ENGN2770: Atomistic Reaction Engineering

September 17, 2020

1 Computational Tutorial

[2]:

This is the pdf version. The interactive version (which includes necessary environment files) can be accessed from using the link provided:

(Note: This can take a minute or two to load and download required environment files)

Once you open the link, you can download the "engn2770_computational_tutorial.ipynb" and save it on your local computer.

Note, you will need jupyter notebook along with other necessary packages if you wish to use it on your local machine(ASE, python, matplotlib, numpy).

1.1 A) Intro to linux bash commands (https://files.fosswire.com/2007/08/fwunixref.pdf)

```
[1]: from IPython.display import Image
[2]: Image(filename='extras/linux_tutorial_page1.png')
```

Job	Command
identify the directory you are in	pwd
see the files in the directory	Is
see the files in the directory with hidden files	ls -a
go to your home directory	cd (or, cd ~/)
change directory to <directory_name></directory_name>	cd <directory_name></directory_name>
move one directory up	cd
move two directories up	cd//
make directory <directory_name></directory_name>	mkdir <directory_name></directory_name>
copy a file	cp
remove a file / remove a file recursively (careful no way to revert if you delete a file/directory)	rm <file_name> / rm -r <file_name></file_name></file_name>
create an empty file	touch <file_name></file_name>

```
[3]: Image(filename='extras/linux_tutorial_page2.png')
```

[3]:

Job	Command
move a file	mv
display the contents of the file	cat <file_name></file_name>
copy file from your local computer to CCV	scp -r <username@ssh.ccv.brown.edu: file_name="" location="" new="" path="" to=""></username@ssh.ccv.brown.edu:>
text editors	vim/ vi/ nano/ gedit

```
Environment Setup:
$ cc
(vi/nano/gedit) ~/.modules

Inside your .modules, type
$ source /gpfs/data/ap31/ap31/teach/2020-engn2770/settings/engn2770packages

Save your .modules, and type
$ source ~/.modules
(This needs to be done only once - i.e whenever you change your ~/.modules)
$ engn2770 loadase
$ python3
$ import ase
$ ase. file (The output should point to '/users/ap31/data/ap31/teach/2020-engn2770/packages/ase/ase/__init__.py'

Note: The first line in your python script should be always be #!/usr/bin/env python3
```

1.2 B) Python

1.2.1 Basic arithmetic

```
[4]: # Basic arithmetic
a = 2.
b = 3.
print(a + b)
```

5.0

```
[5]: # square of a number
sq = a**2
print(sq)
```

4.0

```
[6]: # division & % print(4./2)

# remainder of a division
```

```
print(15.%8)
print(8.%2) # If this is 0, then 8 is divisible by 2!
```

2.0

7.0

0.0

1.2.2 Functions

```
[7]: # functions

# define add function
def add(a,b):
    return a+b

# define power function
def power(a,pow=2):
    return a**pow

# call add function
add_result = add(2.,3.)
print(add_result)

# call power function
power_result = power(5.,3)
print(power_result)
```

5.0 125.0

```
[8]: sum_result = sum((2.,2.))
print(sum_result)

pow_result = pow(5.,4)
print(pow_result)
```

4.0 625.0

1.2.3 numpy, list & arrays

```
[9]: # using numpy (Numerical Python)
import numpy as np

sum_result = np.sum((2.,-3.))
print(sum_result)

power_result = np.power(2.,8)
print(power_result)
```

```
-1.0
256.0
```

```
[10]: # list
things = ['a',7]

for thing in things:
    print(thing)
```

a 7

```
[11]: # Basic list commands

_list = [1,2,3,4]

#_list.append(5)

#_list.pop(2)

#_list.reverse()

print(_list)
```

[1, 2, 3, 4]

```
[12]: import numpy as np

# return evenly spaced numbers over a specified interval

a = np.linspace(1,12,12)
print(a)
print(a.shape)

# convert a 1D array into 2D
a.reshape(3,4)
```

```
[ 1. 2. 3. 4. 5. 6. 7. 8. 9. 10. 11. 12.] (12,)
```

1.2.4 For loop & conditional statements

```
[13]: # for loops
for element in range(5):
    print(element, element**2)
```

```
1 1
     2 4
     3 9
     4 16
[14]: # For loops with conditional statements
      dice = [1,2,3,4,5,6]
      type_list = []
      for roll in dice:
          if roll % 2 == 0:
              type_list.append('even')
          else:
              type_list.append('odd')
      print(type_list)
     ['odd', 'even', 'odd', 'even', 'odd', 'even']
[15]: num = np.linspace(0,10,11)
     print(num)
      doubled_list = []
      for element in num:
          if element > 5:
              doubled_list.append(element*2)
      print(doubled_list)
     [0. 1. 2. 3. 4. 5. 6. 7. 8. 9. 10.]
     [12.0, 14.0, 16.0, 18.0, 20.0]
[16]: # For loop with conditional statement in one line
      num = np.linspace(0,10,11)
      print(num)
      doubled = [element * 2 for element in num if element>5]
      print(doubled)
     [0. 1. 2. 3. 4. 5. 6. 7. 8. 9. 10.]
     [12.0, 14.0, 16.0, 18.0, 20.0]
[17]: long_words = ['platinum', 'hi', 'carbon', 'welcome', 'hydrogen', 'neptune', 'covid', __
      # checks and stores into new list based on length of a word
```

0 0

```
short_words = [word for word in long_words if len(word) < 6]
short_words</pre>
```

[17]: ['hi', 'covid']

1.3 C) Atomic Simutlation Environment

```
import os
import numpy as np
import ase
from ase.parallel import paropen
from ase.io import read, write
from ase.visualize import view

# Plotting and image visualizing in Jupyter Notebook
import matplotlib
import matplotlib.pyplot as plt
from IPython.display import Image

%matplotlib inline
```

2 Module 1: Atom / Atoms

```
[19]: # import the module (use tab tab for auto-completion)
      from ase import Atom, Atoms
[20]: # use ? to know more
      #atom = Atom?
[21]: atom = Atom('0')
      atom.mass = 16
      atom.charge = -2
      atom.position = (0,2,3)
      #atom.symbol
      #atom.x
      #atom.y
[22]: # use ? to know more
      #Atoms?
[23]: d = 1.104 # N2 bondlength
      # The following three are equivalent
      N2 = Atoms('N2', [(0, 0, 0), (0, 0, d)])
```

```
N2 = Atoms(numbers=[7, 7], positions=[(0, 0, 0), (0, 0, d)])
N2 = Atoms([Atom('N', (0, 0, 0)), Atom('N', (0, 0, d))])

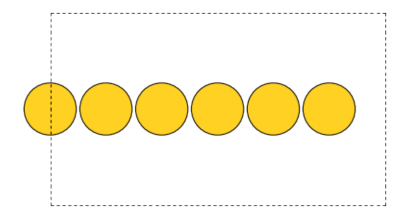
# to save and view a snapshot
write('extras/n2.png', N2, rotation='-80y')
Image(filename='extras/n2.png')

# to have a 3D view in notebook
#view(N2, viewer='x3d')
```

[23]:



[24]:



3 Module 2: Build (molecule, bulk, surfaces)

```
[25]: from ase.build import molecule

m = molecule('CH3CH2OH')
print(m.get_chemical_symbols())
print(m.get_chemical_formula())
print(m.get_positions())

# to save and view a snapshot
write('extras/ethanol.png', m, rotation='-20x')
Image(filename='extras/ethanol.png',width=150)

#view(m, viewer='x3d')

#view(m)
```

[25]:



4 Optimize lattice constant

```
[26]: from ase.build import bulk
  from ase.calculators.emt import EMT
  from ase.io import Trajectory

a0 = 3.6
  cu = bulk('Cu', 'fcc', a=a0, cubic=True)

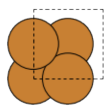
print(cu.get_cell())

# to save and view a snapshot
  write('extras/cu.png', cu)
```

```
Image(filename='extras/cu.png',width=150)
#view(cu, viewer='x3d')
#view(cu)
```

Cell([3.6, 3.6, 3.6])

[26]:



```
[27]: # create an empty list to store energy
energy_list = []

# start with a good guess
# and create a list with fluctuations in a.

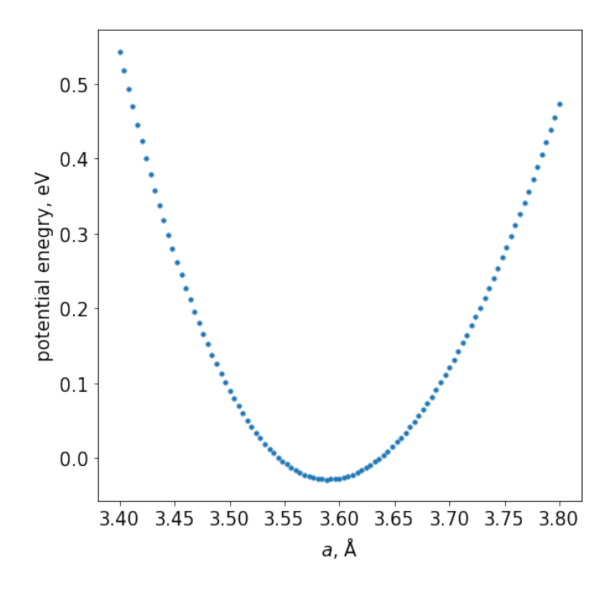
a0 = 3.6
a_list = a0 + np.linspace(-0.2, 0.2, 101)

# For loop to use each and every value of a
# and calculate & store the potential energy

for a in a_list:
    cu = bulk('Cu', 'fcc', a=a, cubic=True)
    cu.set_calculator(EMT())
    e = cu.get_potential_energy()
    energy_list.append(e)
```

```
[28]: # plotting using matplotlib (https://matplotlib.org/)

fig, ax = plt.subplots(figsize=(7,7))
plt.scatter(a_list, energy_list, marker='o', s=10)
plt.xlabel('$a$, $\mathrm{\AA}$', fontsize=15)
plt.xticks(fontsize=15)
plt.ylabel('potential enegry, eV', fontsize=15)
plt.yticks(fontsize=15)
fig.savefig('extras/lattice_constant.png')
plt.show()
```



```
[29]: # index corresponding to the lowest energy
index = energy_list.index(min(energy_list))
lc = a_list[index]
print('lattice constant:%0.3f'%lc)
```

lattice constant:3.588

5 Module 3: Surface, Constraints

```
[30]: from ase.build import fcc111 from ase.constraints import FixAtoms
```

[31]: #fcc111?

```
[32]: # Create a slab using build
slab = fcc111('Cu', size=(3,3,3), a=3.588, vacuum=10)
slab.set_pbc((1,1,0))

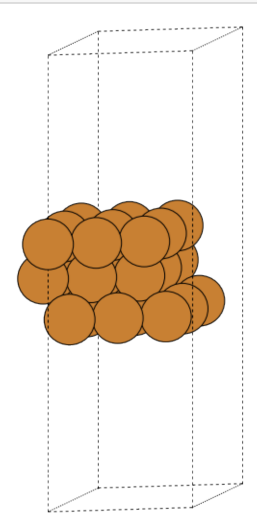
# Use of constraints, here, Fix the bottom most layer
indices=[atom.index for atom in slab if atom.tag == 3]
constraint = FixAtoms(indices)
slab.set_constraint(constraint)

# Note that the constraint is not visible through
# the following method

write('extras/cu_slab.png', slab, rotation='10z,-80x')
Image(filename='extras/cu_slab.png')

#view(slab, viewer='x3d')
#view(slab)
```

[32]:



6 Module 4: Add an adsorbate

method 1

```
[33]: from ase.build import add_adsorbate

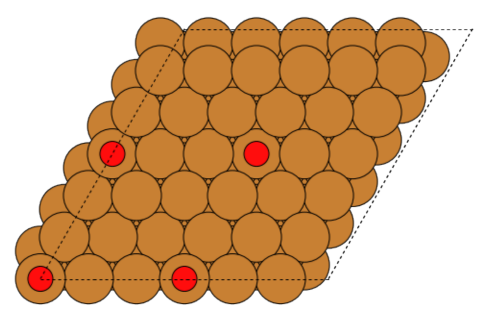
# create a slab
slab = fcc111('Cu', size=(3,3,3), a=3.588, vacuum=10)
slab.set_pbc((1,1,0))

# create an atom and use add_adsorbate to add it to the slab
adsorbate = Atom('0')
add_adsorbate(slab, adsorbate, 1.8, 'ontop')

write('extras/slab_with_adsorbate.png', slab * (2, 2, 1))
Image(filename='extras/slab_with_adsorbate.png')

#view(slab, viewer='x3d')
#view(slab)
```

[33]:



method 2

```
[34]: a = 3.558

# create a slab
slab = fcc111('Cu', size=(3,3,3), a=3.588, vacuum=10)
slab.set_pbc((1,1,0))
```

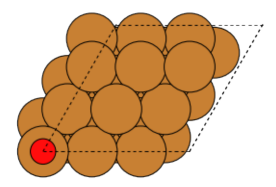
```
# use the slab atom position to add it to the slab
adsorbate = Atom('0')
adsorbate.position = slab[18].position + (0,0,2)

slab_with_add = slab + adsorbate

write('extras/slab_with_adsorbate.png', slab_with_add)
Image(filename='extras/slab_with_adsorbate.png')

#view(slab_with_add, viewer='x3d')
#view(slab_with_add)
```

[34]:



7 Module 5: Structure optimization using EMT

```
[35]: from ase.constraints import FixAtoms
      from ase.build import fcc111, add_adsorbate
      from ase.calculators.emt import EMT
      from ase.optimize import BFGS
      # Construct a slab with optimized lattice constant
      slab = fcc111('Cu', size=(3,3,3), a=3.588, vacuum=10)
      # Periodic boundary conditions
      slab.set_pbc((1,1,0))
      # Construct & add an adosorbate
      adsorbate = Atom('0')
      add_adsorbate(slab, adsorbate, 1.8, 'ontop')
      # Fix atoms
      indices=[atom.index for atom in slab if atom.tag == 3]
      constraint = FixAtoms(indices)
      slab.set_constraint(constraint)
      # Calculator
      calc = EMT()
      slab.set_calculator(calc)
```

```
if os.path.exists('qn.log'):
          os.remove('qn.log')
      # Optimization
      opt = BFGS(slab, logfile='qn.log', trajectory='qn.traj')
      opt.run(fmax=0.01)
[35]: True
[36]: #relaxed_slab = read('qn.traj')
      #view(relaxed_slab)
[37]: myfile = open("qn.log",'r')
      txt = myfile.read()
      print(txt)
           Step
                    Time
                                  Energy
                                                 fmax
                                                0.9096
     BFGS:
             0 09:20:44
                                6.520552
     BFGS:
              1 09:20:44
                                6.505566
                                                0.4225
     BFGS: 2 09:20:44
                                6.501307
                                                0.0559
     BFGS: 3 09:20:44
                                6.501107
                                                0.0437
     BFGS: 4 09:20:44
                                                0.0845
                                6.500731
     BFGS: 5 09:20:44
                                6.500402
                                                0.0943
     BFGS: 6 09:20:44
                                6.500170
                                                0.0693
     BFGS: 7 09:20:45
                                6.500026
                                                0.0385
     BFGS: 8 09:20:45
                                6.499856
                                                0.0668
     BFGS:
            9 09:20:45
                                6.499623
                                                0.0791
     BFGS: 10 09:20:45
                                6.499416
                                                0.0562
     BFGS: 11 09:20:45
                                6.499321
                                                0.0183
     BFGS: 12 09:20:45
                                6.499287
                                                0.0128
     BFGS: 13 09:20:45
                                6.499261
                                                0.0155
     BFGS: 14 09:20:45
                                6.499236
                                                0.0165
     BFGS: 15 09:20:46
                                6.499223
                                                0.0082
[38]: from ase.constraints import FixAtoms
      from ase.build import fcc111, add_adsorbate
      from ase.calculators.emt import EMT
      from ase.optimize import BFGS
      # Construct a slab with optimized lattice constant
      slab = fcc111('Cu', size=(3,3,3), a=3.588, vacuum=10)
      # periodic boundary condition
      slab.set_pbc((1,1,0))
      # Construct & add an adosorbate
      adsorbate = Atom('0')
      add_adsorbate(slab, adsorbate, 1.8, 'hcp')
      # Fix atoms constraint to fix the last atomic layer
      indices=[atom.index for atom in slab if atom.tag == 3]
```

```
constraint = FixAtoms(indices)
     slab.set_constraint(constraint)
     # Calculator
     calc = EMT()
     slab.set_calculator(calc)
     if os.path.exists('qn.log'):
         os.remove('qn.log')
      # Optimization
     opt = BFGS(slab, logfile='qn.log', trajectory='qn.traj')
     opt.run(fmax=0.01)
[38]: True
[39]: myfile = open("qn.log", 'r')
     txt = myfile.read()
     print(txt)
                                                fmax
          Step
                   Time
                                 Energy
     BFGS: 0 09:20:46
                               6.570523
                                               1.7805
     BFGS: 1 09:20:46
                               6.508532
                                               1.5692
     BFGS: 2 09:20:46
                               6.388970
                                               0.6331
     BFGS: 3 09:20:46
                               6.368169
                                               0.3576
     BFGS: 4 09:20:46
                                               0.2807
                               6.363440
     BFGS: 5 09:20:46
                               6.346981
                                               0.1420
     BFGS: 6 09:20:46
                               6.345193
                                               0.1026
     BFGS: 7 09:20:47
                               6.343991
                                               0.0898
     BFGS: 8 09:20:47
                               6.342511
                                               0.0919
     BFGS: 9 09:20:47
                               6.340466
                                               0.0910
     BFGS: 10 09:20:47
                               6.339097
                                               0.1304
     BFGS: 11 09:20:47
                               6.338357
                                               0.1316
     BFGS: 12 09:20:47
                               6.337813
                                               0.1097
     BFGS: 13 09:20:47
                               6.337098
                                               0.0630
     BFGS: 14 09:20:47
                                               0.0408
                               6.336492
     BFGS:
            15 09:20:47
                               6.336258
                                               0.0134
     BFGS: 16 09:20:48
                               6.336218
                                               0.0105
     BFGS: 17 09:20:48
                               6.336205
                                               0.0090
```

8 Module 6: Calculate Barriers using nudged elastic band (NEB)

```
[43]: #!/usr/bin/env python3

import ase
from ase.io import read, write
from ase import Atom
from ase.build import fcc111
from ase.constraints import FixAtoms
```

```
from ase.neb import NEB
from ase.optimize import BFGS
from ase.calculators.emt import EMT
# Create a bare slab
slab = fcc111('Cu', size=(3,3,3), a=3.588, vacuum=10)
slab.set_pbc((1,1,0))
indices=[atom.index for atom in slab if atom.tag == 3]
constraint = FixAtoms(indices)
slab.set_constraint(constraint)
slab.set_calculator(EMT())
opt = BFGS(slab, logfile='bare_slab.log', trajectory='bare_slab.traj')
opt.run(fmax=0.01)
# create an adsorbate
adsorbate = Atom('0')
###### SLAB 1: Cu with 0 in fcc (position 1) ######
initial = slab + adsorbate
initial[-1].position = initial[18].position + (1.3,0.8,1)
initial.set_calculator(EMT())
opt = BFGS(initial, logfile='initial.log', trajectory='initial.traj')
opt.run(fmax=0.01)
###### SLAB 1: Cu with 0 in fcc (position 2) ######
final = slab + adsorbate
final[-1].position = final[19].position + (1.3, 0.8, 1)
final.set_calculator(EMT())
opt = BFGS(final, logfile='final.log', trajectory='final.traj')
opt.run(fmax=0.01)
n = 10
images = [initial]
for i in range(n):
   image = initial.copy()
   image.set_calculator(EMT())
   image.get_potential_energy()
   image.get_forces()
   images.append(image)
images.append(final)
```

```
neb = NEB(images)
neb.interpolate()

qn = BFGS(neb,logfile='neb.log', trajectory='neb.traj')
qn.run(fmax=0.05)
```

[43]: True