
Modeling strong-field ionization in small molecules

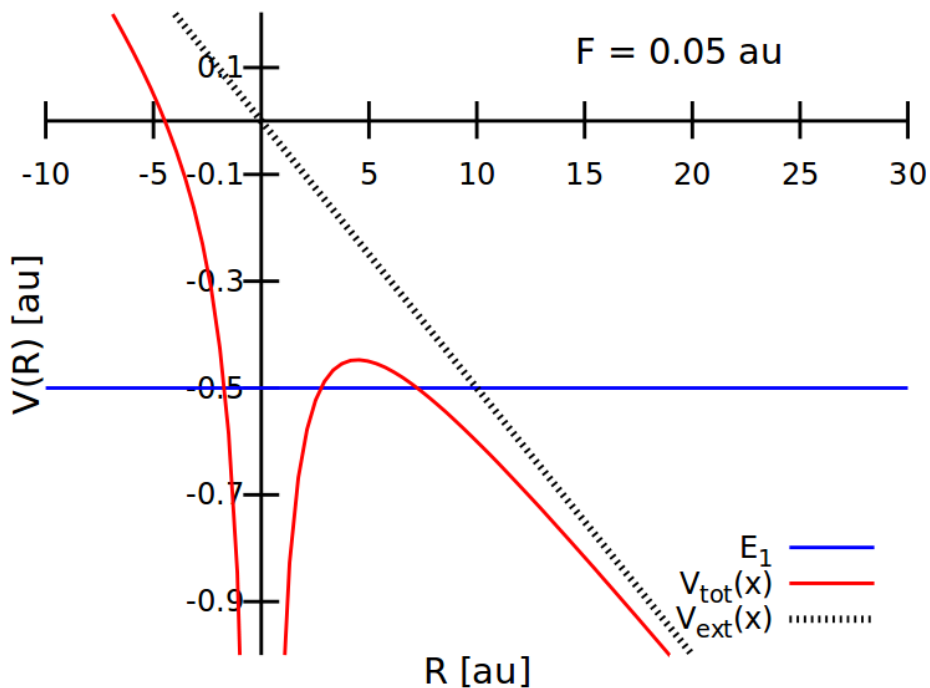
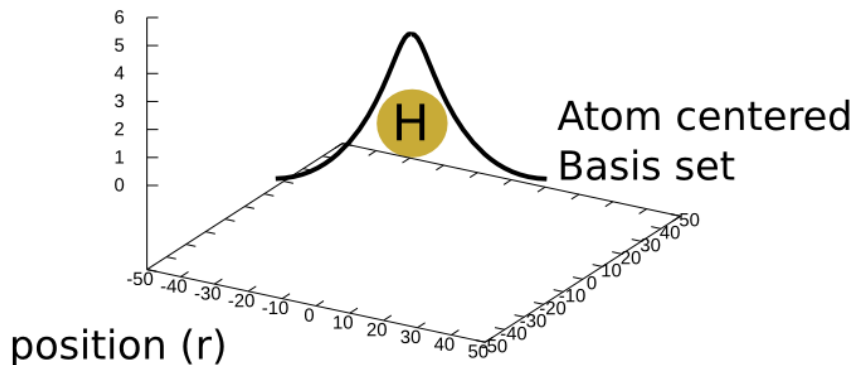
Margarite Picone, LSMSA, Georgia Tech

Dr. Kenneth Lopata, LSU Chemistry Dept.

Adonay Sissay, LSU Chemistry Dept.

Recap

- Strong-field ionization
- Tunneling
- Basis sets



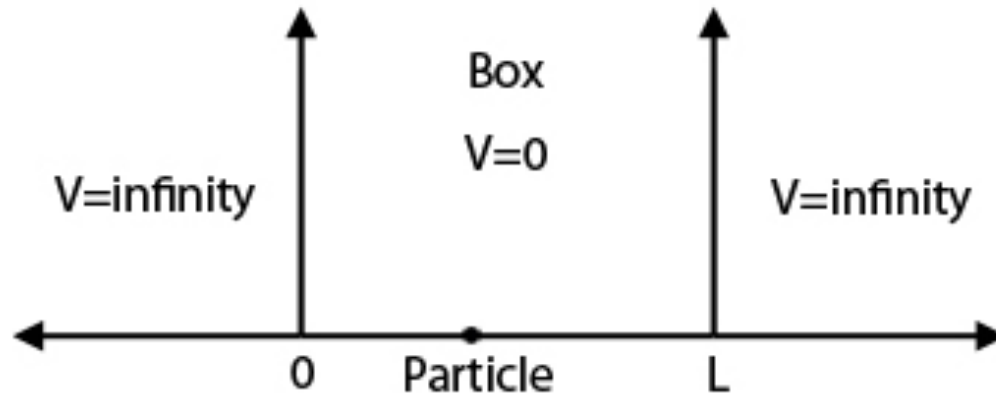
Goals

- learning the approximate wave functions to form a set
- engineering basis sets
- extending the density functional theory to strong-field
- find tunnel lengths in hydrogen atom

Achieved

- derived and understood wave functions for the particle in a box problem (PIB)
- created code to find energy levels of PIB with an electric field
- formed basis sets with wave functions
- studied polarizability of 1D “molecules”

Particle in a box



Particle in a box

- no potential acting on particle (free particle)
- Schrodinger Equation for free particle:

$$\frac{d^2\Psi}{dx^2} + \frac{2m}{\hbar^2}[E - V(x)]\Psi(x) = 0$$

- Solution: $\Psi(x) = A\cos(kx) + B\sin(kx)$
- $\Psi(x)$ is probability a particle is in a region
- $\Psi(x)$ must be zero for barriers

Particle in a box

- $B\sin(ka)$ must be zero, so ka must be $n\pi$ for $n=1, 2, 3, \dots$
- Solving further reveals discrete energy levels

Code

- building phi
- symmetrical box
- new conditions

```
def phi(n, a, x):  
    if (n%2 == 0): #even n  
        val = math.sqrt(2.0/a)*math.sin(2.0*n*math.pi*x/a)  
    else:          # odd n  
        val = math.sqrt(2.0/a)*math.cos(n*math.pi*x/a)  
    return val
```


Linear Combination of Wavefunction

- Basis set of wave functions constructs ψ

$$\psi(x) = \sum_i c_i \phi_i(x)$$

- Optimize combination to minimize energy
- Best ground-state guess will be lowest eigenvalue from eigenvalue problem

Hamiltonian

- Eigenvalue problem

$$H\psi = E\psi$$

- Hamiltonian operator:

$$H\psi = (H + V(x))\psi(x)$$

- with electric field in box $V(x) = -Fqx$:

Code

- Construct matrix

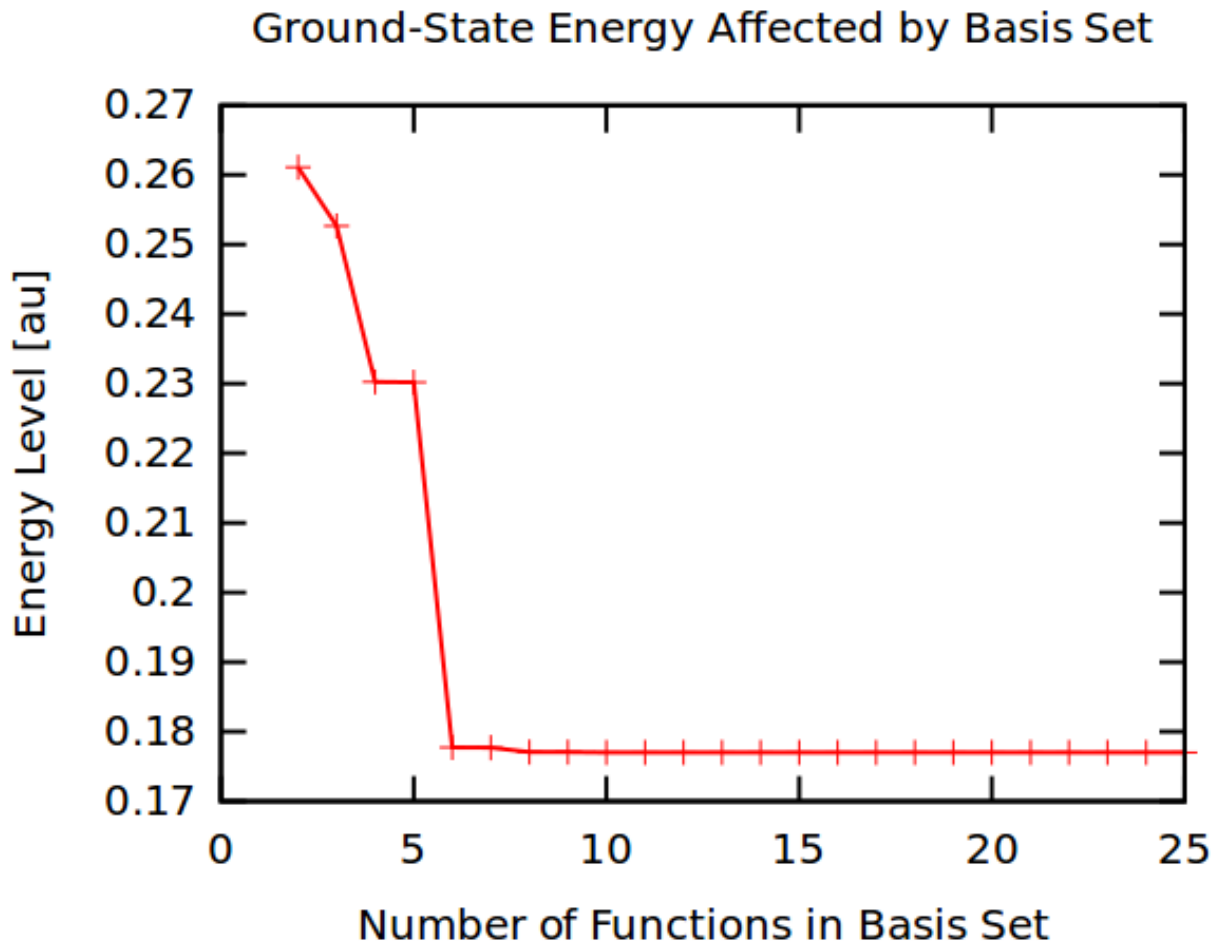
```
H_matrix= np.array( [ [ calc_Hnm(n, m) for m in range(1,10)] for n in range(1,10)] )
```

- Perform matrix operations. Pull lowest value

```
vals,vecs= LA.eigh(H_matrix)
idx = vals.argsort()
vals = vals[idx]
vecs= vecs[:,idx]
bestGuess= vals[0]
```

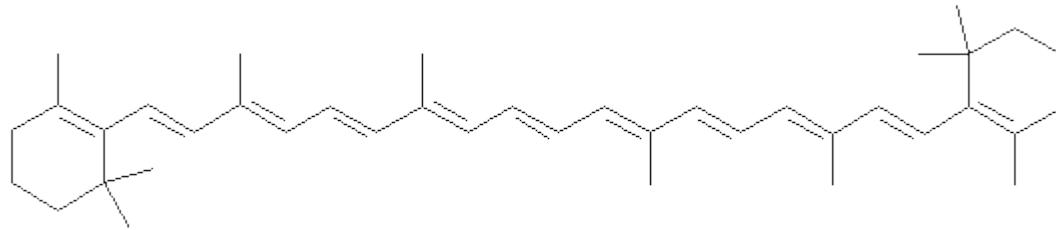
Basis Set

- $A = 18.9$ au
- $F = .3$ au



beta-Carotene

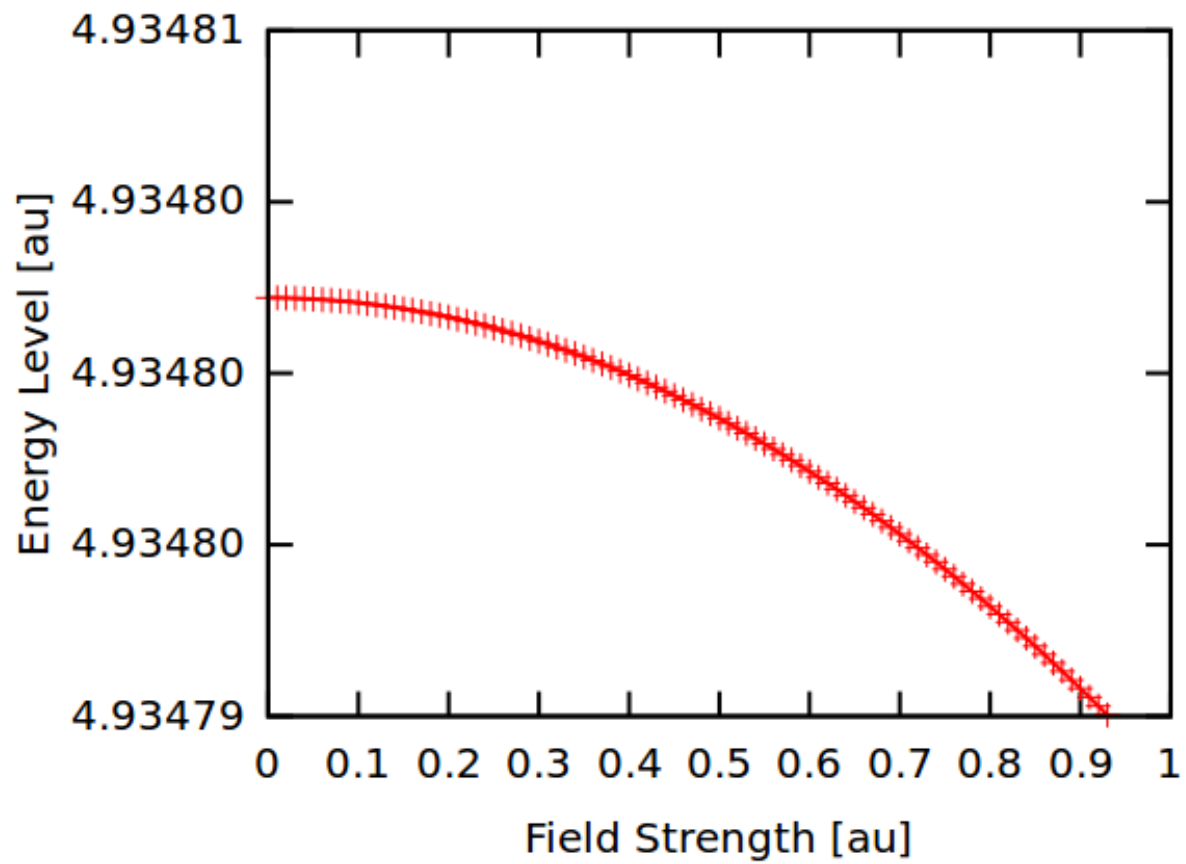
- bad model for molecules
- simple, linear ones do exist



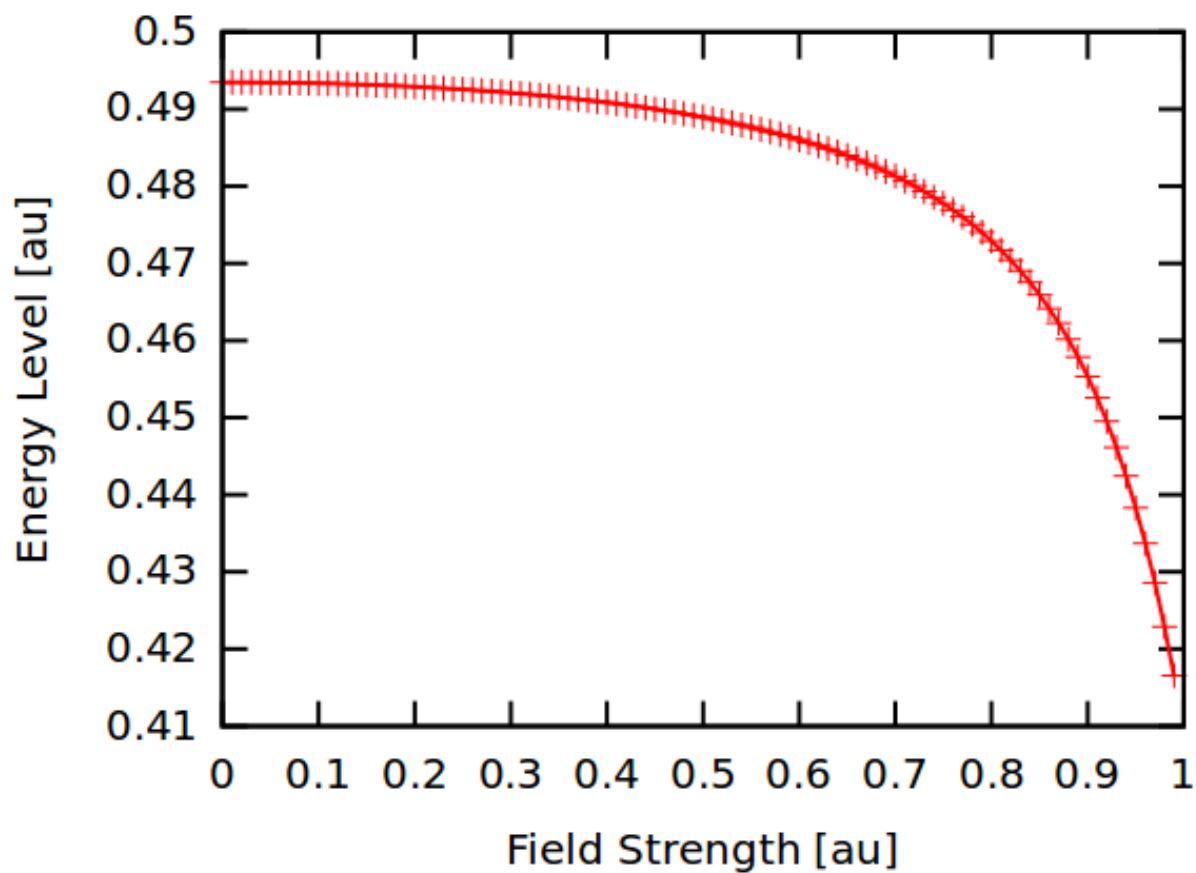
Length of molecules

- set basis at 25 functions
- set electric field at 0.3 au
- size of molecule will affect strength of field needed for ionization

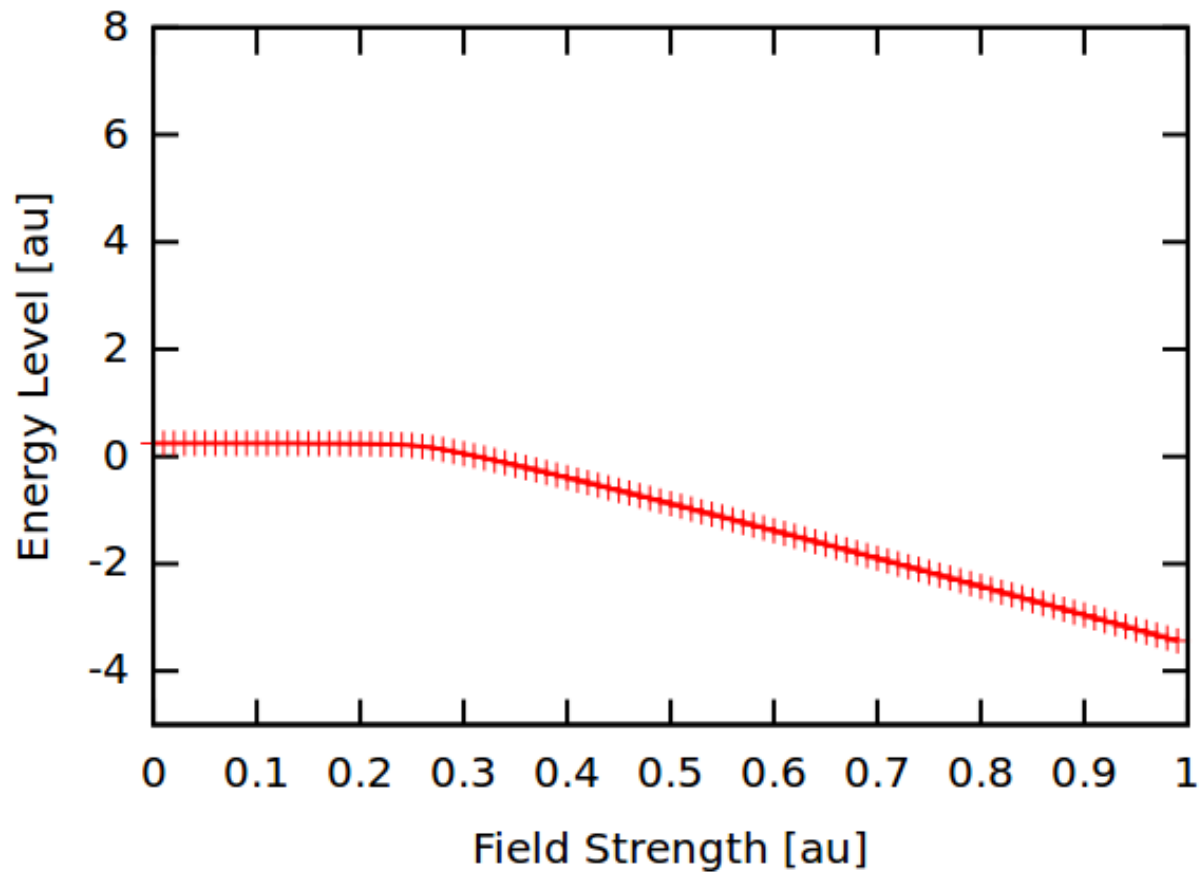
Sensitivity of Small Molecule [1 au]



Sensitivity of Medium Sized Molecule [10 au]



Sensitivity of Large Molecule [20 au]

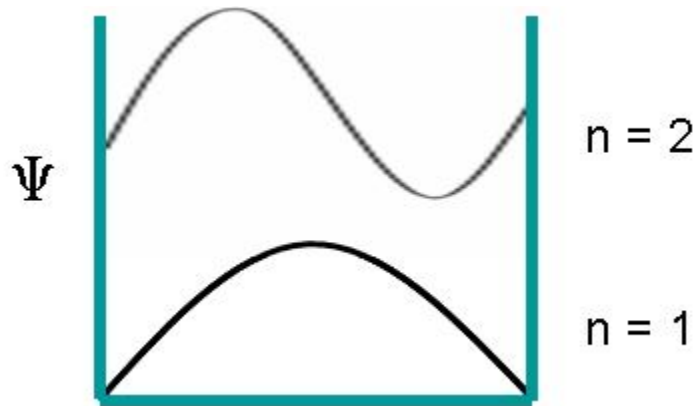


Polarizability

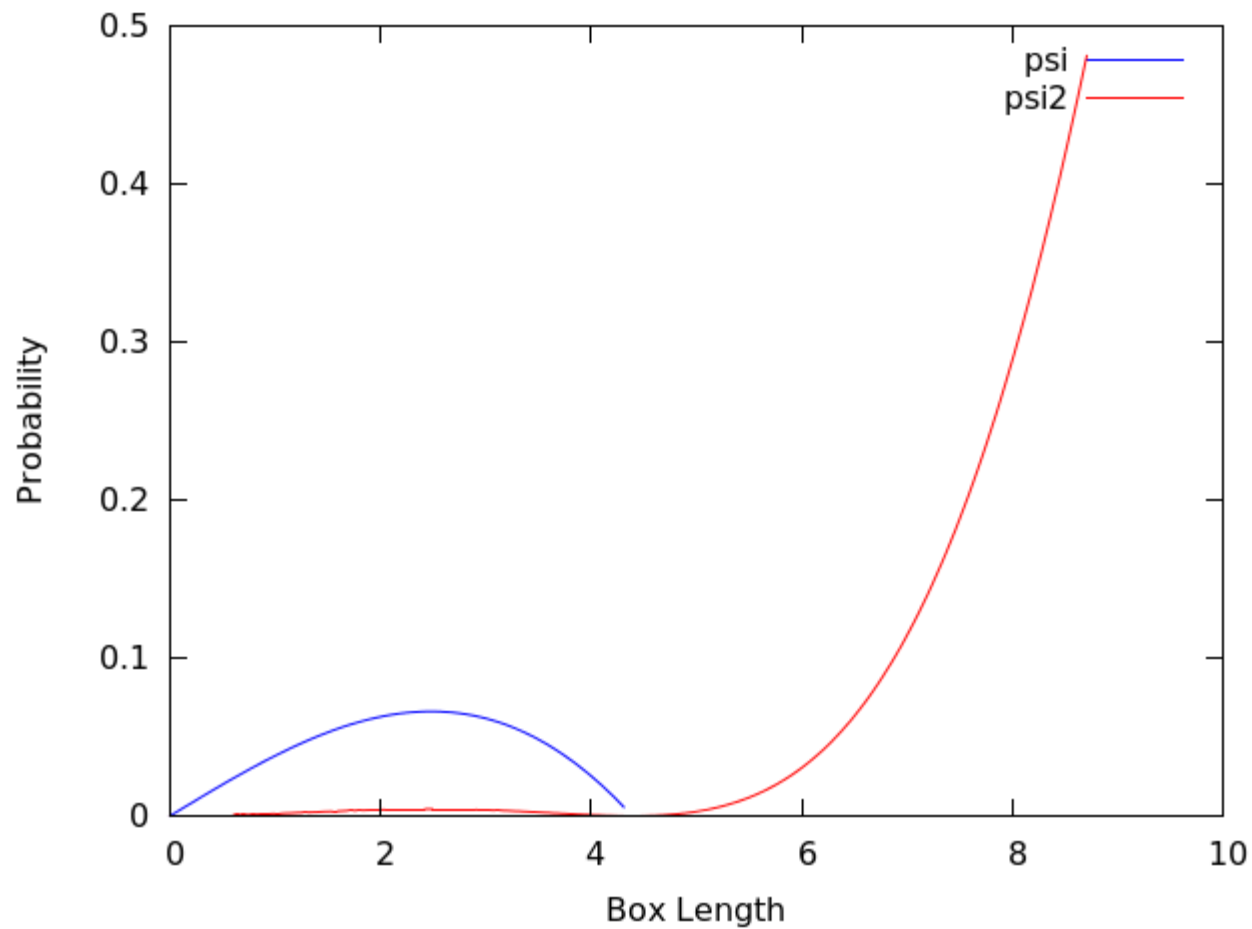
- inclination of a system to respond to an external force
- ground-state energy to be affected by field
- can provide insight to molecular structures

Location of Particle

- probability of location = ψ^2
- probability without an electric field



$n = 1$
 $F = .1 \text{ au}$



Future Work

- modify code for Hydrogen
- radial and angular parameters
- potentially expand for diatomic molecules
- add to code to find tunnel length

Acknowledgements

- Dr. Lopata and Adonay Sissay
- LA-SiGMA
- NSF EPSCoR
- Louisiana Board of Regents