

## **10.009 The Digital World**

Term 3. 2016

Problem Set 7 (for Chemistry Project)

Last update: March 10, 2016

Due dates:

- **Problems:** Following week: Monday, Week 11, 11:59pm.

### **Objectives:**

1. Learn to use Numpy for numerical computations

**Note:** Solve the programming problems listed below using the IDLE or Canopy editor. Make sure you save your programs in files with suitably chosen names and in an newly created directory. In each problem find out a way to test the correctness of your program. After writing each program, test it, debug it if the program is incorrect, correct it, and repeat this process until you have a fully working program. Show your working program to one of the cohort instructors.

## Problems

1. Create a function to calculate the energy level of a given principal quantum number. This function should take 1 integer argument and return the energy level in eV. Round to 5 decimal places. (Pre-requisite: Part 2a) Hint: You can use `import scipy.constants as c` to get the necessary constants.

To test:

```
print 'n = 1'
ans= energy_n(1)
print ans

print 'n = 2'
ans= energy_n(2)
print ans

print 'n = 3'
ans= energy_n(3)
print ans
```

The output should be:

```
n = 1
-13.60569
n = 2
-3.40142
n = 3
-1.51174
```

2. Create two functions to convert degrees to radian and radian to degrees respectively. These functions should take 1 float argument and return the respective conversions each. Round to 5 decimal places.

To Test:

```
print 'degToRad(90)'
ans=degToRad(90)
print ans

print 'degToRad(180)'
ans=degToRad(180)
print ans

print 'degToRad(270)'
ans=degToRad(270)
print ans

print 'radToDeg(3.14)'
ans=radToDeg(3.14)
print ans

print 'radToDeg(3.14/2.0)'
ans=radToDeg(3.14/2.0)
print ans

print 'radToDeg(3.14*3/4)'
```

```
ans=radToDeg(3.14*3/4)
print ans
```

The output should be:

```
degToRad(90)
1.5708
degToRad(180)
3.14159
degToRad(270)
4.71239
radToDeg(3.14)
179.90875
radToDeg(3.14/2.0)
89.95437
radToDeg(3.14*3/4)
134.93156
```

3. Create two functions to convert spherical to cartesian coordinates and cartesian to spherical coordinates. These functions should take 3 float arguments and return the 3 respective conversions. Round to 5 decimal places. (Pre-requisite: Part 2d)

Hint: you can use Numpy trigonometric function by doing `import numpy as np`.

To test:

```
print 'sphericalToCartesian(3,0,np.pi)'
ans=sphericalToCartesian(3,0,np.pi)
print ans

print 'sphericalToCartesian(3,np.pi/2.0,np.pi/2.0)'
ans=sphericalToCartesian(3,np.pi/2.0,np.pi/2.0)
print ans

print 'sphericalToCartesian(3,np.pi, 0)'
ans=sphericalToCartesian(3,np.pi,0)
print ans

print 'cartesianToSpherical(3,0,0)'
ans=cartesianToSpherical(3,0,0)
print ans

print 'cartesianToSpherical(0,3,0)'
ans=cartesianToSpherical(0,3,0)
print ans

print 'cartesianToSpherical(0,0,3)'
ans=cartesianToSpherical(0,0,3)
print ans

print 'cartesianToSpherical(0,-3,0)'
ans=cartesianToSpherical(0,-3,0)
print ans
```

The output should be:

```
sphericalToCartesian(3,0,np.pi)
(-0.0, 0.0, 3.0)
sphericalToCartesian(3,np.pi/2.0,np.pi/2.0)
```

```

(0.0, 3.0, 0.0)
sphericalToCartesian(3,np.pi, 0)
(0.0, 0.0, -3.0)
cartesianToSpherical(3,0,0)
(3.0, 1.5708, 0.0)
cartesianToSpherical(0,3,0)
(3.0, 1.5708, 1.5708)
cartesianToSpherical(0,0,3)
(3.0, 0.0, 0.0)
cartesianToSpherical(0,-3,0)
(3.0, 1.5708, -1.5708)

```

4. Create a function using the while loop that calculates the factorial of the input integer. For example,  $4! = 4 \times 3 \times 2 \times 1 = 24$ . This function should take 1 integer argument and return the calculated factorial. The return value should be an integer.

To test:

```

print 'fact(3) '
ans=fact(3)
print ans

```

```

print 'fact(5) '
ans=fact(5)
print ans

```

```

print 'fact(4) '
ans=fact(4)
print ans

```

```

print 'fact(1) '
ans=fact(1)
print ans

```

The output should be:

```

fact(3)
6
fact(5)
120
fact(4)
24
fact(1)
1

```

5. Create a function that calculates the associated Legendre function. This function should take in 2 integer arguments and return the function that computes the associated Legendre with a variable  $\cos(\theta)$ . You should be able to obtain the function for any combination of  $m=0,1,2,3$  and  $n=0,1,2,3$  (Pre-requisite: Part 2g)

Hint: You may want to create a function that acts as a reference table to a set of recalculated associated Legendre function. For example

```

def p00(theta):
    return 1

def assocLegendre(m,l):
    if m==0 and l==0:
        return p00

To test:

print 'f=assocLegendre(0,0)'
print 'f(1)'
f=assocLegendre(0,0)
ans=f(1)
print ans

print 'f=assocLegendre(1,1)'
print 'f(1)'
f=assocLegendre(1,1)
ans=f(1)
print ans

print 'f=assocLegendre(2,3)'
print 'f(1)'
f=assocLegendre(2,3)
ans=f(1)
print ans

print 'f=assocLegendre(2,3)'
print 'f(0)'
f=assocLegendre(2,3)
ans=f(0)
print ans

```

You can refer to the Tables found in : [this PDF](#).

The output should be:

```

f=assocLegendre(0,0)
f(1)
1
f=assocLegendre(1,1)
f(1)
0.841470984808
f=assocLegendre(2,3)
f(1)
5.73860550926
f=assocLegendre(2,3)
f(0)
0.0

```

6. Create a function that calculates the associated Laguerre function. This function should take in 2 integer arguments and return the function that computes the associated Laguerre with a variable x. You should be able to obtain the function for any combination of  $p = 0, 1, 2, 3$  and  $q - p = 0, 1, 2, 3$  (Pre-requisite: Part 2h)

Hint: You may want to create a function that acts as a reference table to a set of recalculated associated Laguerre function. For example

```
def l00(x):
    return 1

def assocLaguerre(p,qmp):
    if p==0 and qmp==0:
        return l00
```

Note that  $qmp$  is the argument for  $q - p$ . You can refer to the Tables found in : [this PDF](#).

To test:

```
print 'f=assocLaguerre(0,0)'
print 'f(1)'
f=assocLaguerre(0,0)
ans=f(1)
print ans

print 'f=assocLaguerre(1,1)'
print 'f(1)'
f=assocLaguerre(1,1)
ans=f(1)
print ans

print 'f=assocLaguerre(2,2)'
print 'f(1)'
f=assocLaguerre(2,2)
ans=f(1)
print ans

print 'f=assocLaguerre(2,2)'
print 'f(0)'
f=assocLaguerre(2,2)
ans=f(0)
print ans
```

The output should be:

```
f=assocLaguerre(0,0)
f(1)
1
f=assocLaguerre(1,1)
f(1)
2
f=assocLaguerre(2,2)
f(1)
60
f=assocLaguerre(2,2)
f(0)
144
```

7. Create a function that calculates the normalized angular solution. This function should take 4 float arguments and return the value of the normalized angular solution for the specific  $m$ ,  $l$ ,  $\theta$  and  $\phi$ . (Pre-requisite: Part 2g). The return value is a complex number rounded to 5 decimal places for both the real and the imaginary parts. Hint: You may want to use `np.round()` function to round the return value to 5 decimal places. You can use the previous function `assocLegendre(m,1)`, in the following way:

```
pfunc=assocLegendre(m,l)
y=pfunc(theta)
```

where  $m$  and  $l$  are the respective quantum numbers and  $\theta$  is the angle  $\theta$ . This means that `assocLegendre(m,l)` must return a function with one argument  $\theta$ . See the test cases on the `assocLegendre(m,l)` question.

To test:

```
print 'angular_wave_func(0,0,0,0)'
ans=angular_wave_func(0,0,0,0)
print ans

print 'angular_wave_func(0,1,c.pi,0)'
ans=angular_wave_func(0,1,c.pi,0)
print ans

print 'angular_wave_func(1,1,c.pi/2,c.pi)'
ans=angular_wave_func(1,1,c.pi/2,c.pi)
print ans

print 'angular_wave_func(0,2,c.pi,0)'
ans=angular_wave_func(0,2,c.pi,0)
print ans
```

The output should be:

```
angular_wave_func(0,0,0,0)
(0.28209+0j)
angular_wave_func(0,1,c.pi,0)
(-0.4886+0j)
angular_wave_func(1,1,c.pi/2,c.pi)
(0.34549+0j)
angular_wave_func(0,2,c.pi,0)
(0.63078+0j)
```

8. Create a function that calculates the normalized radial solution. This function should take 3 float arguments and return the value of the normalized radial solution. The return value should be normalized to  $a^{-3/2}$ , where  $a$  is the Bohr's radius, and rounded to 5 decimal places. Hint: You may want to use `np.round()` function to round the return value to 5 decimal places. You can use the previous function `assocLaguerre(p, qmp)`, in the following way:

```
lfunc=assocLaguerre(p, qmp)
y=lfunc(x)
```

where the argument  $p$  and  $qmp$  refers to  $p$  and  $q - p$  in the associated Laguerre. This means that `assocLaguerre(n,l,r)` must return a function with one argument. See the test cases on the `assocLaguerre(p, qmp)` question.

To test:

```
a=c.physical_constants['Bohr radius'][0]
print 'radial_wave_func(1,0,a)'
```

```

ans=radial_wave_func(1,0,a)
print ans

print 'radial_wave_func(1,0,a)'
ans=radial_wave_func(1,0,a)
print ans

print 'radial_wave_func(2,1,a)'
ans=radial_wave_func(2,1,a)
print ans

print 'radial_wave_func(2,1,2*a)'
ans=radial_wave_func(2,1,2*a)
print ans

```

The output should be:

```

radial_wave_func(1,0,a)
0.73576
radial_wave_func(1,0,a)
0.73576
radial_wave_func(2,1,a)
0.12381
radial_wave_func(2,1,2*a)
0.15019
radial_wave_func(3,1,2*a)
0.08281

```

9. Create a function that calculates the square of the magnitude of the wave function. The function takes in several arguments:

- $n$ : quantum number  $n$
- $l$ : quantum number  $l$
- $m$ : quantum number  $m$
- $roa$ : maximum distance to plot from the centre, normalized to Bohr radius, i.e.  $r/a$ .
- $N_x$ : Number of points in the positive  $x$  axis.
- $N_y$ : Number of points in the positive  $y$  axis.
- $N_z$ : Number of points in the positive  $z$  axis.

The function should returns:

- $xx$ :  $x$  location of all the points in a 3D Numpy array.
- $yy$ :  $y$  location of all the points in a 3D Numpy array.
- $zz$ :  $z$  location of all the points in a 3D Numpy array.
- $density$ : The square of the magnitude of the wave function, i.e.  $|\Psi|^2$

Hint: You may find the following functions to be useful:



- `fvec=numpy.vectorize(f)`: This function takes in a function and return its vectorized version of the function.
- `xx,yy,zz=numpy.meshgrid(x,y,z)`: This function takes in 1D arrays and returns its 3D arrays to conform to a 3D grid.
- `m=numpy.absolute(c)`: This function takes in a complex number and returns its absolute value or its magnitude.

Hint: You also need to use all the previous functions you have done. Note that some of those functions may round the output to 5 decimal places and the final magnitude output from this function should also be rounded to 5 decimal places.

To test:

```
print 'Test 1'
x,y,z,mag=hydrogen_wave_func(2,1,1,8,2,2,2)
print 'x, y, z:'
print x, y, z
print 'mag:'
print mag

print 'Test 2'
x,y,z,mag=hydrogen_wave_func(2,1,1,5,3,4,2)
print 'x, y, z:'
print x, y, z
print 'mag:'
print mag

print 'Test 3'
x,y,z,mag=hydrogen_wave_func(2,0,0,3,5,4,3)
print 'x, y, z:'
print x, y, z
print 'mag:'
print mag
```

The output should be:

```
Test 1
x, y, z:
[[[-8. -8.]
  [ 8.  8.]]

 [[-8. -8.]
  [ 8.  8.]]] [[[-8. -8.]
 [-8. -8.]]

 [[ 8.  8.]
  [ 8.  8.]]] [[[-8.  8.]
 [-8.  8.]]

 [[-8.  8.]
 [-8.  8.]]]
mag:
[[[ 0.  0.]
  [ 0.  0.]
```

```

[[ 0.  0.]
 [ 0.  0.]]]
Test 2
x, y, z:
[[[-5. -5.]
 [ 0.  0.]
 [ 5.  5.]]

[[[-5. -5.]
 [ 0.  0.]
 [ 5.  5.]]

[[[-5. -5.]
 [ 0.  0.]
 [ 5.  5.]]

[[[-5. -5.]
 [ 0.  0.]
 [ 5.  5.]]] [[[-5.      -5.      ]
 [-5.      -5.      ]
 [-5.      -5.      ]]]

[[-1.66667 -1.66667]
 [-1.66667 -1.66667]
 [-1.66667 -1.66667]]

[[ 1.66667  1.66667]
 [ 1.66667  1.66667]
 [ 1.66667  1.66667]]

[[ 5.      5.      ]
 [ 5.      5.      ]
 [ 5.      5.      ]]] [[[-5.  5.]
 [-5.  5.]
 [-5.  5.]]

[[[-5.  5.]
 [-5.  5.]
 [-5.  5.]]

[[[-5.  5.]
 [-5.  5.]
 [-5.  5.]]

[[[-5.  5.]
 [-5.  5.]
 [-5.  5.]]]

mag:
[[[ 4.00000000e-05  4.00000000e-05]
 [ 1.10000000e-04  1.10000000e-04]
 [ 4.00000000e-05  4.00000000e-05]]

[[ 1.00000000e-04  1.00000000e-04]
 [ 7.00000000e-05  7.00000000e-05]
 [ 1.00000000e-04  1.00000000e-04]]

[[ 1.00000000e-04  1.00000000e-04]
 [ 7.00000000e-05  7.00000000e-05]
 [ 1.00000000e-04  1.00000000e-04]]

[[ 4.00000000e-05  4.00000000e-05]

```

```

[ 1.10000000e-04 1.10000000e-04]
[ 4.00000000e-05 4.00000000e-05]]]
Test 3
x, y, z:
[[-3. -3. -3.]
 [-1.5 -1.5 -1.5]
 [ 0.  0.  0.]
 [ 1.5  1.5  1.5]
 [ 3.  3.  3.]]

[[-3. -3. -3.]
 [-1.5 -1.5 -1.5]
 [ 0.  0.  0.]
 [ 1.5  1.5  1.5]
 [ 3.  3.  3.]]

[[-3. -3. -3.]
 [-1.5 -1.5 -1.5]
 [ 0.  0.  0.]
 [ 1.5  1.5  1.5]
 [ 3.  3.  3.]]

[[-3. -3. -3.]
 [-1.5 -1.5 -1.5]
 [ 0.  0.  0.]
 [ 1.5  1.5  1.5]
 [ 3.  3.  3.]]] [[[-3. -3. -3.]
 [-3. -3. -3.]
 [-3. -3. -3.]
 [-3. -3. -3.]
 [-3. -3. -3.]]

[[-1. -1. -1.]
 [-1. -1. -1.]
 [-1. -1. -1.]
 [-1. -1. -1.]
 [-1. -1. -1.]]

[[ 1.  1.  1.]
 [ 1.  1.  1.]
 [ 1.  1.  1.]
 [ 1.  1.  1.]
 [ 1.  1.  1.]]

[[ 3.  3.  3.]
 [ 3.  3.  3.]
 [ 3.  3.  3.]
 [ 3.  3.  3.]
 [ 3.  3.  3.]]] [[[-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]]

[[-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]]

[[-3.  0.  3.]

```

```

[-3.  0.  3.]
[-3.  0.  3.]
[-3.  0.  3.]
[-3.  0.  3.]]

[[-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]
 [-3.  0.  3.]]]
mag:
[[[ 5.60000000e-04  7.20000000e-04  5.60000000e-04]
 [ 6.90000000e-04  6.40000000e-04  6.90000000e-04]
 [ 7.20000000e-04  5.00000000e-04  7.20000000e-04]
 [ 6.90000000e-04  6.40000000e-04  6.90000000e-04]
 [ 5.60000000e-04  7.20000000e-04  5.60000000e-04]]

 [[ 7.10000000e-04  5.70000000e-04  7.10000000e-04]
 [ 6.80000000e-04  6.00000000e-05  6.80000000e-04]
 [ 5.70000000e-04  3.66000000e-03  5.70000000e-04]
 [ 6.80000000e-04  6.00000000e-05  6.80000000e-04]
 [ 7.10000000e-04  5.70000000e-04  7.10000000e-04]]

 [[ 7.10000000e-04  5.70000000e-04  7.10000000e-04]
 [ 6.80000000e-04  6.00000000e-05  6.80000000e-04]
 [ 5.70000000e-04  3.66000000e-03  5.70000000e-04]
 [ 6.80000000e-04  6.00000000e-05  6.80000000e-04]
 [ 7.10000000e-04  5.70000000e-04  7.10000000e-04]]

 [[ 5.60000000e-04  7.20000000e-04  5.60000000e-04]
 [ 6.90000000e-04  6.40000000e-04  6.90000000e-04]
 [ 7.20000000e-04  5.00000000e-04  7.20000000e-04]
 [ 6.90000000e-04  6.40000000e-04  6.90000000e-04]
 [ 5.60000000e-04  7.20000000e-04  5.60000000e-04]]]

```

10. **Plots:** Submit your plot for your assigned quantum numbers to your Chemistry instructors to get a point for this item.
  11. In the final function to calculate the hydrogen wave function, you are to use the other previous functions you have calculated. However, some of those functions rounds the result to 5 decimal places. The error on the final wave function magnitude is called \_\_\_\_ due to \_\_\_\_.
- (a) floating point error, rounding error.
  - (b) propagation error, rounding error.
  - (c) propagation error, floating point error.
  - (d) rounding error, propagation error.

**Submit your answer on Tutor.**

12. What is the effect when you increase the number of points  $Nx, Ny, Nz$ , while maintaining the values the other parameters?

- (a) increase of accuracy, decrease of computational time.
- (b) decrease of accuracy, increase of computational time.
- (c) increase of accuracy, increase of computational time.
- (d) decrease of accuracy, decrease of computational time.

**Submit your answer on Tutor.**

13. What is the effect of increasing the distance  $r/a$ , while maintaining the values of the other parameters?

- (a) increase of accuracy, no change in computational time.
- (b) decrease of accuracy, change in computational time.
- (c) increase of accuracy, change in computational time.
- (d) decrease of accuracy, no change in computational time.

**Submit your answer on Tutor.**

### Plotting sample codes:

- You can use the following code to save the Python data to a file:

```
import numpy as np

#####
# write all your function definitions here
#####

x,y,z,mag=hydrogen_wave_func(3,1, 0,10,20,20,20)

x.dump('xdata310.dat')
y.dump('ydata310.dat')
z.dump('zdata310.dat')
mag.dump('density310.dat')
```

- You can use the following code to plot using matplotlib:

```
import numpy as np
import matplotlib.pyplot as plt
from mpl_toolkits.mplot3d import Axes3D

x = np.load('xdata310.dat')
y = np.load('ydata310.dat')
z = np.load('zdata310.dat')

mag = np.load('density310.dat')

fig = plt.figure()
ax = fig.add_subplot(111, projection='3d')

for a in range(0,len(mag)):
    for b in range(0,len(mag)):
        for c in range(0,len(mag)):
            ax.scatter(x[a][b][c],y[a][b][c],z[a][b][c], marker='o',
                      alpha=(mag[a][b][c]/np.amax(mag)))

plt.show()
```

- You can use the following code to plot using mlab Mayavi package:

```
import numpy as np
from mayavi import mlab

x = np.load('xdata310.dat')
y = np.load('ydata310.dat')
z = np.load('zdata310.dat')

density = np.load('density310.dat')

figure = mlab.figure('DensityPlot')
mag=density/np.amax(density)

pts = mlab.points3d(mag,opacity=0.5, transparent=True)
# or pts = mlab.contour3d(mag, opacity=0.5)

mlab.colorbar(orientation='vertical')
mlab.axes()
mlab.show()
```

## Plots

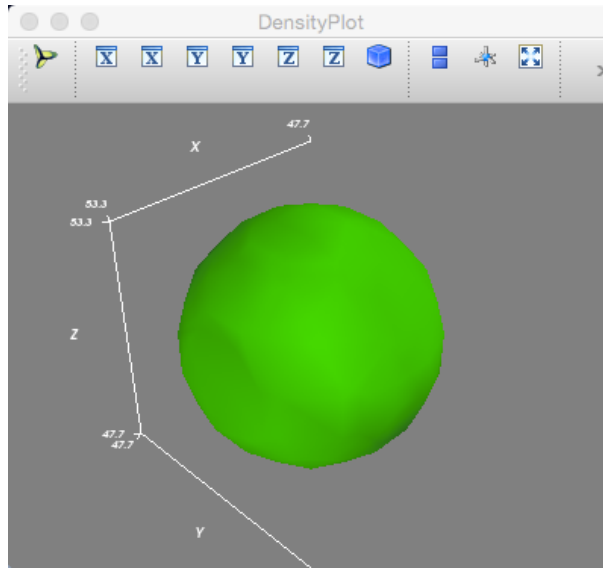


Figure 1: Magnitude plot for  $n = 2, l = 0, m = 0$  using contour3d from mlab Mayavi package.

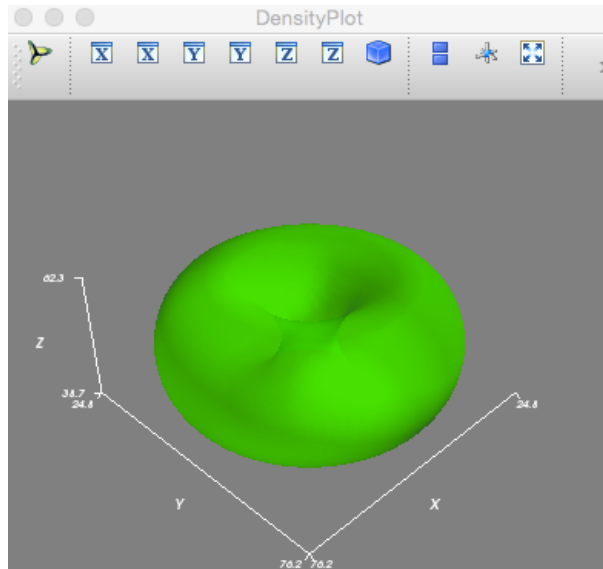


Figure 2: Magnitude plot for  $n = 2, l = 1, m = 1$  using contour3d from mlab Mayavi package.

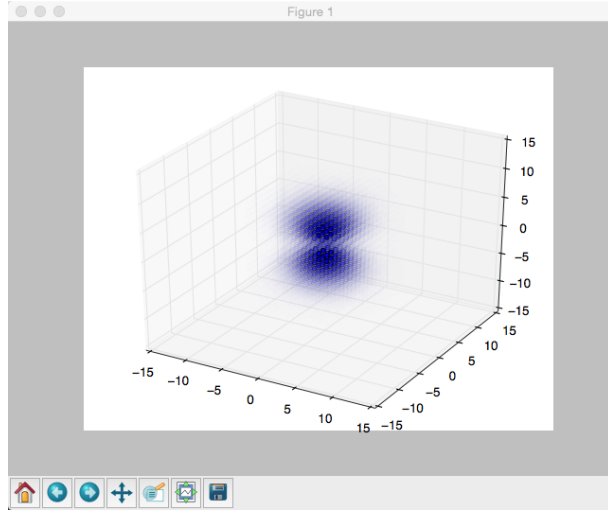


Figure 3: Magnitude plot for  $n = 2, l = 1, m = 0$  using scatter from Matplotlib package.

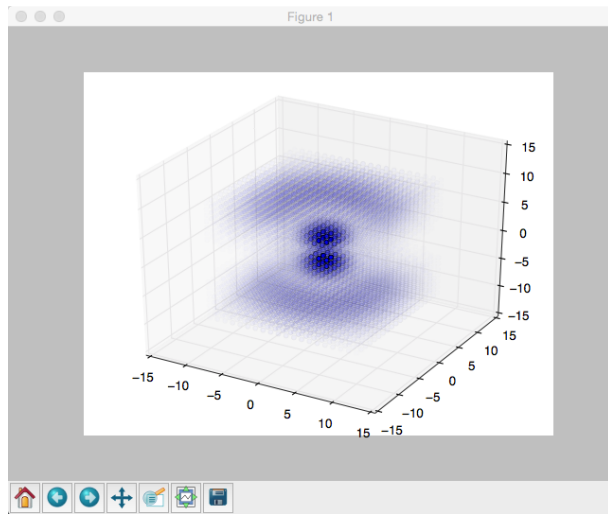


Figure 4: Magnitude plot for  $n = 3, l = 1, m = 0$  using scatter from Matplotlib package.



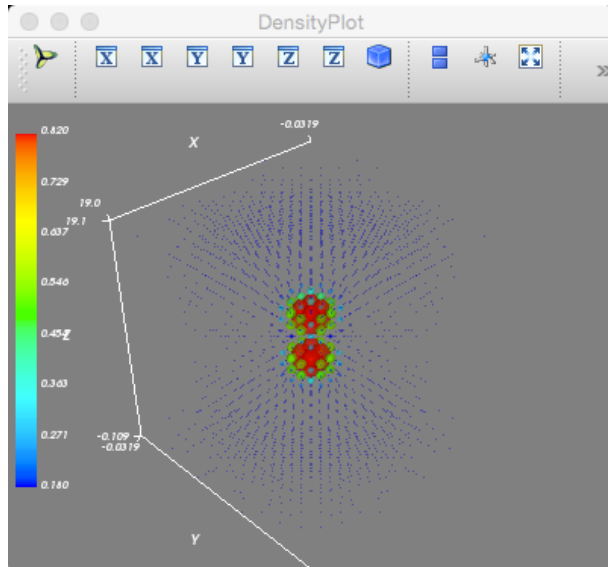


Figure 5: Magnitude plot for  $n = 3, l = 1, m = 0$  using points3d from mlab Mayavi package.