Project 2

Vilde Mari Reinertsen

Abstract

Many theoretical physics systems involve the many-body Schrödinger equation (SE) and modelling a quantum dot is one of them. The problem with the many-body SE is that it is very hard or impossible to solve it analytically. Therefore, numerical solution are necessary to gain insight into the behaviour of many-body systems of particles. In this particular project I am working with quantum Variational Monte Carlo (VMC) methods to solve the many-body SE for fermions in an harmonic oscillator trap, which here represents the quantum dot. I am looking at full-shell systems of two, six and twelve fermions in harmonic oscillator traps of different strengths, i.e. trap frequencies, and investigating the different systems by comparing expectation energies and one-body densities. Additionally, I look at how these properties change by introducing interaction between the particles.

Contents

1	Intro	duction	1	2								
2	The	Theory and method										
	2.1	1 Two particle system										
	2.2	particles	4									
	2.3	Releva	nt topics for this project	5								
		2.3.1	Gradient descent - two parameters	5								
		2.3.2	One-body density	6								
		2.3.3	Virial theorem	6								
		2.3.4	Trap frequency	6								
		2.3.5	Evaluating the error	7								
	2.4	The pr	rogram	7								
		2.4.1	Initialization	7								
		2.4.2	Metropolis steps	7								
		2.4.3	Sampling	8								
	2.5	Improv	ving performance and effiency	8								
		2.5.1	Vectorization	8								
		2.5.2	Parallelizing	8								
		2.5.3	Reducing computational cost	8								
3	Resu	Results and discussion										
	2 1	T	landaria da Arra dimanatana									

		3.1.1	Brute force sampling	9			
		3.1.2	Including importance sampling	10			
		3.1.3	Including optimization	13			
		3.1.4	Including interaction	14			
		3.1.5	One-body density	16			
	3.2	Extend	ling to more particles	18			
	3.3	Six par	ticles	18			
	3.4	Twelve	particles	21			
	3.5	Efficier	псу	23			
	3.6	Perforr	mance analysis	23			
		3.6.1	Vectorization	23			
		3.6.2	Parallellization	23			
4	Conc	clusion		24			
Ар	pendi	ces		25			
Α	A Dealing with the Slater determinant efficiently A.1 Slater determinant						
В	B Energies 27						
С	Hermite polynomials and the wavefunction derivatives 28						
D	More MC cycles 30						

1 Introduction

The main objective of this project is to investigate the many-body problem of simulating a quantum dot. A quantum dot is basically electrons that are trapped in an electrical potential. In this project, the potential is modelled as a harmonic oscillator potential. This assumption is ... (maybe not accurate because ...), but it is an easy potential to start with and the potential can later be expanded and most of the code can be reused as it is. Furthermore, we use a trial wavefunction with one (two) fitting parameters for the case without (with) a Jastrow factor to represent the interaction between the particles. This trial wave function is not the exact wave function, hence the simulation can only approximate the exact values for the ground state energy.

The report starts out by introducing the system with the representative equations and analysis tools. Most of the numerical tools used in the programming in this project has already been described in detail in project 1 [1], but some additional things are explained or more thoroughly elaborated on in the theory and method part of this report.

Furthermore, the results and discussion part first analyses how different important parameters have been chosen, i.e. choise of evaluation of the double derivative, step size of sampling techniques and method and parameters of the optimization. In addition the expectation energies and the one-body densities of the systems involves are compared and evaluated. At last, an evaluation of code's efficiently

is made and some concluding remarks are stated.

2 Theory and method

In this project we investigate a fermionic system of N=2, 6 and 12 electrons, where N is the number of particles. It is a so-called closed shell-system. The Hamiltonian used to model this system is

$$\hat{H} = \sum_{i=1}^{N} \left(-\frac{1}{2} \nabla_i^2 + \frac{1}{2} \omega^2 r_i^2 \right) + \sum_{i < j} \frac{1}{r_{ij}},\tag{1}$$

where

$$\hat{H}_0 = \sum_{i=1}^{N} \left(-\frac{1}{2} \nabla_i^2 + \frac{1}{2} \omega^2 r_i^2 \right)$$

is the single particle part and

$$\hat{H}_1 = \sum_{i < j} \frac{1}{r_{ij}},\tag{2}$$

represent the interaction potential between particles. The Hamiltonian is written in atomic units, which implies that $\hbar=1, m=1$, the unit of length is $a_0={}^{4\pi\epsilon_0\hbar^2}/m_ee^2$ and the unit of energy is $E_h={}^{\hbar^2}/m_ea_0^2$. We also have $r_{ij}=|{\bm r}_i-{\bm r}_j|=\sqrt{(x_i-x_j)^2+(y_i-y_j)^2}$ and ω is the oscillator frequency. Later we will study the dependence of the system on the oscillator frequency.

2.1 Two particle system

The single particle wave function in two dimensions is

$$\phi_{n_x,n_y}(x,y) = AH_{n_x}(\sqrt{\omega}x)H_{n_y}(\sqrt{\omega}y)\exp\left(-\omega(x^2+y^2)/2\right). \tag{3}$$

where the functions $H_{n_x}(\sqrt{\omega}x)$ are Hermite polynomials, while A is a normalization constant. The relevant Hermite polynomials in this project are listed in Appendix C. ω is the trap frequency.

For the lowest-lying state, E_{00} (see Fig. 1), we have $n_x=n_y=0$ and an energy $\epsilon_{n_x,n_y}=\omega(n_x+n_y+1)=\omega$, the total energy of the lowest-lying state is hence 2ω because there is room for two electrons with opposite spins.

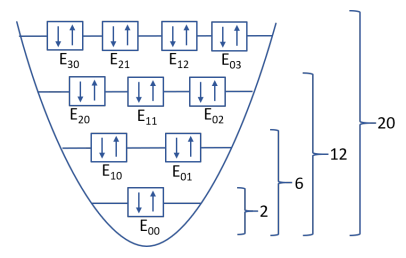


Figure 1: Illustration of the different electron states in a harmonic oscillator trap. The numbers E_{ij} represent the different single-particle states and the states at the same level have the same energy. The arrows show that two electrons with opposite spins can occupy the same state. On the right the full shell systems with the according number of particles are pointed out.

The expectation value can be found by solving the equation

$$\langle E \rangle = \frac{\int d\mathbf{r}_1 d\mathbf{r}_2 \psi_T^*(\mathbf{r}_1, \mathbf{r}_2) \hat{H}(\mathbf{r}_1, \mathbf{r}_2) \psi_T(\mathbf{r}_1, \mathbf{r}_2)}{\int d\mathbf{r}_1 d\mathbf{r}_2 \psi_T^*(\mathbf{r}_1, \mathbf{r}_2) \psi_T(\mathbf{r}_1, \mathbf{r}_2)}.$$
 (4)

We will use Variational Monte Carlo (VMC) methods to evaluate the Eq. 4 [1]. The exact wave function for two not interacting electrons in the ground state is given by

$$\Phi(\mathbf{r}_1, \mathbf{r}_2) = C \exp(-\omega(r_1^2 + r_2^2)/2),$$

where $r_i=\sqrt{x_i^2+y_i^2}$ and C is a normalization constant. The trial wavefunction we use for the not interacting case is

$$\Phi(\mathbf{r}_1, \mathbf{r}_2) = C \exp\left(-\alpha \omega (r_1^2 + r_2^2)/2\right). \tag{5}$$

with the parameter α . From the exact wave function we know that $\alpha=1$ for the situation without interaction. On the other hand, for the interacting case, the trial wave function for the two-electron system is

$$\psi_T(\mathbf{r}_1, \mathbf{r}_2) = C \exp\left(-\alpha \omega (r_1^2 + r_2^2)/2\right) \exp\left(\frac{ar_{12}}{(1 + \beta r_{12})}\right),\tag{6}$$

where we introduce another parameter, β , and a spin factor, a. a is 1 when the two electrons have anti-parallel spins and 1/3 when they have the parallel spins (this is not relevant before we introduce more particles to the system, as can be seen from Fig. 1).

2.2 More particles

Since we are looking at closed shell systems, the next amount of particles are six. We can see this from Fig. 1, there are room for two electrons with opposite spin in two different states, in addition to the two in the lowest lying state. The trial wave function is now given by

$$\psi_T(\boldsymbol{r}_1, \boldsymbol{r}_2, \dots, \boldsymbol{r}_6) = Det(\phi_1(\boldsymbol{r}_1), \phi_2(\boldsymbol{r}_2), \dots, \phi_6(\boldsymbol{r}_6)) \prod_{i < j}^6 \exp\left(\frac{ar_{ij}}{(1 + \beta r_{ij})}\right), \tag{7}$$

where

$$Det(\phi_1(\boldsymbol{r}_1), \phi_2(\boldsymbol{r}_2), \dots, \phi_6(\boldsymbol{r}_6)) = \begin{vmatrix} \phi_1(\boldsymbol{r}_1) & \phi_2(\boldsymbol{r}_1) & \cdots & \phi_6(\boldsymbol{r}_1) \\ \phi_1(\boldsymbol{r}_2) & \phi_2(\boldsymbol{r}_2) & \cdots & \phi_6(\boldsymbol{r}_2) \\ \vdots & \vdots & \ddots & \vdots \\ \phi_1(\boldsymbol{r}_6) & \phi_2(\boldsymbol{r}_6) & \cdots & \phi_6(\boldsymbol{r}_6) \end{vmatrix}$$

is the Slater determinant. This determinant occurs because electron are indistinguishable particles and they are antisymmetric The functions, $\phi_i(r_j)$, are given by Eq. 3 and the notation is explained in Tab. 1.

Table 1: The relation between the notation used in the determinant (left) compared to Eq. 3 (right).

Similarly if we include another shell in our system we get 12 particles and the trial wavefunction is

$$\psi_T(\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_{12}) = Det(\phi_1(\mathbf{r}_1), \phi_2(\mathbf{r}_2), \dots, \phi_{12}(\mathbf{r}_{12})) \prod_{i \le j}^{12} \exp\left(\frac{ar_{ij}}{(1 + \beta r_{ij})}\right).$$
(8)

The determinant have the same structure as for six particles and the relation to the single-particle wave functions are shown in Tab. 1.

2.3 Relevant topics for this project

2.3.1 Gradient descent - two parameters

In this project the gradient descent method also known as steepest descent method was used to evaluate the parameters that makes the trial wavefunction approximate the ground state wavefunction [2]. Compared to Project 1, there are two parameters to be evaluated. The derivative of the trial wavefunction with regards to the parameter α is

$$\frac{\frac{\partial \psi_T}{\partial \alpha}}{\psi_T} = -\frac{\omega}{2} \sum_{i}^{N} r_i^2 \tag{9}$$

and β is

$$\frac{\frac{\partial \psi_T}{\partial \beta}}{\psi_T} = -\sum_{i < j}^N \frac{a_{ij} r_{ij}^2}{(1 + \beta r_{ij})^2}.$$
 (10)

The derivative used in the gradient descent method is

$$\frac{\partial \langle E_L \rangle}{\partial \alpha} = 2 \left(\left\langle \frac{\bar{\psi}_T^{\alpha}}{\psi_T} E_L \right\rangle - \left\langle \frac{\bar{\psi}_T^{\alpha}}{\psi_T} \right\rangle \langle E_L \rangle \right) \tag{11}$$

where $\left\langle \frac{\bar{\psi_T^{\alpha}}}{\psi_T} \right\rangle$ is the expectation value of Eq. 9 and $\left\langle \frac{\bar{\psi_T^{\alpha}}}{\psi_T} E_L \right\rangle$ is the expectation value of Eq. 9 multiplied with the local energy, and the same applies for the parameter β (except using Eq. 10).

2.3.2 One-body density

The radial one-body density is a measure of the spacial distribution of the electrons with respect to the distance from the middle of the harmonic oscillator trap. To calculate the radial one-body density, we want to sample the position of the electrons. The distance from the origin to a set cut-off is separated into bins with a length Δr . For every Monte Carlo step, the distance between the electron's position and the origin is calculated, and the bin that corresponds to the current distance get a count. In the end, you have an array corresponding to the different bins with counts corresponding to how many times an electron was found to have that particular distance to the origin. This array is normalized by dividing by the number of Monte Carlo steps. However, to get the density, we have to divide the number in the bins with the area or volume the bin represents. Because we have two-dimensional problem in this project and we calculate the radial one-body density, we divide bin i with the area $A = \pi(r_i + \Delta r)^2 - \pi r_i^2$ where r_i is distance from the origin to bin i. normalized to the number of particles. Mention. Should happen automatically... Does not happen automatically for me, but I have notice that Even has the same scale on the y-axis.[3]

2.3.3 Virial theorem

The virial theorem gives a relation between the time-average total kinetic energy, $\langle T \rangle$, and the time-average external potential energy, $\langle V_{ext} \rangle$, that is

$$2\langle T \rangle = n \langle V_{ext} \rangle \tag{12}$$

where $V(r) = Br^n$. In our case when we do not include interaction n=2 from the external potential term in Eq. 1, and therefore the average kinetic energy should be equal to the average potential energy. This can be used as a test to see if the simulations executed are correct.

2.3.4 Trap frequency

The trap frequency changes the external potential felt by the electrons. Figure 2 shows how a larger trap frequency, ω , results in a narrower external potential. In this narrow harmonic oscillator trap, the electrons are forced to be closer to each other. Later, we will investigate how ω affects the energy when the electrons are interacting with each other through the potential given in Eq. 2

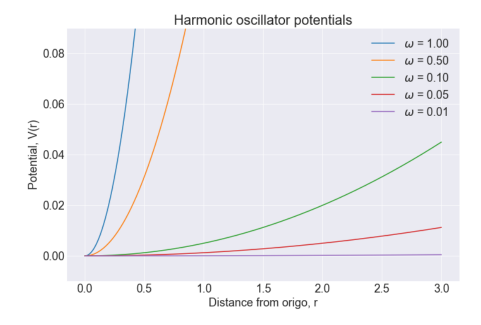


Figure 2: Illustration of how the potential changes when the trap frequency changes.

2.3.5 Evaluating the error

Standard error of the mean (SEM):

$$\mathsf{SEM} = \frac{\sigma}{\sqrt{N}} \tag{13}$$

where N is the number of observations, in our case the number of Monte Carlo cycles.

2.4 The program

A short introduction?

2.4.1 Initialization

First the different parts of the program is initialized based on what Hamiltonian that is used and what wave function is used. In this program, one can choose an Hamiltonian with and without interaction (InteractionHarmonicOscillator and HarmonicOscillator, respectively), and accordingly a wave function with and without a Jastrow factor (SlaterDeterminantInteraction and Slaterdeterminant, respectively). Thereafter, the initial state is set up. The particles are distributed after randomly after a set distribution. Two different distributions are used and chosen based on the sampling method. For importance sampling a Gaussian distribution is used (GaussainDistribution) and for brute force sampling a random uniform distribution is used (RandomUniform). This initialization reflects in the way the sampling of new positions are made by the sampling method, which is explained in project 1 [1].

2.4.2 Metropolis steps

After the initialization the particles are moved one by one, chosen at random, and the new position is found through the sampling method. The metropolis ratio is calculated to determine wether the step is accepted or not. The metropolis ratio is also determined by the sampling method. If the step is

accepted, the energy is sampled and if not, the former energy is sampled again. This is continued until all Monte Carlo (MC) cycles have finished.

2.4.3 Sampling

In this program one can choose to sample all local energies for all MC cycles. This makes it possible to performed resampling techniques on the dataset to improve estimate of the error by estimating the correlation between the different metropolis steps, i.e. the samples made of the local energy. For the other outputs, e.g. kinetic, potential and interaction energy and the mean distance, the average is calculated based on the sum of all the sampled values by dividing by the number of MC cycles.

2.5 Improving performance and efficiency

2.5.1 Vectorization

Solving as vectors instead of scalars - using more of memory - is sort of parallel. Added set (CMAKE CXX FLAGS RELEASE "-O3")

in CMakeLists.

2.5.2 Parallelizing

Doing operations in parallell instead of secquentially. Can use different methods and combine different methods like MPI and openMP. Includes the transfer of information between different processes. In this case the parallelization needed is very simple. Want more MC cycles in less time. In this project I chose to parallelize using a Makefile and cmake. I ran the whole executeable with different seeds for the random number generator and different filenames to save all the local energies in four different threads on my own personal computer which is a quad-core laptop.

2.5.3 Reducing computational cost

Decrease floating point operations (flops) by smarter code and use of ratios:

- Slater determinant split into spin-up determinat and spin down determinant: Only have to update either of them when a spin up/spin down is moved. Detailed information in Appedix A.1.
- Metropolis ratio instead of calculating individual parts and afterwards calculate the ratio, a much simpler expression for the ratio is used and then calculated. Details in Appendix A.2
- Updating the inverse instead of calculating the inverse for every step.
- Another structure in metropolis step (importance sampling):
 - Pick particle at random and find new position
 - Update Slater determinant row (spin up or spin down one based on the particle that was picked)
 - Calculate both old and new quantum force (gradient in a spesific position) using the old inverse Salter determinant.
 - Calculate the metropolis ratio using the new Slater determinant and the old inverse determinant.
 - Update the inverse Slater determinant (only spin up or spin down one based on the particle that was moved) only if the move is accepted.

- Using this structure saves many flops. The inverse update takes e.g. $O(N^2)$ instead of $O(N^3)$.

3 Results and discussion

3.1 Two electrons in two dimensions

We start with the simple case of two electrons in a harmonic oscillator trap. These electrons do not interact with each other and the trial wavefunction is given by Eq. 5.

3.1.1 Brute force sampling

First, brute force sampling was used to calculate the new position and evaluate the metropolis ratio. The double derivative of the wavefunction, used to calculate the kinetic energy part of the expectation energy, was evaluated both analytically and numerically. Table 2 shows the energy for different values for the parameter α . The numbers show that the standard error of the mean (SEM) is underestimating the deviations. From Tab. 2 one can observe that including the correlations, i.e. mainly correlations between one state and the next where only one particle is moved, increases the deviation, giving us σ_b . This value is also an estimate of the error, but probably a more true estimate of the error.

From Tab. 2 one can also observe that $\alpha=1.0$ gives zero standard deviation and is therefore the optimal parameter. By comparing the results for the analytical and the numerical cases one can observe that the SEM and σ_b is similar for both cases, especially around the ground state ($\alpha=1.0$). If he expectation energies from the analytical case and the numerical cases are compared, they differ with values at the scale of 10^{-3} , which is reasonable with a σ_b around 10^{-3} to 10^{-2} . At last one can observe, both from the individual CPU time measurements and the mean CPU time of these 10 measurements (though with different α s), that the analytical solution of the double derivative is much faster and more efficient than the numerical case.

Table 2: Comparing the results for analytical/numerical evaluation of the double derivative. Here $\langle E_L \rangle$ is the expectation value for the energy given in atomic units (a.u.) and CPU time is in units of seconds. σ_B is the standard deviation after resampling with the blocking method and SEM is the standard deviation of the mean. Number of MC cycles are 2^{21} .

Analyt	ical:				Numer	ical:			
α :	$\langle E_L \rangle$:	SEM:	σ_B :	CPU time:	α :	$\langle E_L \rangle$:	SEM:	σ_B :	CPU time:
0.50	2.49402	0.00073	0.01022	5.57812	0.50	2.49991	0.00073	0.01093	18.20310
0.60	2.26441	0.00052	0.00690	5.76562	0.60	2.26412	0.00053	0.00727	18.21880
0.70	2.13118	0.00035	0.00448	5.92188	0.70	2.13039	0.00036	0.00436	18.62500
0.80	2.05016	0.00022	0.00263	5.67188	0.80	2.04993	0.00022	0.00269	18.46880
0.90	2.01015	0.00010	0.00116	5.96875	0.90	2.01160	0.00010	0.00118	18.46880
1.00	2.00000	0.00000	0.00000	5.62500	1.00	2.00000	0.00000	0.00000	18.31250
1.10	2.00871	0.00009	0.00102	6.20312	1.10	2.00825	0.00009	0.00097	18.35940
1.20	2.03402	0.00018	0.00175	6.34375	1.20	2.03308	0.00018	0.00170	18.31250
1.30	2.07259	0.00026	0.00244	5.95312	1.30	2.06460	0.00026	0.00243	20.23440
1.40	2.11041	0.00034	0.00311	6.15625	1.40	2.11803	0.00033	0.00308	19.00000
Mean	CPU time:	5.91875			Mean	CPU time:	18.62033		

3.1.2 Including importance sampling

Figure 3 compare the expectation value of the energy and the acceptance rate of brute force sampling and importance sampling. It can be observed from the right part of the figure that the acceptance rate of both methods increase with decreasing step size, but one can also observe that the acceptance is lower for importance sampling than brute force sampling at large step sizes. These observations could indicate that a small step size would be ideal for both methods.

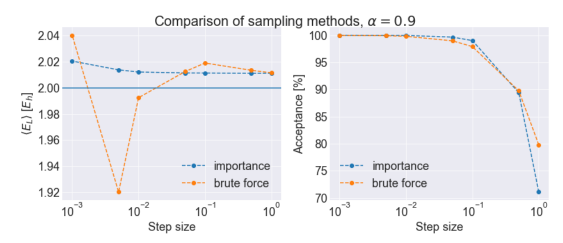


Figure 3: Left: Expectation energies after 2^{21} MC cycles for different step sizes. Right: Percentage of accepted steps for different step sizes. Here importance sampling and brute force sampling is compared.

From the left part of the figure it can be observed that one of the expectation values for the energies are lower than the ground state energy (dl = 0.005 with brute force sampling) when these calculations where done with $\alpha = 0.9$. However, in Tab. 3 which compare the result of brute force sampling and

importance sampling for different step sizes one can observe that the standard deviation from the blocking method is larger for the case of brute force sampling with a step size of 0.005. However, the SEM does not indicate anything to be special about this result.

Table 3: Comparing the results for importance/brute force sampling. Here the parameter α is set to 0.9 and number of MC cycles are 2^{21} . Acc. is short for acceptance and is here given in % and t_{CPU} is the CPU time used by the program in units of seconds. The rest of the values are as described in Tab. 2.

	Brute fo	rce:				Importa	nce:			
dl:	$\langle E_L \rangle$:	SEM:	σ_B :	Acc.:	t_{CPU} :	$\langle E_L \rangle$:	SEM:	σ_B :	Acc.:	t_{CPU} :
1.000	2.010	0.00010	0.00066	79.832	6.625	2.011	0.00010	0.00022	71.078	8.531
0.500	2.011	0.00010	0.00112	89.794	7.266	2.011	0.00010	0.00023	89.343	8.672
0.100	2.015	0.00011	0.00549	97.919	7.078	2.011	0.00010	0.00047	99.049	8.859
0.050	2.007	0.00011	0.00946	99.001	7.016	2.011	0.00010	0.00068	99.662	8.188
0.010	2.002	0.00007	0.01510	99.788	6.953	2.007	0.00010	0.00136	99.968	8.281
0.005	1.942	0.00006	0.02140	99.916	6.578	2.010	0.00010	0.00199	99.989	8.547
0.001	2.118	0.00002	0.00371	99.974	6.531	2.007	0.00010	0.00410	99.999	8.016
	Mean C	PU time: (5.86384			Mean C	PU time: 8	8.44197		

I took a closer look at the actual local energies for the brute force sampling method. Figure 4 shows how the energy is not stable for steps sizes smaller than 0.01, so even though the step sizes 0.001 and 0.01 seems to give reasonable expectation values for the energy (see Tab. 3 and Fig. 3), Fig. 4 seems to show that that is sort of a lucky shot. I also saw this by running the calculation with brute force sampling and the step size, 0.005, with different seeds for the random number generator. The expectation energy for five different runs where $\langle E_L \rangle = 1.91487, 2.03452, 1.90805, 1.88356$ and 1.9284. From Fig. 4 one can observe that even dl=0.1 seems to be too small since it also results in the local energy varying slowly and taking longer "trips" to higher energies and using many steps to get back down again, but for this step size the "trips" to higher energies are more frequent than for the smaller step sizes. I concluded that a step size of 0.5 is the best choice for the brute force sampling because it gives reasonable changes of the local energy and an acceptance rate of \sim 90 % (see Fig. 3).

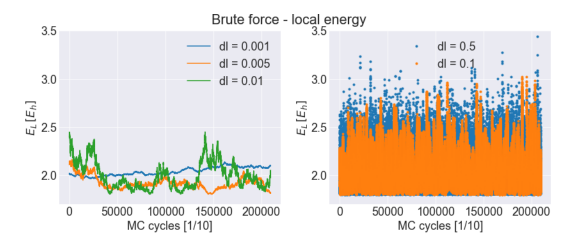


Figure 4: The local energy for every tenth MC cycle for brute force sampling at different step sizes, dl. a) shows the smaller step sizes and b) some that are a bit larger.

Proceeding to evaluate the energy, Table 4 shows how the energy changes with different trap frequencies, ω . From Tab. 4 one can observe that the mean distance is increasing with decreasing trap frequency. This is as expected from Fig. 2, where the potential is broadened with decreasing trap frequency and hence is not forcing the particles closer together. The mean distance is a bit different for brute force sampling compared to importance sampling, but the similarity might improve if more MC cycles are used. The other value show, however, no large difference between the results from brute force sampling and importance sampling. But to be able to compare the sampling methods more thoroughly it is better to look at the case where the system is not in the ground state.

Table 4: Ground state energy of two electrons in a harmonic oscillator trap. Here \bar{r}_{12} is the mean distance between the two electrons at positions r_1 and r_2 given in units of a_0 and $\langle E_L \rangle$, $\langle T \rangle$, $\langle V_{ext} \rangle$ and $\langle V_{int} \rangle$ are the expectation value of the local energy, kinetic energy, potential energy and interaction energy, respectively, given in units of E_h . The rest of the values are as explained in Tab. 2. Number of MC cycles are 2^{23} .

	Brut	te force	:			Imp	ortance	:		
ω	α	$\langle E_L \rangle$	\overline{r}_{12}	$\langle T \rangle$	$\langle V_{ext} \rangle$	α	$\langle E_L \rangle$	\overline{r}_{12}	$\langle T \rangle$	$\langle V_{ext} \rangle$
1.00	1	2.00	1.250	1.0008	0.9992	1	2.00	1.254	0.9982	1.0018
0.50	1	1.00	1.775	0.4971	0.5029	1	1.00	1.781	0.4973	0.5027
0.10	1	0.20	3.967	0.0996	0.1004	1	0.20	4.046	0.0987	0.1013
0.05	1	0.10	5.638	0.0497	0.0503	1	0.10	5.534	0.0512	0.0488
0.01	1	0.02	12.631	0.0099	0.0101	1	0.02	12.488	0.0101	0.0099

Virial theorem! + energy is given by $\hbar\omega$, so energy change according to $\hbar\omega$.

Table 5 shows the expectation value for the energy for various parameters, alpha. The calculated expectation values for the energy for the two different sampling methods are similar, especially close to the correct parameter α , varying only by $\pm 10^{-2}$. The SEM is here underestimating the error compared to σ_b from the blocking resampling technique for both sampling methods. However, what is different is that σ_b is larger for importance sampling than for brute force sampling. Why is that? Is it just estimating the error more correctly (but the blocking code is the same), or is there a larger error resulting from importance sampling? At last, one can observe that the CPU time of the the

importance sampling method is larger than brute force sampling which is expected because of the calculation of the gradient and the quantum force. But we know from Fig. 3 that the acceptance rate of brute force sampling is around 90 % compared to importance sampling which should be close to 100 % and this makes the importance sampling technique more effective in terms of MC cycles.

Table 5: Comparing the results for brute force sampling/importance sampling. Values are as explained in Tab. 2. Number of MC cycles are 2^{21} .

	Brute forc	e:			Importance:			
α :	$\langle E_L \rangle$:	SEM:	σ_B :	CPU time:	$\langle E_L \rangle$:	SEM:	σ_B :	CPU time:
0.50	2.49074	0.00072	0.01030	6.26562	2.52950	0.00075	0.01952	8.53125
0.60	2.27645	0.00053	0.00721	6.53125	2.26531	0.00052	0.01281	8.62500
0.70	2.12481	0.00036	0.00451	6.59375	2.12449	0.00036	0.00813	8.76562
0.80	2.04873	0.00022	0.00265	6.68750	2.04797	0.00022	0.00454	9.37500
0.90	2.01135	0.00010	0.00115	6.82812	2.01171	0.00010	0.00203	8.81250
1.00	2.00000	0.00000	0.00000	6.75000	2.00000	0.00000	0.00000	8.76562
1.10	2.00863	0.00009	0.00097	6.59375	2.00954	0.00009	0.00171	8.40625
1.20	2.03343	0.00018	0.00174	7.29688	2.03276	0.00018	0.00305	8.50000
1.30	2.07103	0.00026	0.00246	6.75000	2.07363	0.00026	0.00440	8.67188
1.40	2.11725	0.00033	0.00317	6.70312	2.10697	0.00034	0.00536	8.67188
	Mean CPl	J time: 6.7	7000		Mean CPU	J time: 8.7	7125	

3.1.3 Including optimization

Because there are two parameters to optimize, I chose, in this project compared to the previous one, to experiment with the minimization rate during the actual optimization. I started out using a minimization rate, $\gamma=0.5$. It resulted in the fewest steps until the parameter value stabilized both for guesses close to the optimal value and for guesses far away from the optimal value, but for the smallest trap frequencies I had to use $\gamma=0.1$ or 0.2. For the case of two interacting fermions, the parameters were optimized by trying out different first guesses for α and β and tuning γ so that the parameters stabilized during the first 200 iterations. The optimal parameters were extracted from the mean of the last 50 iterations. An example run is shown in Fig. 5 for $\omega=0.5$.

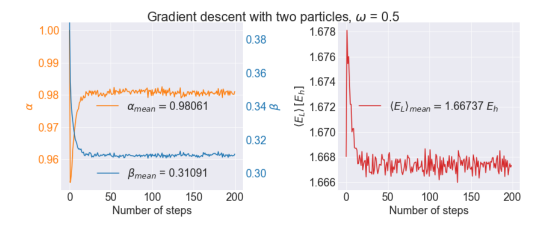


Figure 5: Left: The development of the parameters during the steps of gradient descent. The values α_{mean} and β_{mean} printed on the figure is the mean of the last 50 values. Right: The expectation value after 2^{19} number of MC cycles. Here also, the value printed on the figure is the mean of the last 50 iterations.

For the system with six interacting fermions, the method described above was used for the largest ω s (i.e. $\omega=1.0,\ 0.5,\ 0.1$), but for the smaller ones I had to use a smaller minimization rate (i.e. 0.01-0.05) and I also had to move step by step from $\omega=0.1$ to $\omega=0.01$ with the step size of $\Delta\omega=0.01$. I found the parameters for the current ω and used that as a guess for the next ω . I tried to do it in a more efficient way and let the simple gradient descent method find the minimum on its own, but with this unguided method, the optimization ended up in local minima at higher energies or going to infinite energies. To improve the code, I attempted to use the extended gradient descent method, described in project 1, which utilize the previous gradient to find the new parameter [1]. But the attempt did not improve the behaviour described earlier. Compared to project 1, this systems local energy dependence on the parameters ($\langle E_L \rangle (\alpha, \beta)$), is more complicated, and also involve two parameters instead of one, which makes it harder to optimize the parameters with this simple gradient descent method.

Is the value calculated from the mean of 50 2^{19} with slightly different α s and β s, which values are oscillating around the optimal value, better than one calculation with the mean values of α and β for 2^{21} number of MC cycles? Maybe if I also could calculate the standard deviation and also save the local energies so that I get σ_b ?

3.1.4 Including interaction

Here is the results of the expectation values approximating the ground state energy for two interacting electrons in a harmonic oscillator trap. The equations used to model the system is as described in the theory part of this report. That includes the trial wavefunction (see Eq. 6) with the Jastrow factor that is used to model the many-body wavefunction. Table 6 and 7 show the results of the calculations with the optimal parameters found with the gradient descent method for brute force and importance sampling, respectively.

For two-dimensional dots, he found the energy to be E=3 for the frequency w=1 and E=2/3 with w=1/6 as the frequency. [4]

Table 6: Ground state energy of two interacting electrons in harmonic oscillator trap found with brute force sampling. Here \bar{r}_{12} , $\langle T \rangle$, $\langle V_{ext} \rangle$ and $\langle V_{int} \rangle$ are as explain in Tab. 4. The rest of the values are as explained in Tab. 2 . Number of MC cycles are 2^{23} .

ω	α	β	$\langle E_L \rangle$	SEM	σ_B	\overline{r}_{12}	$\langle T \rangle$	$\langle V_{ext} \rangle$	$\langle V_{int} \rangle$
1.00	0.98847	0.39965	3.0068	0.00001	0.00009	1.636	0.8944	1.2990	0.8135
0.50	0.98061	0.31091	1.6674	0.00001	0.00010	2.481	0.4488	0.7051	0.5135
0.10	0.94693	0.17764	0.4486	0.00001	0.00011	6.695	0.1003	0.1767	0.1716
0.05	0.92747	0.13815	0.2609	0.00000	0.00011	10.389	0.0533	0.0997	0.1076
0.01	0.88398	0.07287	0.0777	0.00000	0.00006	29.177	0.0129	0.0284	0.0364

Table 7: Ground state energy of two interacting electrons in harmonic oscillator trap found with importance sampling. Here \bar{r}_{12} , $\langle T \rangle$, $\langle V_{ext} \rangle$ and $\langle V_{int} \rangle$ as explain in Tab. 4. The rest of the values are as explained in Tab. 2. Number of MC cycles are 2^{23}

ω	α	β	$\langle E_L \rangle$	SEM	σ_B	\overline{r}_{12}	$\langle T \rangle$	$\langle V_{ext} \rangle$	$\langle V_{int} \rangle$
1.00	0.98846	0.39954	3.0069	0.00001	0.00018	1.643	0.8931	1.3052	0.8086
0.50	0.98082	0.31068	1.6674	0.00001	0.00019	2.481	0.4547	0.6997	0.5130
0.10	0.94734	0.17810	0.4486	0.00001	0.00023	6.724	0.0989	0.1787	0.1710
0.05	0.92262	0.14090	0.2610	0.00000	0.00021	10.333	0.0495	0.1024	0.1091
0.01	0.88305	0.07366	0.0776	0.00000	0.00010	29.293	0.0131	0.0283	0.0362

From Tab. 6 and 7 one can observe that the two sampling methods give approximately the same results. The different expectation values for the different energies are very similar and the mean distance is also very similar. However, the standard deviation from the blocking resampling method are different. Importance sampling seems to result in larger σ_B here as well as for the case without interaction (see Tab. 5).

Have to compare with Tout's work. Say that it should be 3. Can maybe also compare that other value for another omega.

The results above show that the energy of the system is decreasing with decreasing trap frequency, ω . One can also observe that the potential energy dominates for large trap frequencies ($\omega=1.0$ and $\omega=0.5$), but for the smaller ω s the interaction energy and the potential energy is approximately equal. For all ω , the kinetic energy is the smallest one. Does this make sense? What is the potential energy? The energy stored in the force from the trap - keeping the electrons together - stronger trapping force - stronger potential energy. What is kinetic energy? Momentum? Why does it decrease with trap frequency? Because potential energy is less? Without interaction - energy oscillates between potential energy and kinetic energy - energy is conserved.

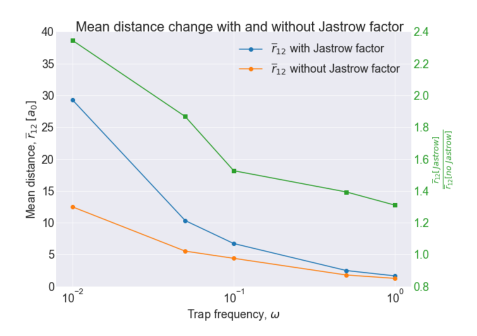


Figure 6: The mean distance between the two particles, calculated with importance sampling, compared for the situation with and without the Jastrow factor and at different trap frequencies.

Figure 6 shows the combined results from Tab. 4 and 7. One can observe that the mean distance is larger for the case with interaction. This is expected since the interaction potential (see Eq. 2) is a repulsive potential, and will hence force the electrons further apart. One can also observe that the ratio of the two different cases increase with decreasing trap frequency, i.e. the mean distance increases more for smaller trap frequencies. is that as expected? Why? Does that mean that the trap frequency is more important than the interaction?

3.1.5 One-body density

The one-body density can tell how the particles are distributed in space, and Fig. 7 shows the result of the one-body density of the system of two not interacting fermions for different trap frequencies. One can observe, as one saw in the development of the mean distance (see Fig. 6), that the particle distribution or density is spread out in space for smaller trap frequencies. From Fig. 7 one can also observe that the electrons spend most of their time around the origin of the harmonic oscillator trap. The shape of the curve seems not to change for the different trap frequencies, only the height and width.

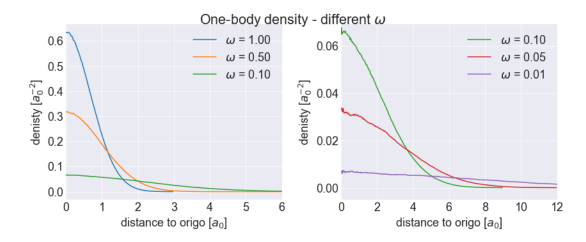


Figure 7: One-body density of the two electron system without the Jastrow factor, but with different trap frequencies. Left: The larger trap frequencies. Right: The smaller trap frequencies. The system with $\omega=0.1$ is in both plots to show the difference in scale for the two plots. Used 2^{24} MC cycles.

To get a smooth curve, I found out that you either have to use larger bin sizes or more MC cycles. Since the one-body density is found by counting how many times a particle is situated within a certain bin, i.e. with a certain distance to origin, more samples of the positions will give a more true distribution. So if the bin size is small, you need more MC cycles to smooth out the random difference between the two bins. On the other hand, if the bin size is too large real differences or fast changes in the distribution might not be detected.

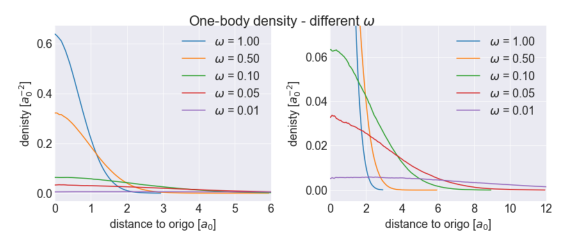


Figure 8: One-body density of the two electron system without the Jastrow factor, but with different trap frequencies. Left: The larger trap frequencies. Right: The smaller trap frequencies. The system with $\omega=0.1$ is in both plots to show the difference in scale for the two plots. Here showed for a case with larger bins then Fig. 7. Used 2^{24} MC cycles.

Furthermore, Fig. 9 shows the one-body densities with and without the Jastrow factor. Here, one can also observe the same as was seen with the mean distance, that the Jastrow factor spreads out the distribution of the particles, but the difference is a lot clearer in the one-body density plot. Another interesting observation is that, for the interacting case, the maximum does not seem to lay at the origin. It is not that easy to see for the larger trap frequencies, but one can easily see it for $\omega=0.1$ in the right part of Fig. 9. This seems to be reasonable because there is a repulsive force between the particles, they spend more time further away from the origin. Could it be that for larger trap frequencies there is a larger energy-wise gain (i.e. the energy is lower) to be situated closer to origin, hence the maximum is not shifted as far from origin for those frequencies, but for the lower trap

frequencies the gain obtained closer to origin does not compensate enough of the repulsive forces (or interaction energy), so the maximum is shifted away from origin. That should imply that the position of the maximum reflects a balance between the potential energy and the interaction energy.

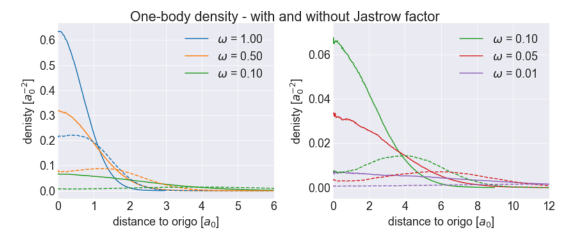


Figure 9: One-body density of the two electron system with and without the Jastrow factor and with different trap frequencies (indicated by the color of the lines). Dashed lines are with the Jastrow factor and filled lines are without Jastrow factor. Left: The larger trap frequencies. Right: The smaller trap frequencies. The system with $\omega=0.1$ is in both plots to show the difference in scale for the two plots. Used 2^{24} MC cycles.

3.2 Extending to more particles

To investigate the quantum dot system further, the next shell (illustrated in Fig. 1) is filled which results in a system of six particles. Furthermore, the third shell is filled resulting in a system of twelve particles.

3.3 Six particles

The expectation value for the total energy (i.e. the local energy), the kinetic energy and the potential energy for the different trap frequencies and without interaction are listed in Tab. 8. One can observe that for all the trap frequencies, the kinetic and potential energy seems to follow the virial theorem (see Eq. 12), if one allows for a small difference of maximum ± 0.03 a.u.. The energies are as expected from the calculated exact energies in Tab. 15 in Appendix B.

Table 8: Ground state energy of six electrons in a harmonic oscillator trap. Here $\langle E_L \rangle$, $\langle T \rangle$ and $\langle V_{ext} \rangle$ are as explain in Tab. 4. Number of MC cycles are 2^{23} .

ω	$\langle E_L \rangle$	$\langle T \rangle$	$\langle V_{ext} \rangle$
1.00	10.00	4.9865	5.0135
0.50	5.00	2.4973	2.5027
0.10	1.00	0.4861	0.5139
0.05	0.50	0.2483	0.2517
0.01	0.10	0.0454	0.0546

Furthermore, the result after including interaction is showed in Tab. 9. One can observe, if one compares the result with Tab. 7, that the both the SEM and σ_b are larger for the system with six

electrons. This could be explained by the fact that the total energies are larger for this system, and hence the errors will be larger. If one compare the error in as a fraction for the case with $\omega=1.0$ $\sigma_b/\langle E_L\rangle=5.99\cdot 10^{-5}$ for two electrons and $3.98\cdot 10^{-4}$ for six electrons and one can see that the error is larger for the six electron case as a fraction as well. Would the blocking error be worse if the optimal parameters where worse? I guess not because it sort of says how good the expectation value is for the parameters given. It tells if the energy that comes out can be relied upon. It does not tell us how far we are from the real energy.

Table 9: Ground state energy of six interacting electrons in harmonic oscillator trap found with importance sampling. Number of MC cycles are 2^{23} .

ω	α	β	$\langle E_L \rangle$	SEM	σ_B	$\langle T \rangle$	$\langle V_{ext} \rangle$	$\langle V_{int} \rangle$
1.00	0.71567	0.49372	20.4492	0.00022	0.00813	2.3429	10.7076	7.3988
0.50	0.75823	0.34260	11.9868	0.00011	0.00522	1.3226	5.8094	4.8548
0.10	0.78852	0.15041	3.6542	0.00003	0.00416	0.2951	1.7035	1.6556
0.05	0.76518	0.10733	2.2223	0.00003	0.00436	0.1178	1.0882	1.0162
0.01	0.64907	0.05085	0.7191	0.00001	0.00586	0.0021	0.3803	0.3367

Compare with Even's values?

I wanted to take a closer look at the optimized parameters for the two different systems too, especially since I had so much trouble finding the parameters for the six electron system. Figure 10 compares the optimized parameters for the two systems with different amounts of particles at the different trap frequencies. One can observe that the β parameter which is involved in the Jastrow factor is similar for both systems and also show a similar development from one trap frequency to the next. This is expected, I guess, since the interaction between the particles do not change that much by increasing the number of particles. The other parameter α , however, which is involved in the Slater determinant, shows completely different values and development for the two systems. This is also reasonable because the wave function, i.e. the Slater determinant part, changes a lot when we increase the number of particles. Because when the number of particles is increased higher states are filled and the Slater determinant is changed a lot. The development of α though is strange, I would have expected it to either increase or decrease with an increasing trap frequency, but In the right part of Fig. 10 one can observe that the largest α is found for $\omega=0.1$ which is the middle ω investigated. But my expectation might be wrong, maybe the strange development is due to the single-particle states that are introduced with the higher energy states.

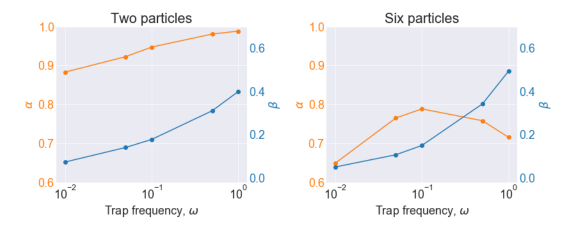


Figure 10: Left: The optimized parameters for the system with two particles and with the different trap frequencies. Right: The optimized parameters for for the system with six particles and with the different trap frequencies. All found with the simple gradient descent method.

Next, Fig. 11 compares the energy per particle of the two systems when interaction is included. The left part of the figure shows that the magnitude of kinetic energy per particle and the development of the kinetic energy with trap frequency is very similar for both systems (two and six particles). The right part of the figure shows that both the potential and interaction energy per particle is larger for the six particle system at the same trap frequency, than the two particle system. This is as expected since the trap frequency determine the "size" of the trap. The six particle system's potential energy is hence larger for the same size system when there are more particles involved. The interaction energy is also larger when more particles, that are repulsed by each other, are forced together in the "same amount of space" (it is not exactly the same amount of space since the space is not strictly limited, but it is intuitive to image it like that). Apart from this, the six particle system show the same behaviour as the two particle system, energy wise.

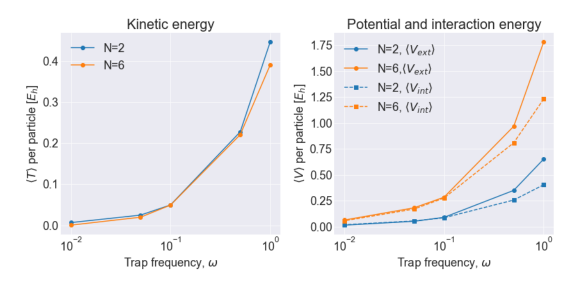


Figure 11: Left: The expectation value for the kinetic energy per particle for the two different systems examined at the different trap frequencies. Right: The expectation value of the potential and interaction energy per particle for the two different systems at the different trap frequencies.

Proceeding to the one-body density, Fig. 12 shows the one-body density for the six electron system. First, one can notice that the shape is different from the two electron system. Here the maximum

denisty is not at the origin of the trap, but approximately a_0 away from the origin for $\omega=1.0$ without the Jastrow factor. Furthermore, one can observe that the maximum moves further and further away from the origin when ω increases. Regarding the inclusion of interaction, the same behavior is seen here, compared to the two electron one-body denisty, that including interaction broadens the whole curve and lowers the maximum density. The shape of the one-body density curve seems to be similar with and without interaction except for $\omega=0.1$ and maybe also smaller ω where there is another maximum closer to the origin. Maybe the broadening of the trap with decreasing ω opens up for a redistribution of the particles when the repulsive interaction is included.

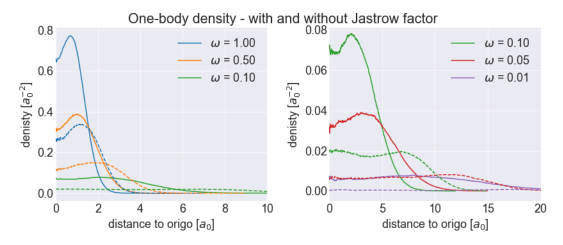


Figure 12: One-body density of the six electron system with and without the Jastrow factor and with different trap frequencies (indicated by the color of the lines). Dashed lines are with the Jastrow factor and filled lines are without Jastrow factor. Left: The larger trap frequencies. Right: The smaller trap frequencies. The system with $\omega=0.1$ is in both plots to show the difference in scale for the two plots. Used 2^{24} MC cycles.

3.4 Twelve particles

By including another shell in the two dimensional harmonic oscillator trap, the system consists of twelve particles. The ground state energy at different ω s are listed in Tab. 10 and the energies are the same as the exact energies calculate based on the different energy states (see Tab. 15 in Appendix B).

Table 10: Ground state energy of twelve electrons in a harmonic oscillator trap. Here $\langle E_L \rangle$, $\langle T \rangle$ and $\langle V_{ext} \rangle$ are as explain in Tab. 4. Number of MC cycles are 2^{23} .

ω	$\langle E_L \rangle$	$\langle T \rangle$	$\langle V_{ext} \rangle$
1.00	28.00	14.0117	13.9883
0.50	14.00	7.0463	6.9537
0.10	2.80	1.4084	1.3916
0.05	1.40	0.6901	0.7099
0.01	0.28	0.1419	0.1381

I tried to use the same method, the simple gradient descent method, to evaluate the parameters that gives me the closest thing to the ground state I can get with the trial wavefunction used in this project, but it seems like the system have gotten too complicated with twelve particles. An example of the behaviour of the gradient descent method is showed in Fig. 13. One can observe, from the

right part of the figure that the energy seems to be converging towards a minimum, but it is a very slow process. However, the left part of the figure shows how much the parameters are varying and that the parameter α is negative. I do not think that a negative α is a good sign especially after comparing my results with Ref. [3] where $\langle E_L \rangle = 65.7026(4)$ was found for the same system.

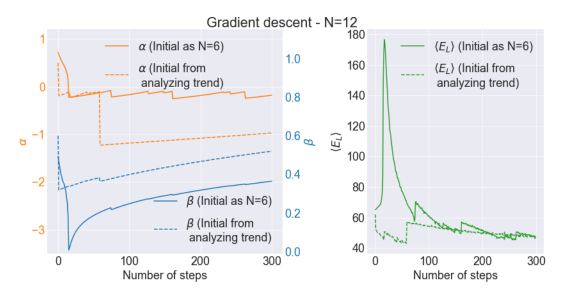


Figure 13: Two examples of the gradient descent method for twelve particles with different initial guesses for the two parameters. One of them is a guess based on the parameters found for the six electron system and the other is a guess based on the difference and trend found when the two electron and the six electron parameters was compared. Left: The development of the parameters. Right: The corresponding expectatino value for the energy.

Continuing to the one-body density, Fig. 14 shows the one-body denisty for the twelve electron system, but only without interaction since a ground state for the interaction case was not found. Compared to the two smaller systems, this system shows another curve for the one-body density. Here, the maximum is once angain at the origin of the trap, but there is also another local maximum approximately $1.5a_0$ away from the origin. The scaling of the density seems to be similar to the former two systems where one see that the distribution broadens and the maximum is hence lowered, but the shape remains similar when ω is decreased.

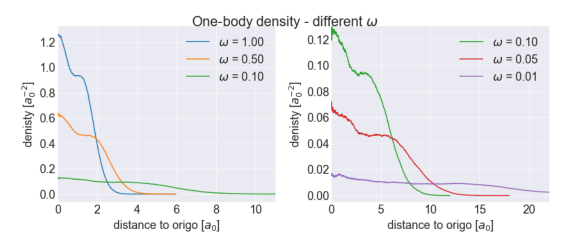


Figure 14: One-body density of the twelve electron system without the Jastrow factor, but with different trap frequencies. Left: The larger trap frequencies. Right: The smaller trap frequencies. The system with $\omega=0.1$ is in both plots to show the difference in scale for the two plots. Used 2^{24} MC cycles.

3.5 Efficiency

In this project I wanted to write an efficient code, but I ran into some problems and after a while I figured out that it was best to use the unefficeint, but working code to get some results and use time to analyse them. Therefore, I have two branches in my repository at github. One called no_optimization and one master branch. The master branch includes my attempts to make the code more efficient as described in the theory part of this report, and the other branch is the code used to generate the results in this proejct.

3.6 Performance analysis

At last, I include a small analysis of the performance of the code with and without vectorization and with and without parallelizing.

3.6.1 Vectorization

Table 11 lists the average CPU time of ten runs of the code with and without vectorization. The mean was found four times for both cases at different times. There is not a clear difference in CPU time, and one can also see from the four differnt attempts that the CPU time changes based on the computer. I think one can conclude that the program is just as efficient with and without the vectorization. But this is just comapring the CPU time. As I have understood, vectorization can also help exploit the memory in a more efficient way.

Table 11: The average CPU time in seconds after 10 runs. The calculation is of the six electron system with interaction and involves importance sampling, saving and printing all local energies to file. Here I used 2^{22} MC cycles.

	CPU time	
Run	With vectorization	Without vectorization
1	145.930	148.531
2	149.653	142.881
3	155.731	153.519
4	146.769	146.834

3.6.2 Parallellization

In this project I chose to parallelize using a Makefile and cmake. I ran the whole executeable with different seeds for the random number generator and different filenames to save all the local energies in four different threads on my own personal computer which is a quad-core laptop. The CPU times for the different threads for five different runs are listed in Tab. 12. The maximum CPU time of the four different threads is counted as the total CPU time. From the different runs, one can observe that the CPU time can vary based on the laptop and other processes on it, and it shows that it is important to time several runs to get a proper comparison of the CPU time of different methods.

Table 12: Four threads 2^{22} MC cycles on each.

Thread	1	2	3	4	Total:
CPU time	206.520	207.370	209.260	210.400	210.400
	228.270	229.930	229.930	230.110	230.110
	231.880	234.690	235.030	235.730	235.730
	231.330	234.060	234.170	234.300	234.300
	230.200	231.280	231.770	232.070	232.070
				Mean:	228.522

Table 13 compare the CPU time of parallelized runs and the not parallelized runs. One can see clearly that the parallelized runs are faster, but even though the parellelized run is done on four threads it is only between two and three times faster. One reason might be that the number of MC cycles used to reach an equilibrium is the same $(1\cdot 10^5)$ for both cases. For the single thread case it is done only once and afterwards all the local energeis are sampled, but for the four thread case this is done independently on all four threads. At last I compared the expectation value, the SEM and σ_B for one of the runs with and without parallelizing. The results are listed in Tab. 14 and shows a difference in the expectation value of the energy at approximatly ± 0.01 which is larger than the calculated σ_B . One would expect a difference because of the different seeds in the random number generator, but maybe not that large. This might improve with a larger number of MC cycles though.

Table 13: Not parallelized: 2^{24} MC cycles and one thread. Parallelized: 2^{22} MC cycles and four threads. The CPU times are given in seconds and as an average over five runs.

	Not parallelized	Parallelized	Times faster
CPU time	656.210	228.522	2.87
	602.816	257.922	2.34

Table 14: A comparison of the expectation value, the SEM and σ_B for one of the runs with and without parallelizing. Units are as described in Tab. 4.

	$\langle E_L \rangle$	SEM	σ_B
Parallelized			
Not parallelized	20.4572	0.00016	0.00629

4 Conclusion

 Better optmization technique - maybe conjugate gradient - took a long time and work + could not really find the parameters for the N=12 case. Maybe, if I had enough time, I could have found the parameters giving a correct minimum in the energy, but after trying out a lot of different starting guesses for the parameters, different number of MC cycles for each step and different minimization rates without finding a minimum in the energy I figured it was too difficult with the simple gradient descent method.

- Not efficient code new how to do it, but I could not do it. Did not get the correct energies (below ground state energy at omega not 1 - no interaction).
- Not efficent to which to Eigen notation for matrix in inverse matrix function, should have whiched all code to eigen, but I used the time I had left for analysis etc instead

•

Appendices

A Dealing with the Slater determinant efficiently

Determining a determinant numerically is a costly operation, so we want to do some alteration to increase the efficiency of the code.

A.1 Slater determinant

The Slater determinant is contain the single-particle wave function of the number of particles included in the system evaluated at the posistion for all particles included because electrons are indistinguishable. The determinant is written as

$$D = Det\left(\phi_1(\boldsymbol{r}_1), \phi_2(\boldsymbol{r}_2), \dots, \phi_N(\boldsymbol{r}_N)\right) = \begin{vmatrix} \phi_1(\boldsymbol{r}_1) & \phi_2(\boldsymbol{r}_1) & \cdots & \phi_N(\boldsymbol{r}_1) \\ \phi_1(\boldsymbol{r}_2) & \phi_2(\boldsymbol{r}_2) & \cdots & \phi_N(\boldsymbol{r}_2) \\ \vdots & \vdots & \ddots & \vdots \\ \phi_1(\boldsymbol{r}_N) & \phi_2(\boldsymbol{r}_N) & \cdots & \phi_N(\boldsymbol{r}_N). \end{vmatrix}$$

Hence the rows represent different positions, r_i , and the coloumns represent different states. To simplify the calcualtions we want to have all states with spin up in one determinant and all states with spin down in another determinant. For six particles we then get

$$D = D_{\uparrow}D_{\downarrow} = egin{array}{c|ccc} \phi_1(m{r}_1) & \phi_3(m{r}_1) & \phi_5(m{r}_1) \ \phi_1(m{r}_2) & \phi_3(m{r}_2) & \phi_5(m{r}_2) \ \phi_1(m{r}_3) & \phi_3(m{r}_3) & \phi_5(m{r}_3) \ \end{array} egin{array}{c|cccc} \phi_2(m{r}_4) & \phi_4(m{r}_4) & \phi_6(m{r}_4) \ \phi_2(m{r}_5) & \phi_4(m{r}_5) & \phi_6(m{r}_5) \ \phi_2(m{r}_6) & \phi_4(m{r}_6) & \phi_6(m{r}_6) \ \end{array} .$$

We see this from Tab. 1 and Eq. 3. Mister anti-symmertrien, men expectation value er lik.

The trial wavefunction can therefore be rewritten to

$$\psi_T = D_{\uparrow} D_{\downarrow} \psi_C$$

where ψ_C is the correlation part of the trial wavefunction. Now we only have to update one of these matrises when we move a particle, depending on which spin the particle has.

A.2 The Metropolis ratio

In the metropolis test we calculate the ratio bewteen the wavefunction before and after a proposed move, but now the wavefunction includes a determinant which is costly to calculate. We therefore want to utilize some relations from linear algebra to simplify the ratio and make the algorithm more efficient. The ratio between the Slater determinant part of the wavefunction, ψ_{SD} , is

$$R = \frac{\psi_{SD}(\boldsymbol{r}^{new})}{\psi_{SD}(\boldsymbol{r}^{old})} = \frac{\sum_{i}^{N} d_{ij}(\boldsymbol{r}^{new}) C_{ij}(\boldsymbol{r}^{new})}{\sum_{i}^{N} d_{ij}(\boldsymbol{r}^{old}) C_{ij}(\boldsymbol{r}^{old})}.$$
(14)

where $d_{ij} = \psi_i(j)$

Here we have used the fact that when you calculate a determinant, you break it down into a sum of smaller determinants times a factor:

$$D = \begin{vmatrix} d_{11} & d_{12} & \cdots & d_{1N} \\ d_{21} & d_{22} & \cdots & d_{2N} \\ \vdots & \vdots & \ddots & \vdots \\ d_{N1} & d_{N2} & \cdots & d_{NN} \end{vmatrix} = \sum_{i}^{N} d_{ij} C_{ij}.$$

So if $dij = d_{11}$ then

$$C_{11} = \begin{vmatrix} d_{22} & d_{23} & \cdots & d_{2N} \\ d_{32} & d_{33} & \cdots & d_{3N} \\ \vdots & \vdots & \ddots & \vdots \\ d_{N2} & d_{N3} & \cdots & d_{NN} \end{vmatrix}.$$

We observe in Eq. 14 that if we move particle j from r_j^{old} to r_j^{new} the matrix C_{ij} is unchanged, we have only changed the d_{ij} in the original determinant D that is not included in C_{ij} . Equation 14 is then

$$R = \frac{\sum_{i}^{N} d_{ij}(\boldsymbol{r}^{new})}{\sum_{i}^{N} d_{ij}(\boldsymbol{r}^{old})}$$
(15)

We can simplify this even further with the relation

$$\sum_{k=1}^{N} d_{ik} d_{kj}^{-1} = \delta_{ij} = \begin{cases} 0 & \text{if } i \neq j \\ 1 & \text{if } i = j \end{cases}$$
 (16)

The ratio can be rewritten as

$$R = \frac{\sum_{i}^{N} d_{ij}(\boldsymbol{r}^{new}) d_{ij}(\boldsymbol{r}^{old})^{-1}}{\sum_{i}^{N} d_{ij}(\boldsymbol{r}^{old}) d_{ij}(\boldsymbol{r}^{old})^{-1}} = \sum_{i}^{N} d_{ij}(\boldsymbol{r}^{new}) d_{ij}(\boldsymbol{r}^{old})^{-1}.$$
(17)

The consequence of these calculations are that we now only have to calculate the invers values of the determinant once to know the values for $d_{ij}(\mathbf{r}^{old})^{-1}$ and then update only the row of the position that was changed in the Slater determinant and calculate the invers of the determinant again only if the move is accepted.

A.3 Updating the inverse of the Slater determinant

After a move is accepted in the Metropolis test, the row in the Slater determinant representing that particle is updated, but the inverse of the Slater determinant also needs to be updated because the Slater determinant has changed. This could be done by simply calculating the inverse of the determinant, but this is costly and there is a more efficient way. The elements of the determinant d_{kj}^{-1} (hva betyr den $^{-1}$? At den skal opphøyes i minus 1 eller er det notasjon på at det er et element i den inverse matrisen?) can be found through

$$d_{kj}^{-1}(\boldsymbol{r}^{new}) = \begin{cases} d_{kj}^{-1}(\boldsymbol{r}^{old}) - \frac{d_{kj}^{-1}(\boldsymbol{r}^{old})}{R} \sum_{l=1}^{N} d_{il}^{-1}(\boldsymbol{r}^{new}) d_{lj}^{-1}(\boldsymbol{r}^{old}) & \text{if } i \neq j \\ \frac{d_{kj}^{-1}(\boldsymbol{r}^{old})}{R} \sum_{l=1}^{N} d_{il}^{-1}(\boldsymbol{r}^{old}) d_{lj}^{-1}(\boldsymbol{r}^{old}) & \text{if } i = j \end{cases},$$

where i is the number of the row representing the particle that was moved.

B Energies

$$E_{n_x n_y} = \hbar \omega (n_x + n_y + \frac{d}{2}) \tag{18}$$

where d is the number of dimensions. In this project d=2.

Table 15: The exact energies for the non-interacting case with different number of particles in a closed shell system.

Energies	
E_{00}	$\hbar\omega$
$E_{10} = E_{01}$	$2\hbar\omega$
$E_{20} = E_{02} = E_{11}$	$3\hbar\omega$
$E_{30} = E_{03} = E_{21} = E_{12}$	$4\hbar\omega$
$E_{N=2} = 2E_{00}$	$2\hbar\omega$
$E_{N=6} = E_{N=2} + 2E_{10} + 2E_{01}$	$10\hbar\omega$
$E_{N=12} = E_{N=6} + 2E_{20} + 2E_{02} + 2E_{11}$	$28\hbar\omega$
$E_{N=20} = E_{N=12} + 2E_{30} + 2E_{03} + 2E_{21} + 2E_{12}$	$60\hbar\omega$

C Hermite polynomials and the wavefunction derivatives

The relevant Hermite polynomials

$$H_0(\sqrt{\omega}x)$$
 1
 $H_1(\sqrt{\omega}x)$ 2 $\sqrt{\omega}x$
 $H_2(\sqrt{\omega}x)$ 4 $\omega x^2 - 2$
 $H_3(\sqrt{\omega}x)$ 8 $\omega \sqrt{\omega}x^3 - 12\sqrt{\omega}x$

$$\phi_{n_x,n_y}(x,y) = AH_{n_x}(\sqrt{\omega}x)H_{n_y}(\sqrt{\omega}y)\exp\left(-\omega(x^2+y^2)/2\right).$$

Table 16: $\psi_{n_x n_y}$

Trial wavefunctions for the different states

ψ_{00}	$A \exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{01}	$2\sqrt{\omega}xA\exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{10}	$2\sqrt{\omega}yA\exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{20}	$(4\omega x^2 - 2)A \exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{02}	$(4\omega y^2 - 2)A \exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{11}	$4\omega xyA\exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{30}	$\left((8\omega\sqrt{\omega}x^3 - 12\sqrt{\omega}x)A\exp\left(\frac{-\alpha\omega r^2}{2}\right) \right)$
ψ_{03}	$\left(8\omega\sqrt{\omega}y^3 - 12\sqrt{\omega}y\right)A\exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{21}	$\left(8\omega\sqrt{\omega}x^2y - 4\sqrt{\omega}y\right)A\exp\left(\frac{-\alpha\omega r^2}{2}\right)$
ψ_{12}	$\left (8\omega\sqrt{\omega}xy^2 - 4\sqrt{\omega}x)A\exp\left(\frac{-\alpha\omega r^2}{2}\right) \right $

Table 17: $\psi_{n_x n_y}$

The derivative of the trial wavefunctions for the different states

$$\nabla \psi_{00} \begin{vmatrix} (-\alpha \omega x, -\alpha \omega y)A \exp\left(\frac{-\alpha \omega r^2}{2}\right) \\ \nabla \psi_{01} \end{vmatrix} = \frac{(-\alpha \omega x, -\alpha \omega y)A \exp\left(\frac{-\alpha \omega r^2}{2}\right)}{-(\sqrt{\omega}(a\omega x^2-1), \alpha \omega^{3/2}xy)2A \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{10} \end{vmatrix} = \frac{(-\alpha \omega^2 x^2, \sqrt{\omega}(a\omega y^2-1))2A \exp\left(\frac{-\alpha \omega r^2}{2}\right)}{-(2\alpha \omega^2 x^3 - \alpha \omega x - 4\omega x, 2\alpha \omega^2 x^2y - \alpha \omega y)2A \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{02} \end{vmatrix} = \frac{(-2\alpha \omega^2 xy^2 - \alpha \omega x, 2\alpha \omega^2 y^3 - \alpha \omega y - 4\omega y)2A \exp\left(\frac{-\alpha \omega r^2}{2}\right)}{-(4\omega y(\alpha \omega x^2-1), -4\omega x(\alpha \omega y^2-1))A \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{11} \end{vmatrix} = \frac{(-4\omega y(\alpha \omega x^2-1), -4\omega x(\alpha \omega y^2-1))A \exp\left(\frac{-\alpha \omega r^2}{2}\right)}{-(4\sqrt{\omega}(2\alpha \omega^2 x^4 - 3(\alpha + 2)\omega x^2 + 3), -4\alpha \omega^{3/2}xy(2\omega x^2 - 3))A \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{03} \end{vmatrix} = \frac{(-4\sqrt{\omega}(-4\alpha \omega^{3/2}xy(2\omega y^2 - 3), 2\alpha \omega^2 y^4 - 3(\alpha + 2)\omega y^2 + 3))A \exp\left(\frac{-\alpha \omega r^2}{2}\right)}{-(4\sqrt{\omega}(\alpha \omega x^2(2\omega xy - 1) - 4\omega xy + 1), -4\omega^{3/2}x(2x(\alpha \omega y^2 - 1) - \alpha y))A \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{12} \end{vmatrix} = \frac{(-4\omega^{3/2}y(2y(\alpha \omega x^2 - 1) - \alpha x), -4\sqrt{\omega}(\alpha \omega y^2(2\omega xy - 1) - 4\omega xy + 1)}{-(2\omega x^2 x^2 + 3)} + \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{12} \end{vmatrix} = \frac{(-4\omega^{3/2}y(2y(\alpha \omega x^2 - 1) - \alpha x), -4\sqrt{\omega}(\alpha \omega y^2(2\omega xy - 1) - 4\omega xy + 1)}{-(2\omega x^2 x^2 + 3)} + \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{12} \end{vmatrix} = \frac{(-4\omega^{3/2}y(2y(\alpha \omega x^2 - 1) - \alpha x), -4\sqrt{\omega}(\alpha \omega y^2(2\omega xy - 1) - 4\omega xy + 1)}{-(2\omega x^2 x^2 + 3)} + \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{13} \end{vmatrix} = \frac{(-4\omega^{3/2}y(2y(\alpha \omega x^2 - 1) - \alpha x), -4\sqrt{\omega}(\alpha \omega y^2(2\omega xy - 1) - 4\omega xy + 1)}{-(2\omega x^2 x^2 + 3)} + \exp\left(\frac{-\alpha \omega r^2}{2}\right)} \\ \nabla \psi_{13} \end{vmatrix} = \frac{(-4\omega^{3/2}y(2y(\alpha \omega x^2 - 1) - \alpha x), -4\sqrt{\omega}(\alpha \omega y^2(2\omega xy - 1) - 4\omega xy + 1)}{-(2\omega x^2 x^2 + 3)} + \exp\left(\frac{-\alpha \omega r^2}{2}\right)}$$

Table 18: $\psi_{n_x n_y}$

The double derivative of the trial wavefunctions for the different states

$\nabla^2 \psi_{00}$	$(\alpha^2 \omega^2 r^2 - \alpha \omega) A \exp$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$\nabla^2 \psi_{01}$	$2\alpha\omega^{3/2}x(\alpha\omega r^2-4)A\exp$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$\nabla^2 \psi_{10}$	$2\alpha\omega^{3/2}y(\alpha\omega r^2-4)A\exp{-i\omega r^2}$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$ abla^2 \psi_{20}$	$2\omega(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 1)r^2 + \alpha(2\omega x^2 - 12\omega x^2) + 4))A\exp(\alpha^2\omega(2\omega x^2 - 12\omega x^2) + 4)$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$ abla^2 \psi_{02}$	$2\omega(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2 - 12\omega y^2) + 4))A\exp(\alpha^2\omega(2\omega y^2 - 1)r^2 + \alpha(2\omega y$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$\nabla^2 \psi_{11}$	$4\alpha\omega^2 xy(\alpha\omega r^2 - 6)A\exp$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$ abla^2 \psi_{30}$	$4\omega^{3/2}x(\alpha^2\omega(2\omega x^2 - 3)r^2 - 4\alpha(4\omega x^2 - 3) + 12)A\exp(-\frac{\pi}{2})$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$ abla^2 \psi_{03}$	$4\omega^{3/2}y(\alpha^2\omega(2\omega y^2 - 3)r^2 - 4\alpha(4\omega y^2 - 3) + 12)A\exp(-2\omega y^2 - 3)r^2 + 4\alpha(4\omega y^2 - 3) + 12A\exp(-2\omega y^2 - 3)r^2 + 4\alpha(4\omega y^2 -$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$ abla^2 \psi_{21}$	$4\omega^{3/2}(\alpha^2\omega xr^2(2\omega xy - 1) + 4\alpha x(1 - 4\omega xy) + 4y)A\exp(-2\omega xy - 1)$	$\left(\frac{-\alpha\omega r^2}{2}\right)$
$\nabla^2 \psi_{12}$	$4\omega^{3/2}(\alpha^2\omega yr^2(2\omega xy-1)+4\alpha y(1-4\omega xy)+4x)A\exp(-4\omega xy)$	$\left(\frac{-\alpha\omega r^2}{2}\right)$

D More MC cycles

Table 19: Ground state energy of two interacting electrons in harmonic oscillator trap found with importance sampling. Here \overline{r}_{12} , $\langle T \rangle$, $\langle V_{ext} \rangle$ and $\langle V_{int} \rangle$ as explain in Tab. 4. The rest of the values are as explained in Tab. 2. Number of MC cycles are 2^{24}

ω	α	β	$\langle E_L \rangle$	SEM	σ_B	\overline{r}_{12}	$\langle T \rangle$	$\langle V_{ext} \rangle$	$\langle V_{int} \rangle$
1.00	0.98846	0.39954	3.0068	0.00001	0.00013	1.631	0.9002	1.2907	0.8160
0.50	0.98082	0.31068	1.6674	0.00001	0.00014	2.471	0.4511	0.7003	0.5160
0.10	0.94734	0.17810	0.4483	0.00000	0.00015	6.744	0.0985	0.1788	0.1709
0.05	0.92262	0.14090	0.2612	0.00000	0.00015	10.369	0.0536	0.1003	0.1073
0.01	0.88305	0.07366	0.0775	0.00000	0.00008	28.997	0.0131	0.0277	0.0367

Table 20: Ground state energy of six interacting electrons in harmonic oscillator trap found with importance sampling. Number of MC cycles are 2^{24} .

ω	α	β	$\langle E_L \rangle$	SEM	σ_B	$\langle T \rangle$	$\langle V_{ext} \rangle$	$\langle V_{int} \rangle$
1.00	0.71567	0.49372	20.4580	0.00016	0.00605	2.4011	10.6658	7.3911
0.50	0.75823	0.34260	11.9850	0.00008	0.00430	1.3360	5.7919	4.8572
0.10	0.78852	0.15041	3.6447	0.00003	0.00438	0.2837	1.7094	1.6517
0.05	0.76518	0.10733	2.2214	0.00002	0.00529	0.1324	1.0619	1.0272
0.01	0.64907	0.05085	0.7078	0.00001	0.00427	-0.0211	0.3998	0.3290

References

- [1] V. M. Reinertsen. Project 1, 2020.
- [2] Wikipedia contributors. Gradient descent, Jun 2020.
- [3] E. M. Nordhagen. Studies of quantum dots using machine learning. Master's thesis, University of Oslo, 2019.
- [4] M. Taut. Two electrons in an external oscillator potential: Particular analytic solutions of a coulomb correlation problem. *Physical Review A*, 48(5):3561, 1993.