Project 3

Benedicte Allum Pedersen, Emil Heland Broll Fredrik Oftedal Forr

Abstract

We have computed the integral for the expectation value of the correlation energy between to electrons in a helium atom by using Gauss-Legendre and Gauss-Laguerre quadrature as well as Monte Carlo integration. We experienced that the methods that were brute force gave a less satisfactory resulst compared to the methods that required some more thinking. The exact value of the integral should be 0.192765. The best value we got from Gauss-Legendre was 0.192651, while the best value from Gauss-Laguerre was 0.194779. We also experienced that Gauss-Laguerre gave better results when we increased the number of integration points, while this was not the case for Gauss-Legendre.

Introduction

In this report there were given a brief rundown on different numerical integration methods. The methods that have been used is Gauss-Legendre and Gauss-Laguerre quadrature as well as Monte Carlo integration. There will be adjustments to these to show the different strengths of the methods. Under this section you can find the method on how the program works, results from the program, discussion where we analyze the results and ending with a conclusion.

The integral we have worked with is a six-dimensional integral which determines the ground state correlation energy between two electrons in a helium atom.

The single-particle wave function for an electron i in the 1s state in a hydrogen atom is as followed:

$$\psi_{1s}(\mathbf{r}_i) = e^{-\alpha r_i}$$

where $\mathbf{r}_i = x_i \mathbf{e}_x + y_i \mathbf{e}_y + z_i \mathbf{e}_z +$, and the value of r_i is given by:

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

We set $\alpha=2$ which corresponds to the charge of the helium atom, Z=2. We assume that the wave function for each electron in the helium atom can be modelled like the single-particle wavefunction above. The wavefunction for two electron is then given by the product of two single-particle wavefunctions:

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

We need to solve the integral for the expectation value of the correlation energy between two electrons:

$$\left\langle \frac{1}{|\mathbf{r}_1 - \mathbf{r}_2|} \right\rangle = \int_{-\infty}^{\infty} d\mathbf{r}_1 d\mathbf{r}_2 e^{-\alpha(r_1 + r_2)} \frac{1}{|\mathbf{r}_1 - \mathbf{r}_2|}$$

The exact value of this integral is $5\pi^2/16^2 = 0.192765$.

Method

In this problem there have been used c++ as programming language and mainly a variety of functions to solve the integral.

Gaussian Quadrature

The first method to be used is Gaussian Quadrature. To compute the integral numerically we do the approximation:

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

Unlike other more basic methods for numerical integrations where the mesh points x_i are equidistantly spaced, the mesh points in the gaussian quadrature are not equidistantly spaced. In the sum above w_i corresponds to the weights. Gaussian quadrature uses some orthonogonal Legendre- and Laguerre-polynomials, to obtain the mesh points and weights.

Gauss-Legendre is a brute-force method and to obtain the value of the integral using this method we have to set up the mesh points and weights that corresponds to some finite integrations limits [a, b]. To set up the mesh points and weights we haved used the function gauleg from Morten's example program.cpp. We can replace the intigration limits $-\infty$ and ∞ with $-\lambda$ and $+\lambda$.

While the Legendre polynomials are defined for $x \in [-1, 1]$ the Laguarre polynomials are defined for $x \in [0, \infty)$. To improve our result, we rewrite the integral by changeing to spherical coordinates.

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

Thus the integral becomes:

$$I = \int_0^\infty r_1^2 dr_1 \int_0^\infty r_2^2 dr_2 \int_0^\pi dcos(\theta_1) \int_0^\pi dcos(\theta_2) \int_0^{2\pi} d\phi_1 \int_0^{2\pi} d\phi_2 \frac{e^{-2\alpha(r_1 + r_2)}}{r_{12}}$$

where

$$\frac{1}{r_{12}} = \frac{1}{\sqrt{r_1^2 + r_2^2 - 2r_1r_2\cos(\beta)}}$$

with

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

We perform the integration over $\theta \in [0, \pi]$, $\phi \in [0, 2\pi]$ and $r \in [0, \infty)$. To calculate the mesh points and weight for the angles we use the same function as for the Gauss-Legendre quadrature. To calculate the mesh points and weights corresponding to r_1 and r_2 we use Morten's function gauss_laguerre.cpp.

Monte Carlo Integration

We'll firstly perform a brute force Monte Carlo integration of our system. We use the same function as in a) and b), but in the Monte Carlo method, we "guess" at the values for r1 and r2 using a simple uniform distribution function. To approximate the infinite limits of our integral, we again choose the limits to be [-3,3] because we know our function flattens out when the variables approach these limits. This lets us approximate the integral as:

$$I \approx (3 - (-3))^6 \frac{1}{N} \sum_{i=1}^{N} f(x_{1,(i)}, y_{1,(i)}, z_{1,(i)}, x_{2,(i)}, y_{2,(i)}, z_{2,(i)})$$

For the next part of the Monte Carlo strategy, we will try to adapt our method to the actual integrand and integration variables. Looking at our function, we can transform our integral into polar coordinates, as in b), and using an exponential distribution of the form $p(x) = \frac{1}{\lambda} e^{-x/\lambda}$. This results in the following new function to be integrated:

$$f(r_1, r_2, \phi_1, \phi_2, \theta_1, \theta_2) = \frac{r_1^2 r_2^2 \sin \theta_1 \sin \theta_2}{\sqrt{r_1^2 + r_2^2 - 2r_1 r_2 \cos b}}$$
$$b = \cos \theta_1 \cos \theta_2 + \sin \theta_1 \sin \theta_2 \cos(\phi_1 - \phi_2)$$

The integral can then be approximated like this:

$$I \approx \frac{(2\pi)^2 \pi^2}{16N} \sum_{i=1}^{N} f(r_1, r_2, \phi_1, \phi_2, \theta_1, \theta_2)$$

In order to be able to get the best possible results with the Monte Carlo method, we want N to be as big as possible. To achieve this, we use various compiler flags, and parallelisation, with OpenMP.

Results

Gaussian Quadrature

We have plotted the wave function for the electrons in the Helium atom in figure 1. From this plot we can tell that the integral converges to zero for x < -2 and x > 2. Therefore the integral would not need to be evaluated for those values of x.

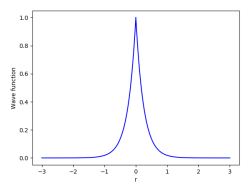


Figure 1: Plot of the wave function for to electron in a Helium atom.

Table 1 and table 2 shows the results for the two Gaussian quadratures with different values of N. The tables also shows the difference from the exact value of the integral, namely 0.192765. The integral values for both of the Gaussian quadratures are plotted against different number of integration points in figure 2 You can also see a plot of the difference from the exact value for Gauss-Legendre and Gauss-Laguerre in figure 3.

The Gauss-Legendre quadrature gave different results when we changed the limits of the integral, $-\lambda$ and $+\lambda$. We found that we got the best result for Gauss-Legendre using $\lambda=2.89$ and N=27. The value of the integral then where 0.192651 with a difference 0.000114 from the exact value.

Monte Carlo Integration

The results for the brute force Monte Carlo integration are showed in Table 3, and the integration values for the improved Monte Carlo method are showed inn Table 4. In Table 3 and 4 also lists the time used for calculating the integral. When we parallelised our code we experienced a quicker calculation of the integral.

Table 1: Integration using Gauss-Legendre, variying N, with $\lambda=3$

| Ν | Legendre | Diff. from exact |
|----|----------|------------------|
| 10 | 0.071980 | 0.120785 |
| 15 | 0.239088 | 0.046323 |
| 20 | 0.156139 | 0.036626 |
| 25 | 0.196817 | 0.003052 |
| 27 | 0.193524 | 0.000759 |
| 30 | 0.177283 | 0.015482 |

Table 2: Integration using Gauss-Laguerre, variying N

| N | Laguerre | Diff. from exact |
|----|----------|------------------|
| 10 | 0.177081 | 0.0156840 |
| 15 | 0.193285 | 0.000520 |
| 20 | 0.194786 | 0.002021 |
| 25 | 0.194804 | 0.002039 |
| 27 | 0.193524 | 0.002030 |
| 30 | 0.194779 | 0.002014 |

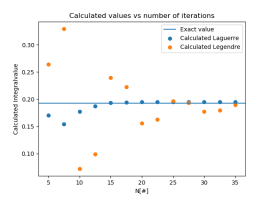


Figure 2: Calculated values for the integral against number of iterations using Gauss-Legendre and Gauss-Laguerre.

Table 3: Integration using Monte Carle integration, variying N

| N | Laguerre | Diff. from exact | Time used |
|----------|----------|------------------|-------------------|
| 10^{5} | 0.244992 | 0.052227 | 0 |
| 10^{6} | 0.183088 | 0.009676 | 0 |
| 10^{7} | 0.193923 | 0.001159 | $1 \cdot 10^{-6}$ |
| 10^{8} | 0.192657 | 0.000108 | $8 \cdot 10^{-6}$ |

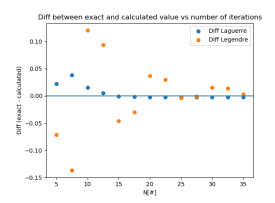


Figure 3: Difference from the exact value of the integral for Gauss-Legendre and Gauss-Laguerre against number of iterations.

Table 4: Integration using imporoved Monte Carle integration, variying N

| N | Laguerre | Diff. from exact | Time used (without parallization) | Time used (with parallization) |
|----------|----------|------------------|-----------------------------------|--------------------------------|
| 10^{5} | 0.192894 | 0.000129 | 0 | 0 |
| 10^{6} | 0.193056 | 0.000291 | 0 | |
| 10^{7} | 0.193333 | 0.000568 | $1 \cdot 10^{-6}$ | |
| 10^{8} | 0.193324 | 0.000049 | $9 \cdot 10^{-6}$ | |

Discussion

The Gauss-Legendre quadrature is a brute force method and we can see from our results that is is not a very good method. We had to try with different integration values and test which values that gave the best result. With increasing values of N the integrand got less exact insted of more close to the correct value. When we use the Gauss-Laguerre quadrature we have to do some more thinking by changing to spherical coordinates. The results got better and our we got the best result for N=15, with difference 0.0005020 from the exact value.

Conclution

Bibliography

Link to our GitHub repository.