Molecular Simulation HOMEWORK 3

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1 Reaction energy of CO oxidation

1.1 Compute the reaction energy for CO + 1/2 O₂ \rightarrow CO₂

Use a cutoff energy of 250 eV. The molecules should all be relaxed to their lowest energy geometry (perform a geometry optimization). Demonstrate that all the forces on the molecule are less than $0.05~{\rm eV/\mathring{A}}$.

```
from ase import Atoms, Atom
     from jasp import *
     import numpy as np
    np.set_printoptions(precision=3, suppress=True)
4
    CO = Atoms([Atom('C', (0, 0, 0)),
Atom('0', (1.2, 0, 0))],
cell=(10, 10, 10))
7
     02 = Atoms([Atom('0', (0, 0, 0)),
9
                  Atom('0', (1.23, 0, 0))], cell=(9.8, 9.9, 10))
10
11
     CO2 = Atoms([Atom('0', (0, 0, 0)),
12
                    Atom('C', (1.16, 0, 0)),
13
                   Atom('0', (2.32, 0, 0))],
14
                  cell=(10, 10, 10))
15
     with jasp('probla/CO',
16
                xc='PBE', lreal=False,
17
                encut=250, prec='Accurate',
18
                kpts=(1, 1, 1), ismear=1, sigma=0.05,
```

```
20
             ibrion=1, nsw=50, ediffg=-0.05, isif=2,
21
             atoms=CO) as COcalc:
22
23
            eC0 = C0.get_potential_energy()
           fC0 = C0.get_forces()
24
25
        except:
26
           pass
    with jasp('probla/02',
27
             xc='PBE', lreal=False,
             encut=250, prec='Accurate',
29
             kpts=(1, 1, 1), ismear=1, sigma=0.05,
30
             ibrion=1, nsw=50, ediffg=-0.05, isif=2,
31
             atoms=02) as 02calc:
32
33
34
           eO2 = O2.get_potential_energy()
           f02 = 02.get_forces()
35
36
        except:
           pass
37
38
    with jasp('prob1a/CO2',
             xc='PBE', lreal=False,
39
             encut=250, prec='Accurate',
40
             kpts=(1, 1, 1), ismear=1, sigma=0.05,
41
42
             ibrion=1, nsw=50, ediffg=-0.05, isif=2,
             atoms=CO2) as CO2calc:
43
44
           eC02 = C02.get_potential_energy()
45
           fCO2 = CO2.get_forces()
46
47
48
           pass
49
    re = eC02 - eC0 - 0.5*e02
50
    print 'The total energy of CO is {0:1.3f}'.format(eCO)
51
    print 'The forces (eV/angstrom) on the atoms in CO are'
    print 'C: {0}'.format(fCO[0])
53
    print '0: {0}\n'.format(fCO[1])
54
    print 'The total energy of 02 is {0:1.3f}'.format(e02)
    print 'The forces (eV/angstrom) on the atoms in O2 are'
56
   print '0: {0}'.format(f02[0])
    print '0: {0}\n'.format(f02[1])
   print 'The total energy of CO2 is {0:1.3f}'.format(eCO2)
59
   print 'The forces (eV/angstrom) on the atoms in CO2 are'
    print '0: {0}'.format(fCO2[0])
61
    print 'C: {0}'.format(fCO2[1])
62
   print '0: {0}\n'.format(fCO2[2])
    print 'The reaction energy is {0:1.3f}'.format(re)
    The total energy of CO is -15.168
    The forces (eV/angstrom) on the atoms in CO are
    C: [-0.033 0.
                            0.
                                   ]
                                   ٦
    0: [ 0.033 0.
                            0.
    The total energy of O2 is -8.719
    The forces (eV/angstrom) on the atoms in O2 are
    0: [ 0.019 0.
                            0.
                                   ]
    0: [-0.019 0.
                            0.
                                   1
    The total energy of CO2 is -23.508
    The forces (eV/angstrom) on the atoms in CO2 are
    0: [-0.015 0.
                            0.
                                   ]
    C: [ 0. 0. 0.]
```

```
0: [ 0.015 0. 0. ]
```

The reaction energy is -3.980

1.2 Convergence test

Repeat the previous problem at 350, 450, and 500 eV. Reoptimize the geometry at each ENCUT value. Compare (in a graph) the convergence of the total energy of each species with the convergence of the reaction energy. Which converges faster?

```
from ase import Atoms, Atom
1
2
    from jasp import *
3
    import numpy as np
    np.set_printoptions(precision=3, suppress=True)
4
    CO = Atoms([Atom('C', (0, 0, 0)),
6
                 Atom('0', (1.2, 0, 0))],
                cell=(10, 10, 10))
    02 = Atoms([Atom('0', (0, 0, 0)),
Atom('0', (1.23, 0, 0))],
9
10
                cell=(9.8, 9.9, 10))
11
    CO2 = Atoms([Atom('0', (0, 0, 0)),
12
                  Atom('C', (1.16, 0, 0))
13
                  Atom('0', (2.32, 0, 0))],
14
                 cell=(10, 10, 10))
15
16
    dirs = ('/e350', '/e450', '/e500')
    cuts = (350, 450, 500)
17
    eCOs = []
18
    e02s = []
19
    eCO2s = []
20
21
    for d, cut in zip(dirs, cuts):
22
         with jasp('prob1b' + d + '/CO'.
23
                   xc='PBE', lreal=False,
24
                   encut=cut, prec='Accurate',
25
                   kpts=(1, 1, 1), ismear=1, sigma=0.05,
26
                   ibrion=1, nsw=50, ediffg=-0.05, isif=2,
                   atoms=CO) as COcalc:
28
29
                 eC0 = C0.get_potential_energy()
30
                 {\tt eCOs.append(eCO)}
31
32
             except:
                 pass
33
         with jasp('prob1b' + d + '/02'),
34
35
                   xc='PBE', lreal=False,
                   encut=cut, prec='Accurate',
36
37
                   kpts=(1, 1, 1), ismear=1, sigma=0.05,
                   ibrion=1, nsw=50, ediffg=-0.05, isif=2,
38
                   atoms=02) as 02calc:
39
                 e02 = 02.get_potential_energy()
41
42
                 e02s.append(e02)
43
             except:
44
                 pass
         with jasp('prob1b' + d + '/C02'),
45
                   xc='PBE', lreal=False,
46
                   encut=cut, prec='Accurate',
47
                   kpts=(1, 1, 1), ismear=1, sigma=0.05,
                   ibrion=1, nsw=50, ediffg=-0.05, isif=2,
49
                    atoms=CO2) as CO2calc:
50
51
             try:
                 eCO2 = CO2.get_potential_energy()
52
```

```
eCO2s.append(eCO2)
53
            except:
54
55
                pass
56
    import matplotlib.pyplot as plt
57
58
    from matplotlib.ticker import ScalarFormatter
59
    import numpy as np
60
    eCOs = np.array(eCOs)
61
    e02s = np.array(e02s)
62
    eCO2s = np.array(eCO2s)
63
64
    fig = plt.figure(1)
65
    axC0 = fig.add_subplot(221)
66
    axCO.plot(cuts, eCOs, marker='o')
67
    axCO.set_title('$\mathdefault{CO}$')
68
69
    axCO.set_xlim((300, 550))
    axCO.set_ylim((-14.81, -14.75))
70
71
    axCO.set_xticklabels(())
    axCO.yaxis.set_major_formatter(ScalarFormatter(useOffset=False))
72
    axCO.set_ylabel('Total Energy (eV/atom)')
73
74
75
    ax02 = fig.add_subplot(222)
    ax02.plot(cuts, e02s, marker='o')
76
    ax02.set_title('$\mathdefault{0_{2}}$')
    ax02.set_xlim((300, 550))
78
    ax02.set_ylim((-8.76, -8.70))
79
    ax02.set_xticklabels([])
    ax02.yaxis.set_major_formatter(ScalarFormatter(useOffset=False))
81
82
    axCO2 = fig.add_subplot(223)
83
    axC02.plot(cuts, eC02s, marker='o')
84
    axC02.set_title('$\mathdefault{C0_{2}}$')
85
    axCO2.set_xlim((300, 550))
86
    axCO2.set_xlabel('Kinetic Energy Cutoff (eV)')
87
     axCO2.set_ylabel('Total Energy (eV/atom)')
    axC02.xaxis.set_major_formatter(ScalarFormatter(useOffset=False))
89
90
    axCO2.yaxis.set_major_formatter(ScalarFormatter(useOffset=False))
91
    axrxn = fig.add_subplot(224)
92
    axrxn.plot(cuts, eCO2s - eCOs - 0.5*eO2s, marker='o')
93
    94
    axrxn.set xlim((300, 550))
95
    axrxn.set_ylim((-3.84, -3.78))
    axrxn.set_xlabel('Kinetic Energy Cutoff (eV)')
97
    axrxn.xaxis.set_major_formatter(ScalarFormatter(useOffset=False))
98
    axrxn.yaxis.set_major_formatter(ScalarFormatter(useOffset=False))
    fig.tight_layout()
100
101
    plt.savefig('1b.png')
    plt.show()
102
```

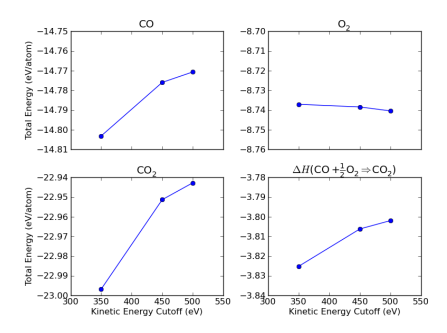


Figure 1: Convergence of CO, O_2 , CO_2 , and the reaction enthalpy of $CO + 1/2O_2 \rightarrow CO_2$ with respect to plane wave cutoff energy

The total energy of the oxygen molecule converges the fastest. Note, all y-axis tick spacings are the same.

2 Zero-point energy corrections

2.1 Compute vibrational modes for CO, CO_2 and O_2

Compute the vibrational modes of each molecule in the CO oxidation reaction. Do this at 350 eV cutoff energy only. Prepare a table of the vibrational modes for molecule.

```
import os
    import sys
    from ase.calculators.vasp import Vasp
4
    import ase.units
    from jasp import *
    # Since we wanted relaxed molecules for these calculations, we can take
    # these geometries from the previous problem.
8
    with jasp('prob1b/e350/CO') as calc:
10
        CO = calc.get_atoms()
11
    with jasp('prob1b/e350/02') as calc:
12
        02 = calc.get_atoms()
13
    with jasp('prob1b/e350/C02') as calc:
14
         CO2 = calc.get_atoms()
15
16
    # Now we're ready to perform the vibrational calculations
17
    with jasp('prob2a/CO',
18
               xc='PBE', lreal=False,
19
               encut=350, prec='Accurate', ediff=1e-8,
20
               kpts=(1, 1, 1), ismear=0, sigma=0.05,
21
```

```
22
               ibrion=6, nsw=1, potim=0.015, nfree=2,
23
               atoms=CO) as calcCO:
24
        try:
25
             CO.get_potential_energy()
            energies, modes = calcCO.get_vibrational_modes()
26
27
             print 'Energies of CO\n======'
28
            for i, e in enumerate(energies):
                print '{0:02d}: {1} eV'.format(i, e)
29
         except:
            pass
31
    with jasp('prob2a/02',
32
              xc='PBE', lreal=False,
               encut=350, prec='Accurate', ediff=1e-8,
34
35
               kpts=(1, 1, 1), ismear=0, sigma=0.05,
               ibrion=6, nsw=1, potim=0.015, nfree=2,
36
               atoms=02) as calc02:
37
38
        try:
            02.get_potential_energy()
39
40
             energies, modes = calc02.get_vibrational_modes()
            print '\nEnergies of O2\n======
41
            for i, e in enumerate(energies):
42
                print '{0:02d}: {1} eV'.format(i, e)
43
44
        except:
            pass
45
    with jasp('prob2a/CO2',
              xc='PBE', lreal=False,
47
               encut=350, prec='Accurate', ediff=1e-8,
48
               kpts=(1, 1, 1), ismear=0, sigma=0.05,
               ibrion=6, nsw=1, potim=0.015, nfree=2,
50
               atoms=CO2) as calcCO2:
51
52
            CO2.get_potential_energy()
53
54
             energies, modes = calcCO2.get_vibrational_modes()
55
            print '\nEnergies of CO2\n=====
            for i, e \underline{in} enumerate(energies):
56
57
                print '{0:02d}: {1} eV'.format(i, e)
        except:
58
            pass
```

Energies of CO

```
======
```

00: 0.261840727 eV 01: 0.003767323 eV 02: 0.003767323 eV 03: (3.0739e-05+0j) eV 04: (0.000943898+0j) eV 05: (0.000943898+0j) eV

Energies of 02

======

00: 0.189490603 eV 01: 0.004093929 eV 02: 1e-09 eV 03: 0.0 eV 04: (1e-09+0j) eV 05: (0.006638148+0j) eV

Energies of CO2

```
00: 0.291924562 eV
01: 0.16318552 eV
02: 0.078492458 eV
03: 0.078492458 eV
04: 0.004836504 eV
05: 0.004836504 eV
06: (4.1677e-05+0j) eV
07: (5.9833e-05+0j) eV
08: (5.9833e-05+0j) eV
```

=====

2.2 Compute the CO oxidation reaction energy with zero-point energy corrections.

Compare the reaction energy with and without the zero-point energy correction.

```
from jasp import *
2
    import numpy as np
    c = 3e10 # speed of light cm/s
3
4
    h = 4.135667516e-15 \# eV/s
    # Get the vibrational energies from problem 2a. Get the total energies from
6
7
    # problem 1b at 350 eV.
8
    with jasp('prob2a/CO') as calc:
9
10
        COfreq = calc.get_vibrational_frequencies()
    with jasp('prob1b/e350/CO') as calc:
11
12
        atoms = calc.get_atoms()
        COe = atoms.get_potential_energy()
13
    for f in COfreq:
14
        if not isinstance(f, float):
15
           continue
16
17
        nu = f*c
18
        COe += 0.5*h*nu
    with jasp('prob2a/02') as calc:
19
20
        02freq = calc.get_vibrational_frequencies()
21
    with jasp('prob1b/e350/02') as calc:
        atoms = calc.get_atoms()
22
23
        02e = atoms.get_potential_energy()
24
    for f in O2freq:
        if not isinstance(f, float):
25
26
            continue
27
        nu = f*c
        02e += 0.5*h*nu
28
    with jasp('prob2a/CO2') as calc:
        CO2freq = calc.get_vibrational_frequencies()
30
    with jasp('prob1b/e350/CO2') as calc:
31
32
        atoms = calc.get_atoms()
        CO2e = atoms.get_potential_energy()
33
34
    for f in CO2freq:
        if not isinstance(f, float):
35
36
            continue
        nu = f*c
37
        CO2e += 0.5*h*nu
38
39
    \mathbf{s} = 'The reaction energy for CO oxidation with zero point contributions is {0:1.3f}'
    print s.format(CO2e - COe - 0.5*O2e)
```

The reaction energy for CO oxidation with zero point contributions is -3.697

2.3 Compare your computed energy to a value from the literature.

Provide a reference for your literature value.

All values are taken from the NIST-JANAF Thermochemical Tables at kinetics.nist.gov/janaf.

```
# Values of heats of formation at 0 K in kJ/mol

Hf_CO = -113.805

Hf_CO2 = -393.151

Hf_O2 = 0 # Pure component is reference

Hf_rxn = -393.151 + 113.805

s = 'The experimental heat of reaction is {0:1.3f} eV/atom'
print s.format(Hf_rxn * 0.010364)
```

The experimental heat of reaction is -2.895 eV/atom

Our computed heat of reaction is too exothermic. This means that either the products (CO_2) are too stable, or the reactants $(O_2$ and CO) are unstable.

3 Plot the electron density of the CO2 molecule.

Include the figure in your homework.

```
from jasp import *
1
    from enthought.mayavi import mlab
    from ase.data import vdw_radii
    from ase.data.colors import cpk_colors
    from ase import Atom, Atoms
    # Lets first get the relaxed CO at 500 eV plane wave cutoff, center it,
    # and recalculate the electron density in the centered cell
8
9
    with jasp('prob1b/e500/CO') as calc:
10
        CO = calc.get_atoms()
11
12
        CO.center()
    with jasp('prob3a/CO-centered',
13
              xc='PBE', lreal=False,
14
15
               encut=500, prec='Accurate',
               kpts=(1, 1, 1), ismear=1, sigma=0.05,
16
               atoms=CO) as calc:
17
        CO.get_potential_energy()
18
        x, y, z, cd = calc.get_charge_density()
19
20
21
    mlab.figure(bgcolor=(1, 1, 1))
    \# plot the atoms as spheres
22
    for atom in CO:
23
        mlab.points3d(atom.x,
24
                       atom.y,
25
                       scale_factor=vdw_radii[atom.number]/5.,
27
28
                       resolution=20,
                       # a tuple is required for the color
                       {\tt color=tuple(cpk\_colors[atom.number]),}
30
31
                       scale_mode='none')
32
    # draw the unit cell - there are 8 corners, and 12 connections
33
    a1, a2, a3 = C0.get_cell()
34
    origin = [0, 0, 0]
35
    cell_matrix = [[origin, a1],
```

```
a2],
37
                       [origin,
                       [origin,
38
                                   a3],
                       [a1,
                                   a1 + a2],
39
                                   a1 + a3],
a2 + a1],
40
                       [a1,
                       [a2,
41
                                   a2 + a3],
^{42}
                       [a2,
                                   a1 + a3],
a2 + a3],
43
                       [a3,
                       [a3,
44
                       [a1 + a2, a1 + a2 + a3],
                       [a2 + a3, a1 + a2 + a3],
[a1 + a3, a1 + a3 + a2]]
46
47
48
     for p1, p2 in cell_matrix:
49
          mlab.plot3d([p1[0], p2[0]], # x-positions
50
                        [p1[1], p2[1]], # y-positions
[p1[2], p2[2]], # z-positions
51
52
53
                        tube_radius=0.02)
54
55
     {\it \# Now plot the charge density}
56
     mlab.contour3d(x, y, z, cd, transparent=True)
57
     # this view was empirically found by iteration
58
59
     mlab.view(azimuth=-90, elevation=90, distance='auto')
60
     mlab.savefig('co-density.png')
```

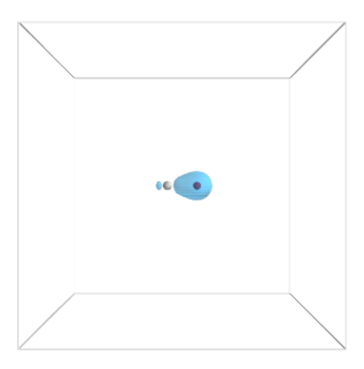


Figure 2: Charge density of CO