arguments described in the previous examples, we can now provide MAI additional geometric parameters if desired. Namely, the new features are now that we can include the X2-X3 bond length and the X1-X2-X3 bond angle. The arguments used here are described below:

- 1. The ads, d\_MX1X2, d\_X1X2, and ang\_MX1X2 are the same as before.
- 2. Now, we have the option to add the d\_X2X3 keyword argument, which specifies the X2-X3 distance. It defaults to d\_X2X3=d\_X1X2 if not specified.
- 3. We can also add the ang\_triads keyword argument, which specifies the X1-X2-X3 bond angle. It defaults to ang\_triads=180 if not specified.

That takes care of initializing the *adsorbate\_constructor* object. With this, we provide the object with the path to the MOF, and it will initialize the adsorbate. Now let's see what happens as a result of running this code! The initialized structures are shown below:

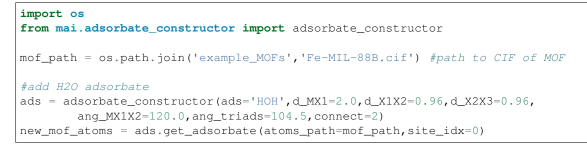


Exactly what we'd expect once more! You can see that in the first example, N2O is bound in an  $\eta$ 1-N mode, whereas the second is bound in an  $\eta$ 1-O mode, as specified. Feel free to play around with the bond distance and bond angle arguments to get a feel for how MAI works.

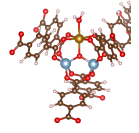
### Noncontiguous Adsorbate

The last bit of trickery comes into play when dealing with what I'll call "noncontiguous" adsorbates. These are adsorbates like water, where it is triatomic, but it is not bound in a sequential fashion. As with water, you will have a central atom of the adsorbate bound to the metal (instead of an M-O-H-H adsorption mode). To tell MAI about this desired adsorption mode, *adsorbate\_constructor* has a keyword argument named connect, which is the atom number in ads that should be bound to the adsorption site. For the case of ads='HOH', we should set connect=2 to have the second atom (i.e. the O atom) bound to the adsorption site. Note that for this reason we cannot use ads='H2O' here.

An example code is shown below. The main thing to keep in mind is that now the connecting atom of the adsorbate is X2 instead of X1. If not specified, ang\_triads will default to ang\_triads=ang\_MX1X2 if connect=2, but here we set it to 104.5°, as is standard for H2O.



This results in the following initialized structure:



That concludes our tutorial for triatomic adsorbates.

#### **Open Metal Detection**

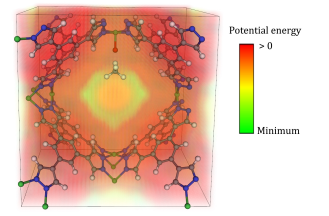
In the previous examples, we manually specified the desired adsorption site via the site\_idx argument to  $get_adsorbate()$ . In practice, it is often desirable to have an automated approach for determining which atom in a MOF is the desired adsorption site. By default, if the user does not specify site\_idx, MAI will assume that you want an automated approach for determining this parameter. By default, MAI will attempt to read in the results from the OpenMetalDetector (OMD) code, which can be used to output information about the number, type, and properties of various open metal sites in a set of MOFs. Instructions for using OMD can be found elsewhere, but the main idea is to call OMD prior to calling MAI. An example is shown below for adding an O atom adsorbate to a bunch of MOFs stored in your current working directory.

```
import os
from omsdetector import MofCollection
from mai.adsorbate constructor import adsorbate constructor
mofs_path = 'example_MOFs' #path to folder of CIFs
oms_analysis_folder = os.getcwd() #path to store the OMS results
oms_data_path = os.path.join(oms_analysis_folder,'oms_results') #path to oms_results...
⇔folder
#Run the Open Metal Detector
mof_coll = MofCollection.from_folder(collection_folder=mofs_path,
        analysis_folder=oms_analysis_folder)
mof_coll.analyse_mofs()
#add adsorbate for every CIF in mofs_path
for file in os.listdir(mofs_path):
        if '.cif' not in file:
                continue
       mof_path = os.path.join(mofs_path,file)
        ads = adsorbate_constructor(ads='0', d_MX1=1.75)
        new_mof_atoms = ads.get_adsorbate(atoms_path=mof_path,
                omd_path=oms_data_path)
```

You can see that this workflow is nearly identical to that in the monatomics tutorial, except we did not specify site\_idx in get\_adsorbate(), and we ran OMD's MofCollection.from\_folder() command prior to calling MAI's adsorbate\_constructor. Generally, the only arguments you'll need to provide to MofCollection.from\_folder() is the collection\_folder and analysis\_folder, which is where the CIF files are located and where you'd like to store the OMD results, respectively. Then, you'll run OMD's analyse\_mofs() command to run the analysis. From there, you're ready to run MAI as usual! The only new MAI-related keyword is that get\_adsorbate() can take an omd\_path keyword argument, which should be where the results from the OMD run are stored. This is generally located at analysis\_folder/oms\_results.

### **Potential Energy Grids**

Some adsorbates do not lend themselves well to the geometric approaches laid out thus far. This is particularly the case for adsorbates that are physisorbed relatively far away from an adsorption site, as opposed to chemisorbed nearby. In these cases, an alternative way of initializing the adsorbate can be considered by mapping out a potential energy grid (PEG) of each MOF and putting the adsorbate in a low-energy site within some cutoff radius of the proposed adsorption site. A visualized example of such a PEG is shown below for a methane adsorbate near a proposed Ni-O active site.



MAI supports two different formats for PEGs. The first is the cube file format, such as that generated from PorousMaterials.jl. Details of how to generate such PEGs can be found here. The second accepted format is a space-delimited file with four columns of (x,y,z,E) entries, where E is the potential energy and (x,y,z) are the coordinates. Each new line represents a new (x,y,z,E) vector. We'll refer to this file as an ASCII grid.

Currently, only single-site CH4 adsorbates are supported with PEGs, although in principle it is trivial to consider other adsorbates as well. When using PEGs to initialize the position of CH4 adsorbates, the C atom of the CH4 molecule will be placed in the low-energy site, and the four remaining H atoms will be arranged to form the tetrahedral structure of CH4, with one of the H atoms pointed directly toward the adsorption site.

Unlike with the monatomics, diatomics, and triatomics tutorials, when dealing with PEGs, one must use the  $get\_adsorbate\_grid()$  function instead of  $get\_adsorbate()$ . The  $get\_adsorbate\_grid()$  function is quite simple. An example code is shown below to initialize an adsorbate based on an example PEG and example MOF.

```
import os
from mai.adsorbate_constructor import adsorbate_constructor
from ase.io import read
grid_path = os.path.join('example_MOFs','energy_grids_ASCII') #path to PEG
mof_path = os.path.join(grid_path,'AHOKIR01-0.cif') #path to CIF of MOF
#Select the last 0 index in the MOF as the adsorption site
```

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The new arguments used in this code are described below:

- 1. The atoms\_path and site\_idx are the same as for get\_adsorbate().
- 2. The grid\_path keyword argument is the path to the PEG file.
- 3. The grid\_format keyword argument can be either grid\_format='cube' or grid\_format='ASCII'.

The result of running the previous example code is the following structure.



#### **Advanced Skills**

The previous tutorials should get you up and running so you can automate the construction of adsorbate geometries for a wide range of MOFs. However, for more advanced users, there are some additional features that may be of use. Let's walk through some more complex examples.

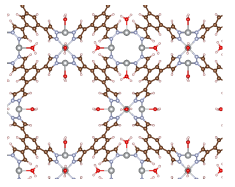
#### Adding Multiple Adsorbates

Instead of adding a single adsorbate, you may want to add an adsorbate to every metal site in a given MOF. We will consider the Ni-BTP MOF from the diatomic tutorial, except this time we will add an H2O molecule to every Ni site. This can be done using the code below.

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There are a few changes we've made to the usual workflow. The first is that we needed to identify all of the possible adsorption site indices, which we have defined as Ni\_idx. This can be done using ASE's built-in tools for working with ASE Atoms ojbects. We then iterate over each site and add an H2O molecule. Since MAI adds adsorbates sequentially, we don't want to write out all the intermediate structures, so we also set write\_file=False in get\_adsorbate(). The last new aspect to introduce is that, instead of passing a CIF file via atoms\_path in get\_adsorbate(), we can directly pass in an ASE Atoms object via the atoms keyword argument. This is useful when you make modifications to the same MOF in a loop, as done here. The result of running this code is shown below.



### **Neighbor Algorithms**

When specifying site\_idx, the get\_adsorbate() function will automatically use Pymatgen's built-in nearest neighbor algorithms to determine the atoms in the first coordination sphere of the adsorption site. By default, MAI uses Pymatgen's crystalNN algorithm, but additional algorithms are available as listed in get\_NNs\_pm() and described here. Through iterative testing, we have found crystalNN to be the best-performing algorithm for MOFs in general, but other algorithms can be considered if desired.

#### **Pre-specification of Indices**

In addition to the site\_idx keyword in the *get\_adsorbate()* function, it is also possible to manually specify the indices of the atoms in the first coordination sphere via the NN\_indices keyword argument. This flexibility makes it possible to use a wide range of workflows in determining the desired adsorption site and coordinating atoms.

### 1.3 Citing MAI

If you use MAI in your work, please cite A.S. Rosen, J.M. Notestein, R.Q. Snurr. "Identifying Promising Metal-Organic Frameworks for Heterogeneous Catalysis via High-Throughput Periodic Density Functional Theory", J. Comput. Chem. (2019). DOI: [10.1002/jcc.25787](https://onlinelibrary.wiley.com/doi/10.1002/jcc.25787). If you feel so inclined, you can also reference the corresponding Zenodo DOI for the MAI code: 10.5281/zenodo.1451875.

### 1.4 Code documentation

### 1.4.1 mai.ads\_sites module

class	<pre>mai.ads_sites.ads_pos_optimizer(adsorbate_constructor,</pre>	
	new_mofs_path=None,	
	$log\_stats=True)$	

write\_file=True,
error\_path=None,

Bases: object

This identifies ideal adsorption sites

Args: adsorbate\_constructor (class): adsorbate\_constructor class containing many relevant defaults

write\_file (bool): if True, the new ASE atoms object should be written to a CIF file (defaults to True)

new\_mofs\_path (string): path to store the new CIF files if write\_file is True (defaults to /new\_mofs)

error\_path (string): path to store any adsorbates flagged as problematic (defaults to /errors)

log\_stats (bool): print stats about process

check\_and\_write(new\_mof, new\_name)

Check for overlapping atoms and write CIF file

Args: new\_mof (ASE Atoms object): the new MOF-adsorbate complex

new\_name (string): the name of the new CIF file to write

**Returns:** overlap (boolean): True or False for overlapping atoms

construct\_mof(mof, ads\_pos, site\_idx)

Construct the MOF-adsorbate complex

Args: ads\_pos\_optimizer (class): see ads\_sites.py for details

ads (string): adsorbate species

ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

Returns: mof (ASE Atoms object): ASE Atoms object with adsorbate

#### get\_NNs (ads\_pos, site\_idx)

Get the number of atoms nearby the proposed adsorption site within r\_cut

Args: ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

site\_idx (int): ASE index for adsorption site

Returns: NN (int): number of neighbors within r\_cut

min\_dist (float): distance from adsorbate to nearest atom

n\_overlap (int): number of overlapping atoms

- get\_bi\_ads\_pos (normal\_vec, scaled\_sum\_dist, center\_coord, site\_idx)
  Get adsorption site for a 2-coordinate site
  - Args: normal\_vec (numpy array): 1D numpy array for the normal vector to the line

scaled\_sum\_dist (numpy array): 2D numpy array for the scaled Euclidean distance vectors between each coordinating atom and the central atom (i.e. the adsorption site)

center\_coord (numpy array): 1D numpy array for adsorption site

site\_idx (int): ASE index of adsorption site

Returns: ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

**get\_dist\_planar** (*normal\_vec*) Get distance vector for planar adsorption site

Args: normal\_vec (numpy array): 1D numpy array for normal vector

Returns: dist (float): distance vector scaled to d\_MX1

get\_new\_atoms (ads\_pos, site\_idx)

Get new ASE atoms object with adsorbate from pymatgen analysis

Args: ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

Returns: new\_mof (ASE Atoms object): new ASE Atoms object with adsorbate

**get\_new\_atoms\_grid** (*site\_pos*, *ads\_pos*) Get new ASE atoms object with adsorbate from energy grid

**Args:** ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

Returns: new\_mof (ASE Atoms object): new ASE Atoms object with adsorbate

- get\_nonplanar\_ads\_pos (scaled\_sum\_dist, center\_coord) Get adsorption site for non-planar structure
  - **Args:** scaled\_sum\_dist (numpy array): 2D numpy array for the scaled Euclidean distance vectors between each coordinating atom and the central atom (i.e. the adsorption site)

center\_coord (numpy array): 1D numpy array for adsorption site

Returns: ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

- get\_opt\_ads\_pos (mic\_coords, site\_idx)
  Get the optimal adsorption site
  - **Args:** mic\_coords (numpy array): 2D numpy array for the coordinates of each coordinating atom using the central atom (i.e. adsorption site) as the origin

site\_idx (int): ASE index of adsorption site

**Returns:** ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

#### **get\_planar\_ads\_pos** (*center\_coord*, *dist*, *site\_idx*) Get adsorption site for planar structure

Args: center\_coord (numpy array): 1D numpy array for adsorption site (i.e. the central atom)

site\_idx (int): ASE index for adsorption site

Returns: ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

get\_tri\_ads\_pos (normal\_vec, scaled\_sum\_dist, center\_coord, site\_idx)
Get adsorption site for a 3-coordinate site

Args: normal\_vec (numpy array): 1D numpy array for the normal vector to the line

scaled\_sum\_dist (numpy array): 2D numpy array for the scaled Euclidean distance vectors between each coordinating atom and the central atom (i.e. the adsorption site)

center\_coord (numpy array): 1D numpy array for adsorption site

site\_idx (int): ASE index of adsorption site

Returns: ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

### 1.4.2 mai.adsorbate\_constructor module

Bases: object

This class constructs an ASE atoms object with an adsorbate Initialized variables

Args: ads (string): string of element or molecule for adsorbate (defaults to 'X')

d\_MX1 (float): distance between adsorbate and surface atom. If used with get\_adsorbate\_grid, it represents the maximum distance (defaults to 2.0)

eta (int): denticity of end-on (1) or side-on (2) (defaults to 1)

connect (int): the connecting atom in the species string (defaults to 1)

d\_X1X2 (float): X1-X2 bond length (defaults to 1.25)

 $d_X2X3$  (float): X2-X3 bond length for connect == 1 or X1-X3 bond length for connect == 2 (defaults to  $d_bond1$ )

ang\_MX1X2 (float): site-X1-X2 angle (for diatomics, defaults to 180 degrees except for side-on in which it defaults to 90 or end-on O2 in which it defaults to 120; for triatomics, defaults to 180 except for H2O in which it defaults to 104.5)

ang\_triads (float): X3-X1-X2 angle (defaults to 180 degrees for connect == 1 and 90 degrees for connect == 2)

r\_cut (float): cutoff distance for calculating nearby atoms when ranking adsorption sites

sum\_tol (float): threshold to determine planarity. when the sum of the Euclidean distance vectors of coordinating atoms is less than sum\_tol, planarity is assumed

rmse\_tol (float): second threshold to determine planarity. when the root mean square error of the best-fit plane is less than rmse\_tol, planarity is assumed

overlap\_tol (float): distance below which atoms are assumed to be overlapping

get\_adsorbate (atoms\_path=None, site\_idx=None, omd\_path=None, NN\_method='crystal', allowed\_sites=None, write\_file=True, new\_mofs\_path=None, error\_path=None,

NN\_indices=None, atoms=None, new\_atoms\_name=None)

Add an adsorbate using PymatgenNN or OMD

Args:

atoms\_path (string): filepath to the CIF file

site\_idx (int): ASE index for the adsorption site

omd\_path (string): filepath to OMD results folder (defaults to '/oms\_results')

NN\_method (string): string representing the desired Pymatgen nearest neighbor algorithm. options include 'crystal', vire', 'okeefe', and others. See NN\_algos.py (defaults to 'crystal')

allowed\_sites (list of strings): list of allowed site species for use with automatic OMS detection

write\_file (bool): if True, the new ASE atoms object should be written to a CIF file (defaults to True)

new\_mofs\_path (string): path to store the new CIF files if write\_file is True (defaults to /new\_mofs within the directory containing the starting CIF file)

error\_path (string): path to store any adsorbates flagged as problematic (defaults to /errors within the directory containing the starting CIF file)

NN\_indices (list of ints): list of indices for first coordination sphere (these are usually automatically detected via the default of None)

atoms (ASE Atoms object): the ASE Atoms object of the MOF to add the adsorbate to (only include if atoms\_path is not specified)

new\_atoms\_name (string): the name of the MOF used for file I/O purposes (defaults to the basename of atoms\_path if provided)

**Returns:** new\_atoms (Atoms object): ASE Atoms object of MOF with adsorbate

get\_adsorbate\_grid (atoms\_path=None, site\_idx=None, grid\_path=None, grid\_format='ASCII', write\_file=True, new\_mofs\_path=None, error\_path=None, atoms=None,

new\_atoms\_name=None)

This function adds a molecular adsorbate based on a potential energy grid

Args: atoms\_path (string): filepath to the CIF file

site\_idx (int): ASE index for the adsorption site

grid\_path (string): path to the directory containing the PEG (defaults to /energy\_grids)

grid\_format (string): accepts either 'ASCII' or 'cube' and is the file format for the PEG (defaults to 'ASCII')

write\_file (bool): if True, the new ASE atoms object should be written to a CIF file (defaults to True)

new\_mofs\_path (string): path to store the new CIF files if write\_file is True (defaults to /new\_mofs)

error\_path (string): path to store any adsorbates flagged as problematic (defaults to /errors)

atoms (ASE Atoms object): the ASE Atoms object of the MOF to add the adsorbate to (only include if atoms\_path is not specified)

new\_atoms\_name (string): the name of the MOF used for file I/O purposes (defaults to the basename of atoms\_path if provided)

**Returns:** new\_atoms (Atoms object): ASE Atoms object of MOF with adsorbate

### 1.4.3 mai.grid\_handler module

mai.grid\_handler.cube\_to\_xyzE(cube\_file)

Converts cube to ASCII file Adopted from code by Julen Larrucea Original source: https://github.com/julenl/molecular\_modeling\_scripts

**Args:** cube\_file (string): path to cube file

**Returns:** pd\_data (Pandas dataframe): dataframe of (x,y,z,E) grid

mai.grid\_handler.get\_best\_grid\_pos (atoms, max\_dist, site\_idx, grid\_filepath)
Finds minimum energy position in grid dataframe

Args: atoms (ASE Atoms object): Atoms object of structure

max\_dist (float): maximum distance from active site to consider

site\_idx (int): ASE index of adsorption site

grid\_filepath (string): path to energy grid

Returns: ads\_pos (array): 1D numpy array for the ideal adsorption position

mai.grid\_handler.grid\_within\_cutoff (df, atoms, max\_dist, site\_pos, partition=1000000.0)
Reduces grid dataframe into data within max\_dist of active site

**Args:** df (pandas df object): df containing energy grid details (x,y,z,E)

atoms (ASE Atoms object): Atoms object of structure

max\_dist (float): maximum distance from active site to consider

site\_pos (array): numpy array of the adsorption site

partition (float): how many data points to partition the df for. This is used to prevent memory overflow errors. Decrease if memory errors arise.

**Returns:** new\_df (pandas df object): modified df only around max\_dist from active site and also with a new distance (d) column

mai.grid\_handler.read\_grid(grid\_filepath)

Convert energy grid to pandas dataframe

Args: grid\_filepath (string): path to energy grid (must be .cube or .grid)

Returns: df (pandas df object): df containing energy grid details (x,y,z,E)

### 1.4.4 mai.NN\_algos module

mai.NN\_algos.get\_NNs\_pm(atoms, site\_idx, NN\_method)
Get coordinating atoms to the adsorption site

Args: atoms (Atoms object): atoms object of MOF

site\_idx (int): ASE index of adsorption site

NN\_method (string): string representing the desired Pymatgen nearest neighbor algorithm: refer to http: //pymatgen.org/\_modules/pymatgen/analysis/local\_env.html

Returns: neighbors\_idx (list of ints): ASE indices of coordinating atoms

### 1.4.5 mai.oms\_handler module

mai.oms\_handler.get\_ase\_NN\_idx (atoms, coords)
 Get the ASE indices for the coordinating atoms

Args: atoms (Atoms object): ASE Atoms object for the MOF

coords (numpy array): coordinates of the coordinating atoms

**Returns:** ase\_NN\_idx (list of ints): ASE indices of the coordinating atoms

mai.oms\_handler.get\_ase\_oms\_idx (atoms, coords)
 Get the ASE index of the OMS

Args: atoms (Atoms object): ASE Atoms object for the MOF

coords (numpy array): coordinates of the OMS

Returns: ase\_oms\_idx (int): ASE index of OMS

mai.oms\_handler.get\_omd\_data (oms\_data\_path, name, atoms)
Get info about the open metal site from OpenMetalDetector results files

Args: oms\_data\_path (string): path to the OpenMetalDetector results

name (string): name of the MOF

atoms (ASE Atoms object): Atoms object for the MOF

Returns: omsex\_dict (dict): dictionary of data from the OpenMetalDetector results

### 1.4.6 mai.regression module

mai.regression.OLS\_fit (xyz) Make ordinary least squares fit to z=a+bx+cy and return the normal vector Args: xyz (numpy array): 2D numpy array of XYZ values (N rows, 3 cols) Returns: normal\_vec (numpy array): 1D numpy array for the normal vector mai.regression.TLS\_fit (xyz) Make total least squares fit to ax+by+cz+d=0 and return the normal vector Args: xyz (numpy array): 2D numpy array of XYZ values (N rows, 3 cols)

Returns: rmse (float): root mean square error of fit

normal\_vec (numpy array): 1D numpy array for the normal vector

### 1.4.7 mai.species\_rules module

mai.species\_rules.add\_CH4\_SS (mof, site\_idx, ads\_pos)
 Add CH4 to the structure from single-site model

Args: mof (ASE Atoms object): starting ASE Atoms object of structure

site\_idx (int): ASE index of site based on single-site model

ads\_pos (array): 1D numpy array for the best adsorbate position

**Returns:** mof (ASE Atoms object): ASE Atoms object with adsorbate

Add diatomic to the structure

Args: mof (ASE Atoms object): starting ASE Atoms object of structure

ads\_species (string): adsorbate species

ads\_pos (array): 1D numpy array for the best adsorbate position

site\_idx (int): ASE index of site

d\_X1X2 (float): X1-X2 bond length (defaults to 1.25)

ang\_MX1X2 (float): site-X1-X2 angle (defaults to 180 degrees except for side-on in which it defaults to 90 degrees prior to the centering)

eta (int): denticity of end-on (1) or side-on (2) (defaults to 1)

connect (int): the connecting atom in the species string (defaults to 1)

r\_cut (float): cutoff distance for calculating nearby atoms when ranking adsorption sites (defualts to 2.5)

overlap\_tol (float): distance below which atoms are assumed to be overlapping (defualts to 0.75)

Returns: mof (ASE Atoms object): ASE Atoms object with adsorbate

mai.species\_rules.add\_monoatomic (mof, ads\_species, ads\_pos)
Add adsorbate to the ASE atoms object

Args: mof (ASE Atoms object): starting ASE Atoms object of structure

ads\_species (string): adsorbate species

ads\_pos (numpy array): 1D numpy array for the proposed adsorption position

Returns: mof (ASE Atoms object): ASE Atoms object with adsorbate

Add triatomic to the structure

Args: mof (ASE Atoms object): starting ASE Atoms object of structure

ads\_species (string): adsorbate species

ads\_pos (array): 1D numpy array for the best adsorbate position

site\_idx (int): ASE index of site

d\_X1X2 (float): X1-X2 bond length (defaults to 1.25)

 $d_X2X3$  (float): X2-X3 bond length for connect == 1 or X1-X3 bond length for connect == 2 (defaults to  $d_X1X2$ )

ang\_MX1X2 (float): site-X1-X2 angle (defaults to 180 degrees)

ang\_triads (float): triatomic angle (defaults to 180 degrees for connect == 1 and ang\_MX1X2 for connect == 2)

connect (int): the connecting atom in the species string (defaults to 1)

r\_cut (float): cutoff distance for calculating nearby atoms when ranking adsorption sites (defualts to 2.5)

overlap\_tol (float): distance below which atoms are assumed to be overlapping (defualts to 0.75)

Returns: mof (ASE Atoms object): ASE Atoms object with adsorbate

### 1.4.8 mai.tools module

mai.tools.get\_refcode (atoms\_filename)
Get the name of the MOF

Args: atoms\_filename (string): filename of the ASE Atoms object (accepts CIFS, POSCARs, and CONTCARs)

**Return:** refcode (string): name of MOF (defaults to 'mof' if the original filename is just named CONTCAR or POSCAR)

mai.tools.**string\_to\_formula** (*species\_string*) Convert a species string to a chemical formula

**Args:** species\_string (string): string of atomic/molecular species

Return: formula (string): stoichiometric chemical formula

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