

HW2_Gabriel

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1 Homework 2 - Hückel Theory and Argus Lab

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In this homework, we had to do the following: 1. Calculate energy levels of HOMO/LUMO levels 2. Determine HOMO/LUMO wavefunctions and their electron density (and plot them) 3. Compare results with ArgusLab

```
In [27]: import numpy as np
import matplotlib.pyplot as plt
import os
%matplotlib inline
```

```
In [99]: alpha=-6.6 #eV
beta=-2.7 #eV
N= 10 #available e- in naphthalene
```

Here we determine the Hamiltonian of the system, defined by:

$$H_{ij} = \alpha \quad \text{if } i = j \quad (1)$$

$$H_{ij} = \beta \quad \text{if } i \text{ and } j \text{ are next neighbors} \quad (2)$$

$$H_{ij} = 0 \quad \text{otherwise} \quad (3)$$

$$(4)$$

```
In [47]: # determine hamiltonian => alpha if i=j, beta if first neighbours, 0 otherwise
H = np.zeros((N,N))
for i in range(N):
    for j in range(N):
        if i==j:
            H[i,j] = alpha
        elif j==(i-1) or j==(i+1):
            H[i,j] = beta

#additional neighbors, according to image from homework 1
#C5 neighbor of C10 and vice-versa
H[4,-1] = beta
H[-1,0] = beta
```

```

H[-1,4] = beta
H[0,-1] = beta
print H

[[-6.6 -2.7  0.   0.   0.   0.   0.   0.   0.  -2.7]
 [-2.7 -6.6 -2.7  0.   0.   0.   0.   0.   0.   0. ]
 [ 0.  -2.7 -6.6 -2.7  0.   0.   0.   0.   0.   0. ]
 [ 0.   0.  -2.7 -6.6 -2.7  0.   0.   0.   0.   0. ]
 [ 0.   0.   0.  -2.7 -6.6 -2.7  0.   0.   0.  -2.7]
 [ 0.   0.   0.   0.  -2.7 -6.6 -2.7  0.   0.   0. ]
 [ 0.   0.   0.   0.   0.  -2.7 -6.6 -2.7  0.   0. ]
 [ 0.   0.   0.   0.   0.   0.  -2.7 -6.6 -2.7  0. ]
 [ 0.   0.   0.   0.   0.   0.   0.  -2.7 -6.6 -2.7]
 [-2.7  0.   0.   0.  -2.7  0.   0.   0.  -2.7 -6.6]]

```

Below we calculate the eigenvalues and eigenvectors of the hamiltonian in order to get the energies and the corresponding wave functions (molecular orbitals) as coefficients ck (coeff_k)

```

In [104]: Ek, coeff_k = np.linalg.eig(H) # c_ik eigenvectors, E_k their eigenvalues

indices_sorted = np.argsort(Ek) #sort in ascending energies
Ek.sort()

sorted_coeff_k = coeff_k[indices_sorted]
print 'energies sorted: ' + str(Ek)

energies sorted: [-12.81749422 -10.96869177 -10.11749422  -9.3          -8.26869177
 -4.93130823  -3.9          -3.08250578  -2.23130823  -0.38250578]

```

Since we have 10 available electrons, we can fill up until the 5th orbital. We have the 1s₂, 2s₂ and 2p₆ totally filled. Therefore, our top most filled level (HOMO) is the 5th level (Ek[4]) and our lowest unoccupied level (LUMO) is the 6th level (Ek[5])

```

In [108]: E_gap = Ek[np.int(N/2)] - Ek[np.int(N/2 - 1)]
print 'Egap : ' +str(E_gap)+' eV'

Egap : 3.33738353925 eV

```

2 Calculate wave functions

From the website <http://winter.group.shef.ac.uk/orbitron/AOs/1s/equations.html>, we have that the wave functions for the 2p_z orbital can be constructed as follows

$$\phi_s = R_{1s} Y_{1s} = \frac{1}{2\sqrt{6}} \rho Z^{3/2} \exp(-\rho/2) \left(\frac{\sqrt{3} * z/r}{\sqrt{4\pi}} \right) \quad (5)$$

and the electron density is given by

$$n = \phi_{1s}^2 \quad (6)$$

```
In [109]: #Calculate wave function
def wavefunction(x,y,z):

    Z_eff = 3.136 #orbital specific, got a mean number for Carbon from wikipedia
    n = 2 #2p orbitals
    r = np.sqrt(x**2 + y**2 + z**2) #radial coordinate
    rho = 2*Z_eff*r/n / 52.9 #conversion between Bohr radius and atomic units
    R = 1./(2.*np.sqrt(6))*rho*Z_eff**(1.5)*np.exp(-rho/2)
    Y = np.sqrt(3)*z/r /np.sqrt(4.*np.pi)
    Psi_pz = R * Y

    return Psi_pz
```

Below we initialize the xy grid where the wavefunctions are going to be defined, and also the molecule is localized.

```
In [110]: width = 1000
length = 500
step = 0.5

gridX = np.arange(0,width,step)
gridY = np.arange(0,length,step)
x,y = np.meshgrid(gridX,gridY,sparse = True)

N = 10 #number of atoms

r = np.zeros((N,2)) #holds [x,y] position of each atom

d = 140
h = 0.5*d
w = np.sqrt(3)*0.5*d

#positions of atoms in molecule
r[0] = [w, 2*h+d]
r[1] = [0,h+d]
r[2] = [0,h]
r[3] = [w,0]
r[4] = [2*w,h]
r[5] = [3*w,0]
r[6] = [4*w,h]
r[7] = [4*w,h+d]
r[8] = [3*w,2*h+d]
r[9] = [2*w,h+d]
```

```

#we recenter the molecule so that the image is better localized in the grid and the pl
r[:,0] += 100
r[:,1] += 100

In [166]: rho = np.zeros((length/step,width/step))
#list_energies = []
list_z=[]
list_rhos=[]

for z in (-70,-30,1,10,50,80,100):
    list_energies=[]
    rho = np.zeros((length/step,width/step))
    for E in range(6):

        Psi_k = np.zeros((length/step,width/step))

        for n in range(N):

            xnew = x - r[n,0]
            ynew = y - r[n,1]
            Psi_k += coeff_k[n,indices_sorted[E]] * wavefunction(xnew, ynew, z)

        #if E <= 5:
        rho += 2 * (Psi_k)**2

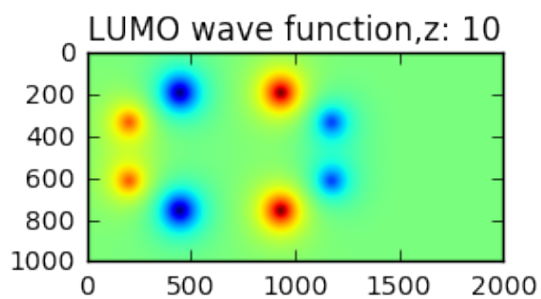
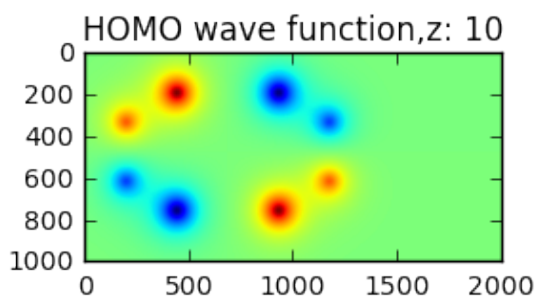
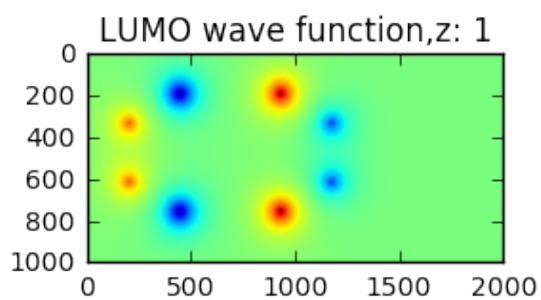
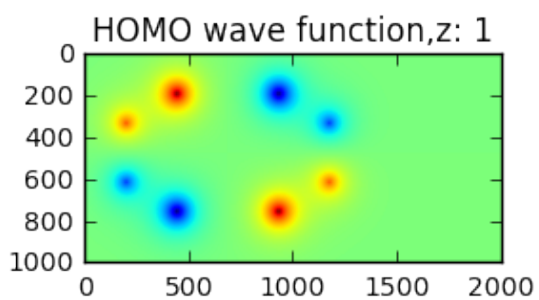
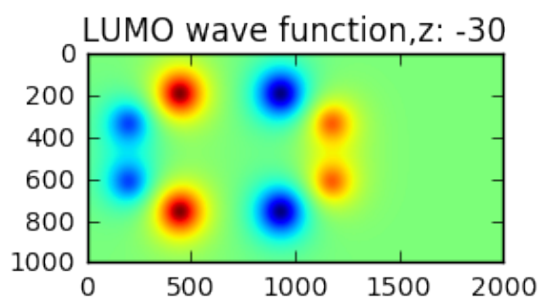
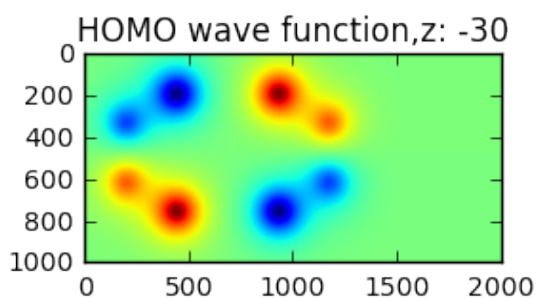
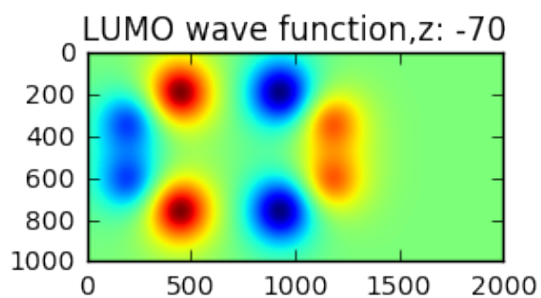
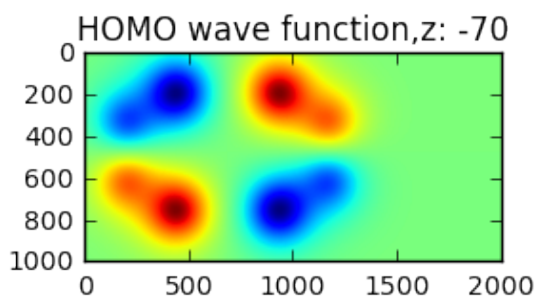
        list_energies.append([z,E,Psi_k])
    list_z.append([z,list_energies])
    list_rhos.append(rho)

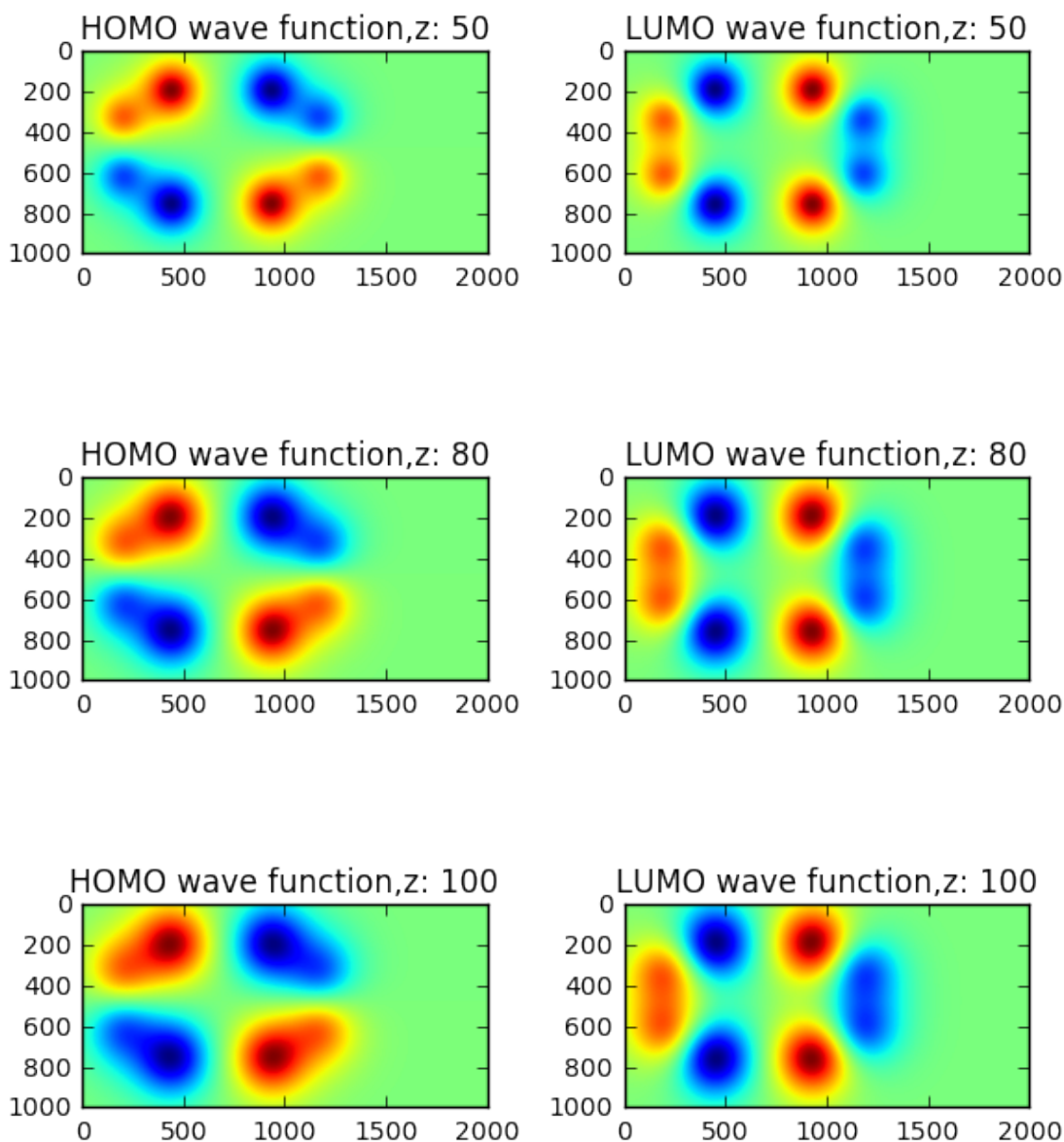
/Users/gabrielfior/anaconda/lib/python2.7/site-packages/ipykernel/__main__.py:1: VisibleDeprecat
if __name__ == '__main__':
/Users/gabrielfior/anaconda/lib/python2.7/site-packages/ipykernel/__main__.py:8: VisibleDeprecat
/Users/gabrielfior/anaconda/lib/python2.7/site-packages/ipykernel/__main__.py:11: VisibleDepreca

In [167]: #list_energies
for i in range(len(list_z)):
    plt.figure(i)

    plt.subplot(121)
    plt.imshow(list_z[i][1][-2][2])
    plt.title('HOMO wave function,z: '+str(list_z[i][1][-2][0]))
    plt.subplot(122)
    plt.imshow(list_z[i][1][-1][2])
    plt.title('LUMO wave function,z: '+str(list_z[i][1][-2][0]))
    plt.tight_layout()
    plt.show()
#print len(list_z[0][1][-1])

```

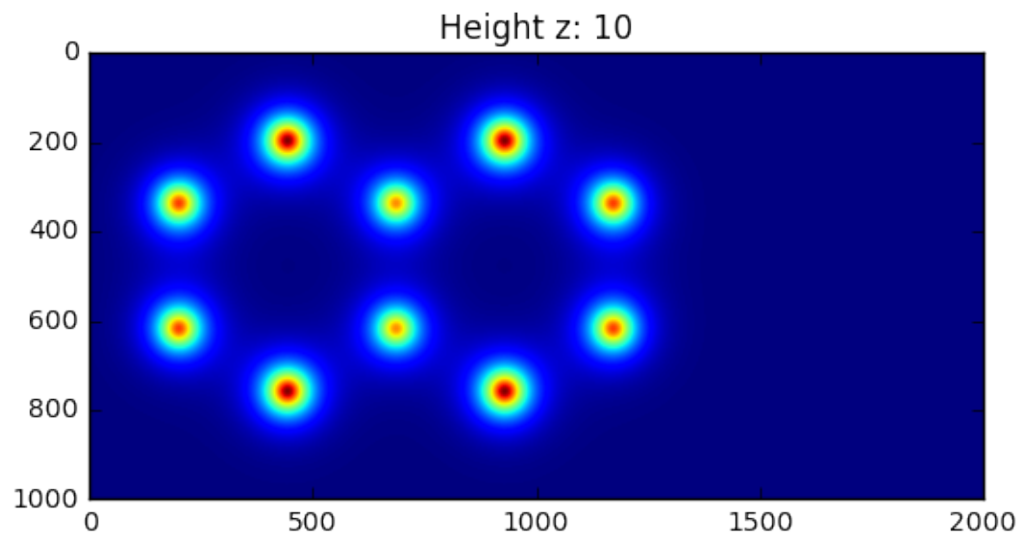
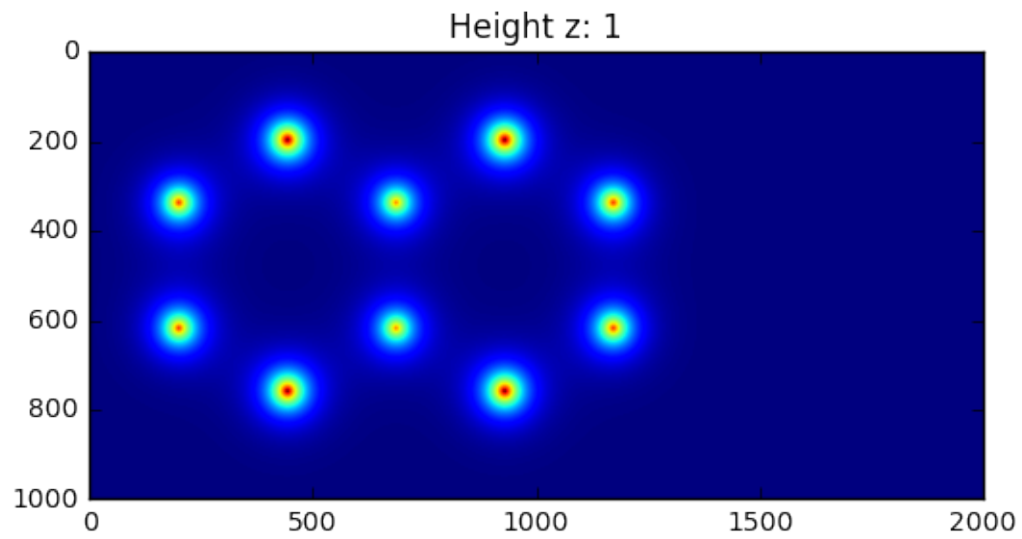


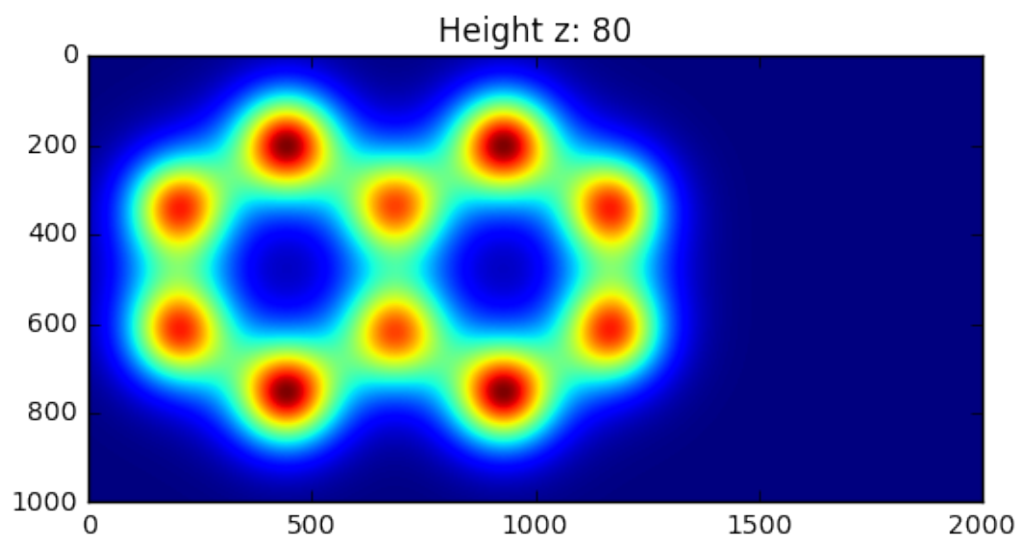
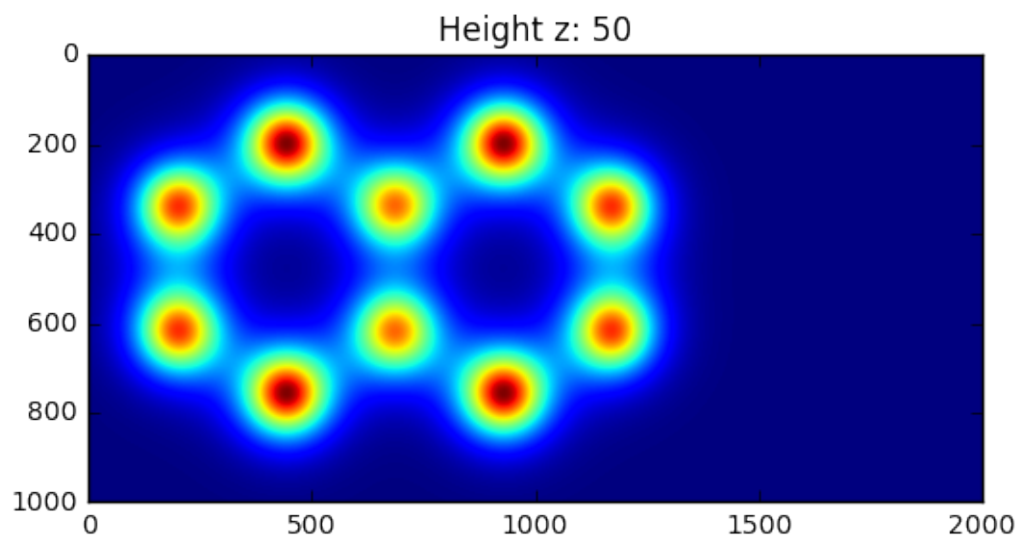


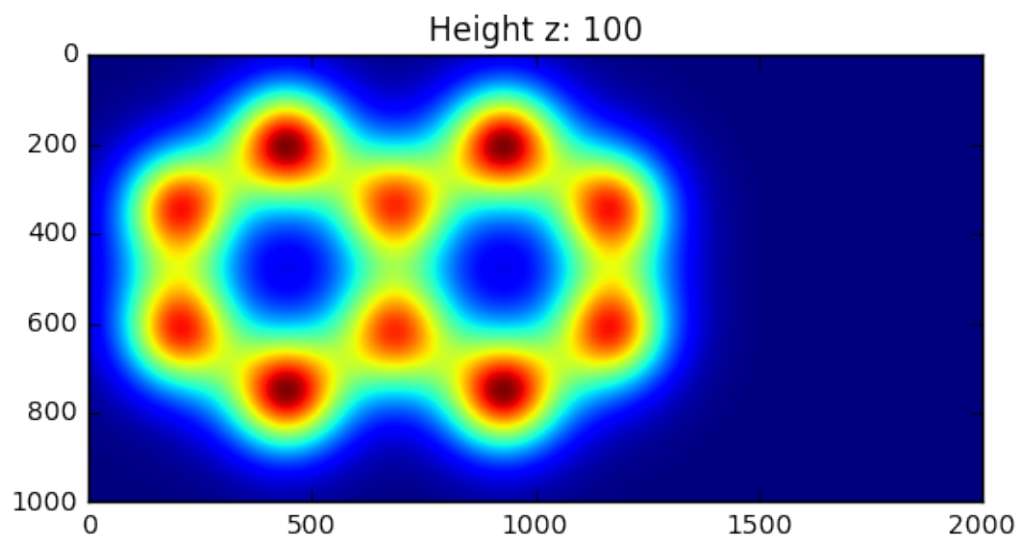
In the images above, one can notice that, for the low heights z (1,10) the atom positions can be directly noticed. As the height increases, the wave function get smeared out.

Below I plot the electron charge density ρ , which was obtained by summing up the absolute value of the wave function up to the HOMO level.

```
In [165]: print len(list_rhos)
          for i,j in zip(range(len(list_rhos)),(1,10,50,80,100)):
              plt.imshow(list_rhos[i])
              plt.title('Height z: '+str(j))
              plt.show()
```







The final step is to compare my results with ArgusLab. As can be seen in the images below, the plots of the HOMO/LUMO orbitals agree very well, specially if you see that in both calculations the sign of the wave function changes (going from blue to red, when z goes from a negative value to a positive value), and vice-versa.

LUMO orbital calculated for Naphtalene by Argus

HOMO orbital calculated for Naphtalene by Argus

In []: