## pm-project-1

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### 1 Introduction

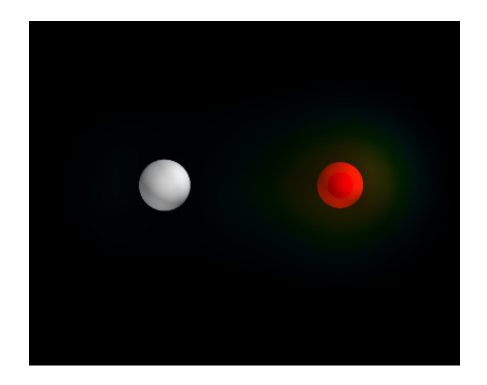
This project aims to plot electron density as a "fog" rather than with isosurfaces using jasp. The code has been adapted from the examples in dft-book and the example given on http://docs.enthought.com/mayavi/mayavi/auto/example\_chemistry.html. The mayavi documentation on volume rendering (http://docs.enthought.com/mayavi/mayavi/mlab.htm volumetric-scalar-data) and the jasp documentation for plotting charge density (https://github.com/jkitchin/jasp/blob/master/CHG.py) have been used as reference. Examples of python code have been written for the CO and the  $\rm H_2O$  molecule.

# 2 CO Example

```
from jasp import *
from enthought.mayavi import mlab
from ase.data import vdw_radii
from ase.data.colors import cpk_colors
from ase import Atom, Atoms

# creating atoms object
# creating atoms object
```

```
atoms = Atoms([Atom('C', [2.422, 0.0, 0.0]),
9
                    Atom('0', [3.578, 0.0, 0.0])],
10
                    cell=(10,10,10))
11
    atoms.center()
12
13
    # running dft calculation
14
15
    with jasp('molecules/co-fog',
              encut=350,
16
              xc='PBE',
17
              atoms=atoms) as calc:
18
         calc.calculate()
19
20
         atoms = calc.get_atoms()
        x, y, z, cd = calc.get_charge_density()
21
22
    # making a black background
23
    mlab.figure(bgcolor=(0, 0, 0))
24
25
    # plotting the atoms as spheres,
26
27
    for atom in atoms:
28
29
        mlab.points3d(atom.x,
30
                       atom.y,
31
                       atom.z,
32
                       scale_factor=vdw_radii[atom.number]/5.,
33
                       resolution=20,
                       # a tuple is required for the color
                       color=tuple(cpk_colors[atom.number]),
35
36
                       scale_mode='none')
37
    # plotting the charge density
38
39
    source = mlab.pipeline.scalar_field(x,y,z,cd)
    min=cd.min()
40
    max=cd.max()
41
    \# vmin and vmax were adjusted by iteration
42
    vol = mlab.pipeline.volume(source, vmin=min+0.1*(max-min),
43
                                         vmax=max)
45
    # view adjusted by iteration
46
    mlab.view(azimuth=-90, elevation=90, distance='auto')
47
48
49
    mlab.savefig('molecules/co-fog.png')
    mlab.show()
50
```



## 3 H<sub>2</sub>O Example

```
from jasp import *
   from enthought.mayavi import mlab
3 from ase.data import vdw_radii
    from ase.data.colors import cpk_colors
    from ase import Atom, Atoms
5
     \# a similar approach to the one shown for the CO molecule has been taken
    atoms = Atoms([Atom('H', [0.5960812, -0.7677068, Atom('O', [0.000000, 0.0000000, Atom('H', [0.5960812, 0.7677068,
                                                               0.0000000]),
9
10
                                                                0.0000000]),
11
                                                              0.0000000])],
                      cell=(8, 8, 8))
12
13
     atoms.center()
14
15
    with jasp('molecules/water-fog',
16
17
                encut=350,
                xc='PBE',
18
19
                atoms=atoms) as calc:
20
         calc.calculate()
         atoms = calc.get_atoms()
21
22
         x, y, z, cd = calc.get_charge_density()
23
24
```

```
25
   n1, n2, n3=cd.shape
26
    mlab.figure(bgcolor=(0, 0, 0))
27
28
    for atom in atoms:
29
30
        {\tt mlab.points3d(atom.x,}
31
                       atom.y,
32
                       atom.z,
33
                       scale_factor=vdw_radii[atom.number]/5.,
34
                       resolution=20,
                       color=tuple(cpk_colors[atom.number]),
35
                       scale_mode='none')
36
37
38
    source = mlab.pipeline.scalar_field(x,y,z,cd)
39
    min = cd.min()
max = cd.max()
40
41
   vol = mlab.pipeline.volume(source, vmin=min+.1*(max-min),
42
43
44
    mlab.savefig('molecules/water-fog.png')
45
    mlab.show()
46
```

