1 Theory

1.1 Wavefunction of Helium

The single-particle wave function of an electron i in the 1s state is given in terms of a dimensionless variable (the wave function is not normalized) as

$$\psi_{1s}(\vec{r_i}) = e^{-\alpha r_i}$$

Where the electron position \vec{r}_i is

$$\vec{r}_i = x_i \vec{e}_x + y_i \vec{e}_y + z_i \vec{e}_z$$

and its distance from the origin r_i is

$$r_i = \sqrt{x_i^2 + y_i^2 + z_i^2}$$

 α is a parameter set to 2, which corresponds to the carge of the Helium atom, Z=2.

For our system with two electrons, we have the product of the two 1s wave functions defined as

$$\Psi(\vec{r}_1, \vec{r}_2) = e^{-\alpha(r_1 + r_2)}$$

This leads to the integral which will be solved nummerically with the different methods mentioned earlier. The value of the integral corresponds to the expectation value of the energy between the two electrons repelling each other due to Columb interactions.

$$\langle \frac{1}{|\vec{r_1} - \vec{r_2}|} \rangle = \int_{\infty}^{\infty} d\vec{r_1} d\vec{r_2} e^{-2\alpha(r_1 + r_2)} \frac{1}{\vec{r_1} - \vec{r_2}}$$
(1)

This is the integration that will be performed numerically in multiple ways in this paper. The analytical result is $5\pi/16^2$.

1.2 Gaussian Quadrature

The main idea of Gaussian quadrature is to integrate over a set of points x_i not equally spaced with weights w_i , which are calculated in /code/Gauss-Quadrature/src/gauleg.cpp. The weights are found through orthogonal polynomials(Laguerre and Legendre polynomials) in a set interval. The points x_i are chosen in a optimal sense and lie in the interval.

The intgral is approximated as

$$\int_{a}^{b} W(x)f(x) \approx \sum_{i=1}^{n} \omega_{i}f(x_{i})$$

For a more detalled derivation and explanation of Gaussian quadrature see [1].

1.2.1 Gauss-Legendre

Using Gauss-Legendre quadrature with Legendre polynomials will make it possible to solve the integral numerically. The first step is to change the integration limits from $-\infty$ and ∞ to $-\lambda$ and λ . The λ 's are found by inserting it for r_i in the expression $e^{-\alpha r_i}$ because $r_i \approx \lambda$ when $e^{-\alpha r_i} \approx 0$. From figure 1, $\lambda \in [-5, 5]$ is therefor a good approximation for the integration limits.

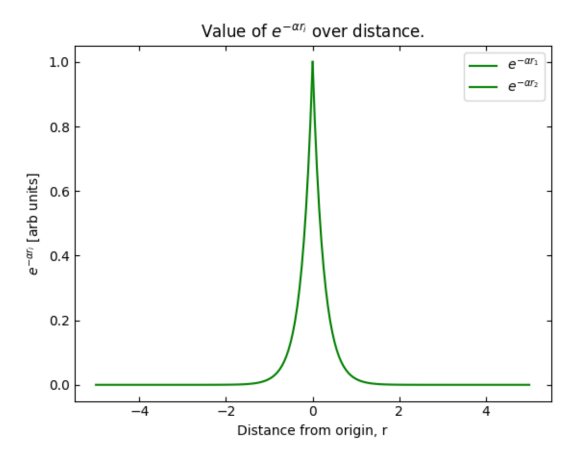


Figure 1: Plot of wavefunction in one dimension

The weights and mesh points are computed using /code/Gauss-Quadrature/src/gauleg.cpp. Eventually ending up with a sixdimensional integral, where all six integration limits are the same.

$$\int_a^b \int_a^b \int_a^b \int_a^b \int_a^b \int_a^b e^{-x} f(x) dx \approx \sum_{i=1}^n w_i f(x_i)$$

1.2.2 Improved Gauss-Quadrature- Laguerre

Gauss-Legendre quadrature gets the job done, but it is unstable and unsatisfactory. By changing to spherical coordinates and replacing Legendre- with Laguerre polynomials an improvement in accuracy is expected. The Laguerre polynomials are defined for $x \in [0, \infty)$, and in spherical coordinates:

$$d\vec{r}_1 d\vec{r}_2 = r_1^2 dr_1 r_2^2 dr_2 d\cos(\theta_1) d\cos(\theta_2) d\phi_1 d\phi_2$$

with

$$\frac{1}{r_{12}} = \frac{1}{\sqrt{r_1^2 + r_2^2 - 2r_1r_2cos(\beta)}}$$

and

$$cos(\beta) = cos(\theta_1)cos(\theta_2) + sin(\theta_1)sin(\theta_2)cos(\phi_1 - \phi_2)$$

For numerical integration, the deployment of the following relation is nessecary:

$$\int_0^\infty e^{-x} f(x) dx \approx \sum_{i=1}^n w_i f(x_i)$$

where x_i is the *i*-th root of the Laguerre polynomial $L_n(x)$ and the weight w_i is given by

$$w_i = \frac{x_i}{(n+1)^2 [L_{n+1}(x_i)]^2}$$

The Laguerre polynomials are defined by Rodrigues formula:

$$L_n(x) = \frac{e^x}{n!} \frac{d^n}{dx^n} \left(e^{-x} x^n \right) = \frac{1}{n!} \left(\frac{d}{dx} - 1 \right)^n x^n$$

or the recursive relation:

$$L_0(x) = 1$$

$$L_1(x) = 1 - x$$

$$L_{n+1}(x) = \frac{(2n+1-x)L_n(x) - nL_{n-1}(x)}{n+1}$$

1.2.1

1.3 Monte Carlo

1.3.1 Generalized

Monte Carlo integration is based on the idea of finding the mean of a function in a domain by sampling random function values. This mean multiplied by the volume of the domain will be an approximation of the integral.

Say we have an integral I of $f(\mathbf{x})$ we want to find:

$$I = \int_D f(\mathbf{x}) d\mathbf{x}$$

where \mathbf{x} is in the domain D. This integral can be approximated by using random numbers distributed on D by the probability distribution function (PDF) $p(\mathbf{x})$. Discretizing, the approximated integral now becomes

$$I \approx \langle I \rangle = \frac{1}{N} \sum_{i=0}^{N} \frac{f(\mathbf{x}_i)}{p(\mathbf{x}_i)},$$
 (2)

where N is the number of sampled values.

1.3.2 Naïve approach (uniform PDF)

To solve our six-dimensional integral, we first take the naïve approach and distribute our randomly chosen variables on the uniform distribution

$$\theta(x) = \begin{cases} \frac{1}{b-a}, & \text{for } x \in [a, b], \\ 0 & \text{else} \end{cases}$$

and keep our variables \mathbf{r}_1 and \mathbf{r}_2 in cartesian coordinates. Putting the uniform distribution into (2), we get the naïve approximation of an integral:

$$\langle I \rangle = \frac{V}{N} \sum_{i=0}^{N} f(\mathbf{x}_i).$$
 (3)

Here V is the integration volume (for d dimensions in cartesian coordinates $V = (b-a)^d$, with b and a being the integration limits for each dimension). Going back to our original integral (1), our approximation of it using this method is

1.3.3 Paralellization

To run the computations faster, openMP will be used to paralellize the code. This shares the workload across multiple processor threads and results in a substantional decrease in time spent for the same amount of operations. Some important remarks when doing Monte-Carlo integration in paralell is:

• Create a random number generator in earch thread.

- \bullet Keep the summations private for each thread.
- Sum the private summations from each thread together after the calculations are completed.

By doing this we avoid having the threads wait for the random number generator and writing to the same memory, thereby achieving optimal speedup.

The code is commented in for example /code/Monte-Carlo/src/naiveMC.cpp.