# Variational Monte Carlo studies of Atoms

# by HÅVARD SANDSDALEN

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Department of Physics
University of Oslo

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# Contents

Ι	Th	neory	7
1	Intr	roduction	9
2	Qua	antum Physics	11
	2.1	Quantum Mechanics in one dimension	11
		2.1.1 Probability and statistics	11
		2.1.2 The time-independent Schrödinger Equation	13
	2.2	Bra-Ket notation	15
	2.3	Quantum Mechanics in three dimensions	15
		2.3.1 Separation of variables - quantum numbers $l$ and $m$	16
		2.3.2 The Angular equation	17
		2.3.3 The Radial equation and solution for the hydrogen atom	18
3	Ma	ny-particle Systems	23
	3.1	Atoms	23
		3.1.1 Two-particle systems	23
		3.1.2 Wave functions for N-particle atoms	25
		3.1.3 Electron configuration	26
		3.1.4 Hamiltonian and scaling	27
	3.2	Hartree-Fock Theory	28
		3.2.1 Hartree-Fock equations	31
4	Qua	antum Monte Carlo	37
	4.1	Markov chains	38
	4.2	Random numbers	39
	4.3	The Variational Principle	40
	4.4	Variational Monte Carlo	41
		4.4.1 VMC and the simple Metropolis algorithm	42
		4.4.2 Metropolis-Hastings algorithm and importance sampling	44
	4.5	Blocking	48
5	Wa	ve Functions	51
	5.1	Cusp conditions and the Jastrow factor	51
		5.1.1 Single particle cusp conditions	51
		5.1.2 Correlation cusp conditions	53
	5.2	Rewriting the Slater determinant	53

## CONTENTS

	5.3 5.4	Variational Monte Carlo wave function	55 55
		5.4.1 S-orbitals	55
		5.4.2 P-orbitals	57
	5.5	Roothaan-Hartree-Fock with Slater-type orbitals	60 62
II	In	nplementation and results	65
6	Imp	plementation of the Variational Monte Carlo Method	67
	6.1	1 0	68
		6.1.1 Optimizing the ratio - $\Psi_T(r^{new})/\Psi_T(r^{old})$	69
	6.2	6.1.2 Derivative ratios	74 80
7	Har	tree-Fock implementation	83
	7.1	Interaction matrix elements	83
	7.2	Algorithm and HF results	84
8	VM	IC Results	87
	8.1	Validation runs	87
		8.1.1 Hydrogen-like wave functions	87
	0.0	8.1.2 Slater-type orbitals	88
	8.2	Variational plots	89
		8.2.1 Hydrogenic single particle wave functions	89
	8.3	8.2.2 Roothaan-Hartree-Fock wave functions	90
	0.0	Optimal parameters with DFP	90 90
		8.3.1 Hydrogenic wave-functions	90
		8.3.3 Neon	93
	8.4	Time-step interpolation	93
	8.5	Magnesium	93
	8.6	Comparison with experimental data	94
$\mathbf{A}$	Roc	othaan-Hartree-Fock results	95
	A.1	Helium	95
	A.2	Beryllium	95
	A.3		96
	A.4	Magnesium	96
	A.5	Silicon	96
В		tistics	99
	B.1	Correlated measurements	100
$\mathbf{C}$	DFI	P and energy minimization	105

Bibliography 108

Part I

Theory

# Chapter 1 Introduction

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# Chapter 2

# Quantum Physics

In the 1920's european physicists developed quantum mechanics in order to describe the physical phenomena they had been discovering for some years. Such phenomena as the photoelectric effect and the diffraction patterns from the double-slit experiment indicated that physicists needed a new set of tools when handling systems on a very small scale, e.g. the behavior of single particles and isolated atoms.

This chapter will give an introduction to the relevant topics in quantum physics needed to describe the atomic systems in this project. Some parts will closely follow the discussions in the books [1] and [2], while other parts only contain elements from the sources listed in the bibliography.

# 2.1 Quantum Mechanics in one dimension

The general idea and goal of quantum mechanics is to solve the complex, time-dependent Schrödinger-equation (S.E.) for a specific physical system which cannot be described by classical mechanics. Once solved, the S.E. will give you the quantum mechanical wave function,  $\Psi$ , a mathematical function which contains all information needed about such a non-classical system. It introduces probabilities and statistical concepts which contradicts with the deterministic nature of a classical system. The full one-dimensional S.E. for an arbitrary potential, V, reads:

$$i\hbar \frac{\partial \Psi}{\partial t} = -\frac{\hbar^2}{2m} \frac{\partial^2 \Psi}{\partial x^2} + V\Psi. \tag{2.1}$$

The Schrödinger equation is the classical analogy to Newton's second law in classical physics, and describes the dynamics of virtually any physical system, but is only useable in small scale systems, quantum systems.

# 2.1.1 Probability and statistics

The wave function,  $\Psi$ , is now a function of both position, x, and time, t. Quantum mechanics uses the concept of probability and statistics via  $\Psi$ , and these solutions of the S.E. may be complex as the equation Eq. (2.1) is complex itself. To comply

with the statistical interpretation we must have a function that is both real and non-negative. As described in [1], Born's statistical interpretation takes care of this problem by introducing the complex conjugate of the wave function. The product  $\Psi^*\Psi = |\Psi|^2$  is interpreted as the probability density for the system state. If the system e.g. consists of only one particle, the integral

$$\int_{a}^{b} |\Psi(x,t)|^{2} dx \tag{2.2}$$

is then interpreted as the probability of finding the particle between positions a and b at an instance t. To further have a correct correspondence with probability, we need the total probability of finding the particle anywhere in the universe to be 1. That is

$$\int_{-\infty}^{\infty} |\Psi(x,t)|^2 dx = 1.$$
(2.3)

In quantum mechanics, operators represent the observables we wish to find, given a wave function,  $\Psi$ . The operator representing the position variable,  $\hat{\mathbf{x}}$ , is just x itself, while the momentum operator is  $\hat{\mathbf{p}} = -i\hbar(\partial/\partial x)$ . All classical dynamic variables are expressed in terms of just momentum and position, as said on p. 17 in [1]. Another important operator is the Hamilton operator. The Hamilton operator gives the time evolution of the system and is the sum of the kinetic energy operator  $\hat{\mathbf{T}}$  and the potential energy operator  $\hat{\mathbf{V}}$ ,  $\hat{\mathbf{H}} = \hat{\mathbf{T}} + \hat{\mathbf{V}}$ .  $\hat{\mathbf{V}}$  is represented by the function V from Eq. (2.1), while the kinetic energy operator is

$$\widehat{\mathbf{T}} = \frac{\widehat{\mathbf{p}}^2}{2m} = \frac{(i\hbar)^2}{2m} \left(\frac{\partial}{\partial x}\right)^2 = -\frac{\hbar^2}{2m} \frac{\partial^2}{\partial x^2}$$
 (2.4)

For an arbitrary operator,  $\hat{\mathbf{Q}}$ , the expectation values,  $\langle Q \rangle$ , is found by the formula

$$\langle Q \rangle = \int \Psi^* \widehat{\mathbf{Q}} \Psi \mathrm{d}x. \tag{2.5}$$

The same goes for expectation values of higher moments, e.g.  $\langle p^2 \rangle$ :

$$\langle p^2 \rangle = \int \Psi^* \hat{\mathbf{p}}^2 \Psi dx. \tag{2.6}$$

The so-called variance of an operator or observable,  $\sigma_Q^2$ , can be calculated by the following formula:

$$\sigma_Q^2 = \langle Q^2 \rangle - \langle Q \rangle^2. \tag{2.7}$$

This quantity determines the standard deviation,  $\sigma_Q = \sqrt{\sigma_Q^2}$ . The standard deviation describes the spread around the expectation value. The smaller the standard deviation, the smaller the variation between the possible values of Q.

The wave functions,  $\Psi$ , exist in the so-called *Hilbert space*, a mathematical vector space of square-integrable functions.

#### 2.1.2 The time-independent Schrödinger Equation

However, in any cases, we are only interested in a time-independent version of Eq. (2.1). A crucial point is then to demand the potential, V, to be a time-independent potential as well, viz. V = V(x) (in one spatial dimension). This equation can be obtained by the well-known method of **separation of variables**. The trick is to write our wave function,  $\Psi(x,t)$ , as a product of a purely spatial function,  $\psi(x)$ , and another function,  $\phi(t)$ , depending only of time. That is, we assume:  $\Psi(x,t) = \psi(x)\phi(t)$ . By inserting  $\Psi$  into the full S.E., Eq. (2.1), remembering that  $\psi$  and  $\phi$  only depend on one variable each, then dividing by  $\Psi = \psi \phi$ , we get

$$i\hbar \frac{1}{\phi} \frac{\mathrm{d}\phi}{\mathrm{d}t} = -\frac{\hbar^2}{2m} \frac{1}{\psi} \frac{\mathrm{d}^2\psi}{\mathrm{d}x^2} + V(x). \tag{2.8}$$

By inspecting Eq. (2.8), we see that the left side of the equation depends on t alone, while the right side depends only on x. This means that both sides must be **constant** and of course equal. By varying t and changing the left side, the right side would change without ever varying x, which says it must a constant. We call this separation constant E, giving us the two equations

$$\frac{\mathrm{d}\phi}{\mathrm{d}t} = -\frac{iE}{\hbar}\phi,\tag{2.9}$$

and

$$-\frac{\hbar^2}{2m}\frac{\mathrm{d}^2\psi}{\mathrm{d}x^2} + V\psi = E\psi. \tag{2.10}$$

Equation (2.9) can be solved quite easily, and will give an exponential form of the time-dependent part.

$$\phi(t) = e^{-iEt/\hbar}. (2.11)$$

The second equation, Eq. (2.10), is called the **time-independent Schrödinger** equation. By inspecting Eq. (2.11), we see that expectation values will be constant in time because the time-dependent part from  $\Psi$ ,  $\phi(t)$ , will only give a factor 1 when it is multiplied with its complex conjugate from  $\Psi^*$ . That is:

$$e^{-iEt/\hbar}e^{+iEt/\hbar} = 1. (2.12)$$

The expectation values depend solely on the spatial parts,  $\phi$ . We call these separable solutions **stationary states**.

#### Stationary states and expectation values

Another point about the stationary solutions is the close relation to classical mechanics. The Hamilton function determines the total energy of the system, and is the sum of the kinetic and potential energy. The classical Hamiltonian function for any system with a time-independent potential is

$$H(x,p) = \frac{p^2}{2m} + V(x). \tag{2.13}$$

By using the canonical substitution  $p \to (\hbar/i)(\mathrm{d}/\mathrm{d}x)$  for the quantum mechanical momentum operator, we get

$$\widehat{\mathbf{H}} = -\frac{\hbar^2}{2m} \frac{\partial^2}{\partial x^2} + V(x). \tag{2.14}$$

This is identical to the time-independent Schrödinger equation, Eq. (2.10), and we can then obtain a much simplified Schrödinger equation:

$$\widehat{\mathbf{H}}\psi = E\psi. \tag{2.15}$$

The expectation value of the Hamilton operator, the total energy, is now given as

$$\langle H \rangle = \int \psi^* \widehat{\mathbf{H}} \psi dx = E \int |\psi|^2 dx = E \int |\Psi|^2 dx = E.$$
 (2.16)

Calculating  $\langle H^2 \rangle$  gives

$$\langle H^2 \rangle = \int \psi^* \widehat{\mathbf{H}}^2 \psi dx = E \int \psi^* \widehat{\mathbf{H}} \psi dx = E^2 \int |\psi|^2 dx = E^2 \int |\Psi|^2 dx = E^2.$$
(2.17)

This means the variance and standard deviation of  $\hat{\mathbf{H}}$  are both zero

$$\sigma_H^2 = \langle H^2 \rangle - \langle H \rangle^2 = E^2 - E^2 = 0.$$
 (2.18)

For these separable solutions, every measurement of the total energy will return the same value, E. Thus the spread around the expectation value is exactly zero.

#### General solutions

The general solution for these systems is a linear combination of different separable solutions. Different solutions with different separation constants, e.g.

$$\Psi_1(x,t) = \psi_1(x)e^{-iE_1t/\hbar}, \quad \Psi_2(x,t) = \psi_2(x)e^{-iE_2t/\hbar}$$
 (2.19)

which both are solutions of Eq. (2.10), can be used to construct the general solution

$$\Psi(x,t) = \sum_{n=1}^{\infty} c_n \psi_n(x) e^{-iE_n t/\hbar} = \sum_{n=1}^{\infty} c_n \Psi_n(x,t),$$
 (2.20)

where the factors  $c_n$  are probability weights for its corresponding stationary state.

#### Spin

An important, but difficult consept in quantum physics is spin. Spin is an intrinsic property of every elementary particle. Also composite systems will have a certain value of spin when imposing addition rules on the single particles that make up such a system. In this project we will deal with fermions, i.e. half integer spin particles (see section 3.1.1).

A particle will either have *spin up* or *spin down*. This is denoted by tacking on a factor to the wave function itself, namely the factors

$$\chi_{+} = \uparrow \tag{2.21}$$

and

$$\chi_{-} = \downarrow . \tag{2.22}$$

These spin states,  $\chi_{\pm}$ , are mutually orthogonal, but exist in another Hilbert space than the spatial wave functions,  $\Psi$ , and will not interfere with the integration  $\int dx$ .

This presentation of spin is very short due to the fact that we don't need much information about spin in this thesis. As shown later, the Hamiltonian will not depend on spin values, so the spin states  $\chi_{\pm}$  will only be used as a label to indicate which states are occupied or not. This will become more apparent in the discussion of many particle systems.

### 2.2 Bra-Ket notation

As we will be dealing a lot with expectiation values, which are integrals, it can be smart to introduce a compact way to describe the wave functions and the integrals. The so-called **ket**,  $|\Psi\rangle$ , represents our  $\Psi$ , while the **bra**,  $\langle\Psi|$ , represents the  $\Psi^*$ . The product

$$\langle \Psi | \, \widehat{\mathbf{H}} \, | \Psi \rangle$$
 (2.23)

is defined as

$$\langle \Psi | \, \widehat{\mathbf{H}} \, | \Psi \rangle = \int \Psi^* \widehat{\mathbf{Q}} \Psi \mathrm{d}x,$$
 (2.24)

where x now represents all spatial dimensions and quantum numbers. Now we can write the expectiation values of  $\widehat{\mathbf{H}}$  as

$$\langle H \rangle = \langle \Psi | \, \widehat{\mathbf{H}} \, | \Psi \rangle \tag{2.25}$$

# 2.3 Quantum Mechanics in three dimensions

While discussing quantum mechanics in one dimension is useful for getting in the basics, most real life systems occur in three dimensions. This section closely follows the discussion in [1].

In three dimensions the one-dimensional Hamiltonian,  $H(x, p_x) = p_x^2/2m_e + V(x)$  is replaced by

$$H(x, y, z, p_x, p_y, p_z) = \frac{1}{2m_e} \left( p_x^2 + p_y^2 + p_z^2 \right) + V(x, y, z).$$
 (2.26)

with  $m_e$  being the electron mass. For quantum mechanical systems the momentum operators are substituted by

$$p_x \to \frac{\hbar}{i} \frac{\partial}{\partial x}, \ p_y \to \frac{\hbar}{i} \frac{\partial}{\partial y}, \ p_z \to \frac{\hbar}{i} \frac{\partial}{\partial z}.$$
 (2.27)

We could also write this more compact in vector form

$$\mathbf{p} = \frac{\hbar}{i} \nabla. \tag{2.28}$$

Introducing the **Laplacian** in Cartesian coordinates,  $\nabla^2 = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2}$ , we can write the full Hamiltonian as

$$i\hbar \frac{\partial \Psi}{\partial t} = -\frac{\hbar^2}{2m_e} \nabla^2 \Psi + V \Psi. \tag{2.29}$$

The normalization integral in three dimensions changes using the infinitesimal volume element  $d^3\mathbf{r} = dx \, dy \, dz$ . We now have

$$\int |\Psi(x, y, z, t)|^2 dx dy dz = \int |\Psi(\mathbf{r}, t)|^2 d^3 \mathbf{r} = 1.$$
 (2.30)

The general solutions in three dimensions can be expressed as

$$\Psi(\mathbf{r},t) = \sum c_n \psi_n(\mathbf{r}) e^{-iE_n t/\hbar}.$$
 (2.31)

The spatial wave functions,  $\psi_n$ , satisfy the time-independent Schrödinger equation:

$$-\frac{\hbar^2}{2m_e}\nabla^2\psi + V\psi = E\psi. \tag{2.32}$$

# 2.3.1 Separation of variables - quantum numbers l and m

For a central symmetrical potential where the function V only depends on the distance,  $V = V(|\mathbf{r}|) = V(r)$ , it is common to introduce **spherical coordinates**,  $(r, \theta, \varphi)$  (see figure 2.1), and try the approach of separation of variables. That means solutions of the form

$$\psi(r,\theta,\varphi) = R(r)Y(\theta,\varphi). \tag{2.33}$$

In spherical coordinates the time-independent Schrödinger equation, Eq. (2.32), is

$$-\frac{\hbar^2}{2m_e} \left[ \frac{1}{r^2} \frac{\partial}{\partial r} \left( r^2 \frac{\partial \psi}{\partial r} \right) + \frac{1}{r^2 \sin \theta} \frac{\partial}{\partial \theta} \left( \sin \theta \frac{\partial \psi}{\partial \theta} \right) + \frac{1}{r^2 \sin^2 \theta} \left( \frac{\partial^2 \psi}{\partial \varphi^2} \right) \right] + V\psi = E\psi.$$
(2.34)

By performing the same exercise as in section 2.1.2, calling the separation constant l(l+1), this will give rise to the radial and angular equations for a single particle in a three dimensional central symmetrical potential:

$$\frac{1}{R}\frac{d}{dr}\left(r^{2}\frac{dR}{dr}\right) - \frac{2m_{e}r^{2}}{\hbar^{2}}\left[V(r) - E\right] = l(l+1); \tag{2.35}$$

$$\frac{1}{Y} \left[ \frac{1}{\sin \theta} \frac{\partial}{\partial \theta} \left( \sin \theta \frac{\partial Y}{\partial \theta} \right) + \frac{1}{\sin^2 \theta} \frac{\partial^2 Y}{\partial \varphi^2} \right] = -l(l+1). \tag{2.36}$$

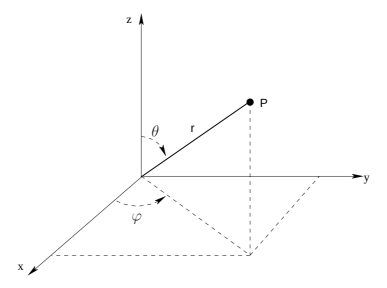


Figure 2.1: Visualizing the spherical coordinates of some point P; radius r, polar angle  $\theta$  and azimuthal angle  $\varphi$ .

#### 2.3.2 The Angular equation

If we multiply Eq. (2.36) by  $Y \sin^2 \theta$ , again using separation of variables, with  $Y(\theta, \varphi) = T(\theta)F(\varphi)$ , and divide by Y = TF, we get

$$\frac{1}{T} \left[ \sin \theta \frac{d}{d\theta} \left( \sin \theta \frac{dT}{d\theta} \right) \right] + l(l+1) \sin^2 \theta = -\frac{1}{F} \frac{d^2 F}{d\varphi^2}. \tag{2.37}$$

This time we call the separation constant  $m^2$ .

#### The $\varphi$ equation

The equation for  $\varphi$  is calculated quite easily, with

$$\frac{d^2F}{d\varphi^2} = -m^2F \Rightarrow F(\varphi) = e^{im\varphi} \tag{2.38}$$

letting the value m take both positive and negative values. Since the angle  $\varphi$  represents a direction in space, we must require that

$$F(\varphi + 2\pi) = F(\varphi). \tag{2.39}$$

It then follows that

$$e^{im(\varphi+2\pi)} = e^{im\varphi} \rightarrow e^{2\pi im} = 1. \tag{2.40}$$

For this to be fulfilled, m has to be an integer. Viz.,  $m = 0, \pm 1, \pm 2, \ldots$ 

#### The $\theta$ equation

The differential equation for the polar angle,  $\theta$ , reads

$$\sin\theta \frac{d}{d\theta} \left( \sin\theta \frac{dT}{d\theta} \right) + [l(l+1)\sin^2\theta - m^2]T = 0. \tag{2.41}$$

This equation is more difficult to solve by standard mathematics. In short, the solution is given by

$$T(\theta) = AP_I^m(\cos \theta), \tag{2.42}$$

where A is a constant, and  $P_l^m$  is the associated Legendre function

$$P_l^m(x) = (1 - x^2)^{|m|/2} \left(\frac{d}{dx}\right)^{|m|} P_l(x). \tag{2.43}$$

 $P_l(x)$  is the *l*th degree **Legendre polynomial**, defined by the so-called **Rodrigues** formula:

$$P_l(x) = \frac{1}{2^l l!} \left(\frac{d}{dx}\right)^l (x^2 - 1)^l.$$
 (2.44)

We see that l must be a positive integer for the differentiations to make sense, and furthermore, |m| must be smaller or equal to l for  $P_l^m(x)$  to be non-zero.

These solutions are the physically acceptable ones from the differential equation Eq. (2.36). Another set of non-physical solutions also exist, but these are not interesting for us.

Taking into account the normalization of the angular wave functions, a quite general expression for the functions  $Y_l^m$  is then

$$Y_l^m(\theta,\varphi) = \epsilon \sqrt{\frac{(2l+1)}{4\pi} \frac{(l-|m|)!}{(l+|m|)!}} e^{im\varphi} P_l^m(\cos\theta). \tag{2.45}$$

These functions are called the **spherical harmonics**. Here  $\epsilon = (-1)^m$  for  $m \ge 0$  and  $\epsilon = 1$  for  $m \le 0$ . Examples of these are:

$$Y_0^0 = \left(\frac{1}{4\pi}\right)^{1/2},\tag{2.46}$$

$$Y_1^0 = \left(\frac{3}{4\pi}\right)^{1/2} \cos\theta \tag{2.47}$$

and

$$Y_1^{\pm 1} = \pm \left(\frac{3}{8\pi}\right)^{1/2} \sin\theta e^{\pm i\varphi}.$$
 (2.48)

The quantum number l is commonly called the **azimuthal quantum number**, while m is called the **magnetic quantum number**. We observe that the spherical harmonics do not depend on the potential V, but we have required that the potential is spherically symmetric for the derivations.

# 2.3.3 The Radial equation and solution for the hydrogen atom

The radial equation reads

$$\frac{d}{dr}\left(r^2\frac{dR}{dr}\right) - \frac{2m_e r^2}{\hbar^2}[V(r) - E]R = l(l+1)R. \tag{2.49}$$

The first simple step is to change variables by using

$$u(r) = rR(r), (2.50)$$

which will give the radial part of the Schrödinger equation on a much simpler form

$$-\frac{\hbar^2}{2m_e}\frac{d^2u}{dr^2} + \left[V + \frac{\hbar^2}{2m_e}\frac{l(l+1)}{r^2}\right]u = Eu.$$
 (2.51)

The only difference between this Eq. (2.51) and the one-dimensional Schrödinger equation, Eq. (2.10), is the potential. Here we have an effective potential

$$V_{eff} = V + \frac{\hbar^2}{2m_e} \frac{l(l+1)}{r^2} \tag{2.52}$$

as opposed to the simple V in Eq. (2.10). The extra term in  $V_{eff}$ 

$$\frac{\hbar^2}{2m_e} \frac{l(l+1)}{r^2},\tag{2.53}$$

is called the centrifugal term.

#### The Hydrogen atom and the principal quantum number - n

An important quantum mechanical system is the hydrogen atom. It consists of an electron, charged -e, and a proton, charged e. When dealing with atomic systems, it is common to invoke the Born-Oppenheimer approximation (BOA). The BOA says that the kinetic energy of the nucleus is so small compared to the kinetic energy of the electron(s), that we can freeze out the nucleus' kinetic degrees of freedom. More on the Born-Oppenheimer approximation in section 3.1.4.

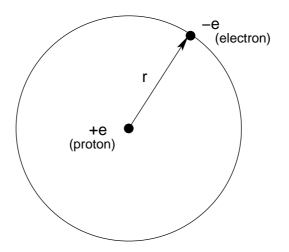


Figure 2.2: A shell model of the hydrogen atom.

For the hydrogen atom, the potential function is given by Coulomb's law between charged particles:

$$V(r) = -\frac{e^2}{4\pi\epsilon_0} \frac{1}{r},\tag{2.54}$$

where  $\epsilon_0$  is the vacuum permittivity. The radial equation for the hydrogen atom is then

$$-\frac{\hbar^2}{2m_e}\frac{d^2u}{dr^2} + \left[ -\frac{e^2}{4\pi\epsilon_0} \frac{1}{r} + \frac{\hbar^2}{2m_e} \frac{l(l+1)}{r^2} \right] u = Eu.$$
 (2.55)

Figure 2.2 gives a schematic overview of the simple hydrogen atom.

In these derivations we will consentrate on the bound states, viz. E < 0.

The radial differential equation, Eq. (2.55), is quite tedious and difficult to solve, so in this section I will just state the important results. From solving the equation we will obtain a certain quantum number, n, also called the **principal quantum** number. This number determines the allowed energies for the system given by

$$E = -\left[\frac{m_e}{2\hbar^2} \left(\frac{e^2}{4\pi\epsilon_0}\right)^2\right] \frac{1}{n^2} = \frac{E_1}{n^2}$$
 (2.56)

for  $n = 1, 2, 3, \dots$ . The ground state of hydrogen is then given as

$$E_1 = -\left[\frac{m_e}{2\hbar^2} \left(\frac{e^2}{4\pi\epsilon_0}\right)^2\right] = -13.6 \text{ eV}.$$
 (2.57)

The radial part of the wave function is now labeled by quantum numbers n and l, giving the full wave function the form

$$\psi_{nlm}(r,\theta,\varphi) = R_{nl}(r)Y_l^m(\theta,\varphi). \tag{2.58}$$

By solving the equation, we also find a constraint on l, namely

$$l = 0, 1, 2, \dots, n - 1. \tag{2.59}$$

The mathematical form of radial wave function is expressed as

$$R_{nl} = \sqrt{\left(\frac{2}{na}\right)^3 \frac{(n-l-1)!}{2n[(n+l)!]^3}} e^{-r/na} \left(\frac{2r}{na}\right)^l \left[L_{n-l-1}^{2l+1}(2r/na)\right], \tag{2.60}$$

where a is the Bohr radius

$$a \equiv \frac{4\pi\epsilon_0 \hbar^2}{m_e e^2} = 0.529 \cdot 10^{-10} \text{m},$$
 (2.61)

and

$$L_{q-p}^{p}(x) \equiv (-1)^{p} \left(\frac{d}{dx}\right)^{p} L_{q}(x)$$
(2.62)

is the associated Laguerre polynomial. The function  $L_q(x)$  is the qth Laguerre polynomial and is defined by

$$L_q(x) \equiv e^x \left(\frac{d}{dx}\right)^q (e^{-x}x^q) \tag{2.63}$$

with p = 2l + 1 and q = n + l. The radial wave functions needed in this project are

$$R_{10} = 2a^{-3/2}e^{-r/a}, (2.64)$$

$$R_{20} = \frac{1}{\sqrt{2}} a^{-3/2} \left( 1 - \frac{1}{2} \frac{r}{a} \right) \frac{r}{a} e^{-r/2a}, \tag{2.65}$$

$$R_{21} = \frac{1}{\sqrt{24}} a^{-3/2} \frac{r}{a} e^{-r/2a}, \tag{2.66}$$

$$R_{30} = \frac{2}{\sqrt{27}} a^{-3/2} \left( 1 - \frac{2}{3} \frac{r}{a} + \frac{2}{27} \left( \frac{r}{a} \right)^2 \right) e^{-r/3a}, \tag{2.67}$$

$$R_{31} = \frac{8}{27\sqrt{6}}a^{-3/2}\left(1 - \frac{1}{6}\frac{r}{a}\right)\left(\frac{r}{a}\right)e^{-r/3a}.$$
 (2.68)

In sections 5.4 and 3.1.2 we will see how we can use these functions to construct the wave functions needed in this project.

# Chapter 3

# Many-particle Systems

While single-particle systems are instructive and are fine examples to examine the properties of quantum mechanical systems, most real life systems consist of many particles that interact with each other and/or an external potential. The aim of this chapter is to describe the quantum mechanics of such systems, or more precisely, atomic systems.

The Hamiltonian of the atomic system will be discussed in section 3.1.4. This is the most important quantity when discussing the energy of a quantum mechanical system. In this part we will see how we can scale the Hamiltonian to have a much cleaner and more comfortable setup for our calculations.

We will also present the Hartree-Fock method (HF), a much used technique used in studies of many-particle systems. The HF method will mainly be used as a way to improve the single particle wave functions used in the Variational Monte Carlo calculations, which will be discussed in chapters 4 and 5.

## 3.1 Atoms

In quantum mechanics an atom can be viewed as a many-particle system. While the wave function for a single particle system is a function of only the coordinates of that particular particle and time,  $\Psi(\mathbf{r},t)$ , a many-particle system will depend on the coordinates of all the particles.

# 3.1.1 Two-particle systems

The simplest example of a many-particle system is of course a two-particle system. The wave function will now be the function

$$\Psi(\mathbf{r}_1, \mathbf{r}_2, t), \tag{3.1}$$

and the Hamiltonian will take the form

$$\widehat{\mathbf{H}} = -\frac{\hbar^2}{2m_1} \nabla_1^2 - \frac{\hbar^2}{2m_2} \nabla_2^2 + V(\mathbf{r}_1, \mathbf{r}_2, t) + V_{external}(\mathbf{r}_1, \mathbf{r}_2).$$
(3.2)

where  $V_{external}$  depends on the system, e.g. the Coulomb interaction between electrons and the nucleus when dealing with atoms. The indices on the Laplacian

operators indicate which electron/coordinates the derivative is taken with respect to. The normalization integral is now

$$\int |\Psi(\mathbf{r}_1, \mathbf{r}_2, t)|^2 d^3 \mathbf{r}_1 d^3 \mathbf{r}_2 = 1$$
(3.3)

with the integrand

$$|\Psi(\mathbf{r}_1, \mathbf{r}_2, t)|^2 d^3 \mathbf{r}_1 d^3 \mathbf{r}_2 \tag{3.4}$$

being the probability of finding particle 1 in the infinitesimal volume  $d^3\mathbf{r}_1$ , and particle 2 in  $d^3\mathbf{r}_2$ . For a time-independent potential, this will give the stationary solutions from separation of variables analogous to the single particle case in section 2.1.2. The full wave function will be

$$\Psi(\mathbf{r}_1, \mathbf{r}_2, t) = \psi(\mathbf{r}_1, \mathbf{r}_2)e^{-iEt/\hbar},\tag{3.5}$$

and the probability distribution function (PDF) will be time-independent

$$P(\mathbf{r}_1, \mathbf{r}_2) = |\psi(\mathbf{r}_1, \mathbf{r}_2)|^2. \tag{3.6}$$

#### Antisymmetry and wavefunctions

For general particles in a composite system, a wave function can be constructed by using the single particle wave functions the particles currently occupy. An example is if electron 1 is in state  $\psi_a$  and electron 2 is in state  $\psi_b$ . The wave function can be constructed as

$$\psi(\mathbf{r}_1, \mathbf{r}_2) = \psi_a(\mathbf{r}_1)\psi_b(\mathbf{r}_2). \tag{3.7}$$

The particles that make up an atom are electrons and the nuclei. The nuclei are the protons and neutrons that the atomic nucleus consists of. As we will see in section 3.1.4, the interesting parts are the electrons. Electrons and the nuclei are so-called fermions, that means particles with *half integer* spin, while particles with *integer* spin are called bosons (e.g. photons). An important point with respect to our quantum mechanical approach is how to construct the wave function for a composite system of particles. Quantum physics tells us that the particles are identical, and cannot be distinguished by some classic method like labelling the particles. This is taken care of by constructing a wave function that opens for the possibility of both electrons being in both states. That is,

$$\psi_{\pm}(\mathbf{r}_1, \mathbf{r}_2) = A[\psi_a(\mathbf{r}_1)\psi_b(\mathbf{r}_2) \pm \psi_b(\mathbf{r}_1)\psi_a(\mathbf{r}_2)], \tag{3.8}$$

with A being a normalization constant. The plus sign is for bosons, while the minus sign applies for fermions. This also shows the **Pauli exclusion principle** which states: No two identical fermions can occupy the same state at the same time. E.g. if  $\psi_a = \psi_b$  the wave function automatically yields zero for fermions, that is

$$\psi_{-}(\mathbf{r}_1, \mathbf{r}_2) = A[\psi_a(\mathbf{r}_1)\psi_a(\mathbf{r}_2) - \psi_a(\mathbf{r}_1)\psi_a(\mathbf{r}_2)] = 0, \tag{3.9}$$

and so the PDF will be zero everywhere. It also tells you that two bosons in **can** occupy the same state at the same time. In fact, any number of bosons may occupy the same state at the same time. Equation (3.8) also underlines an important property of fermionic wave function. That is the antisymmetric property when interchanging two particles. By switching particles 1 and 2 in  $\psi_-$ , you will get  $-\psi_-$  in return, illustrated by

$$\psi_{-}(\mathbf{r}_{2}, \mathbf{r}_{1}) = A[\psi_{a}(\mathbf{r}_{2})\psi_{b}(\mathbf{r}_{1}) - \psi_{b}(\mathbf{r}_{2})\psi_{a}(\mathbf{r}_{1})]$$

$$= -A[\psi_{a}(\mathbf{r}_{1})\psi_{b}(\mathbf{r}_{2}) - \psi_{b}(\mathbf{r}_{1})\psi_{a}(\mathbf{r}_{2})] = -\psi_{-}(\mathbf{r}_{1}, \mathbf{r}_{2}). \tag{3.10}$$

In atoms however, two partices are allowed to occupy the same spatial wave function state, but only if these have different spin states (ref. section 2.1.2). An example is helium, an atom consisting of two electrons and and a core with charge Z=2. Since we can't solve the helium problem, we use an ansatz for the wave function, with the two electrons occupying the  $\psi_{100}$ -functions found when solving the hydrogen problem. The two electrons in the helium case will have the wave functions

$$\Psi = 2a^{-3/2}e^{-r/a}\left(\frac{1}{4\pi}\right)^{1/2}\chi_{\pm} \tag{3.11}$$

with  $\chi_{+}=\uparrow$  as the spin up state, and  $\chi_{-}=\downarrow$  as the spin down state.

## 3.1.2 Wave functions for N-particle atoms

The many particle systems of most interest in this thesis are atomic systems. We can consider both the neutral atom, with the number of electrons equalling the number of protons in the nuclues, or as an isotope, where some of the outermost electrons have been excited. Examples which will be dealt with in this thesis are the neutral atoms helium, beryllium, neon, magnesium and silicon. As mentioned above, we have no closed form solution to any atomic system other than the hydrogen atom, and we have to make an ansatz for the wave function.

#### Slater Determinants

The two-electron wave function

$$\psi_{-}(\mathbf{r_1}, \mathbf{r_2}) = A[\psi_a(\mathbf{r_1})\psi_b(\mathbf{r_2}) - \psi_b(\mathbf{r_1})\psi_a(\mathbf{r_2})]$$
(3.12)

can be expressed as a determinant

$$\psi_{-}(\mathbf{r_1}, \mathbf{r_2}) = A \begin{vmatrix} \psi_a(r_1) & \psi_a(r_2) \\ \psi_b(r_1) & \psi_b(r_2) \end{vmatrix}. \tag{3.13}$$

This is called the **Slater determinant** for the two particle case where the available single particle states are  $\psi_a$  and  $\psi_b$ . The factor A is in this case  $1/\sqrt{2}$ . The general

expression for a Slater determinant,  $\Phi$ , with N electrons and N available single particle states is as given in [3]:

$$\Phi = \frac{1}{\sqrt{N!}} \begin{vmatrix} \psi_1(r_1) & \psi_1(r_2) & \cdots & \psi_1(r_N) \\ \psi_2(r_1) & \psi_2(r_2) & \cdots & \psi_2(r_N) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_N(r_1) & \psi_N(r_2) & \cdots & \psi_N(r_N) \end{vmatrix} .$$
(3.14)

This will automatically comply with the antisymmetry principle and is suited as a wave function for a fermionic many-particle system. The general structure is to take the determinant of an  $N \times N$ -matrix with available particle states in increasing order in the columns. The states are functions of the particle coordinates which are in increasing order in the rows.

More properties of the many-particle Slater determinant will be discussed in section 3.2.

#### 3.1.3 Electron configuration

The electron configuration of an atom describes how the electrons are distributed in the given orbitals of the system. The rules for quantum numbers obtained in sections 2.3.1 and 2.3.3 give us the allowed combinations for the configuration. As shown, n can be  $n = 1, 2, 3, \ldots$ , while l has the allowed values  $l = 0, 1, \ldots, n - 1$ . The quantum number m can take the values  $m = -l, -l + 1, \ldots, l - 1, l$ .

A common way to label the orbitals is to use the spectroscopic notation (see section 5.2.2 in [1]) which labels states with l=0 as s-states, l=1 as p-states and so on. An orbital with n=1 and l=0 is labelled 1s, and an orbital with n=2 and l=1 is labelled 2p. Table 3.1 shows the full list for the azimuthal quantum number, l. The letters s, p, d and f are historical names origining from the words

Quantum number $l$	Spectroscopic notation
0	S
1	p
2	d
3	f
4	g
5	h
6	i
7	k
:	

Table 3.1: The table shows the spectroscopic notation for the azimuthal quantum number l.

sharp, principal, diffuse and fundamental. From g and out the letters are just given alphabetically, but actually skipping j for historical reasons.

The neon atom consists of N=10 electrons and its orbital distribution is given as follows:

- When n = 1, l (and therefore m) can only assume the value 0. But an electron can either be in a spin up, or a spin down state. This means that two electrons can be in the 1s-orbital. This is written as  $(1s)^2$ .
- For n=2 and l=0 there is only m=0, but again two spin states, i.e.  $(2s)^2$ .
- For n=2 and l=1, we can have m=-1,0,1, and both spin states. This means 6 electrons in the 2p-orbital, i.e.  $(2p)^6$ .
- The electron configuration in neon can then be written short and concise as:  $(1s)^2(2s)^2(2p)^6$ .

We can now use this notation to describe the distribution of electrons in the different orbitals for the atoms examined in this thesis. Table 3.2 shows a few important examples. All this tells us is which orbitals to put our electrons in, but it doesn't say anything about what the orbitals in fact are. This is taken care of in sections 5.4 and 5.5.

Atom	Full spectroscopic notation
Hydrogen	(1s)
Helium	$(1s)^2$
Beryllium	$(1s)^2(2s)^2$
Neon	$(1s)^2(2s)^2(2p)^6$
Magnesium	$(1s)^2(2s)^2(2p)^6(3s)^2$
Silicon	$(1s)^2(2s)^2(2p)^6(3s)^2(3p)^2$

Table 3.2: The table shows the electron configuration for hydrogen, helium, beryllium, neon, magnesium and silicon.

# 3.1.4 Hamiltonian and scaling

The Hamiltonian,  $\hat{\mathbf{H}}$ , for an N-electron atomic system is subject to the Born-Oppenheimer approximation (BOA). The BOA effectively freezes out the nucleus' degrees of freedom given the electron mass is small compared to the mass of the nucleus. As a result, we approximate the kinetic energy of the nucleus to be zero during the calculations. The Hamiltonian consists of only the kinetic energy of electrons in addition to the potential energy between electrons and the core, and between the electrons themselves. The Borh-Oppenheimer Hamiltionian is given as

$$\widehat{\mathbf{H}} = \sum_{i}^{N} \frac{\widehat{\mathbf{p}}_{i}^{2}}{2m_{e}} - \sum_{i}^{N} \frac{Ze^{2}}{4\pi\epsilon_{0}r_{i}} + \sum_{i < j} \frac{e^{2}}{4\pi\epsilon_{0}r_{ij}}.$$
(3.15)

The operator  $\hat{\mathbf{p}}$  is the kinetic energy operator,  $m_e$  is the electron mass, Z is the number of protons in the nucleus (equal to the number of electrons in neutral atoms) and  $\epsilon_0$  the vacuum permitivity. The variables  $r_i$  and  $r_{ij}$  represent the distances between an electron and the nucleus and the distance between to different electrons respectively.

By substituting  $\hat{\mathbf{p}}$  with the quantum mechanical operator  $\hat{\mathbf{p}} = -i\hbar\nabla$ , *i* being the imaginary unit and  $\hbar$  Planck's original constant divided by  $2\pi$ , we get

$$\widehat{\mathbf{H}} = -\frac{\hbar^2}{2m_e} \sum_{i}^{N} \nabla_i^2 - \sum_{i}^{N} \frac{Ze}{4\pi\epsilon_0 r_i} + \sum_{i < i} \frac{e^2}{4\pi\epsilon_0 r_{ij}}.$$
 (3.16)

It is more desireable to work with a scaled version of the Hamiltonian. By defining

$$r_0 \equiv \frac{4\pi\epsilon_o \hbar^2}{m_e e^2} \tag{3.17}$$

and

$$\Omega \equiv \frac{m_e e^4}{(4\pi\epsilon_0 \hbar)^2},\tag{3.18}$$

we can write dimensionless variables such as  $H' = H/\Omega$ ,  $\nabla' = \nabla/r_0$  and  $r' = r/r_0$ . The  $r_0$ 's in  $\nabla'$  comes from the factors  $d/dx = d/d(x'r_o)$ . By analyzing each term in Eq. (3.15) and writing out the original variables in terms of constants  $r_0$  and  $\Omega$ , and the dimensionless quantities r' and  $\hat{\mathbf{H}}'$  we get

$$\widehat{\mathbf{H}} = \widehat{\mathbf{H}}'\Omega = -\frac{\hbar^2}{2m_e r_0^2} \sum_{i}^{N} (\nabla_i')^2 + \frac{e^2}{4\pi\epsilon_0 r_o} \sum_{i}^{N} \frac{1}{r_i'} - \frac{Ze^2}{4\pi\epsilon_0 r_0} \sum_{i < i}^{N} \frac{1}{r_{ij}'}.$$
 (3.19)

By inserting Eqs. (3.17) and (3.18) into Eq. (3.19) we obtain

$$\widehat{\mathbf{H}}'\Omega = -\frac{\Omega}{2} \sum_{i}^{N} (\nabla_i')^2 - \Omega \sum_{i}^{N} \frac{Z}{r_i'} + \Omega \sum_{i < j} \frac{1}{r_{ij}'}$$
(3.20)

which divided by  $\Omega$  will give the final dimensionless Hamiltonian

$$\widehat{\mathbf{H}} = -\frac{1}{2} \sum_{i}^{N} \nabla_{i}^{2} - \sum_{i}^{N} \frac{Z}{r_{i}} + \sum_{i < j} \frac{1}{r_{ij}}.$$
(3.21)

Now the energy will be measured in the units of Hartree,  $E_h$ , which converts as,  $1E_h = 2 \cdot 13.6$  eV. These are also called atomic units. We recognize the number 13, 6 as the ground state energy of the hydrogen atom (see section 2.3.3).

# 3.2 Hartree-Fock Theory

A popular and well used many-particle method is the Hartree-Fock method. Hartree-Fock theory assumes the Born-Oppenheimer approximation and can be

used to approximate the ground state energy and ground state wave function of a quantum many body system. In Hartree-Fock theory we assume the wave function,  $\Phi$ , to be a single N-particle Slater determinant. A Hartree-Fock calculation will result in obtaining the single-particle wave functions which minimizes the energy. These wave functions can then be used as optimized wave functions for a variational Monte Carlo machinery (see section 4.4. The single particle orbitals for an atom are given as

$$\psi_{nlm_lsm_s} = \phi_{nlm_l}(\mathbf{r})\chi_{m_s}(s) \tag{3.22}$$

where  $m_l$  is the projection of the quantum number l, given as merely m in section 3.1.3. The quantum number s represents the intrinsic spin of the electrons, which is s = 1/2. The quantum number  $m_s$  now represents the two projection values an electron can have, namely  $m_s = \pm 1/2$ . For a simple description, we label the single particle orbitals as

$$\psi_{nlm_lsm_s} = \psi_{\alpha}, \tag{3.23}$$

with  $\alpha$  containing all the quantum numbers to specify the orbital. The N-particle Slater determinant is given as

$$\Phi(\mathbf{r}_{1}, \mathbf{r}_{2}, \dots, \mathbf{r}_{N}, \alpha, \beta, \dots, \nu) = \frac{1}{\sqrt{N!}} \begin{vmatrix} \psi_{\alpha}(\mathbf{r}_{1}) & \psi_{\alpha}(\mathbf{r}_{2}) & \cdots & \psi_{\alpha}(\mathbf{r}_{N}) \\ \psi_{\beta}(\mathbf{r}_{1}) & \psi_{\beta}(\mathbf{r}_{2}) & \cdots & \psi_{\beta}(\mathbf{r}_{N}) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_{\nu}(\mathbf{r}_{1}) & \psi_{\nu}(\mathbf{r}_{2}) & \cdots & \psi_{\nu}(\mathbf{r}_{N}) \end{vmatrix} .$$
(3.24)

The Hamiltonian (see section 3.1.4) is given as

$$\widehat{\mathbf{H}} = \widehat{\mathbf{H}}_1 + \widehat{\mathbf{H}}_2 = \sum_{i=1}^{N} \left( -\frac{1}{2} \nabla_i^2 - \frac{Z}{r_i} \right) + \sum_{i < j} \frac{1}{r_{ij}}.$$
 (3.25)

or even as

$$\widehat{\mathbf{H}} = \sum_{i=1}^{N} \hat{h}_i + \sum_{i < j} \frac{1}{r_{ij}},\tag{3.26}$$

with  $\hat{h}_i$  being the one-body Hamiltonian

$$\hat{h}_i = -\frac{1}{2}\nabla_i^2 - \frac{Z}{r_i}. (3.27)$$

The variational principle (section 4.3) tells us that the ground state energy for our Hamiltonian,  $E_0$ , is always less or equal to the expectation value of the Hamiltonian with a chosen trial wave function,  $\Phi$ . That is

$$E_0 \le E[\Phi] = \int \Phi^* \widehat{\mathbf{H}} \Phi d\tau \tag{3.28}$$

where the brackets in  $E[\Phi]$  tells us that the expectation value is a functional, i.e. a function of a function, here an integral-function. The label  $\tau$  represents a shorthand notation for  $d\tau = d\mathbf{r}_1 d\mathbf{r}_2 \dots d\mathbf{r}_N$ . We also assume the trial function  $\Phi$  is normalized

$$\int \Phi^* \Phi d\tau = 1. \tag{3.29}$$

By introducing the so-called anti-symmetrization operator,  $\mathcal{A}$ , and the Hartreefunction,  $\Phi_H = \psi_{\alpha}(\mathbf{r}_1)\psi_{\beta}(\mathbf{r}_2)\dots\psi_{\nu}(\mathbf{r}_N)$ , we can write the Slater determinant (Eq. 3.24) more compactly as

$$\Phi(\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N, \alpha, \beta, \dots, \nu) = \sqrt{N!} \mathcal{A} \Phi_H.$$
(3.30)

The Hartree-function is as shown a simple product of the available single particles states and the operator A is given as

$$\mathcal{A} = \frac{1}{N!} \sum_{P} (-)^{P} P, \tag{3.31}$$

with P being the permutation operator of two particles, and the sum spanning over all possible permutations of two particles. The operators  $\widehat{\mathbf{H}}_1$  and  $\widehat{\mathbf{H}}_2$  themselves do not depend on permutations and will commute with the anti-symmetrization operator  $\mathcal{A}$ , namely

$$[\widehat{\mathbf{H}}_1, \mathcal{A}] = [\widehat{\mathbf{H}}_2, \mathcal{A}] = 0. \tag{3.32}$$

The operator  $\mathcal{A}$  also has the property

$$\mathcal{A} = \mathcal{A}^2 \tag{3.33}$$

which we can use in addition to Eq. (3.32) to show that

$$\int \Phi^* \widehat{\mathbf{H}}_1 \Phi d\tau = N! \int \Phi_H^* \mathcal{A} \widehat{\mathbf{H}}_1 \mathcal{A} \Phi_H d\tau \tag{3.34}$$

$$= N! \int \Phi_H^* \widehat{\mathbf{H}}_1 \mathcal{A} \Phi_H d\tau. \tag{3.35}$$

By using Eq. (3.30) and inserting the expression for  $\hat{\mathbf{H}}_1$  we arrive at

$$\int \Phi^* \widehat{\mathbf{H}}_1 \Phi d\tau = \sum_{i=1}^N \sum_P (-)^P \int \Phi_H^* \widehat{\mathbf{h}}_i P \Phi_H d\tau.$$
 (3.36)

Orthogonality of the single-particle functions ensures us that we can remove the sum over P, as the integral will disappear when the two Hartree-Functions,  $\Phi_H^*$  and  $\Phi_H$  are permuted differently. As the operator  $\hat{\mathbf{h}}_i$  is a single-particle operator, all factors in  $\Phi_H$  exept one,  $\phi_{\mu}(\mathbf{r}_i)$ , will integrate out under orthogonality conditions, and the expectation value of the  $H_1$ -operator is written as

$$\int \Phi^* \widehat{\mathbf{H}}_1 \Phi d\tau = \sum_{\mu=1}^N \int \psi_\mu^*(\mathbf{r}_i) \widehat{\mathbf{h}}_i \psi_\mu(\mathbf{r}_i) d\mathbf{r}_i.$$
 (3.37)

The two-body Hamiltonian,  $H_2$ , can be treated similarly:

$$\int \Phi^* \widehat{\mathbf{H}}_2 \Phi d\tau = N! \int \Phi_H^* \mathcal{A} \widehat{\mathbf{H}}_2 \mathcal{A} \Phi_H d\tau$$
 (3.38)

$$= \sum_{i < j=1}^{N} \sum_{P} (-)^{P} \int \Phi_{H}^{*} \frac{1}{r_{ij}} P \Phi_{H} d\tau.$$
 (3.39)

In order to reduce this to a simpler form, we cannot use the same argument as for  $H_1$  for get rid of the P-sum, because of the form of the interation,  $1/r_{ij}$ . The permutations of two electrons will not vanish and must be accounted for by writing

$$\int \Phi^* \widehat{\mathbf{H}}_2 \Phi d\tau = \sum_{i < j=1}^N \int \Phi_H^* \frac{1}{r_{ij}} (1 - P_{ij}) \Phi_H d\tau.$$
 (3.40)

 $P_{ij}$  is the operator which permutes electrons i and j. However, by inspecting the paranthesis  $(1 - P_{ij})$  we can again take advantage of the orthogonality condition in order to further simplify the expression:

$$\int \Phi^* \widehat{\mathbf{H}}_2 \Phi d\tau = \frac{1}{2} \sum_{\mu=1}^N \sum_{\nu=1}^N \left[ \int \psi_{\mu}^*(\mathbf{r}_i) \psi_{\nu}^*(\mathbf{r}_j) \frac{1}{r_{ij}} \psi_{\mu}(\mathbf{r}_i) \psi_{\nu}(\mathbf{r}_j) d\mathbf{r}_i d\mathbf{r}_j - \int \psi_{\mu}^*(\mathbf{r}_i) \psi_{\nu}^*(\mathbf{r}_j) \frac{1}{r_{ij}} \psi_{\mu}(\mathbf{r}_j) \psi_{\nu}(\mathbf{r}_i) d\mathbf{r}_i d\mathbf{r}_j \right].$$
(3.41)

The first term is the *Hartree* term, while the second term is called the *Fock* term. Alternatively they are called the *direct* and *exchange* terms. We see the exchange term has permuted the particles i and j in the  $\Phi_H$ -function. The factor of 1/2 is due to the fact that we sum freely over  $\mu$  and  $\nu$  instead of using  $\sum_{i < j}$ .

Combining the results from  $\widehat{\mathbf{H}}_1$  and  $\widehat{\mathbf{H}}_2$  you get the full Hartree-Fock energy functional

$$E[\Phi] = \sum_{\mu=1}^{N} \int \psi_{\mu}^{*}(\mathbf{r}_{i}) \widehat{\mathbf{h}}_{i} \psi_{\mu}(\mathbf{r}_{i}) d\mathbf{r}_{i} + \frac{1}{2} \sum_{\mu}^{N} \sum_{\nu}^{N} \left[ \int \psi_{\mu}^{*}(\mathbf{r}_{i}) \psi_{\nu}^{*}(\mathbf{r}_{j}) \frac{1}{r_{ij}} \psi_{\mu}(\mathbf{r}_{i}) \psi_{\nu}(\mathbf{r}_{j}) d\mathbf{r}_{i} d\mathbf{r}_{j} - \int \psi_{\mu}^{*}(\mathbf{r}_{i}) \psi_{\nu}^{*}(\mathbf{r}_{j}) \frac{1}{r_{ij}} \psi_{\nu}(\mathbf{r}_{i}) \psi_{\nu}(\mathbf{r}_{j}) d\mathbf{r}_{i} d\mathbf{r}_{j} \right]$$
(3.42)

where now  $\hat{\mathbf{h_i}}$  are the one body Hamiltonians, that is the sum of the kinetic energy and the one-body interaction with the nucleus. The states  $\psi_{\mu}$  and  $\psi_{\nu}$  are the Hartree-Fock orbitals which minimizes the energy when we solve the Hartree-Fock equations. Using Bra-Ket notation, the energy functional can be written more compactly as

$$E[\Phi] = \sum_{\mu=1}^{N} \langle \mu | h | \mu \rangle + \frac{1}{2} \sum_{\mu=1}^{N} \sum_{\nu=1}^{N} \left[ \langle \mu \nu | \frac{1}{r_{ij}} | \mu \nu \rangle - \langle \mu \nu | \frac{1}{r_{ij}} | \nu \mu \rangle \right].$$
 (3.43)

## 3.2.1 Hartree-Fock equations

There are basically two approaches when it comes to minimizing the energy functional in Eq. (3.43). The most straight forward, but a bit more demanding method is to vary the single particle orbitals. The other approach is to expand the single particle orbitals in a well known basis as

$$\psi_a = \sum_{\lambda} C_{a\lambda} \psi_{\lambda},\tag{3.44}$$

where now the  $\psi_{\lambda}$  are in a known orthogonal basis (e.g. hydrogen-like wave functions, harmonic oscillator functions etc.), and we vary the functions  $\psi_a$  with respect to the expansion coefficients  $C_{a\lambda}$ .

#### Varying the single particle wave functions

The background for these calculations are the principles of variational calculus as described in section 18.4.1 in [4]. By introducing so-called Lagrangian multipliers we can minimize a multi-variable functional with constraints. For this particular problem, we introduce  $N^2$  such Lagrange multipliers,  $\epsilon_{\mu\nu}$ . The variational equation (section 18.4.1 in [4]) for the energy functional in Eq. (3.42) is written as

$$\delta E - \sum_{\mu=1}^{N} \sum_{\nu=1}^{N} \epsilon_{\mu\nu} \delta \int \psi_{\mu}^{*} \psi_{\nu} = 0.$$
 (3.45)

We still assume the wave functions are orthogonal, so the variational equation can be written

$$\delta E - \sum_{\mu=1}^{N} \epsilon_{\mu} \int \psi_{\mu}^* \psi_{\mu} = 0. \tag{3.46}$$

The next step is to perform variation with respect to the single particle orbitals  $\psi_{\mu}$ , which gives

$$\sum_{\mu}^{N} \int \delta \psi_{\mu}^{*} \widehat{\mathbf{h}}_{i} \psi_{\mu} d\mathbf{r}_{i} + \frac{1}{2} \sum_{\mu=1}^{N} \sum_{\nu=1}^{N} \left[ \int \delta \psi_{\mu}^{*} \psi_{\nu}^{*} \frac{1}{r_{ij}} \psi_{\mu} \psi_{\nu} d\mathbf{r}_{i} d\mathbf{r}_{j} - \int \delta \psi_{\mu}^{*} \psi_{\nu}^{*} \frac{1}{r_{ij}} \psi_{\nu} \psi_{\mu} d\mathbf{r}_{i} d\mathbf{r}_{j} \right] 
+ \sum_{\mu}^{N} \int \psi_{\mu}^{*} \widehat{\mathbf{h}}_{i} \delta \psi_{\mu} d\mathbf{r}_{i} + \frac{1}{2} \sum_{\mu=1}^{N} \sum_{\nu=1}^{N} \left[ \int \psi_{\mu}^{*} \psi_{\nu}^{*} \frac{1}{r_{ij}} \delta \psi_{\mu} \psi_{\nu} d\mathbf{r}_{i} d\mathbf{r}_{j} - \int \psi_{\mu}^{*} \psi_{\nu}^{*} \frac{1}{r_{ij}} \delta \psi_{\nu} \psi_{\mu} d\mathbf{r}_{i} d\mathbf{r}_{j} \right] 
- \sum_{\mu=1}^{N} \epsilon_{\mu} \int \delta \psi_{\mu}^{*} \psi_{\mu} d\mathbf{r}_{i} - \sum_{\mu=1}^{N} \epsilon_{\mu} \int \psi_{\mu}^{*} \delta \psi_{\mu} d\mathbf{r}_{i} = 0.$$
(3.47)

The variations  $\delta\psi$  and  $\delta\psi^*$  are obviously not independent as they are only a complex transformation apart, but they can indeed be treated independently by replacing  $\delta\psi$  by the imaginary  $i\delta\psi$  and correspondingly  $\delta\psi^*$  by  $i\delta\psi^*$ . The terms depending on  $\delta\psi$  and  $\delta\psi^*$  respectively can now be set equal to zero, yielding two independent sets of equations. By again combining them we will obtain the Hartree-Fock equations

$$\left[ -\frac{1}{2} \nabla_i^2 - \frac{Z}{r_i} + \sum_{\nu=1}^N \int \psi_{\nu}^*(\mathbf{r}_j) \frac{1}{r_{ij}} \psi_{\nu}^*(\mathbf{r}_j) d\mathbf{r}_j \right] \psi_{\mu}(\mathbf{r}_i) 
- \left[ \sum_{\nu=1}^N \int \psi_{\nu}^*(\mathbf{r}_j) \frac{1}{r_{ij}} \psi_{\mu}^*(\mathbf{r}_j) d\mathbf{r}_j \right] \psi_{\nu}(\mathbf{r}_i) = \epsilon_{\mu} \psi_{\mu}(\mathbf{r}_i),$$
(3.48)

where the integral  $\int d\mathbf{r}_j$  also includes a sum over spin quantum numbers for electron j. The first two terms in the first square bracket is the one-body hamiltonian while

the third term is the direct term, representing a mean field of all the other electrons as seen by electron i. The term in the second square bracket is the exchange term, resulting from our antisymmetric wave function ansatz. This term also takes care of the so-called *self interaction* from the first term by cancellation when i = j. We can now define the direct and exchange operators

$$V_{\mu}^{d}(\mathbf{r}_{i}) = \int \psi_{\mu}^{*}(\mathbf{r}_{i}) \frac{1}{r_{ij}} \psi_{\mu}(\mathbf{r}_{j} d\mathbf{r}_{j})$$
(3.49)

and

$$V_{\mu}^{ex}(\mathbf{r}_i)\psi_{\nu}(\mathbf{r}_i) = \left(\int \psi_{\mu}^* \frac{1}{r_{ij}} \psi_{\nu}(\mathbf{r}_j) d\mathbf{r}_j\right) \psi_{\mu}(\mathbf{r}_i). \tag{3.50}$$

Then the Hartree-Fock equations may be written as

$$H_i^{HF} \psi_{\nu}(\mathbf{r}_i) = \epsilon_{\nu} \psi_{\nu}(\mathbf{r}_i) \tag{3.51}$$

with

$$H_i^{HF} = \hat{\mathbf{h}}_i + \sum_{\mu=1}^N V_{\mu}^d(\mathbf{r}_i) - \sum_{\mu=1}^N V_{\mu}^{ex}(\mathbf{r}_i)$$
 (3.52)

as the Hartree-Fock matrix.

In section 5.5, we will discuss the Roothaan-Hartree-Fock approach, which is based on the method discussed in this section, with

$$\psi = \sum_{p} \chi_p C_p \tag{3.53}$$

with  $C_p(\xi)$  being some coefficient and  $\chi$  chosen to be *Slater type functions* on the form

$$\chi(r) \propto r^{n-1} \exp(-\xi r) \tag{3.54}$$

where  $\xi$  is some factor to be determined during the minimization. We see that we must both vary the parameters and the exponents. The results are given in section 5.5 and the explicit expressions and results in [5].

#### Varying the coefficients

We see how the single particle functions in general, e.g.  $\phi$ , are expressed as an expansion of known functions,  $\psi$ , on the form

$$\phi = \sum_{i} C_i \psi_i. \tag{3.55}$$

In this section we will describe a method where we only vary the coefficients, and let the actual basis functions remain unchanged. This will in time give us a more time-efficient way to perform a Hartree-Fock calculation as we will show at a later stage.

The single particle basis functions are here denoted as  $\psi_{\lambda}$ , where  $\lambda = 1, 2, ...$  and represents the full set of quantum numbers in order to describe a single particle

orbital. Our new single particle orbitals will then be an expansion of these functions  $\psi_{\lambda}$  and will have roman indices  $a=1,2,\ldots$  to distinguish them from the chosen basis functions with greek indices. That is

$$\psi_a = \sum_{\lambda} C_{a\lambda} \psi_{\lambda},\tag{3.56}$$

where  $C_{a\lambda}$  are the expansion coefficients that will be varied. By using the Bra-Ket notation we can introduce a more compact way to write the **direct** and **exchange** terms from the Hartree-Fock energy functional. They are

$$\langle \mu\nu|V|\mu\nu\rangle = \int \psi_{\mu}^{*}(\mathbf{r}_{i})\psi_{\nu}^{*}(\mathbf{r}_{j})V(r_{ij})\psi_{\mu}(\mathbf{r}_{i})\psi_{\nu}(\mathbf{r}_{j})d\mathbf{r}_{i}d\mathbf{r}_{j}$$
(3.57)

and

$$\langle \mu \nu | V | \nu \mu \rangle = \int \psi_{\mu}^*(\mathbf{r}_i) \psi_{\nu}^*(\mathbf{r}_j) V(r_{ij}) \psi_{\nu}(\mathbf{r}_i) \psi_{\mu}(\mathbf{r}_j) d\mathbf{r}_i d\mathbf{r}_j$$
(3.58)

respectively. We have now only written V as a general potential instead of the explicit  $V = 1/r_{ij}$ .

The interaction does not change under a permutation of two particles, which means that

$$\langle \mu \nu | V | \mu \nu \rangle = \langle \nu \mu | V | \nu \mu \rangle.$$
 (3.59)

This is also true for the general case

$$\langle \mu \nu | V | \sigma \rho \rangle = \langle \nu \mu | V | \rho \sigma \rangle.$$
 (3.60)

We can have an even more compact notation by defining an antisymmetrized matrix element

$$\langle \mu \nu | V | \mu \nu \rangle_{AS} = \langle \mu \nu | V | \mu \nu \rangle - \langle \mu \nu | V | \nu \mu \rangle, \qquad (3.61)$$

or for the general case

$$\langle \mu\nu | V | \sigma\rho \rangle_{AS} = \langle \mu\nu | V | \sigma\rho \rangle - \langle \mu\nu | V | \rho\sigma \rangle.$$
 (3.62)

The antisymmetrized matrix element has the following symmetry properties:

$$\langle \mu \nu | V | \sigma \rho \rangle_{AS} = -\langle \mu \nu | V | \rho \sigma \rangle_{AS} = -\langle \nu \mu | V | \sigma \rho \rangle_{AS}. \tag{3.63}$$

It is also hermitian, which means

$$\langle \mu \nu | V | \sigma \gamma \rangle_{AS} = \langle \sigma \gamma | V | \mu \nu \rangle_{AS}.$$
 (3.64)

Using these definitions we can have a short-hand notation for the expectiation value of the  $H_2$  operator (see Eq. (3.41)

$$\int \Phi^* \widehat{\mathbf{H}}_2 \Phi d\tau = \frac{1}{2} \sum_{\mu=1}^N \sum_{\nu=1}^N \langle \mu \nu | V | \mu \nu \rangle_{AS}.$$
 (3.65)

which in turn will give the full energy functional

$$E[\Phi] = \sum_{\mu=1}^{N} \langle \mu | h | \mu \rangle + \frac{1}{2} \sum_{\mu=1}^{N} \sum_{\nu=1}^{N} \langle \mu \nu | V | \mu \nu \rangle_{AS}$$
 (3.66)

similar to the one in Eq. (3.43).

The above functional is valid for any choice of single particle functions, so by using the new single particle orbitals introduced above in Eq. (3.56), and by now calling our Hartree-Fock Slater determinant,  $\Psi$ , we then have

$$E[\Psi] = \sum_{a=1}^{N} \langle a | h | a \rangle + \frac{1}{2} \sum_{a=1}^{N} \sum_{b=1}^{N} \langle ab | V | ab \rangle_{AS}.$$
 (3.67)

By inserting the expressions for  $\psi_a$  we will then have

$$E[\Psi] = \sum_{a=1}^{N} \sum_{\alpha\beta} C_{a\alpha}^* C_{a\beta} \langle \alpha | h | \beta \rangle + \frac{1}{2} \sum_{a=1}^{N} \sum_{b=1}^{N} \sum_{\alpha\beta\gamma\delta} C_{a\alpha}^* C_{b\beta}^* C_{a\gamma} C_{b\delta} \langle \alpha\beta | V | \gamma\delta \rangle_{AS}$$
(3.68)

Similarly as for the previous case where we minimized with respect to the wave functions themselves, we now introduce  $N^2$  Lagrange multipliers. As a consequence of choosing orthogonal basis functions, i.e.  $\langle \alpha | | \beta \rangle = \delta_{\alpha,\beta}$ , where  $\delta_{x,y}$  is the Kroenecker delta function, the new basis functions are orthogonal, and we have

$$\langle a | | b \rangle = \delta_{a,b} = \sum_{\alpha\beta} C_{a\alpha}^* C_{a\beta} \langle \alpha | | \beta \rangle = \sum_{\alpha} C_{a\alpha}^* C_{a\alpha}.$$
 (3.69)

The functional to be minimized will now be

$$E[\Psi] - \sum_{a} \epsilon_a \sum_{\alpha} C_{a\alpha}^* C_{a\alpha}$$
 (3.70)

giving a variational equation

$$\frac{d}{dC_{k\lambda}^*} \left[ E[\Psi] - \sum_a \epsilon_a \sum_{\alpha} C_{a\alpha}^* C_{a\alpha} \right] = 0, \tag{3.71}$$

where we have minimized with respect to some coefficient  $C*_{k\lambda}$  since  $C*_{k\lambda}$  and  $C_{k\lambda}$  are independent. Inserting the energy functional from Eq. (3.66) we will get

$$\sum_{\beta} C_{k\beta} \langle \lambda | h | \beta \rangle - \epsilon_k C_{k\lambda} + \frac{1}{2} \sum_{b=1}^{N} \sum_{\beta,\gamma,\delta} C_{b\beta}^* C_{k\gamma} C_{b\delta} \langle \lambda \beta | V | \gamma \delta \rangle_{AS}$$

$$+ \frac{1}{2} \sum_{a=1}^{N} \sum_{\alpha,\gamma,\delta} C_{a\alpha}^* C_{a\gamma} C_{k\delta} \langle \alpha \lambda | V | \gamma \delta \rangle_{AS} = 0$$
(3.72)

as

$$\frac{\mathrm{d}}{\mathrm{d}C_{k\lambda}} \left[ \sum_{a=1}^{N} \sum_{\alpha,\beta} C_{a\alpha}^* C_{a\beta} \langle \alpha | h | \beta \rangle \right] = \sum_{a=1}^{N} \sum_{\alpha,\beta} C_{a\beta} \langle \alpha | h | \beta \rangle \, \delta_{ka} \delta_{\lambda\alpha} \tag{3.73}$$

$$= \sum_{\beta} C_{k\beta} \left\langle \lambda | h | \beta \right\rangle. \tag{3.74}$$

In the last term in Eq. (3.72) we can use that  $\langle \alpha \lambda | V | \gamma \delta \rangle_{AS} = -\langle \lambda \alpha | V | \gamma \delta \rangle_{AS} = \langle \lambda \alpha | V | \delta \gamma \rangle_{AS}$  and change summation variable in both sums from b to a which gives

$$\sum_{\beta} C_{k\beta} \langle \lambda | h | \beta \rangle - \epsilon_k C_{k\lambda} + \sum_{a=1}^{N} \sum_{\alpha, \gamma, \delta} C_{a\alpha}^* C_{k\gamma} C_{a\delta} \langle \lambda \alpha | V | \gamma \delta \rangle_{AS} = 0, \qquad (3.75)$$

or by using

$$h_{\lambda\gamma}^{HF} = \langle \lambda | h | \gamma \rangle + \sum_{a=1}^{N} \sum_{\alpha,\delta} C_{a\alpha}^* C_{a\delta} \langle \lambda \alpha | V | \gamma \delta \rangle_{AS},$$

we can write the Hartree-Fock equations for this particular approach as

$$\sum_{\gamma} h_{\lambda\gamma}^{HF} C_{k\gamma} = \epsilon_k C_{k\lambda}. \tag{3.76}$$

This is now a non-linear eigenvalue problem which can be solved by an iterative process, or a standard eigenvalue method. The advantage of this approach is the fact that we can calculate the matrix elements  $\langle \alpha | h | \beta \rangle$  and  $\langle \alpha \beta | V | \gamma \delta \rangle$  once and for all, since we don't change the basis functions, but only their coefficients. In the first approach, where we vary the actual wave functions, we have to calculate these matrix elements for each iteration. When we just vary the coefficients, we can calculate the matrix elements and store them in a table before we start the iterative process. This will lead to a quicker algorithm, but we will see that the method of varying both coefficients and basis functions actually produces the best results.

# Chapter 4

# Quantum Monte Carlo

Monte Carlo methods are designed to simulate a mathematical system, or in our case, a quantum mechanical system. By using random numbers, these methods are considered stochastic, i.e. non-deterministic unlike other simulation techniques, such as Hartree Fock, Coupled Cluster Theory and Configuration Interaction, ref. [6]. Monte Carlo methods can be used to simulate quantum mechanical systems, but are also well suited for calculating integrals, especially high-dimensional integrals. There are in fact several Quantum Monte Carlo techniques such as Diffusion Monte Carlo, Green's function Monte Carlo and Variational Monte Carlo, see [7]. In this thesis we will focus on the Variational Monte Carlo method.

The Quantum Monte Carlo calculations are so called *ab initio* methods, which are first principle calculations These methods have their basis on probability and statistics, and in order to get good expectation values and variances, the quantity in question must be sampled millions of times. Each such sample is called a Monte Carlo cycle, and can be mistaken for being linked with the dynamics of the system. However, the systems we will examine are stationary, meaning that they are time-independent systems.

To use the Quantum Monte Carlo technique, we will need a Probability Distribution Function (PDF), P(x), to guide our sampling and which characterizes the system, with x being some set of variables, and of course a quantity to sample, in our case the energy of the system, E. The expectation value of the quantity of interest, e.g. an operator  $\hat{\mathbf{Q}}$ , will now be

$$\langle Q \rangle = \int P(x) \widehat{\mathbf{Q}} dx.$$
 (4.1)

We will see in section 4.4.1 that the operator we are looking for in our case will be the local energy operator

$$\widehat{\mathbf{E}}_L(\mathbf{R};\alpha) = \frac{1}{\psi_T} \widehat{\mathbf{H}} \psi_T(\mathbf{R};\alpha), \tag{4.2}$$

where **R** is the set of all spatial variables,  $\psi_T$  a trial wave function since we don't have the exact solution  $\Psi$  (if we did, we wouldn't have to go through all of this), and  $\alpha$  a set of variational parameters which will be discussed in the following section.

Since the Monte Carlo method is a statistical method involving average values etc., a certain degree of uncertainty and error will always occur. A method to minimize the error estimate, the so-called *blocking* technique, will be discussed as a way to get the results as correct as possible. In addition to this, we will discuss the *Davidon-Fletcher-Powell*-algorithm (DFP), which is an improvement of the Conjugate Gradient Method (CGM), see [8]. DFP is a method which finds the minimum of a multivariable function.

This chapter will also discuss the variational principle and the Metropolis algorithm for the Variational Monte Carlo method. The Metropolis-Hastings algorithm will also be discussed including importance sampling which will improve the method further, with more relevant sample points.

# 4.1 Markov chains

A Markov chain is a way to model a system which evolves in time, and depends solely on it's previous state. We can consider a system that is described by its constituents having different probabilities to occupy some physical state. That is, the system is described by a PDF,  $w_j(t)$ , at an instance t, where the index j indicates that the system is discretized (for simplicity). We now have a transition probability,  $W_{ij}$ , which gives us the probability for the system going from a state given by  $w_j(t)$  to a state  $w_i(t+\epsilon)$ , where  $\epsilon$  is a chosen time step. The relation between the PDFs  $w_i$  and  $w_j$  will be

$$w_i(t+\epsilon) = \sum_j W_{ij} w_j(t), \qquad (4.3)$$

where  $w_i$  and  $W_{ij}$  are probabilites and must fulfill

$$\sum_{i} w_i(t) = 1,\tag{4.4}$$

and

$$\sum_{i} W_{ij} = 1. \tag{4.5}$$

We can also write this as a matrix equation

$$\mathbf{w}(t+\epsilon) = \mathbf{W}\mathbf{w}(t). \tag{4.6}$$

The equilibrium state of the system can be described by

$$\mathbf{w}(\infty) = \mathbf{W}\mathbf{w}(\infty),\tag{4.7}$$

which implies a steady state, since the transition matrix now has no effect on the state vector,  $\mathbf{w}$ . Eq. (4.7) now reads as an eigenvalue equation. By solving this equation, the eigenvectors would then be  $\mathbf{w}_1, \mathbf{w}_2, \mathbf{w}_3, \ldots$  and so on, while the corresponding eigenvalues would be  $\lambda_1, \lambda_2, \lambda_3, \ldots$  etc. If the largest eigenvalue is  $\lambda_1$ , viz.

$$\lambda_1 > \lambda_2, \lambda_3, \lambda_4, \dots, \tag{4.8}$$

the steady state would then be  $\mathbf{w}_1$ , as its eigenvalue would cause this state to dominate as time progresses.

However, we do not know the form of the transition probability,  $W_{ij}$ , and need a way to examine the conditions at equilibrium. We start with the so-called *Master* equation,

$$\frac{dw_i(t)}{dt} = \sum_{i} \left[ W(j \to i)w_j - W(i \to j)w_i \right],\tag{4.9}$$

which expresses the time rate of the PDF as a balance between state transitional probabilities from states j to i and from state i to states j. At equilibrium this time derivative is zero,  $dw(t_{eq})/dt = 0$ , implying that the sums of the transitions must be equal, namely

$$\sum_{j} W(j \to i) w_j = \sum_{j} W(i \to j) w_i. \tag{4.10}$$

By using Eq. (4.5) on the right-hand side of the equation, we get

$$w_i = \sum_j W(j \to i) w_j. \tag{4.11}$$

We do, however, need an even more stringent requirement to obtain the correct solutions, which is the so-called *detailed balance principle*:

$$W(j \to i)w_j = W(i \to j)w_i, \tag{4.12}$$

saying that the transition rate between any two states i and j must be equal as well. This gives the condition

$$\frac{w_i}{w_j} = \frac{W(j \to i)}{W(i \to j)}. (4.13)$$

The detailed balance principle is imposed on the system to avoid cyclic solutions. This principle ensures us that once the most probable state is reached, the one corresponding to the lowest eigenvalue, the system will not traverse back to any state, corresponding to higher eigenvalues, see ref. [9].

A Markov chain must also comply with the *ergodic principle*, which says that a system should have a non-zero probability of visiting all possible states during the process. A Markov chain does only depend on its previous step, and will fulfill ergodicity. The Metropolis algorithm is based on a Markovian process and fulfills the ergodic principle.

Our physical system, an atom consisting of N electrons, is a system that can be described by a PDF and which only depends on its previous system state, the PDF being the absolute square of the wave function, namely  $|\psi|^2$ . A wise choice would then be to use the Metropolis algorithm in order to simulate the system.

### 4.2 Random numbers

A main component of a Monte Carlo calculation is the use of random numbers. The most common random number distribution is the uniform distribution, which returns a random number r between zero and one, that is

$$r \in (0,1)$$
.  $(4.14)$ 

The name *uniform* (chapter 7.1 in [8]) tells us that every number between 0 and 1 could be returned with equal probability. Another much used distribution is the Gaussian, or normal distribution (see chapters 7.3.4 in [8]).

As we saw in section 4.1, a Markov chain is a mathematical model which relies only on it's previous step in the calculation. The true random numbers used to calculate i.e. new positions for particles etc., should be uncorrelated. In computers however, we cannot get true random numbers, and must therefore settle for so-called Pseudo-Random Number Generators (PRNG). A problem with these number generators is that they have a certain period, and this sequence of pseudo-random numbers will repeat itselff after completing a cycle of the sequence. If this period is not sufficiently large, the numbers will be correlated. However, the pseudo-random number generators we use have quite large periods, and we will not suffer noticeably.

# 4.3 The Variational Principle

The main target for us is to solve the time independent Schrödinger equation

$$\widehat{\mathbf{H}}\Psi = E\Psi \tag{4.15}$$

where  $\hat{\mathbf{H}}$  is the Hamiltonian, the operator representing the energy of the system,  $\Psi$  the quantum mechanical wave function, and E the total energy of the system. The variational principle tells us that for any given choice of trial wave function,  $\psi_T$ , the total energy

$$E = \frac{\langle \psi_T | \, \hat{\mathbf{H}} \, | \psi_T \rangle}{\langle \psi_T | \psi_T \rangle} \tag{4.16}$$

will always be larger or equal to the true ground state energy,  $E_0$ , for the chosen Hamiltonian  $\widehat{\mathbf{H}}$ ,

$$E_0 \le \frac{\langle \psi_T | \widehat{\mathbf{H}} | \psi_T \rangle}{\langle \psi_T | \psi_T \rangle}.$$
 (4.17)

To show this, we exploit the fact that  $\hat{\mathbf{H}}$  is Hermitian, and that the Hamiltonian will have a set of exact eigenstates forming a complete basis set,  $|\Psi_i\rangle$ . Our trial function can then be expanded as

$$\langle \psi_T | = \sum_i c_i | \Psi_i \rangle , \qquad (4.18)$$

and the expectation value for  $\hat{\mathbf{H}}$  using the trial function  $\psi_T$  will be

$$E = \frac{\langle \psi_T | \widehat{\mathbf{H}} | \psi_T \rangle}{\langle \psi_T | \psi_T \rangle} = \frac{\sum_{ij} c_i^* c_j \langle \Psi_i | \widehat{\mathbf{H}} | \Psi_j \rangle}{\sum_{ij} c_i^* c_j \langle \Psi_i | \Psi_j \rangle}$$
$$= \frac{|c_i|^2 E_i \langle \Psi_i | \Psi_i \rangle}{|c_i|^2 \langle \Psi_i | \Psi_i \rangle} = \frac{\sum_i |c_i|^2 E_i}{\sum_i |c_i|^2}. \tag{4.19}$$

The crucial point is to know that every excited state energy,  $E_i$ , is greater or equal to the ground state energy,  $E_0$ , that is

$$E = \frac{|c_i|^2 E_i \langle \Psi_i | \Psi_i \rangle}{|c_i|^2 \langle \Psi_i | \Psi_i \rangle} = \frac{\sum_i |c_i|^2 E_i}{\sum_i |c_i|^2} \ge \frac{|c_i|^2 E_0 \langle \Psi_i | \Psi_i \rangle}{|c_i|^2 \langle \Psi_i | \Psi_i \rangle} = \frac{\sum_i |c_i|^2 E_i}{\sum_i |c_i|^2} = E_0. \quad (4.20)$$

This inequality gives an equality when you find the true ground state wave function,  $\Psi_0$ , that is

$$E_0 = \langle \Psi_0 | \widehat{\mathbf{H}} | \Psi_0 \rangle. \tag{4.21}$$

As our main goal is to approximate the ground state energy, this is the basis of our calculations. This gives us an idea to introduce variational parameters to our trial wave function in search for a good approximation. Instead of just having the wave functions depending on the spatial coordinates as  $\psi_T = \psi_T(x, y, z)$ , we can write  $\psi_T$  as

$$\psi_T = \psi_T(x, y, z, \alpha, \beta, \dots). \tag{4.22}$$

Our goal is to find the minimum of functional E in Eq. (4.16), viz. the set of parameters that gives the lowest energy when calculating E. A common approach in VMC is too also search for the minimum of the variance. A priori, we know that when the variance is exactly zero, we have found the true ground state. However, as we have no knowledge of the true form of the wave function, we must make an ansatz for it. This ansatz will in general not have a minimum for both the energy and the variance for the same set of parameters. As mentioned in [10], the most efficient wave function is found if we use a linear combination of the parameters that minimize the energy and the variance respectively.

In this work however, I have focused on finding the minimum of the energy only.

#### 4.4 Variational Monte Carlo

The main exercise with the Variational Monte Carlo process for atoms is to move the particles in the system guided by a probability distribution and sample the energy at these states in order to calculate various expectation values (mainly  $\langle E \rangle$ and  $\langle E^2 \rangle$ ). In our case we have a known Hamiltonian,  $\hat{\mathbf{H}}$ , and a chosen manyparticle trial wave function,  $\psi_T$ . The expectation value of the Hamiltonian is

$$\langle H \rangle = \frac{\int d\mathbf{R} \psi_T^*(\mathbf{R}; \alpha) \widehat{\mathbf{H}}(\mathbf{R}) \psi_T(\mathbf{R}; \alpha)}{\int d\mathbf{R} \psi_T^*(\mathbf{R}; \alpha) \psi_T(\mathbf{R}; \alpha)},$$
(4.23)

where  $\mathbf{R} = (\mathbf{R_1}, \mathbf{R_2}, \dots, \mathbf{R_N})$  are all the coordinates for the N particles, and  $\alpha$  is the set of all the variational parameters in question. The general procedure will now be

- 1. Construct a trial wave function,  $\psi_T$  as a function of the N particles' coordinates and the chosen variational parameters
- 2. Calculate  $\langle H \rangle$  using Eq. (4.23)

#### 3. Vary the parameters according to some minimization technique

We will first focus on the second item in the list above, with a focus on the Metropolis algorithm.

#### 4.4.1 VMC and the simple Metropolis algorithm

We assume first that our trial wave function,  $\psi_T$ , is not normalized so the quantum mechanical probability distribution is then given as

$$P(\mathbf{R}; \alpha) = \frac{|\psi_T(\mathbf{R}; \alpha)|^2}{\int |\psi_T(\mathbf{R}; \alpha)|^2 d\mathbf{R}}.$$
 (4.24)

Equation (4.24) gives the PDF for our system. We now define a new operator, called the local energy operator as

$$\widehat{\mathbf{E}}_L(\mathbf{R};\alpha) = \frac{1}{\psi_T} \widehat{\mathbf{H}} \psi_T(\mathbf{R};\alpha), \tag{4.25}$$

and the expectation value of the local energy is given as

$$\langle E_L(\alpha) \rangle = \int P(\mathbf{R}; \alpha) \widehat{\mathbf{E}}_L(\mathbf{R}; \alpha) d\mathbf{R}.$$
 (4.26)

The dimensionless Hamiltonian in section 3.1.4 is given as

$$\widehat{\mathbf{H}} = -\frac{1}{2} \sum_{i}^{N} \nabla_{i}^{2} - \sum_{i}^{N} \frac{1}{r_{i}} + \sum_{i < j} \frac{1}{r_{ij}}.$$
(4.27)

Inserting the Hamiltonian from Eq. (4.27) into Eq. (4.25), the local energy operator will be

$$\widehat{\mathbf{E}}_{L} = \frac{1}{\psi_{T}(\mathbf{R}, \alpha)} \left( -\frac{1}{2} \sum_{i}^{N} \nabla_{i}^{2} - \sum_{i}^{N} \frac{1}{r_{i}} + \sum_{i < j} \frac{1}{r_{ij}} \right) \psi_{T}(\mathbf{R}, \alpha).$$
(4.28)

By observing that the last two terms don't affect the wave function in any way, we have

$$\widehat{\mathbf{E}}_L = \frac{1}{\psi_T(\mathbf{R}, \alpha)} \left( -\frac{1}{2} \sum_{i=1}^{N} \nabla_i^2 \right) \psi_T(\mathbf{R}, \alpha) - \sum_{i=1}^{N} \frac{1}{r_i} + \sum_{i < j} \frac{1}{r_{ij}}.$$
 (4.29)

This will be quantity we sample in each Monte Carlo cycle, following our PDF from Eq. (4.24). The more we improve the trial wave function the closer the expectation value of the local energy,  $\langle E_L \rangle$ , gets to the exact energy E.

As we sample the local energy  $E_L$  we also sample  $E_L^2$  in order to get  $\langle E_L^2 \rangle$  for the variance

$$\sigma_{E_L}^2 = \langle E_L^2 \rangle - \langle E_L \rangle^2. \tag{4.30}$$

We see from appendix B that we now can calulcate the

$$\langle E_L \rangle = \frac{1}{n} \sum_{k=1}^n E_L(x_k), \tag{4.31}$$

where  $x_k$  are the points at which the local energy is sampled, and n are the number of sample points.

An important note is that in Metropolis algorithm only involves ratios between probabilities, so the denominator in Eq. (4.24) actually never needs to be calculated. We will see this in the next section.

#### The Metropolis algorithm

The Metropolis algorithm uses ratios between probabilities to determine whether or not a chosen particle is to be moved to a proposed position or not. When an electron is moved, the set of positions  $\mathbf{R}$  change to positions  $\mathbf{R}'$ . The ratio  $w = P(\mathbf{R}')/P(\mathbf{R})$  is now the transition probability from the state with particles being in positions  $\mathbf{R}$  to a state where particles are in positions  $\mathbf{R}'$ . This ratio w will now be

$$w = \left| \frac{\psi_T(\mathbf{R}')}{\psi_T(\mathbf{R})} \right|^2. \tag{4.32}$$

The Metropolis algorithm tells us that if

$$w > 1, \tag{4.33}$$

we automatically accept the new positions. If w < 1, we compare w with a random number r, with  $r \in (0,1)$ . If

$$r \le w,\tag{4.34}$$

we also accept the new positions. If neither of the two inequalities are true, the chosen electron remains at its position before the proposed move. The most effective way to use this method is to move one particle at a time before running the tests for w.

We now have enough information to describe the elements needed to calculate the expectation value for the local energy using the Metropolis algorithm. The specifics of the implementation will be discussed in the next chapter, but the following points still show how it's done.

- 1. Set all electrons in random positions initially, that is a set of positions R
- 2. Start a Monte Carlo cycle and a loop over electrons
- 3. Calculate the new position,

$$\mathbf{R}'_i = \mathbf{R}_i + s * r$$

for electron i using a uniform random number, r and a finite stepsize, s. This will change the entire set of positions from  $\mathbf{R}$  to  $\mathbf{R}$ .

4. Calculate the ratio

$$w = |\psi_T(\mathbf{R}')/\psi_T(\mathbf{R})|^2$$

5. Use the checks in Eqs. (4.33) and (4.34) to decide whether to accept new positions,  $\mathbf{R} = \mathbf{R}'$ , or to stay in initial positions  $\mathbf{R}$ .

- 6. If accepted, update positions
- 7. Repeat steps 3 through 6 for each electron
- 8. After looping over all particles, calculate the local energy and update  $\langle E_L \rangle$  and  $\langle E_L^2 \rangle$ .
- 9. Repeat steps 3 through 8 for the chosen number of Monte Carlo cycles

The transition rules being used here is called a uniform symmetrical transition rule, and it is common to keep the acceptance ratio around 0.5 (this is achieved by choosing the step length  $s \approx 1$ ). The acceptance ratio is the number of accepted steps divided by total number of particle moves. However, this simple Metropolis algorithm does not seem very efficient, due to the way the new positions are calculated. They depend solely on the previous position and not on any guiding mechanism. In order to make this method more efficient, we introduce the Metropolis-Hastings algorithm with so-called importance sampling. With importance sampling we will get a much higher acceptance ratio since the calculation of new positions and the Metropolis-test will depend on the gradient of the wave function. Figure 4.1 shows a float chart of the Metropolis method.

# 4.4.2 Metropolis-Hastings algorithm and importance sampling

As an improvement to our Metropolis algorithm, we introduce the Metropolis-Hastings algorithm, which involves importance sampling. A new term that has to be implemented is the **quantum force** of the system:

$$F = 2\frac{1}{\psi_T} \nabla \psi_T \tag{4.35}$$

A physical interpretation of the quantum force, and therefore a good reason to implement it, is that the gradient of the wavefunction tells us which direction the electrons are moving. If we rather use the quantum force to steer us in the direction of the electron movement, rather than sampling random points around the nucleus, the points we sample will be more relevant and therefore our acceptance ratio increases to about 0.8-0.9. Despite the fact that computing the quantum force will increase the calculation time for each cycle, this will in time decrease our thermalization period, the period for the system to reach a most likely state. This will result in a more efficient algorithm.

#### Importance sampling

The one-dimensional diffusion equation is written as

$$\frac{\partial P(x,t)}{\partial t} = D \frac{\partial^2 P(x,t)}{\partial x^2} \tag{4.36}$$

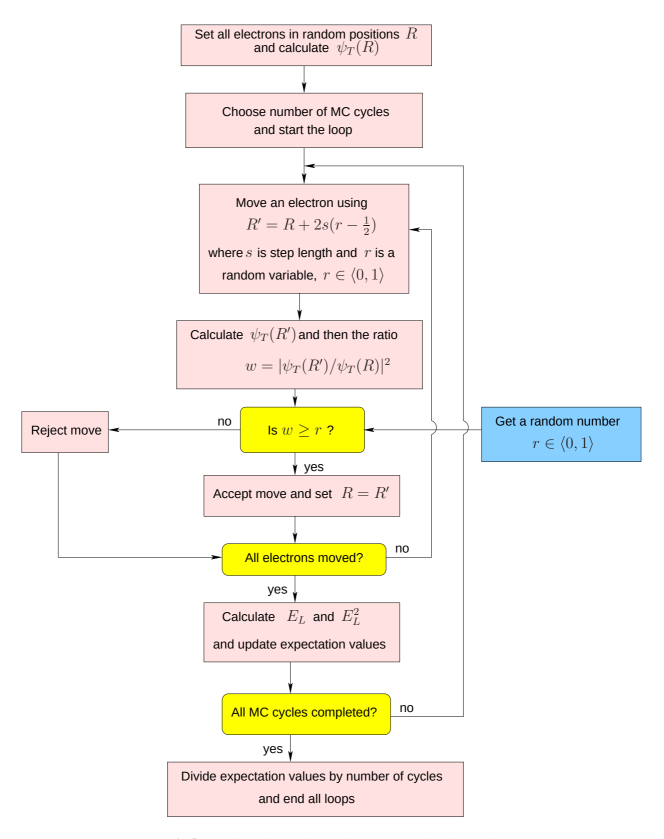


Figure 4.1: A float chart describing the Metropolis algorithm

and describes how the probability distribution, P, of a random walker evolves in time. D is the diffusion constant (see chapter 14 in [4] or chapter 1 in [7]). The Green's function

$$G(y, x, t) = \frac{1}{(4\pi D\Delta t)^{3N/2}} \exp\left(-(y - x)^2/4Dt\right)$$
(4.37)

is a general solution to the diffusion equation for a single walker which starts off at position x at time t=0. By using discrete time steps, it can be shown that a Markov process models a diffusion process, as the transition probability in a time step  $\Delta t$  from position x to position y,  $G(y, x, \Delta t)$ , only depends on the previous position, or state. The Fokker-Planck equation:

$$\frac{\partial P}{\partial t} = D \frac{\partial}{\partial x} \left( \frac{\partial}{\partial x} - F \right) P(x, t), \tag{4.38}$$

describes the time evolution of a PDF. As opposed to the diffusion equation, the Fokker-Planck equation also includes a drift term, F, yielding a solution given as

$$G(y, x, \Delta t) = \frac{1}{(4\pi D\Delta t)^{3N/2}} \exp\left(-(y - x - D\Delta t F(x))^2 / 4D\Delta t\right). \tag{4.39}$$

This is very similar to the solution of the diffusion equation but includes the external "force", F. This can be interpreted as a probability distribution of a single random walker with a drift force term starting off at position x at time t = 0 [11]). This Green's function serves as a transition probability for an electron when it is under influence of the drift term, F. This new propability must be multiplied with the probability from Eq. (4.24) and the ratio of probabilities from Eq. 4.32 is then replaced by the modified ratio

$$w = \frac{G(x, y, \Delta t)|\psi_T(y)|^2}{G(y, x, \Delta t)|\psi_T(x)|^2},$$
(4.40)

or more explicitely

$$w = \frac{|\psi_T(y)|^2}{|\psi_T(x)|^2} \exp\left(-(y - x - D\Delta t F(x))^2 - (x - y - D\Delta t F(y))^2\right). \tag{4.41}$$

The Langevin equation is a stochastic differential equation describing the dynamics of random walkers as

$$\frac{\partial x(t)}{\partial t} = DF(x(t)) + \eta, \tag{4.42}$$

where  $\eta$  is a random variable following the uniform distribution between 0 and 1. Integrating this equation using the consepts of stochastic integration, will yield the equation

$$y = x + DF(x)\Delta t + \xi\sqrt{\Delta t}$$
(4.43)

where  $\xi$  now is a Gaussian distributed random variable and  $\Delta t$  is a chosen timestep. This equation replaces Eq. (3) as the equation that calculates the new position for a particle. We see that this depends on the drift term, the quantum force, F(x). This drift term "pushes" the walkers in the direction of more probable states because the gradient of our PDF will point in the direction of higher values of probability.

By using the Fokker-Planck equation (where P is a time-dependent probability density),

$$\frac{\partial P}{\partial t} = D \frac{\partial}{\partial x} \left( \frac{\partial}{\partial x} - F \right) P(x, t), \tag{4.44}$$

and restraining a system to be in a stationary state, that is  $\partial P/\partial t = 0$ , we find the form of F as

$$F(x) = \frac{1}{P} \frac{dP}{dx},\tag{4.45}$$

or

$$F(\mathbf{r}) = \frac{1}{P} \nabla P \tag{4.46}$$

in three dimension, giving us a constraint for the form of the quantum force. Our quantum mechanical system has a probability distribution given as

$$P(\mathbf{r}) = |\psi_T(\mathbf{r})|^2, \tag{4.47}$$

and the quantum force of our system will be

$$F(x) = 2\frac{1}{\psi_T} \nabla \psi_T. \tag{4.48}$$

This algorithm will speed up the process as more important or relevant states are sampled (or visited in state space), hence the name, importance sampling.

#### Metropolis-Hastings algorithm

The Metropolis-Hastings algorithm can be summarized in the same way as for the simple Metropolis algorithm:

- 1. Set all electrons in random positions as an initial point, x
- 2. Start a Monte Carlo cycle and a loop over electrons
- 3. Calculate the new position,

$$y_i = x_i + DF(x)\Delta t + \xi \sqrt{\Delta t}$$

for the chosen electron i using a Gaussian distributed number,  $\xi$  and a chosen timestep,  $\Delta t$ 

- 4. Calculate the quantum force for the new set of positions, y, and then the Green's function
- 5. Calculate the ratio w from equation Eq. (4.41)
- 6. Use the checks in Eqs. (4.33) and (4.34) to decide whether to accept new positions, y, or to stay in initial positions x

- 7. If accepted, update positions and quantum force
- 8. Repeat steps 3 through 7 for each electron
- 9. After looping over all particles, calculate the local energy and update  $\langle E_L \rangle$  and  $\langle E_L^2 \rangle$ .
- 10. Repeat steps 3 through 9 for the chosen number of Monte Carlo cycles

Figure 4.2 shows a float chart of the Metropolis-Hastings algorithm. The detailed implementation as a computer program will be discussed in the next chapter.

Both the simple Metropolis and the Metropolis-Hastings algorithms demand 4N random numbers per Monte Carlo cycle, where N is the number of particles. Per particle move, we need 3 normal distributed random numbers to make a move in each of the spatial dimensions. We also need one uniformly distributed random number for the Metropolis-test itself.

## 4.5 Blocking

The simulation of physical systems using Monte Carlo methods, involve pseudorandom numbers and will produce correlated results. That is, as the pseudorandom numbers themselves will show correlated behaviour, functions evaluated with respect to variables dependent on these pseudo-random numbers will become correlated as well. As discussed in appendix B, the statistical error for set of correlated stochastic values is given by

$$\operatorname{err}_{X}^{2} = \frac{1}{n}\operatorname{Cov}(x),\tag{4.49}$$

while for uncorrelated values this will reduce to

$$\operatorname{err}_{X}^{2} = \frac{1}{n} \operatorname{Var}(x) = \frac{1}{n^{2}} \sum_{k=1}^{n} (x_{k} - \bar{x}_{n})^{2}.$$
 (4.50)

As given in appendix B, we can write the error for correlated cases as

$$\operatorname{err}_{X}^{2} = \frac{\tau}{n} \operatorname{Var}(x), \tag{4.51}$$

where  $\tau$  is the autocorrelation time

$$\tau = 1 + 2\sum_{d=1}^{n-1} \kappa_d,\tag{4.52}$$

describing the correlation between measurements. However, calculating  $\kappa_d$ , as given in Eqs. (B.34) and (B.35), will be rather time consuming.

By using the blocking technique, or 'data bunching', as described in [12] and [13], we actually have a way to make the measurements 'uncorrelated'. By grouping the data in equally sized blocks, and treating the mean, or sample mean,

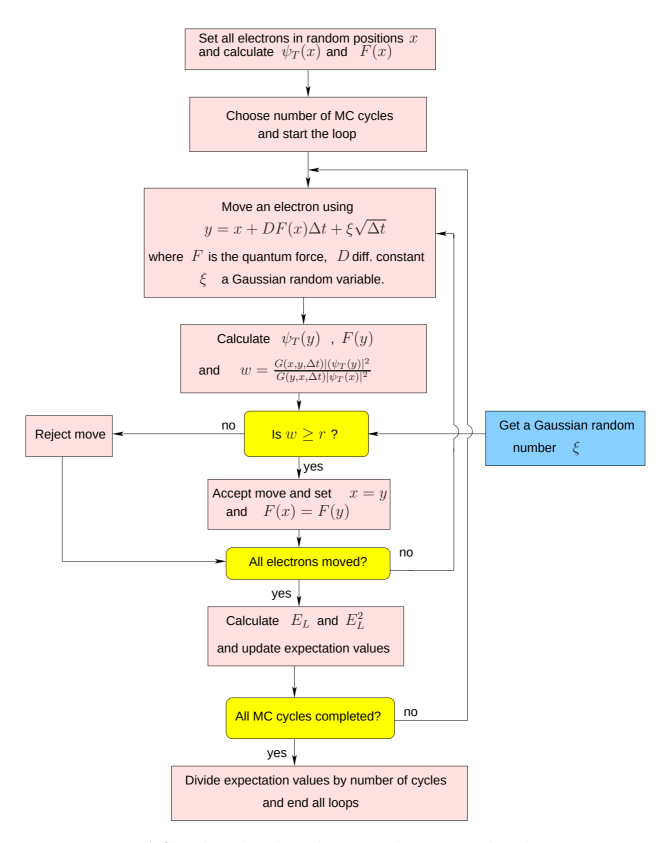


Figure 4.2: A float chart describing the Metropolis-Hastings algorithm

of each block as a measurement, we can find an error estimate by merely plotting the error as a function of block size as if the data were uncorrelated.

Suppose we have 1000 measurements in our sample. We can then divide these values into 10 blocks with 100 measurements in each block. Furthermore, we calculate the mean in each block, and treat them as our measurements. Then we calculate the error as if the data were uncorrelated using Eq. (4.50), where the block means are our  $x_k$ 's and n = 10. The total sample mean,

$$\bar{x}_n = \frac{1}{n} \sum_{k=1}^n x_k, \tag{4.53}$$

will of course be identical whether we calculate it directly, that is for block sizes equal 1 and n equal total number of measurements, or if we group them into blocks.

The blocking method tells us that for block sizes smaller than the correlation time, the distance between correlated measurements, calculating the error using Eq. (4.50) will underestimate the error. When the block size is larger than the correlation time, the measurements are no longer correlated, and the error will no longer be affected by varying the block size. This becomes apparent as there will be a clear plateau when the data becomes uncorrelated. When the error reaches the plateau we have found an estimate for the true error,

A possible problem with relying on blocking is if the plateau is reached at a point where the number of blocks is too small to accurately estimate the variance. In this thesis we are lucky enough for that not to occur.

# Chapter 5

# Wave Functions

As the expectation values calculated with VMC depend on the chosen trial function, the results rely heavily on an educated guess. In this chapter we will examine the wave functions used for the Variational Monte Carlo method, and why they are good choices. This chapter will also introduce a correlation part, the Jastrow factor, to our trial wave function, since a pure Slater determinant is not enough to describe the inter-electronic correlations in an atomic system.

# 5.1 Cusp conditions and the Jastrow factor

The **cusp conditions** are constraints we get on the wave function when considering our system when distances between an electron and the core and between the electrons themselves become very small. The problems lie with the Coulomb force which depends on the inverse of the distance between charged particles. When these distances become small, we need some way to cancel the divergences that occur.

## 5.1.1 Single particle cusp conditions

In order to find the cusp conditions for the single particle wave functions, we start by examining the radial Schrödinger equation

$$-\frac{\hbar^2}{2m} \left( \frac{1}{r^2} \frac{d}{dr} r^2 \frac{d}{dr} - \frac{l(l+1)}{r^2} \right) R(r) + V(r)R(r) = ER(r), \tag{5.1}$$

where R(r) is the radial part of the full wave function. The dimensionless radial equation can be written as

$$\left(\frac{d^2}{d\rho^2} + \frac{2}{\rho}\frac{d}{d\rho} + \frac{2Z}{\rho} - \frac{l(l+1)}{\rho^2} + 2E\right)R(\rho) = 0,$$
(5.2)

with Z being the nuclear charge of the core. We now introduce

$$R(\rho) = \frac{1}{\rho}u(\rho) \tag{5.3}$$

as our radial wave function. This will give the radial equation as

$$\frac{d^2u(\rho)}{d\rho^2} + \frac{2Z}{\rho}u(\rho) - \frac{l(l+1)}{\rho^2}u(\rho) + 2Eu(\rho) = 0.$$
 (5.4)

As  $\rho$  approaches zero, the terms that dominate will be

$$\frac{d^2u(\rho)}{d\rho^2} - \frac{l(l+1)}{\rho^2}u(\rho) = 0.$$
 (5.5)

The general solution of this differential equation is

$$u(\rho) = C\rho^{l+1} + D\rho^{-l}. (5.6)$$

As  $\rho^{-l}$  will blow up as  $\rho$  approaches zero, the only physical solution we can have is then

$$u(\rho) \sim \rho^{l+1}.\tag{5.7}$$

In general we must assume that

$$R(\rho) = \rho^{l+1} \gamma(\rho), \tag{5.8}$$

where  $\gamma(\rho)$  is just some radial dependence function of  $\rho$  with the constraint that  $\gamma(0) \neq 0$ . For the kinetic energy not to diverge, we also need

$$\lim_{\rho \to 0} \frac{d^2 \gamma(\rho)}{d\rho^2} = \Omega,\tag{5.9}$$

where  $\Omega$  is finite. The dimensionless radial equation is now

$$\left(\frac{d^2}{d\rho^2} + \frac{2}{\rho}\frac{d}{d\rho} + \frac{2Z}{\rho} - \frac{l(l+1)}{\rho^2} + 2E\right)\rho^l\gamma(\rho) = 0, \tag{5.10}$$

By finding the derivatives

$$\frac{d^2}{d\rho^2} \left[ \rho^l \gamma(\rho) \right] = (l-1)l\rho^{l-2} \gamma(\rho) + 2l\rho^{l-1} \frac{d\gamma(\rho)}{d\rho} + \rho^l \frac{d^2\gamma(\rho)}{d\rho^2}$$
 (5.11)

and

$$\frac{2}{\rho} \frac{d}{d\rho} \left[ \rho^l \gamma(\rho) \right] = 2l\rho^{l-2} \gamma(\rho) + 2\rho^{l-1} \frac{d\gamma(\rho)}{d\rho}, \tag{5.12}$$

and inserting them into Eq. (5.10) while taking the limit when  $\rho \to 0$ , we will get

$$\gamma(\rho) \left[ \frac{l(l+1)}{\rho^2} + \frac{2Z}{\rho} - \frac{l(l+1)}{\rho^2} \right] + \frac{d\gamma(\rho)}{d\rho} \left[ \frac{2(l+1)}{\rho} \right] = 0 \tag{5.13}$$

as the terms that dominate. The second derivative from Eq. (5.9) is finite, as well as the term involving the energy E. By further simplification we will then have the cusp condition as

$$\frac{1}{\gamma(\rho)}\frac{d\gamma(\rho)}{d\rho} = -\frac{Z}{l+1} \tag{5.14}$$

which indicates the exponential behavior a wave function must have. The solution of equation Eq. (5.14) is

$$\gamma(\rho) \sim e^{-Z\rho/(l+1)}.\tag{5.15}$$

The hydrogenic wave functions and the Roothan-Hartree-Fock orbitals both fulfill the cusp condition.

#### 5.1.2 Correlation cusp conditions

To see how the wave function should behave with respect to correlation, we consider the helium system when the two electrons are very close, but far away from the core, that is,  $r_1 \neq 0$ ,  $r_2 \neq 0$  and  $r_{12} \to 0$ . Now the electron-nucleus terms in the Hamiltonian will be finite, while the repulsive electron-electron potential and the kinetic energy of the two electrons will diverge. When using centre-of-mass coordinates for electron 1 and 2, but neglecting the centre of mass motion since  $r_{12} \to 0$ , we will get a cusp condition also for the correlation part by using much of the same derivations as in section 5.1.1. The cusp condition for the two-particle part,  $\mathcal{R}(r_{12})$ , will then be

$$\frac{d\mathcal{R}(r_{12})}{dr_{12}} = \frac{1}{2(l+1)}\mathcal{R}(r_{12})$$
 (5.16)

as given in [7] and [4].

A correlation function that satisfies this cusp condition is the so-called Padé-Jastrow-function given by the two-particle functions

$$f(r_{ij}) = \exp\left(\frac{r_{ij}}{a(1+\beta r_{ij})}\right),\tag{5.17}$$

where  $\beta$  is a variational parameter. The factor a is a result of whether the intrinsic spins of particles i and j are parallel or not. That is

$$a = \begin{cases} 4 & \text{if spins are parallel} \\ 2 & \text{if spins are anti-parallel.} \end{cases}$$
 (5.18)

The full Padé-Jastrow function, J, for a system of N electrons will then be

$$J = \prod_{i < j}^{N} f(r_{ij}) = \prod_{i < j}^{N} \exp\left(\frac{r_{ij}}{a(1 + \beta r_{ij})}\right),$$
 (5.19)

which also satisfies the cusp condition.

# 5.2 Rewriting the Slater determinant

A general Slater determinant for an N-electron system will take the form:

$$\Phi(\mathbf{r}_{1}, \mathbf{r}_{2}, \dots, \mathbf{r}_{N}, \alpha, \beta, \dots, \nu) = \frac{1}{\sqrt{N!}} \begin{vmatrix} \psi_{\alpha}(\mathbf{r}_{1}) & \psi_{\alpha}(\mathbf{r}_{2}) & \cdots & \psi_{\alpha}(\mathbf{r}_{N}) \\ \psi_{\beta}(\mathbf{r}_{1}) & \psi_{\beta}(\mathbf{r}_{2}) & \cdots & \psi_{\beta}(\mathbf{r}_{N}) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_{\nu}(\mathbf{r}_{1}) & \psi_{\nu}(\mathbf{r}_{2}) & \cdots & \psi_{\nu}(\mathbf{r}_{N}) \end{vmatrix}, (5.20)$$

with the  $\psi$ 's being the available single particle orbitals. We could examine this more specifically by using the beryllium atom as an example. The Slater determinant

for beryllium is written as

$$\Phi(\mathbf{r}_{1}, \mathbf{r}_{2}, \mathbf{r}_{3}, \mathbf{r}_{4}, \alpha, \beta, \gamma, \delta) = \frac{1}{\sqrt{4!}} \begin{vmatrix} \psi_{\alpha}(\mathbf{r}_{1}) & \psi_{\alpha}(\mathbf{r}_{2}) & \psi_{\alpha}(\mathbf{r}_{3}) & \psi_{\alpha}(\mathbf{r}_{4}) \\ \psi_{\beta}(\mathbf{r}_{1}) & \psi_{\beta}(\mathbf{r}_{2}) & \psi_{\beta}(\mathbf{r}_{3}) & \psi_{\beta}(\mathbf{r}_{4}) \\ \psi_{\gamma}(\mathbf{r}_{1}) & \psi_{\gamma}(\mathbf{r}_{2}) & \psi_{\gamma}(\mathbf{r}_{3}) & \psi_{\gamma}(\mathbf{r}_{4}) \\ \psi_{\delta}(\mathbf{r}_{1}) & \psi_{\delta}(\mathbf{r}_{2}) & \psi_{\delta}(\mathbf{r}_{3}) & \psi_{\delta}(\mathbf{r}_{4}) \end{vmatrix},$$
(5.21)

with the quantum numbers representing different states as

$$|\mu\rangle = |n, l, m_l, \chi_{\pm}\rangle, \tag{5.22}$$

with  $|\alpha\rangle = |1,0,0\uparrow\rangle$ ,  $|\beta\rangle = |1,0,0\downarrow\rangle$ ,  $|\gamma\rangle = |2,0,0\uparrow\rangle$  and  $|\delta\rangle = |2,0,0\downarrow\rangle$ . The values  $\chi_{\pm}$  are given as  $\chi_{+} = \uparrow$  and  $\chi_{-} = \downarrow$  (see section 2.1.2).

By writing out the determinant in Eq. (5.21) explicitly, we would get zero, as the spatial wave functions for both spin up and spin down states are identical, and all terms will cancel each other out. We could also try to rewrite the Slater determinant as a product of a spin up-Slater determinant and a spin down-Slater determinant as

$$\Phi(\mathbf{r}_1, \mathbf{r}_2, \mathbf{r}_3, \mathbf{r}_4, \alpha, \beta, \gamma, \delta) = Det \uparrow (1, 2)Det \downarrow (3, 4) - Det \uparrow (1, 3)Det \downarrow (2, 4)$$
(5.23)

$$-Det \uparrow (1,4)Det \downarrow (3,2) + Det \uparrow (2,3)Det \downarrow (1,4) - Det \uparrow (2,4)Det \downarrow (1,3)$$

$$(5.24)$$

$$+ Det \uparrow (3,4)Det \downarrow (1,2), \tag{5.25}$$

where we have defined

$$Det \uparrow (1,2) = \frac{1}{\sqrt{2}} \begin{vmatrix} \psi_{100\uparrow}(\mathbf{r}_1) & \psi_{100\uparrow}(\mathbf{r}_2) \\ \psi_{200\uparrow}(\mathbf{r}_1) & \psi_{200\uparrow}(\mathbf{r}_2) \end{vmatrix}, \tag{5.26}$$

as a spin up-determinant, and

$$Det \downarrow (3,4) = \frac{1}{\sqrt{2}} \begin{vmatrix} \psi_{100\downarrow}(\mathbf{r}_3) & \psi_{100\downarrow}(\mathbf{r}_4) \\ \psi_{200\downarrow}(\mathbf{r}_3) & \psi_{200\downarrow}(\mathbf{r}_4) \end{vmatrix}, \tag{5.27}$$

as a spin down-determinant. However, writing out all these smaller determinants, will still give zero, but the strategy of separating the Slater determinant in one spin up and one spin down determinant is still pursued. The Hamiltonian does not depend on spin, so we wish to avoid summation over spin variables. As described in [14], we can actually approximate the Slater determinant for a variational approach with a plain product of a spin up- and a spin down-Slater determinant

$$\Phi(\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N) \propto Det \uparrow Det \downarrow,$$
 (5.28)

where the determinants  $Det \uparrow$  and  $Det \downarrow$  are  $(N/2) \times (N/2)$ -matrices for even numbered electron systems as examined in this project. We place the first N/2 electrons in the spin up determinant and the rest in the spin down determinant. This ansatz for the wave function is not antisymmetric under particle exchange as a wave function should be, but it will give the same expectiation value as the correct Slater determinant for such a spin independent Hamiltonian.

#### 5.3 Variational Monte Carlo wave function

By reviewing the last two sections we can write the full trial wave function for or Monte Carlo experiment. It can be written

$$\psi_T = \Phi \cdot J \tag{5.29}$$

where the Slater determinant  $\Phi$  can be approximated by

$$\Phi(\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N) \propto Det \uparrow Det \downarrow,$$
 (5.30)

and J is

$$J = \prod_{i < j}^{N} \exp\left(\frac{r_{ij}}{a(1 + \beta r_{ij})}\right). \tag{5.31}$$

For an N-particle system with M=N/2 possible spatial orbitals, the spin up determinant,  $Det \uparrow$ , can be written

$$Det \uparrow \propto \begin{vmatrix} \psi_{1\uparrow}(\mathbf{r}_1) & \psi_{1\uparrow}(\mathbf{r}_2) & \cdots & \psi_{1\uparrow}(\mathbf{r}_{N/2}) \\ \psi_{2\uparrow}(\mathbf{r}_1) & \psi_{2\uparrow}(\mathbf{r}_2) & \cdots & \psi_{2\uparrow}(\mathbf{r}_{N/2}) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_{M\uparrow}(\mathbf{r}_1) & \psi_{M\uparrow}(\mathbf{r}_2) & \cdots & \psi_{M\uparrow}(\mathbf{r}_{N/2}) \end{vmatrix},$$
(5.32)

while the spin down determinant can be written

$$Det \downarrow \propto \begin{vmatrix} \psi_{1\downarrow}(\mathbf{r}_{N/2+1}) & \psi_{1\downarrow}(\mathbf{r}_{N/2+2}) & \cdots & \psi_{1\downarrow}(\mathbf{r}_{N}) \\ \psi_{2\downarrow}(\mathbf{r}_{N/2+1}) & \psi_{2\downarrow}(\mathbf{r}_{N/2+2}) & \cdots & \psi_{2\downarrow}(\mathbf{r}_{N}) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_{M\downarrow}(\mathbf{r}_{N/2+1}) & \psi_{M\downarrow}(\mathbf{r}_{N/2+2}) & \cdots & \psi_{M\downarrow}(\mathbf{r}_{N}) \end{vmatrix} .$$
 (5.33)

These are the expressions needed for our trial wave function. The next two sections describe the single particle orbitals, the  $\psi$ 's, we use in our Slater determinant.

#### 5.4 Orbitals for VMC

In order to construct a trial Slater determinant for the Variational Monte Carlo method, our first choice is to use the hydrogenic single particle functions obtained when solving the Schrödinger equation for hydrogen. These functions comply with the cusp conditions described in section 5.1.1.

#### 5.4.1 S-orbitals

An s-orbital is a spatial wave function with quantum number l=0. Introducing the variational parameter  $\alpha$ , the s-orbitals needed for calculating up to at least silicon, with N=14 electrons, are 1s, 2s and 3s. Here we only need the radial

part of the wave functions because the constant angular parts factor out in the Metropolis algorithm (see section 4.4.1). The spherical harmonics for l = 0 is

$$Y_0^0 = \left(\frac{1}{4\pi}\right)^{1/2}. (5.34)$$

We now don't have to consider changing from spherical to Cartesian coordinates because the expression will remain the same using the simple relation  $r = \sqrt{x^2 + y^2 + z^2}$ . The hydrogenic radial functions are found in table 4.7 in [1] where non-vanishing factors(cf. Metropolis algorithm) 1/a have been replaced by  $\alpha$ .

#### The 1s wave function

For the 1s function we only need the exponential function

$$\phi_{1s} = e^{-r\alpha},\tag{5.35}$$

with the first derivative with respect to an arbitrary coordinate,  $x_i$ , given as:

$$\frac{\partial \phi_{1s}}{\partial x_i} = -\frac{\alpha x_i}{r} e^{-\alpha r}.$$
 (5.36)

The Laplacian,  $\nabla^2 = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2}$ , needed for the kinetic energy is given as

$$\nabla^2 \phi_{1s} = \frac{\alpha}{r} (\alpha r - 2) e^{-\alpha r}. \tag{5.37}$$

#### The 2s wave function

The 2s wave function is given by

$$\phi_{2s} = \left(1 - \frac{\alpha r}{2}\right) e^{-\alpha r/2},\tag{5.38}$$

with the first derivative wrt.  $x_i$  is

$$\frac{\partial \phi_{2s}}{\partial x_i} = \frac{\alpha x_i}{2r} \left( 2 - \frac{\alpha r}{2} \right) e^{-\alpha r/2}.$$
 (5.39)

The **second derivative** of the 2s function gives the Laplacian of the function:

$$\nabla^2 \phi_{2s} = \frac{\alpha}{4r} \left( 5\alpha r - \frac{\alpha^2 r^2}{2} - 8 \right) e^{-\alpha r/2}.$$
 (5.40)

#### The 3s wave function

The final s-function is the 3s wave function. It is

$$\phi_{3s} = \left(1 - \frac{2\alpha r}{3} + \frac{2\alpha^2 r^2}{27}\right) e^{-\alpha r/3}.$$
 (5.41)

The first derivative is

$$\frac{\partial \phi_{3s}}{\partial x_i} = \left(-1 + \frac{10\alpha r}{27} - \frac{2\alpha^2 r^2}{81}\right) \alpha x_i e^{-\alpha r/3} \tag{5.42}$$

while the second derivative summed up in all three dimensions is

$$\nabla^2 \phi_{3s} = \left(-2 + \frac{13\alpha r}{9} - \frac{2\alpha^2 r^2}{9} + \frac{2\alpha^3 r^3}{243}\right) \frac{\alpha}{r} e^{-\alpha r/3}.$$
 (5.43)

#### 5.4.2 P-orbitals

The *p*-orbitals are the single particle functions which have **azimuthal quantum number** l = 1. Now the allowed values for the **magnetic quantum number** are m = -l, -l + 1, ..., l - 1, l, i.e. m = -1, 0, 1. This is only possible for functions with quantum numbers n = 1, 2, 3, ... In this thesis we only need *s*- and *p*-orbitals.

The angular parts for any p-orbital in spherical coordinates are

$$Y_1^0 = \left(\frac{3}{4\pi}\right)^{1/2} \cos \theta, \tag{5.44}$$

$$Y_1^{\pm 1} = \pm \left(\frac{3}{8\pi}\right)^{1/2} \sin\theta e^{\pm i\varphi}.$$
 (5.45)

In my program however, I am working with real wave functions and Cartesian coordinates. This is taken care of by introducing real solid harmonics,  $S_{lm}$ , as described in ref. [2]. The single particle wave functions,  $\psi_{nlm}$ , are written on a separable form with spherical coordinates as

$$\psi(r,\theta,\varphi)_{nlm} = R_{nl}(r)Y_{lm}(\theta,\varphi). \tag{5.46}$$

These are general solutions of the one-electron system in a central symmetric potential field. Our radial functions are on the form  $R_{nl}(r) = r^l \mathcal{R}_{nl}(r)$ . The solid harmonics,  $\mathcal{Y}$ , are related to the spherical harmonics by

$$\mathcal{Y}_{lm}(r,\theta,\varphi) = r^l Y_{lm}(\theta,\varphi). \tag{5.47}$$

We can then write the wave function as

$$\psi(r,\theta,\varphi)_{nlm} = R_{nl}(r)Y_{lm}(\theta,\varphi) = \mathcal{R}_{nl}(r)\mathcal{Y}_{lm}(r,\theta,\varphi). \tag{5.48}$$

Now there is an r-dependence in the angular part. We will only need the real-valued solid harmonics,  $S_{lm}$ , related to the solid harmonics by

$$S_{l,0} = \sqrt{\frac{4\pi}{2l+1}} \mathcal{Y}_{l,0} \tag{5.49}$$

$$S_{l,m} = (-1)^m \sqrt{\frac{8\pi}{2l+1}} \text{Re } \mathcal{Y}_{l,m},$$
 (5.50)

and

$$S_{l,-m} = (-1)^m \sqrt{\frac{8\pi}{2l+1}} \text{Im } \mathcal{Y}_{l,m}.$$
 (5.51)

The real solid harmonics needed for us are simply  $S_{1,1} = x$ ,  $S_{1,0} = z$  and  $S_{1,-1} = y$ . As mentioned earlier, the Metropolis algorithm will take care of any multiplicative factors in front of the wave functions, so we only need to consider one single spatial variable from the angular part of the wave function.

#### The 2p wave function

The expression for the 2p wave function in spherical coordinates is proportional to

$$R_{21} \propto re^{-\alpha r/2}. (5.52)$$

By reviewing Eq. (5.47) and using the results from the real solid harmonics, we see the factor  $r^l = r$  is replaced by a coordinate  $x_i$ . That is,  $x_i = x, y, z$  depending on the quantum number m. The general case is then

$$\phi_{2p}(m) = x_i(m)e^{-\alpha r/2}. (5.53)$$

with  $x_i(1) = x$ ,  $x_i(0) = z$  and  $x_i(-1) = y$ .

The first derivative depends on whether the coordinate being differentiated with respect to matches the coordinate  $x_i$  of the wave function or not. If these are the same, e.g.  $m = 1 \rightarrow x_i = x$ , the **first derivative** is

$$\frac{\partial \phi_{2p}}{\partial x} = \left(1 - \frac{\alpha x^2}{2r}\right) \frac{e^{-\alpha r/2}}{r}.$$
 (5.54)

The expressions are analogous for y and z also.

If these coordinates do not match, e.g.  $\phi_{2p}(1) = x \exp{-\alpha r/2}$  and the derivative is with respect to y, the expression is

$$\frac{\partial \phi_{2p}(1)}{\partial y} = -\frac{\alpha xy}{r} e^{-\alpha r/2}.$$
 (5.55)

As for the first derivative, the second derivative also depends on which coordinate we differentiate with respect to. If the coordinates are the same, e.g.  $m = 1 \rightarrow x_i = x$ , the **second derivative** is

$$\frac{\partial^2 \phi_{2p}}{\partial x^2} = \frac{\alpha}{r} \left( \frac{x^3}{2r} \left( \frac{1}{r} + \frac{\alpha}{2} \right) - \frac{3x}{2} \right) e^{-\alpha r/2},\tag{5.56}$$

while for different coordinates, e.g.  $\phi_{2p}(1) = x \exp{-\alpha r/2}$  and the **second** derivative is with respect to y, the expression is

$$\frac{\partial^2 \phi_{2p}(1)}{\partial y^2} = \frac{\alpha x}{2r^3} \left( \frac{\alpha r y^2}{2} - r^2 + y^2 \right) e^{-\alpha r/2}.$$
 (5.57)

The final Laplacian is a sum of Eqs. (5.56) and (5.57) depending on which l-state is being used.

#### The 3p wave function

The 3p radial wave function is proportional to

$$R_{31} \propto \left(1 - \frac{r\alpha}{6}\right) \alpha r e^{-\alpha r/3}.$$
 (5.58)

By using the real solid harmonics for the full single particle wave function, we get our expression for the 3p wave function;

$$\phi_{3p}(m) = x_i(m) \left( 1 - \frac{\alpha r}{6} \right) \alpha e^{-\alpha r/3}, \tag{5.59}$$

with  $x_i(1) = x$ ,  $x_i(0) = z$  and  $x_i(-1) = y$ . The first derivative when differentiating wrt. to the same coordinate  $x_i$ , e.g. x, is

$$\frac{\partial \phi_{3p}(1)}{\partial x} = \left(-\left(1 - \frac{\alpha r}{6}\right)\left(\frac{\alpha^2 x^2}{3r} + \alpha\right) - \frac{\alpha^2 x^2}{6r}\right)e^{-\alpha r/3} \tag{5.60}$$

while for different coordinates, e.g. x and y, it is

$$\frac{\partial \phi_{3p}(1)}{\partial y} = \left(-\left(1 - \frac{\alpha r}{6}\right) \frac{\alpha^2 yx}{3r} - \frac{\alpha^2 xy}{6r}\right) e^{-\alpha r/3} \tag{5.61}$$

and

$$\frac{\partial \phi_{3p}(1)}{\partial z} = \left(-\left(1 - \frac{\alpha r}{6}\right) \frac{\alpha^2 zx}{3r} - \frac{\alpha^2 zx}{6r}\right) e^{-\alpha r/3} \tag{5.62}$$

The **second derivative** with respect to the coordinate from the solid harmonic, e.g. x, is

$$\frac{\partial^2 \phi_{3p}(1)}{\partial x^2} = \left( \left( 1 - \frac{\alpha r}{6} \right) \left( \frac{x^2}{3r^2} + \frac{\alpha x^2}{9r} - 1 \right) + \frac{\alpha x^2}{9r} + \frac{x^2}{6r^2} - \frac{1}{2} \right) \frac{\alpha^2 x}{r} e^{-\alpha r/3}, \quad (5.63)$$

while the **second derivative** with respect to a different coordinate is

$$\frac{\partial^2 \phi_{3p}(1)}{\partial y^2} = \left( \left( 1 - \frac{\alpha r}{6} \right) \left( \frac{y^2}{3r^2} + \frac{\alpha y^2}{9r} - \frac{1}{3} \right) + \frac{\alpha y^2}{9r} + \frac{y^2}{6r^2} - \frac{1}{6} \right) \frac{\alpha^2 x}{r} e^{-\alpha r/3}. \quad (5.64)$$

and

$$\frac{\partial^2 \phi_{3p}(1)}{\partial z^2} = \left( \left( 1 - \frac{\alpha r}{6} \right) \left( \frac{z^2}{3r^2} + \frac{\alpha z^2}{9r} - \frac{1}{3} \right) + \frac{\alpha z^2}{9r} + \frac{z^2}{6r^2} - \frac{1}{6} \right) \frac{\alpha^2 x}{r} e^{-\alpha r/3}. \quad (5.65)$$

By combining these we get the Laplacian of the wave function,  $\nabla^2 \phi_{3p}(1)$ . The expression for this particular single partice wave function is

$$\nabla^2 \phi_{3p}(1) = \frac{\partial^2 \phi_{3p}(1)}{\partial x^2} + \frac{\partial^2 \phi_{3p}(1)}{\partial y^2} + \frac{\partial^2 \phi_{3p}(1)}{\partial z^2}.$$
 (5.66)

These are all the explicit expressions needed for the Slater determinant with hydrogen-like wave functions.

# 5.5 Roothaan-Hartree-Fock with Slater-type orbitals

Another approach for the single particle basis is to use the single particle wave functions obtained using the Hartree-Fock method discussed in section 3.2.1. The Slater determinant is of the form

$$\Phi = \frac{1}{\sqrt{N!}} \begin{vmatrix} \phi_1(r_1) & \phi_1(r_2) & \cdots & \phi_1(r_N) \\ \phi_2(r_1) & \phi_2(r_2) & \cdots & \phi_2(r_N) \\ \vdots & \vdots & \ddots & \vdots \\ \phi_N(r_1) & \phi_N(r_2) & \cdots & \phi_N(r_N) \end{vmatrix},$$
(5.67)

where the functions  $\phi$  are the single particle states. The Roothaan-Hartree-Fock approach (see [5]) is to choose *Slater type orbitals* (STO) as the basis functions in our wave function expansion. The STO are disussed in [2] and in [5]. In this work I have used the exact form from [5]. These are given as

$$\phi_{i\lambda\alpha} = \sum_{p} \psi_{p\lambda\alpha} C_{i\lambda p}, \tag{5.68}$$

where  $\lambda$  corresponds to quantum number l. The index i refers to the  $i^{th}$  orbital of symmetry  $\lambda$ , and p to the  $p^{th}$  orbital of symmetry  $\lambda$ . The index  $\alpha$  corresponds to quantum number m. The **Slater functions**,  $\psi$ , are on the form

$$\psi_{p\lambda\alpha}(r,\theta,\varphi) = R_{\lambda p}(r)Y_{\lambda\alpha}(\theta,\varphi), \tag{5.69}$$

where

$$R_{\lambda p} = [(2n_{\lambda p})!]^{-1/2} (2\xi_{\lambda p})^{n_{\lambda p} + 1/2} r^{n_{\lambda p} - 1} e^{-\xi_{\lambda p} r}$$
(5.70)

and Y are the normalized spherical harmonics. The solutions of this particular approach are given in appendix A, taken from [5].

By using the idea of real solid harmonics (see section 5.4) and the fact that all basis functions  $\psi$  in  $\phi$  will have the same index  $\alpha$ , hence the same spherical harmonics, we can ignore the pre-factor from the spherical harmonics (see section 4.4.1). Now the basis functions  $\psi$  will go from the form

$$\psi_i = N_i r^{n-1} \exp(-\xi r) Y_{\lambda \alpha} \tag{5.71}$$

in spherical coordinates, to

$$\psi_i = N_i x_i r^{n-2} \exp(-\xi r) \tag{5.72}$$

in Cartesian coordinates, where the factor  $N_i$  is the normalization constant from Eq. (5.70). The results from helium, beryllium, neon, magnesium and silicon are listed in appendix A. As an example on how to construct the single particle orbitals from the tables in appendix A, I use the example given in [5]:

$n,\lambda$	Exponent, $\xi$	1s exp.coeff.	2s exp.coeff.	$n,\lambda$	Exponent, $\xi$	2p exp.coeff.
1S	6.56657	0.19030	0.00754	2P	2.21734	0.21526
1S	4.24927	0.82091	-0.25055	2P	1.00551	0.84052
2S	1.41314	-0.00364	0.87099			
2S	0.87564	0.00251	0.18515			

Table 5.1: The table shows a possible solution for the boron atom.

Consider table 5.1 as a solution of the boron atom. Boron has five electrons distributed as  $(1s)^2(2s)^2(2p)$ . This means we need 1s-, 2s- and 2p-orbitals. The first column tells us how many and which s-type Slater functions the 1s- and 2s-orbitals for the Slater determinant will consist of. In this case it will be two of each type. The second column gives us the factors,  $\xi$ , in the exponents of the exponential functions. These will be the same for both 1s- and 2s, and will also contribute to the prefactors  $N_i$ . The third column gives the coefficients for the 1s single particle orbital for the Slater determinant, while the fourth column gives the coefficients for the 2s single particle orbital. The single particle orbitals for 1s and 2s will then only differ by the constants in front of the Slater functions (the  $\psi$ 's from Eq. (5.68)).

The fifth column indicates that there are only two basis functions needed to describe the p-orbital, namely the 2p single particle orbital, while the sixth and seventh give the exponents and 2p-coefficients respectively.

The single particle orbitals are given in table 5.1 as

$$\phi(1s) = 0.19030\psi_1 + 0.82091\psi_2 - 0.00364\psi_3 + 0.00251\psi_4 \tag{5.73}$$

$$\phi(2s) = 0.00754\psi_1 - 0.25055\psi_2 + 0.87099\psi_3 + 0.18515\psi_4 \tag{5.74}$$

$$\phi(2p) = 0.21526\psi_5 + 0.84052\psi_6 \tag{5.75}$$

where now  $\psi_1$  and  $\psi_2$  are Slater functions of type 1s,  $\psi_3$  and  $\psi_4$  of type 2s, and  $\psi_5$  and  $\psi_6$  Slater functions of type 2p. For larger atoms, the p-orbitals will be linear combinations of different p-type Slater functions. The s- and p-type Slater functions do not mix at any point, seen from Eq. (5.68).

The Slater functions themselves written in Cartesian coordinates are given as

$$\psi_1 = N_1 r^0 \exp(-6.56657r) \tag{5.76}$$

$$\psi_2 = N_2 r^0 \exp(-4.24927r) \tag{5.77}$$

$$\psi_3 = N_3 r^1 \exp(-1.41314r) \tag{5.78}$$

$$\psi_4 = N_4 r^1 \exp(-0.87564r) \tag{5.79}$$

$$\psi_5 = N_5 x_i \exp(-2.21734r) \tag{5.80}$$

$$\psi_6 = N_6 x_i \exp(-1.00551r) \tag{5.81}$$

where  $x_i$  corresponds to either x,y or z as described in section 5.4.2, and the factors  $N_i$  are the normalization factors from Eq. (5.70).

#### 5.5.1 Derivatives of Slater type orbitals

The analytic expressions needed for the Slater type orbitals are of the form

$$\psi_{1s} \propto \exp(-ar) \tag{5.82}$$

$$\psi_{2s} \propto r \exp(-ar) \tag{5.83}$$

$$\psi_{3s} \propto r^2 \exp(-ar) \tag{5.84}$$

$$\psi_{2p} \propto x_i \exp(-ar) \tag{5.85}$$

$$\psi_{4p} \propto r^2 x_i \exp(-ar) \tag{5.86}$$

where a is some exponent determined by the Roothan-Hartree-Fock calculations and  $x_i$  the coordinate from the real solid harmonics. In order to calculate the quantum force (see section 4.4.2) and the kinetic energy we need both the first and second derivatives of the wave functions.

#### First derivatives

The s-orbitals have first derivatives proportional to

$$\frac{\partial \psi_{1s}}{\partial x_i} \propto -ax_i \exp(-ar) \tag{5.87}$$

$$\frac{\partial \psi_{2s}}{\partial x_i} \propto (1 - ar) \frac{x_i}{r} \exp(-ar) \tag{5.88}$$

$$\frac{\partial \psi_{3s}}{\partial x_i} \propto (2x_i - rax_i) \exp(-ar), \tag{5.89}$$

while the 2p-orbitals are

$$\frac{\partial \psi_{2p}(x)}{\partial x} \propto \left(1 - \frac{x^2 a}{r}\right) \exp(-ar) \tag{5.90}$$

$$\frac{\partial \psi_{2p}(x)}{\partial y} \propto -\frac{xya}{r} \exp(-ar) \tag{5.91}$$

where the parenthesis, (x), indicates which real solid harmonics the function depends on. Equations (5.90) and (5.91) are examples with the solid harmonics being the x-coordinate, and the derivatives with respect to x and then y (see the discussion in 5.4.2). The 4p-orbitals are proportional to

$$\frac{\partial \psi_{4p}(x)}{\partial x} \propto \left(2x^2 - r^2 - rx^2 a\right) \exp(-ar) \tag{5.92}$$

$$\frac{\partial \psi_{4p}(x)}{\partial y} \propto (2xy - rxya) \exp(-ar) \tag{5.93}$$

#### Second derivatives

The Laplacian of the s-orbitals are on the form

$$\nabla^2 \psi_{1s} \propto \frac{a}{r} (ar - 2) \exp(-ar) \tag{5.94}$$

$$\nabla^2 \psi_{2s} \propto \left(\frac{2}{r} + a^2 r - 4a\right) \exp(-ar) \tag{5.95}$$

$$\nabla^2 \psi_{3s} \propto (6 - 6ar + a^2 r^2) \exp(-ar),$$
 (5.96)

while the Laplacians of the 2p-orbitals consist of the second derivatives

$$\frac{\partial^2 \psi_{2p}(x)}{\partial x^2} \propto \frac{ax}{r} \left( -3 + \frac{x^2}{r^2} + \frac{x^2 a}{r} \right) \exp(-ar) \tag{5.97}$$

$$\frac{\partial^2 \psi_{2p}(x)}{\partial y^2} \propto \frac{x}{r} \left( \frac{y^2}{r^2} - 1 + \frac{ay^2}{r} \right) \exp(-ar). \tag{5.98}$$

The 4p-orbitals have the second derivatives

$$\frac{\partial^2 \psi_{4p}(x)}{\partial x^2} \propto x \left( 6 - \frac{3x^2a}{r} + 3ar + x^2a^2 \right) \exp(-ar) \tag{5.99}$$

$$\frac{\partial^2 \psi_{4p}(x)}{\partial y^2} \propto x \left( 2 - \frac{3y^2 a}{r} - ar + a^2 y^2 \right) \exp(-ar). \tag{5.100}$$

The expressions for  $\partial^2 \psi_{4p}(y)/\partial y^2$ ,  $\partial^2 \psi_{4p}(z)/\partial y^2$  etc. are identical to the ones above and are used to construct e.g.

$$\nabla^2 \psi_{4p}(x) = \frac{\partial^2 \psi_{4p}(x)}{\partial x^2} + \frac{\partial^2 \psi_{4p}(x)}{\partial y^2} + \frac{\partial^2 \psi_{4p}(x)}{\partial z^2}.$$
 (5.101)

All these expressions are needed for the implementation of the Slater determinant, the quantum force and the kinetic energy in our VMC calculation.

# Part II Implementation and results

# Chapter 6

# Implementation of the Variational Monte Carlo Method

The Variational Monte Carlo method was implemented by writing a C++-based code. The first thing was to implement a brute force code to handle the helium atom, and which could easily be reduced to solve the simple hydrogen system, with only one atom and hence no inter-electron potentials. Brute force means here that there was no importance sampling, so the simple Metropolis algorithm (see figure 4.1) was used. The next step was to implement importance sampling, in order to make the sampling more efficient, and hopefully getting better results for both energy and variance. This would mean we had to replace the simple Metropolis algorithm with the Metropolis-Hastings algorithm, which includes importance sampling.

However, the code was not efficient at this stage. By calculating the explicit wave function for each particle move, and using numerical derivatives, there was still room for improvement. For larger systems like beryllium, neon etc., we would need to implement a more efficient wave function with closed-form expressions for the Slater determinant and its derivatives, using the definitions given in section 5.2, in addition to an efficient computation of the Jastrow factor. By doing this we could improve the efficiency of each Monte Carlo cycle, resulting in less computation time. If this optimization of the wave function was made general enough, and we included closed form expressions for the single particle wave functions and their derivatives, we could easily use the program to calculate an approximation to the ground state energy for basically any even-numbered atom.

In this chapter I will describe the optimized version of the program, since this is of most interest, and the part that has taken the most time to develop. I will also focus on the Metropolis-Hastings algorithm and not the more simple Metropolis algorithm.

Initially, the code was developed using object-oriented programming (OOP), but complications and little programming experience prevented me from fully exploiting the strengths of OOP. However, the main Monte Carlo machinery is built within a single class, so at least one can argue that this approach has left me with a little more organized code. I will therefore not emphasize on the code structure, but rather on the key points of importance for the calculations.

# 6.1 Optimizing the calculations

In order to reduce the time of the calculations, we had to find a way to both rewrite the Slater determinant and the Jastrow factor. When increasing the system size, calculating the entire wave function,

$$\psi_T = \Phi \cdot J = Det \uparrow \cdot Det \downarrow \cdot J, \tag{6.1}$$

for each particle move, seemed inefficient. The determinants are defined as

$$Det \uparrow \propto \begin{vmatrix} \psi_{1\uparrow}(\mathbf{r}_1) & \psi_{1\uparrow}(\mathbf{r}_2) & \cdots & \psi_{1\uparrow}(\mathbf{r}_{N/2}) \\ \psi_{2\uparrow}(\mathbf{r}_1) & \psi_{2\uparrow}(\mathbf{r}_2) & \cdots & \psi_{2\uparrow}(\mathbf{r}_{N/2}) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_{M\uparrow}(\mathbf{r}_1) & \psi_{M\uparrow}(\mathbf{r}_2) & \cdots & \psi_{M\uparrow}(\mathbf{r}_{N/2}) \end{vmatrix},$$
(6.2)

and

$$Det \downarrow \propto \begin{vmatrix} \psi_{1\downarrow}(\mathbf{r}_{N/2+1}) & \psi_{1\downarrow}(\mathbf{r}_{N/2+2}) & \cdots & \psi_{1\downarrow}(\mathbf{r}_{N}) \\ \psi_{2\downarrow}(\mathbf{r}_{N/2+1}) & \psi_{2\downarrow}(\mathbf{r}_{N/2+2}) & \cdots & \psi_{2\downarrow}(\mathbf{r}_{N}) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_{M\downarrow}(\mathbf{r}_{N/2+1}) & \psi_{M\downarrow}(\mathbf{r}_{N/2+2}) & \cdots & \psi_{M\downarrow}(\mathbf{r}_{N}) \end{vmatrix}.$$
(6.3)

for a system of N electrons with M=N/2 possible spatial wave functions. The Jastrow factor is given as

$$J = \prod_{i < j}^{N} f(r_{ij}) = \prod_{i < j}^{N} \exp\left(\frac{r_{ij}}{a(1 + \beta r_{ij})}\right),$$
 (6.4)

A crucial point is the fact that by moving one electron, as we do in the Metropolis algorithms (see sections 4.4.1 and 4.4.2), we actually only change one column in only **one** of the two Slater determinants. This should lead us to believe that an optimization should be obtainable.

We use the wave function,  $\psi_T$ , for:

• Calculating the transition probability in the Metropolis-Hastings algorithm as

$$w = \frac{G(x,y)|\psi_T(y)|^2}{G(y,x)|\psi_T(x)|^2}$$
(6.5)

where y are the new positions, and x are the old ones, and G(y, x) is the Green's function in Eq. (4.39).

• Calculating the quantum force,

$$F(x) = 2\frac{\nabla \psi_T}{\psi_T},\tag{6.6}$$

needed for the Green's function involving the gradient of the wave function.

• Calculating the local energy,

$$E_L = \frac{\nabla^2 \psi_T}{\psi T} - V \tag{6.7}$$

involving the Laplacian of the wave function.

This means that we need to optimize the ratios  $\Phi(r^{new})/\Phi(r^{old})$ ,  $\nabla\Phi(r)/\Phi(r)$ ,  $\nabla^2\Phi(r)/\Phi(r)$ ,  $J(r^{new})/J(r^{old})$ ,  $\nabla J/J$  and  $\nabla^2 J/J$ .

Since our wave function is on the form

$$\psi_T = \Phi \cdot J,\tag{6.8}$$

the total expression for the gradient ratio will be

$$\frac{\nabla \psi_T}{\psi_T} = \frac{\nabla \Phi}{\Phi} + \frac{\nabla J}{J}.\tag{6.9}$$

For the kinetic energy term in the local energy we will have to calculate the Laplacian of  $\psi_T$ , divided by  $\psi_T$  itself as

$$\frac{\nabla^2 \psi_T}{\psi_T} = \frac{\nabla^2 \Phi}{\Phi} + \frac{\nabla^2 J}{J} + \left(\frac{\nabla \Phi}{\Phi} \cdot \frac{\nabla J}{J}\right). \tag{6.10}$$

# 6.1.1 Optimizing the ratio - $\Psi_T(r^{new})/\Psi_T(r^{old})$

When optimizing the wave function part of the transition ratio, it's a good idea to split it into a correlation part, and a Slater determinant part. This means

$$R = \frac{\psi_T(r^{new})}{\psi_T(r^{old})} = R_{SD}R_J = \frac{\Phi(r^{new})}{\Phi(r^{old})} \frac{J(r^{new})}{J(r^{old})}$$
(6.11)

where  $\Phi$  is the Slater determinant, and J is the correlation part, the Jastrow factor. In this section I will label the positions of all particles before a particle move as  $r^{old}$ , and the set of particle positions after the move as  $r^{new}$ .

## Slater determinant ratio - $\Phi(\mathbf{r^{new}})\Phi(\mathbf{r^{old}})$

We start by defining the so-called Slater matrix,  $\mathcal{D}$ , the matrix used to calculate the Slater determinant, and its elements as  $D_{ij} = \phi_j(r_i)$ , where  $\phi_j(r_i)$  are electrons i occupying a single particle state j. The inverse of the Slater matrix,  $\mathcal{D}^{-1}$ , is related to the Slater determinant,  $\Phi$  or just |D|, as

$$\mathcal{D}^{-1} = \frac{\operatorname{adj}\mathcal{D}}{|D|} \tag{6.12}$$

or as matrix elements as

$$D_{ij}^{-1} = \frac{C_{ji}}{|D|},\tag{6.13}$$

where  $\operatorname{adj}\mathcal{D}$  is the so-called *adjugate* of a matrix. The adjugate of a matrix  $\mathcal{A}$  is the transposed of the co-factor matrix of  $\mathcal{A}$  (indices are interchanged from Eq. (6.12) to Eq. (6.13)). For the matrix  $\mathcal{D}$  to be invertible, we must have  $\mathcal{D}^{-1}\mathcal{D} = \mathcal{I}$ , where  $\mathcal{I}$  is the identity matrix, or written out as

$$\sum_{k=1}^{N} D_{ik} D_{kj}^{-1} = \delta_{ij}. {(6.14)}$$

Multiplying Eq. (6.13) with  $D_{ii}$  from the left and summing over i, will give

$$\sum_{i=1}^{N} D_{ji} D_{ij}^{-1} = \frac{\sum_{i=1}^{N} D_{ji} C_{ji}}{|D|},$$
(6.15)

resulting in

$$|D| = \sum_{i=1}^{N} D_{ji} C_{ji}. \tag{6.16}$$

We can now express the ratio between the new and old Slater determinants,  $R_{SD}$ , as

$$R_{SD} = \frac{\sum_{j=1}^{N} D_{ij}(r^{new}) C_{ij}(r^{new})}{\sum_{j=1}^{N} D_{ij}(r^{old}) C_{ij}(r^{old})}.$$
(6.17)

When moving a particle i, only the i-th row of the Slater matrix changes, and therefore only the i-th row of the matrix of co-factors remains unchanged, which means

$$C_{ij}(r^{new}) = C_{ij}(r^{old}). (6.18)$$

Using Eq. (6.13) gives

$$R_{SD} = \frac{\sum_{j=1}^{N} D_{ij}(r^{new}) C_{ij}(r^{old})}{\sum_{j=1}^{N} D_{ij}(r^{old}) C_{ij}(r^{old})} = \frac{\sum_{j=1}^{N} D_{ij}(r^{new}) D_{ji}^{-1}(r^{old})}{\sum_{j=1}^{N} D_{ij}(r^{old}) D_{ji}(r^{old})}.$$
 (6.19)

This denominator will only give unity, so the ratio can be expressed as

$$R_{SD} = \sum_{j=1}^{N} D_{ij}(r^{new}) D_{ji}^{-1}(r^{old}) = \sum_{j=1}^{N} \phi_j(r_i^{new}) D_{ji}^{-1}(r^{old}).$$
 (6.20)

When only moving the *i*-th particle, we only need to calculate a dot product,  $\phi(r_i^{new}) \cdot D_i^{-1}$ , where  $\phi(r_i^{new})$  is the vector

$$\phi(r_i^{new}) = (\phi_1(r_i^{new}), \phi_2(r_i^{new}), \dots, \phi_N(r_i^{new}))$$
(6.21)

and  $D_i^{-1}$  is the *i*-th column of the inverse matrix  $\mathcal{D}^{-1}$  where the electron *i* is at the old position. We also have splitted the determinants in two separate determinants, so we actually only change **one** row in **one** of the Slater determinants when a particle *i* is moved.

The following code shows the function that calculates the ratio  $R_{SD}$ :

Here the function phi(r,alpha,beta,j,i) calculates the single particle wave function for electron i in orbital j.

As described in [7] we can use an algorithm to update the inverse of the Slater matrix without calculating the entire Slater matrix again. After moving particle i, and if the move is accepted, we must for each column  $j \neq i$  calculate a quantity

$$S_j = \sum_{l=1}^{N} D_{il}(r^{new}) D_{lj}(r^{old}).$$
 (6.22)

The new elements of  $\mathcal{D}^{-1}$  are as given on page 277 in [7]:

$$D_{kj}^{-1}(r^{new}) = \begin{cases} D_{kj}^{-1}(r^{old}) - \frac{S_j}{R_{SD}} D_{ki}^{-1}(r^{old}) & \text{if } j \neq i \\ \frac{1}{R_{SD}} D_{kj}^{-1} & \text{if } j = i. \end{cases}$$
(6.23)

The following code shows how we first update the spin up determinant, and then the spin down determinant

```
void atom::update(double **D_up, double **D_down, int i, double ratio, double
    alpha, double beta){

//SPIN UP ELECTRONS
if(i<no_of_particles/2){
    for(int j=0; j<no_of_particles/2; j++){
        if(j!=i){
    for(int l=0; l<no_of_particles/2; l++){
        s_j[j]+=phi(r_new,alpha,beta,l,i)*D_up[l][j];
    }
    }
    for(int j=0; j<no_of_particles/2; j++){
        if(j!=i){
        for(int k=0; k<no_of_particles/2; k++){
            D_up[k][j]=D_up[k][j]-s_j[j]*D_up[k][i]/ratio;
    }
    }
}</pre>
```

```
//i'th column, i=j
   for(int k=0; k<no_of_particles/2; k++){</pre>
     D_up[k][i]=D_up[k][i]/ratio;
 }
//SPIN DOWN ELECTRONS
 else{
   i=i-no_of_particles/2;//This is because the spin-down matrix have indices
       0,1,\ldots, N/2
   for(int j=0; j<no_of_particles/2; j++){</pre>
     if(j!=i){
 for(int l=0; l<no_of_particles/2; l++){</pre>
   s_j[j]+=phi(r_new,alpha,beta,1,i+no_of_particles/2)*D_down[1][j];
   }
   for(int j=0; j<no_of_particles/2; j++){</pre>
     if(j!=i){
 for(int k=0; k<no_of_particles/2; k++){</pre>
   D_down[k][j]=D_down[k][j]-s_j[j]*D_down[k][i]/ratio;
   //i'th column, i=j
   for(int k=0; k<no_of_particles/2; k++){</pre>
     D_down[k][i]=D_down[k][i]/ratio;
   i=i+no_of_particles/2;
```

## Jastrow ratio - $J(r^{new})/J(r^{old})$

We now have to find a functional way to store the inter-electron distances, or for simplicity, the individual Jastrow functions,

$$g_{i,j} = g(r_{ij}) = \exp\left(\frac{r_{ij}}{a(1+\beta r_{ij})}\right), \tag{6.24}$$

between the different electrons i and j, where a is given by

$$a = \begin{cases} 4 & \text{if spins are parallel} \\ 2 & \text{if spins are anti-parallel.} \end{cases}$$
 (6.25)

From section 5.1.2, we have our Jastrow factor, J, on the form

$$J = \prod_{i < j} g_{i,j}. \tag{6.26}$$

A way to store these functions is to construct a matrix with elements above the diagonal as:

$$\mathcal{G} = \begin{bmatrix} \cdot & g_{1,2} & g_{1,3} & \cdots & g_{1,N} \\ & \cdot & g_{2,3} & \cdots & g_{2,N} \\ & & \cdot & \ddots & \vdots \\ & & \cdot & g_{N-1,N} \end{bmatrix}.$$
(6.27)

The Jastrow ratio,  $R_J$  is

$$R_J = \frac{J(r^{new})}{J(r^{old})}. (6.28)$$

From section 5.1.2, our Jastrow factor, J, is on the form

$$J = \prod_{i < j} g_{i,j} = \prod_{i < j} \exp f_{i,j}, \tag{6.29}$$

and when moving only particle k, only the elements with k as an index will change, that is N-1 elements. Because of our exponential form of the correlation part the ratio can be written

$$\frac{J(r^{new})}{J(r^{old})} = e^{\Delta J},\tag{6.30}$$

where the intriguing part is  $\Delta J$ , given as

$$\Delta J = \sum_{i=1}^{k-1} (f_{i,k}^{new} - f_{i,k}^{old}) + \sum_{i=k+1}^{N} (f_{k,i}^{new} - f_{k,i}^{old}), \tag{6.31}$$

with the functions  $f_{ij}$  given as

$$f_{ij} = \frac{r_{ij}}{a(1 + \beta r_{ij})}. (6.32)$$

The following code shows the calculation of the Jastrow factor ratio:

```
double atom::getJastrowRatio(double** distance_old, double**
    distance_new,double beta){
    for(int k=0; k<no_of_particles; k++){
        for(int l=0; l<k; l++){
            jastrowRatio += distance_new[l][k]-distance_old[l][k];
        }
    }
    for(int k=0; k<no_of_particles; k++){
        for(int l=k+1; l<no_of_particles; l++){
            jastrowRatio += distance_new[l][k]-distance_old[l][k];
        }
    return jastrowRatio;
}</pre>
```

The two-dimensional arrays distance\_new and distance\_old have been calculated using:

```
void atom::getDistance(double** distance, double** r_old, double beta){
  for(int k=0; k<no_of_particles; k++){
    for(int l=k+1; l<no_of_particles; l++){
       temp = diffR(r_old, k,1);
       //spin up
       if(((k < no_of_particles/2) && (1 <no_of_particles/2)) ||
            ((k>=no_of_particles/2 && l>=no_of_particles/2))){
       a=0.25;
       distance[k][l] = a*temp/(1+beta*temp);
       }
       //spin down
       else{
    a=0.5;
       distance[k][l] = a*temp/(1+beta*temp);
       }
    }
}
```

We see that the names of the arrays are a bit misleading, as we actually store the values  $f(r_{ij})$  (see Eq. (6.32)) and not  $r_{ij}$  themselves.

#### 6.1.2 Derivative ratios

In order to optimize the calculations, we need to find optimized expressions for the ratios  $\nabla \Phi/\Phi$ ,  $\nabla^2 \Phi/\Phi$ ,  $\nabla J/J$  and  $\nabla^2 J/J$ , as seen in Eqs. (6.9) and (6.10). This will lead to a more efficient evaluation of the quantum force and the kinetic energy.

#### Optimizing the ratios - $\nabla\Phi/\Phi$ and $\nabla^2\Phi/\Phi$

We use a similar procedure as in the previous section to get the explicit expressions for the gradient and Laplacian of the Slater determinant. When differentiating with respect to the coordinates of a single particle i, we will get

$$\frac{\nabla_i \Phi}{\Phi} = \sum_{j=1}^N \nabla_i D_{ij}(r) D_{ji}^{-1}(r) = \sum_{j=1}^N \nabla_i \phi_j(r_i) D_{ji}^{-1}(r)$$
 (6.33)

for the gradient, remembering that the operator  $\nabla$  actually is a vector

$$\nabla = \frac{\partial}{\partial x}\hat{i} + \frac{\partial}{\partial y}\hat{j} + \frac{\partial}{\partial z}\hat{k}, \tag{6.34}$$

with  $\hat{\imath}$ ,  $\hat{\jmath}$  and  $\hat{k}$  are the unit vectors for x-, y- and z- dimensions respectively. For the Laplacian-expression we will obtain

$$\frac{\nabla_i \Phi}{\Phi} = \sum_{j=1}^N \nabla_i^2 D_{ij}(r) D_{ji}^{-1}(r) = \sum_{j=1}^N \nabla_i^2 \phi_j(r_i) D_{ji}^{-1}(r), \tag{6.35}$$

where  $\nabla^2$  is a scalar

$$\nabla^2 = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2}.$$
 (6.36)

We see that the expressions only rely on the elements from the inverse Slater matrix,  $\mathcal{D}^{-1}$ , and the closed form expressions for the derivatives of the single particle orbitals (see sections 5.4 and 5.5).

The following code shows the calculation of the ratio  $\nabla^2 \Phi/\Phi$  in the local energy function:

```
//LAPLACE SLATER-DETERMINANT ANALYTIC
for(int i=0; i<no_of_particles; i++){
   if(i<no_of_particles/2){
      for(int j=0; j<no_of_particles/2; j++){
      E_kinetic -= laplace_phi(r,alpha,beta,j,i)*D_up[j][i];
      }
   }
   else{
      for(int j=0; j<no_of_particles/2; j++){
      E_kinetic -= laplace_phi(r,alpha,beta,j,i)*D_down[j][i-no_of_particles/2];
      }
   }
}</pre>
```

The function laplace\_phi(r,alpha,beta,j,i) simply returns the Laplacian of the single particle wave function where electron i is in orbital j.

#### Optimizing the ratios - $\nabla J/J$ and $\nabla^2 J/J$

The **first derivative** results for the Padé-Jastrow function (as described in Eq. (6.26)) are taken from chapter 19 in [4] and is for a single coordinate, x, for particle k, given by

$$\frac{1}{J}\frac{\partial J}{\partial x_k} = \sum_{i=1}^{k-1} \frac{1}{g_{ik}} \frac{\partial g_{ik}}{\partial x_k} + \sum_{i=k+1}^{N} \frac{1}{g_{ki}} \frac{\partial g_{ki}}{\partial x_k}, \tag{6.37}$$

where  $g_{ij}$  is some correlation function between two particles. We have

$$g_{ij} = \exp f(r_{ij}) = \exp \left(\frac{r_{ij}}{a(1+\beta r_{ij})}\right), \tag{6.38}$$

a function of distances between the electrons. We now use the chain rule to get

$$\frac{\partial g_{ij}}{\partial x_j} = \frac{\partial g_{ij}}{\partial r_{ij}} \frac{\partial r_{ij}}{\partial x_j} = \frac{x_j - x_i}{r_{ij}} \frac{\partial g_{ij}}{\partial r_{ij}}.$$
(6.39)

The exponential form of  $g_{ij}$  also gives us

$$\frac{\partial g_{ij}}{\partial r_{ij}} = g_{ij} \frac{\partial f_{ij}}{\partial r_{ij}},\tag{6.40}$$

so that

$$\frac{1}{J}\frac{\partial J}{\partial x_k} = \sum_{i=1}^{k-1} \frac{\partial f_{ik}}{\partial x_k} + \sum_{i=k+1}^{N} \frac{\partial f_{ki}}{\partial x_k}.$$
 (6.41)

It is easy to show that

$$\frac{\partial f_{ij}}{\partial r_{ij}} = \frac{1}{a(1+\beta r_{ij})^2},\tag{6.42}$$

and combining these results we will get

$$\frac{1}{J}\frac{\partial J}{\partial x_k} = \sum_{i=1}^{k-1} \frac{x_k - x_i}{ar_{ik}(1 + \beta r_{ik})^2} + \sum_{i=k+1}^{N} \frac{x_k - x_i}{ar_{ik}(1 + \beta r_{ik})^2},$$
(6.43)

as the final result for the first derivative with respect to a chosen coordinate x, and particle k. The quantum force for a given particle k in coordinate x,

$$F(x_k) = \frac{2}{J} \frac{\partial J}{\partial x_k} = 2 \left( \sum_{i=1}^{k-1} \frac{x_k - x_i}{ar_{ik}(1 + \beta r_{ik})^2} + \sum_{i=k+1}^{N} \frac{x_k - x_i}{ar_{ik}(1 + \beta r_{ik})^2} \right).$$
 (6.44)

The following code shows the calculation of the quantum force with the first part focusing on the Slater determinant part discussed in the previous section, while the second part focuses on the correlation part of the quantum force (see Eq. (6.9)):

```
void atom::quantum_force(double**r,double**qm_force, double **D_up, double
    **D_down, double alpha, double beta){
  //SLATER PART
 for(int p=0; p<no_of_particles; p++){</pre>
   if(p<no_of_particles/2){</pre>
     for(int q=0; q<dimension; q++){</pre>
 for(int l=0; l<no_of_particles/2; l++){</pre>
   qm_force[p][q] += 2*gradient_phi(r,alpha,q,1,p)*D_up[l][p];
   else{
     for(int q=0; q<dimension; q++){</pre>
 for(int l=0; l<no_of_particles/2; l++){</pre>
   qm_force[p][q] +=
       2*gradient_phi(r,alpha,q,1,p)*D_down[1][p-no_of_particles/2];
 }
 //JASTROW PART
for(int p=0; p<no_of_particles; p++){</pre>
   for(int q=0; q<dimension; q++){</pre>
     for(int l=p+1; l<no_of_particles; l++){</pre>
 qm_force[p][q] += 2*gradient_jastrow(r,p,l,q,beta);
     for(int 1=0; 1<p; 1++){
 qm_force[p][q] += 2*gradient_jastrow(r,p,l,q,beta);
}//end quantum_force
```

The function gradient\_phi(r,alpha,q,l,p) returns the first derivative of the single particle orbital  $\phi_l$  with respect to coordinate q of electron p.

The function gradient\_jastrow(r,p,l,q,beta) is given as

The function diffR(r,p,1) simply returns the distance between electrons p and 1.

For the **second derivative**, I refer to [4] where it is shown that for a correlation function of our form

$$g_{ij} = \exp f_{ij}, \tag{6.45}$$

the full expression for the Laplacian of the correlation part can be written as

$$\frac{\nabla^2 J}{J} = \left(\frac{\nabla J}{J}\right)^2 + \sum_{i=1}^{k-1} \left(\frac{(d-1)}{r_{ik}} \frac{\partial f_{ik}}{\partial r_{ik}} + \frac{\partial^2 f_{ik}}{\partial r_{ik}^2}\right) + \sum_{i=k+1}^N \left(\frac{(d-1)}{r_{ki}} \frac{\partial f_{ki}}{\partial r_{ki}} + \frac{\partial^2 f_{ki}}{\partial r_{ki}^2}\right), \tag{6.46}$$

where the first term is just the dot product of the gradient (see the previous section) with itself, and d is the number of spatial dimensions, i.e. d=3. We have the explicit expression for the Padé-Jastrow correlation function

$$f_{ij} = \frac{r_{ij}}{a(1 + \beta r_{ij})},\tag{6.47}$$

where a is either 2 or 4 depending on the spins of particles i and j. The second derivative of  $f_{ij}$  with respect to  $r_{ij}$  is now

$$\frac{\partial^2 f_{ij}}{\partial r_{ii}^2} = -\frac{2\beta}{a(1+\beta r_{ij})^3},\tag{6.48}$$

since the first derivative is

$$\frac{\partial f_{ij}}{\partial r_{ij}} = \frac{1}{a(1+\beta r_{ij})^2},\tag{6.49}$$

as in the previous section. Inserting Eqs. (6.49) and (6.48) into Eq. (6.46) will then give

$$\frac{\nabla^2 J}{J} = \left(\frac{\nabla J}{J}\right)^2 + \sum_{i=1}^{k-1} \left(\frac{2}{r_{ik}} \frac{1}{a(1+\beta r_{ik})^2} - \frac{2\beta}{a(1+\beta r_{ik})^3}\right) + \sum_{i=k+1}^N \left(\frac{2}{r_{ki}} \frac{1}{a(1+\beta r_{ki})^2} - \frac{2\beta}{a(1+\beta r_{ki})^3}\right), \tag{6.50}$$

which can be reduced to

$$\frac{\nabla^2 J}{J} = \left(\frac{\nabla J}{J}\right)^2 + \sum_{i=1}^{k-1} \frac{2}{ar_{ik}} \frac{1}{(1+\beta r_{ik})^3} + \sum_{i=k+1}^N \frac{2}{ar_{ki}} \frac{1}{(1+\beta r_{ki})^3}.$$
 (6.51)

In the local energy function we need to calculate the cross-term

$$\frac{\nabla \Phi}{\Phi} \cdot \frac{\nabla J}{J},\tag{6.52}$$

and  $\nabla^2 J/J$ . The following code shows how I compute these quantities.

The Slater term,  $\nabla \Phi / \Phi$  is computed as

```
//SLATER TERM:
for(int p=0; p<no_of_particles; p++){
   if(p<no_of_particles/2){
      for(int q=0; q<dimension; q++){
      for(int 1=0; 1<no_of_particles/2; 1++){
        temp1[p][q] += gradient_phi(r,alpha,q,1,p)*D_up[1][p];
      }
      }
      else{
      for(int q=0; q<dimension; q++){
      for(int 1=0; 1<no_of_particles/2; 1++){
        temp1[p][q] += gradient_phi(r,alpha,q,1,p)*D_down[1][p-no_of_particles/2];
      }
      }
    }
    }
}</pre>
```

The following code shows how I compute the Jastrow term,  $\nabla J/J$ , using the gradient\_jastrow-functions explained previously.

```
//JASTROW TERM
for(int p=0; p<no_of_particles; p++){
  for(int q=0; q<dimension; q++){
    for(int l=p+1; l<no_of_particles; l++){
    temp2[p][q] += gradient_jastrow(r,p,l,q,beta);
    }
    for(int l=0; l<p; l++){</pre>
```

```
temp2[p][q] += gradient_jastrow(r,p,1,q,beta);
    }
}
```

We can now compute the cross-term by a simply dot product calculation using the following code:

```
for(int p=0; p<no_of_particles; p++){
  for(int q=0; q<dimension; q++){
    tempE += temp1[p][q]*temp2[p][q];
  }
}</pre>
```

The functions gradient\_jastrow(...) and gradient\_phi(...) have been explained earlier. The array temp2[p] [q] now contains the ratio  $\nabla J/J$  and is used to calculate the ratio  $\nabla^2 J/J$  as given in Eq. (6.51). The code for this equation is:

```
//Gradient ratio squared:
for(int p=0; p<no_of_particles; p++){</pre>
   for(int q=0; q<dimension; q++){</pre>
     tempE2 += temp2[p][q]*temp2[p][q];
  //Second term, involving df/dx and d^2f/dx^2
  for(int p=0; p<no_of_particles; p++){</pre>
   for (int k = 0; k < no_of_particles; k++) {</pre>
     if ( k != p) {
  if(((p < no_of_particles/2) && (k <no_of_particles/2)) ||</pre>
      ((p>=no_of_particles/2 && k>=no_of_particles/2))){
    a=0.25;
    tempE3 += 2*a/diffR(r,p,k)/pow((1+beta*diffR(r,p,k)),3);
  else{
    a=0.5;
    tempE3 += 2*a/diffR(r,p,k)/pow((1+beta*diffR(r,p,k)),3);
     }
   }
  }
```

We now have all the pieces to calculate the local energy with E\_kinetic, tempE,tempE2 and tempE3. We only need to multiply the different term by scalar factors, and calculate the potential energy. The following code shows this.

```
//KINETIC ENERGY
E_kinetic -= 2*tempE;
E_kinetic -= tempE2;
E_kinetic -= tempE3;

E_kinetic *= 0.5;

//POTENTIAL ENERGY Coulomb potential
    //electron-proton-interaction
for(int i=0; i<no_of_particles; i++){
    r_single_particle=0;</pre>
```

```
for(int j=0; j<dimension; j++){
    r_single_particle += r[i][j]*r[i][j];//r^2=x^2+y^2+z^2
}

E_potential -=charge/sqrt(r_single_particle);
}
//electron-electron-interaction
for(int i=0; i<no_of_particles-1; i++){
    for(int j=i+1; j<no_of_particles; j++){
        r_12=0;
        for(int k=0; k<dimension; k++){
    r_12 +=(r[i][k]-r[j][k])*(r[i][k]-r[j][k]);
        }
        E_potential +=1/sqrt(r_12);
    }
}//*/
E_local = E_potential + E_kinetic;
return E_local;</pre>
```

# 6.2 Implementation of Metropolis-Hastings algorithm

We can now use the results from the previous sections to show how the Metropolis-Hastings algorithm is implemented. This algorithm is discussed in section 4.4.2, and figure 4.2 sums up the main points quite nicely.

After calculating the initial positions for the electrons, the inverse spin-up and -down Slater matrices and the quantum force F, I start the first Monte Carlo cycle and move the first electron as

```
for(int j=0; j<dimension; j++){
  r_new[i][j]=r_old[i][j]+D*time_step*qm_force_old[i][j]
  +gaussian_deviate(&idum)*sqrt(time_step)</pre>
```

where the function gaussian\_deviate returns a normal distributed random number. I then calculate the distances between the electrons to find both the Slaterand Jastrow-ratios for the full ratio of probabilities,  $\psi_T(r^{new})/\psi_T(r^{old})$  as

```
//SETTING UP THE DISTANCE MATRIX
getDistance(distance_new, r_new, beta);
//SLATER DETERMINANT RATIO
ratio = getRatio(D_up, D_down, i, alpha, beta);

jastrowRatio = getJastrowRatio(distance_old, distance_new, beta);
jastrowRatio = exp(jastrowRatio);
```

The next point is to update the inverse Slater matrices, and then calculate the quantum force for the Green's function in the Metropolis-Hastings test:

```
//temporary update for quantum force
  update(temp_up, temp_down, i, ratio, alpha, beta);
```

```
quantum_force(r_new, qm_force_new, temp_up, temp_down, alpha, beta);

//IMPORTANCE SAMPLING
double greensfunction = 0.0;
for(int k=0; k<no_of_particles; k++){
for(int j=0; j < dimension; j++) {
  greensfunction += 0.5*(qm_force_old[k][j]+qm_force_new[k][j])*
    (D*time_step*0.5*(qm_force_old[k][j]-qm_force_new[k][j])-r_new[k][j]+r_old[k][j]);
}
greensfunction = exp(greensfunction);</pre>
```

I use temporary arrays in case the Metropolis-Hastings test does not accept the particle move. The test itself is implemented as

```
//METROPOLIS TEST
if(ran1(&idum) <= greensfunction*ratio*ratio*jastrowRatio*jastrowRatio){
  for(int k=0; k<no_of_particles; k++){</pre>
   for(int j=0; j<dimension; j++){</pre>
     r_old[k][j]=r_new[k][j];
     qm_force_old[k][j] = qm_force_new[k][j];
  }
  //UPDATE INVERSE SLATER MATRICES
 for(int m=0; m<no_of_particles/2; m++){</pre>
   for(int n=0; n<no_of_particles/2; n++){</pre>
     D_up[m][n]=temp_up[m][n];
     D_down[m][n]=temp_down[m][n];
  //UPDATE THE DISTANCE MATRIX
 for(int m=0; m<no_of_particles; m++){</pre>
   for(int n=0; n<no_of_particles; n++){</pre>
     distance_old[m][n]=distance_new[m][n];
}//end metropolis test
```

The random number generator ran1 is taken from [8]. If now all particles for the cycle have been moved, we can go on to sample the local energy for the expectation values  $\langle E_L \rangle$  and  $\langle E_L^2 \rangle$ :

```
delta_E = local_energy(r_old, D_up, D_down, alpha, beta);
energy +=delta_E;
energy2 +=delta_E*delta_E;
```

At the end of the Monte Carlo loop, we then divide the expectation values by the number of Monte Carlo cycles we have performed.

Furthermore, an important note must be made regarding the calculation of the local energy. At the beginning of the simulation, the system will in general not be in its most probable state. By reviewing section 4.1, we see that a system must evolve before reaching its largest eigenvalue, corresponding to the most probable state. As a result of this, we must include some *thermalization* cycles before we start sampling the local energy. As seen in the algorithm, we initialize the system

in random positions. If we perform a certain number of Monte Carlo cycles before we start sampling the local energy, the system will hopefully be in its most probable state when we actually **do** start sampling the energy. This will improve our results, since we will calculate the energy where the PDF has greater values, leading to more efficient calculations.

The number of thermalization cycles is not know a priori, and must be dealt with experimentally. In this thesis we found that by using 20,000 thermalization cycles, we will safely start sampling the energy when the system has reached its most probable state.

# Chapter 7

# Hartree-Fock implementation

The Hartree-Fock method (HF) is discussed in section 3.2, and the results supply a great improvement to the single particle wave functions. Using HF, we find the optimal single particle wave functions (SPWF) for our chosen basis functions. By using the optimal SPWF, we no longer have to vary the Slater determinant in a VMC calculation, and can consentrate on finding the minimum by varying the parameter(s) in the Jastrow-function. As an example, I have implemented Hartree-Fock for the Beryllium atom using the method of varying coefficients (see section 3.2.1). The basis functions are here chosen to be the hydrogen-like s-orbitals.

As seen in section 3.2.1, in order to get our single particle solutions

$$\psi_a = \sum_{\lambda} C_{a\lambda} \phi_{\lambda},\tag{7.1}$$

we have to solve the Hartree-Fock equations

$$\sum_{\gamma} h_{\lambda\gamma}^{HF} C_{k\gamma} = \epsilon_k C_{k\lambda}, \tag{7.2}$$

where  $h_{\lambda\gamma}^{HF}$  is

$$h_{\lambda\gamma}^{HF} = \langle \lambda | h | \gamma \rangle + \sum_{a=1}^{N} \sum_{\alpha,\delta} C_{a\alpha}^* C_{a\delta} \langle \lambda \alpha | V | \gamma \delta \rangle_{AS}.$$
 (7.3)

We see how the coefficients  $C_{k\gamma}$  are involved on both sides of the equation, and therefore leading to an non-linear eigenvalue problem.

#### 7.1 Interaction matrix elements

As mentioned in section 3.2.1, this particular Hartree-Fock approach gives us the advantage of computing the matrix elements  $\langle \lambda \alpha | V | \gamma \delta \rangle$  once and for all. We have chosen the hydrogenic s-waves as the basis functions  $\phi_{\lambda}$  from Eq. (7.1).

The interaction is given by the Coloumb force between the electrons, and the matrix elements are given by

$$\langle \lambda \alpha | V | \gamma \delta \rangle = \int r_1^2 dr_1 \int r_2^2 dr_2 R_{n_{\alpha}0}^*(r_1) R_{n_{\beta}0}^*(r_2) \frac{1}{r_{<}} R_{n_{\gamma}0}(r_1) R_{n_{\delta}0}(r_2)$$
 (7.4)

This is the direct term of the interaction piece. By calling the innermost integral over  $r_2$  for I,

$$I = \int r_2^2 dr_2 R_{n_{\beta}0}(r_2) \frac{1}{r_{<}} R_{n_{\delta}0}(r_2),$$

we can split this up into two pieces. One where  $r_2 < r_1$  and  $r_< = r_1$  and one where  $r_2 > r_1$  with  $r_< = r_2$ , giving

$$I = \int_0^{r_1} r_2^2 dr_2 R_{n_{\beta}0}(r_2) \frac{1}{r_1} R_{n_{\delta}0}(r_2) + \int_{r_1}^{\infty} r_2 dr_2 R_{n_{\beta}0}(r_2) R_{n_{\delta}0}(r_2)$$

The matrix element  $\langle \lambda \alpha | V | \gamma \delta \rangle$  can then be written as

$$\langle \lambda \alpha | V | \gamma \delta \rangle = \int_0^\infty r_1^2 dr_1 R_{n_{\alpha}0}(r_1) R_{n_{\gamma}0}(r_1) \left[ \int_0^{r_1} r_2^2 dr_2 R_{n_{\beta}0}(r_2) \frac{1}{r_1} R_{n_{\delta}0}(r_2) + \int_{r_1}^\infty r_2 dr_2 R_{n_{\beta}0}(r_2) R_{n_{\delta}0}(r_2) \right]$$
(7.5)

The exchange term  $\langle \lambda \alpha | V | \delta \gamma \rangle$  can be written the same way only by exchanging  $\delta \leftrightarrow \gamma$  which gives

$$\langle \lambda \alpha | V | \delta \gamma \rangle = \int_0^\infty r_1^2 dr_1 R_{n_{\alpha}0}(r_1) R_{n_{\delta}0}(r_1) \left[ \int_0^{r_1} r_2^2 dr_2 R_{n_{\beta}0}(r_2) \frac{1}{r_1} R_{n_{\gamma}0}(r_2) + \int_{r_1}^\infty r_2 dr_2 R_{n_{\beta}0}(r_2) R_{n_{\gamma}0}(r_2) \right]$$
(7.6)

In the equations above the functions  $R_{n_0}$  are defined by the radial part of the s-states, as

$$R_{n_0} = \left(\frac{2Z}{n}\right)^{3/2} \sqrt{\frac{1}{2n^2}} L_{n-1}^1 \left(\frac{2Zr}{n}\right) \exp\left(-\frac{Zr}{n}\right), \tag{7.7}$$

with  $L_{n-1}^1\left(\frac{2Zr}{n}\right)$  being the generalized Laguerre polynomials.

### 7.2 Algorithm and HF results

The Hartree-Fock program calculates the energy and optimized single particle wave functions to plug into the VMC machinery. We see from equation 7.2 that we are dealing with an eigenvalue problem. However, as the  $h^{HF}$ -matrix depends on the matrix elements of C, we have a non-linear problem, and must use an iterative self-consistency approach.

• The first thing to do is to choose the number basis functions we wish to expand the functions  $\psi$  in Eq. (7.1) with.

- We then calculate the matrix elements from Eq. (7.5). The elements were stored in a 4-dimensional array, and each integral was calculated using Gauss-Legendre-quadrature. One should really use Gauss-Laguerre to match the interval, but the Gauss-Legendre approach seemed to work well.
- The next step is to start the iteration process. The first thing we need is an ansatz for the C-matrix and we choose the identity matrix, corresponding to our new basis functions just being the old hydrogen s-waves. We then calculate the  $h^{HF}$ -matrix from Eq. (7.3) which we then diagonalize using library C++-functions. We now put the eigenvectors corresponding to the 4 lowest eigenvalues in the 4 first rows as we only deal with Beryllium. We can now calculate the energy, and use the modified diagonalized matrix as the C-matrix for next the iteration.
- When the energy difference between two iterations is low enough (determined by a preset tolerance  $\sim 10^{-6}$ ) we stop the iterations and we have reached a desired set of coefficients for the new basis.

Table 7.1 shows the Hartree-Fock energy results for different number of s-waves included in the wave function expansion. It also shows the number of iterations needed for the energy difference between two succeeding iterations to be smaller than the chosen tolerance. We see from table 7.1 that the ground state energy for 15 s-waves produces approximately the same result as for 10 and 7 basis functions, but is improved in the 5th digit. The table also presents the exact result as given in [15]. [15]

We can compare the best Hartree-Fock result when only varying coefficients,  $E = -14.5185 E_h$ , with the results from appendix A,  $E = -14.573021 E_h$ , where Slater-type orbitals have been used. If we compare these with the exact result,  $E_0 = -14.6674 E_h$ , it can lead us to believe that the Slater-type orbitals in the Roothaan-Hartree-Fock approach are a better choice for improving the VMC calculations.

In section 6.5.2 in [2], an argument is given for not using hydrogenic functions as basis functions for larger many-body atomic systems. They do not constitute a complete set by themselves, since the hydrogenic functions only describe bound states, and will not describe the unbound continuum states needed for a complete set. They also spread out quickly and become diffuse because of the 1/n-factors in the polynomials and the exponentials. The Slater-type orbitals have a more compact exponential radial form, and we will avoid the problems involving difuseness and the completeness of the basis set.

HF basis functions	Energy	Iterations
2	-13.7159	2
3	-14.5082	9
4	-14.5115	10
5	-14.5141	11
7	-14.5165	10
10	-14.5178	11
15	-14.5185	10
Exact	-14.6674	-

Table 7.1: The table shows results from different runs with different number of basis functions in addition to the exact result. Energies are given in the Hartree unit.

# Chapter 8

# VMC Results

#### HVORDAN FÅ PASSELIGE MATLAB VARIASJONSPLOTT:

- Kjør fil. Se tinghest.m i Neon-mappe for referanse. Husk å skrive riktige parametre
- Tilpass tekstvinduet til den automatiske figurstørrelsen(.fig-fila)
- Endre tekst på fig-fila fra 10 til 12.
- Skalér figuren tilbake til størrelsen på tekstvinduet.
- Lagre figur og eksporter eps-filer.

In this chapter will present most of the results produced by the Variational Monte Carlo code. The first part shows the variational plots for both sets of single particle orbitals, that is hydrogen like and Roothaan-Hartree-Fock orbitals. The second part will present results of the time step extrapolation discussed in section !!!!!!!!!! for the finely tuned parameter(s) produced by the CGM. These results will also include the error estimated by the blocking method.

The last part will compare the obtained results with experimental data.

#### 8.1 Validation runs

In order to be sure that the Slater determinant is working properly, it is nescessary to test the code with something known.

### 8.1.1 Hydrogen-like wave functions

A good check to see that code with the Slater determinant with hydrogenic single particle orbitals is set up correctly is to run the program with no electron interaction. In this case this means using a pure Slater determinant as the trial wave function without the Jastrow factor. If we also neglect the electron-electron interaction energies, we can compare our results with the analytic results for the hydrogen atom.

For non-interacting atomic systems, we can use the hydrogen results to get a closed form expression for the non-interacting energy,  $E_{NI}$ . The energies are

$$E_{NI} = -\frac{Z^2}{2} \sum_{i=1}^{N} \frac{1}{n_i^2},\tag{8.1}$$

where Z is the charge of the nucleus, N is the number of electrons, and  $n_i$  is the principal quantum number for the state occupied by electron i. The energy is in units of Hartree.

Table 8.1 shows how the electrons for our systems of interest are distributed in the different energy orbitals

Atom	Spectroscopic notation
Hydrogen	(1s)
Helium	$(1s)^2$
Beryllium	$(1s)^2(2s)^2$
Neon	$(1s)^2(2s)^2(2p)^6$
Magnesium	$(1s)^2(2s)^2(2p)^6(3s)^2$
Silicon	$(1s)^2(2s)^2(2p)^6(3s)^2(3p)^2$

Table 8.1: The table shows the electron configuration for hydrogen, helium, beryllium, neon, magnesium and silicon.

The analytic non-interacting helium energy is given as

$$E_{NI}^{He} = -\frac{2^2}{2} \left( \frac{1}{1^2} + \frac{1}{1^2} \right) = -4.$$
 (8.2)

By running the non-interacting version of the program for helium with  $4 \cdot 10^6$  samples and time step  $\delta t = 0.05$ , we get an energy

$$E = -4, (8.3)$$

with variance

$$\sigma_E^2 = 1.6 \cdot 10^{-5}. (8.4)$$

Table 8.2 shows the results for all our atoms of interest. Time step  $\delta t = 0.04$  have been used for all calculations.

### 8.1.2 Slater-type orbitals

To test the Slater-type orbitals (STO) found by using Hartree-Fock in [5], I ran the code with STO but no Jastrow factor. Then I could compare the results with the results from ref. [5], to validate my orbitals. The results are shown in table 8.3

Atom	$E_{NI}$	E	MC cycles	$\sigma_E^2$
Helium	-4	-4	$4 \cdot 10^{6}$	$1.6 \cdot 10^{-5}$
Beryllium	-20	-20	$4 \cdot 10^{7}$	$4.0 \cdot 10^{-5}$
Neon	-200	-200	$1 \cdot 10^{7}$	$4.0 \cdot 10^{-3}$
Magnesium	-304	-304	$1 \cdot 10^{8}$	$9.2 \cdot 10^{-3}$
Silicon	-435.5556			

Table 8.2: The table shows the analytic results,  $E_{NI}$ , the computed results, E, number of Monte Carlo cycles and the variance,  $\sigma_E^2$ .

Atom	$E^{HF}$	E	MC cycles	$\sigma_E^2$
Helium	-2.8616799	-2.8617823	$3 \cdot 10^{7}$	0.57097060
Beryllium	-14.573021	-14.573862	$3 \cdot 10^{7}$	2.9477341
Neon	-128.54705	-128.81981	$1 \cdot 10^{8}$	31.003216
Magnesium	-199.61461	-199.84100	$1 \cdot 10^{8}$	41.645320
Silicon	XX	-XX	XX	XX

Table 8.3: The table shows the Roothaan-Hartree-Fock results,  $E^{HF}$  (see appendix A and ref. [5]), compared with the computed energies using the Roothaan STO, E, number of Monte Carlo cycles and the variance,  $\sigma_E^2$ .

### 8.2 Variational plots

This section will present the plots of the variational runs for the different systems. For the hydrogen-like wave functions in section 8.2.1 these plots will be two dimensional, as the total wave function consists of two variational parameters,  $\alpha$  in the Slater determinant and  $\beta$  in the Jastrow function. For the Roothaan-Hartree-Fock single particle wave functions, the plots will only be one-dimensional, as the single particle orbitals have already been optimized by the Roothaan-Hartree-Fock method. For this case, the wave function has only the single variational parameter in the Jastrow function, namely  $\beta$ .

### 8.2.1 Hydrogenic single particle wave functions

The two-dimensional plots will look like the one depicted in figure 8.1. For the different variational calculations I have used between 10 and 100 million samples per parameter-set depending on the size of the system (smaller systems demand less computational time and we can more easily compute smoother plots). However, these plots are rather qualitative with respect to the energy, and the main goal is to find the whereabouts of the optimal parameters. But by merely looking at the plots and where the energy has its minimum, we should be able to find very good starting points for the DFP-method (see appendix C). The DFP-method will supply us with the **optimal** parameters using the parameters found from the figures in this section as initial values. Table 8.4 shows the results for the variational

Atom	$\alpha$	β	MC cycles	E
Helium	1.85	0.35	$10^{8}$	$\approx -2.89$
Beryllium	3.90	0.12	$10^{8}$	$\approx -14.49$
Neon	9.20	0.50	$3 \cdot 10^{7}$	$\approx -127.7$
Magnesium	10.8	0.40	$10^{7}$	$\approx -200$
Silicon	X	X	X	X

Table 8.4: The table shows the apparent optimal parameters from the variational Monte Carlo plots. It also shows the number of Monte Carlo cycles and the approximate energy for the parameter set. The energies are in units of Hartree.

parameters obtained by inspecting figures 8.2, 8.3, 8.4, 8.5. Some of the plots are very flat, especially the magnesium plot with respect to the  $\alpha$ -parameter. It seems very

#### 8.2.2 Roothaan-Hartree-Fock wave functions

### 8.3 Optimal parameters with DFP

#### 8.3.1 Hydrogenic wave-functions

Atom	$\alpha_{start}$	$\beta_{start}$	$\alpha_{min}$	$\beta_{min}$
Helium	1.85	0.35	1.839	0.348
Beryllium	3.90	0.12	3.925	0.109
Neon	9.20	0.50	9.546	0.177
Magnesium	10.8	0.40	X.xxx	x.xxx
Silicon	X.xx	X.xx	X.xxx	X.xxx

Table 8.5: The table lists the apparent optimal parameters obtained from the plots and the finely tuned parameters from the DFP-algorithm.

For helium with 3,3,3,2,2,4 iterations, and starting point  $\alpha = 1.85$ ,  $\beta = 0.35$ , 6 nodes:

- $\alpha = 1.839228333$
- $\beta = 0.3475618333$

The 6 different runs gave an average energy, E = -2.89053.

#### 8.3.2 Beryllium

For beryllium with 4,4,4,3,3,3 iterations, and starting point  $\alpha = 3.90$ ,  $\beta = 0.12$ , 6 nodes:

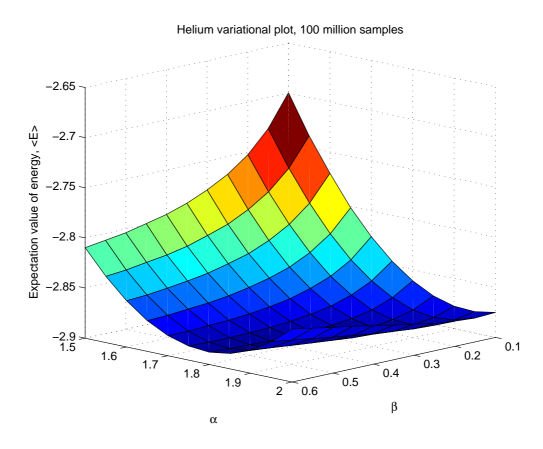


Figure 8.1: A variational plot for helium with 100 million samples

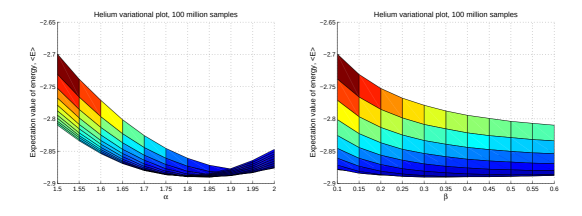


Figure 8.2: The figures show a variational plot of the helium ground state energy with 100 million samples. The figure on the left shows the variation of the  $\alpha$  parameter, while the figure on the right shows the variation of the  $\beta$  parameter.

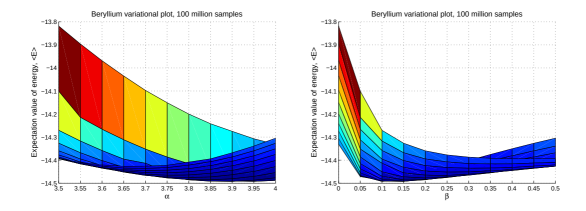


Figure 8.3: The figures show a variational plot of the beryllium ground state energy with 100 million samples. The figure on the left shows the variation of the  $\alpha$  parameter, while the figure on the right shows the variation of the  $\beta$  parameter.

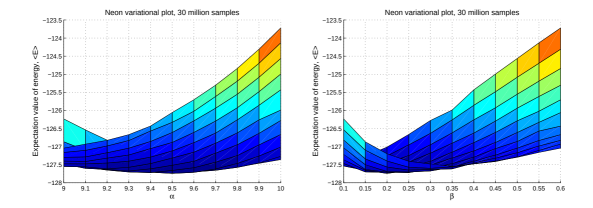
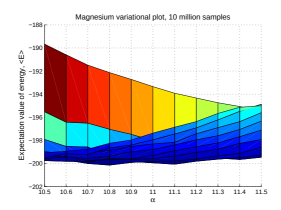


Figure 8.4: The figures show a variational plot of the neon ground state energy with 30 million samples. The figure on the left shows the variation of the  $\alpha$  parameter, while the figure on the right shows the variation of the  $\beta$  parameter.



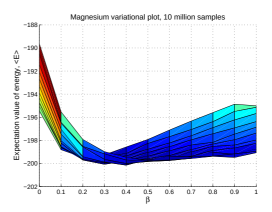


Figure 8.5: The figures show a variational plot of the magnesium ground state energy with 10 million samples. The figure on the left shows the variation of the  $\alpha$  parameter, while the figure on the right shows the variation of the  $\beta$  parameter.

- $\alpha = 3.924768333$
- $\beta = 0.10898885$

The 6 different runs gave an average energy, E = -14.4934125.

#### 8.3.3 Neon

For neon with 3,3,3,4,2,2,3 iterations, and starting point  $\alpha = 9.50$ ,  $\beta = 0.20$ , 7 runs:

- $\alpha = 9.545854286$
- $\beta = 0.1765994286$

The 7 different runs gave an average energy, E = -127.7195714.

### 8.4 Time-step interpolation

### 8.5 Magnesium

For neon with 4,3,3, starting point  $\alpha = 10.8$ ,  $\beta = 0.40$ , 3 runs

- $\alpha = 10.8391$
- $\beta = 0.385164$

The runs gave average energy, E = -199.893.

## 8.6 Comparison with experimental data

The exact results with respect to the Hamiltonian:(XXXXXXXXXXXXXX) for the ground state energy of helium, beryllium, neon and magnesium are taken from [15].

• Helium:  $E_0 = -2.9037$ 

• Beryllium:  $E_0 = -14.6674$ 

• Neon:  $E_0 = -128.9383$ 

• Magnesium:  $E_0 = -200.054$ 

• Silicon: ?

# Appendix A

## Roothaan-Hartree-Fock results

In this part I will present Clementi and Roetti's results from their Roothaan-Hartree-Fock calculations for atoms helium, beryllium, neon, magnesium and silicon. The tables' applications are explained in section 5.5.

#### A.1 Helium

Helium consists of 2 electrons and a core of charge 2e and has the electron distribution  $(1s)^2$ . Table A.1 shows the Roothaan-Hartree-Fock(RHF) solution for Helium. The tabulated Roothaan-Hartree-Fock ground state energy for Helium

$n,\lambda$	Exponent, $\xi$	1s exp.coeff.
1S	1.41714	0.76838
1S	2.37682	0.22356
1S	4.39628	0.04082
1S	6.52699	-0.00994
1S	7.94525	0.00230

Table A.1: The table shows the Roothan-Hartree-Fock results for Helium.

given in [5] is

$$E = -2.8616799 E_h$$

where  $E_h$  is the *Hartree* unit given in section 3.1.4.

### A.2 Beryllium

Beryllium has 4 electrons with a nucleus charged 4e and has electrons distributed as  $(1s)^2(2s)^2$ . Table A.2 shows the RHF-solution for beryllium. In [5] the total ground state energy is calculated as

$$E = -14.573021 E_h$$
.

$n,\lambda$	Exponent, $\xi$	1s exp.coeff.	2s exp.coeff.
1S	3.47116	0.91796	-0.17092
1S	6.36861	0.08724	-0.01455
2S	0.77820	0.00108	0.21186
2S	0.94067	-0.00199	0.62499
2S	1.48725	0.00176	0.26662
2S	2.71830	0.00628	-0.09919

Table A.2: The table shows the Roothan-Hartree-Fock results for Beryllium.

### A.3 Neon

The ten Neon electrons are distributed as  $(1s)^2(2s)^2(2p)^6$ . Table A.3 shows the RHF-solution. The Roothaan-Hartree-Fock energy is

$n,\lambda$	Exponent, $\xi$	1s exp.coeff.	2s exp.coeff.	$n,\lambda$	Exponent, $\xi$	2p exp.coeff.
1S	9.48486	0.93717	-0.23093	2P	1.45208	0.21799
1S	15.56590	0.04899	-0.00635	2P	2.38168	0.53338
2S	1.96184	0.00058	0.18620	2P	4.48489	0.32933
2S	2.86423	-0.00064	0.66899	2P	9.13464	0.01872
2S	4.82530	0.00551	0.30910			
2S	7.79242	0.01999	-0.13871			

Table A.3: The table shows the Roothan-Hartree-Fock results for Neon.

$$E = -128.54705 E_h$$
.

### A.4 Magnesium

The Magnesium atom has 12 electrons in orbitals  $(1s)^2(2s)^2(2p)^6(3s)^2$ . The RHF-solution is given in table A.4. The RHF-energy is given in [5] as

$$E = -199.61461 E_h$$
.

#### A.5 Silicon

The silicon atom consists of a nucleus with charge 14e and 14 electrons distributed in orbitals as  $(1s)^2(2s)^2(2p)^6(3s)^2(3p)^2$ . Table A.5 shows the s-orbitals for the RHF-solution. Table A.6 shows the RHF results for the p-orbitals. The ground state energy calculated with the RHF-method gives

$$E = -288.85431 E_h$$
.

$n,\lambda$	Exponent, $\xi$	1s e.c.	2s e.c.	3s e.c.	$n,\lambda$	Exponent, $\xi$	2 <i>p</i> e.c.
1S	12.01140	0.96430	-0.24357	0.04691	2P	5.92580	0.52391
3S	13.91620	0.03548	-0.00485	0.00144	4P	7.98979	0.07012
3S	9.48612	0.02033	0.08002	-0.01850	4P	5.32964	0.31965
3S	6.72188	-0.00252	0.39902	-0.07964	4P	3.71678	0.20860
3S	4.24466	0.00162	0.57358	-0.13478	4P	2.59986	0.03888
3S	2.53466	-0.00038	0.05156	-0.01906			
3S	1.46920	0.00015	-0.00703	0.48239			
3S	0.89084	-0.00004	0.00161	0.60221			

Table A.4: The table shows the Roothan-Hartree-Fock results for Magnesium.

$n,\lambda$	Exponent, $\xi$	1s exp.coeff.	2s exp.coeff.	3s exp.coeff.
1S	14.01420	0.96800	-0.25755	0.06595
3S	16.39320	0.03033	-0.00446	0.00185
3S	10.87950	0.02248	0.11153	-0.03461
3S	7.72709	-0.00617	0.40339	-0.10378
3S	5.16500	0.00326	0.55032	-0.19229
3S	2.97451	-0.00143	0.03381	-0.06561
3S	2.14316	0.00081	-0.00815	0.59732
3S	1.31306	-0.00016	0.00126	0.55390

Table A.5: The table shows the s-orbital Roothan-Hartree-Fock results for Silicon.

$n,\lambda$	Exponent, $\xi$	2p exp.coeff.	3p exp.coeff.
2P	7.14360	0.54290	-0.11535
4P	16.25720	0.00234	-0.00189
4P	10.79720	0.04228	-00473
4P	6.89724	0.32155	-0.07552
4P	4.66598	0.22474	0.01041
4P	2.32046	0.00732	0.46075
4P	1.33470	-0.00105	0.57665
4P	0.79318	0.00041	0.06274

Table A.6: The table shows the p-orbital Roothan-Hartree-Fock results for Silicon.

# Appendix B

# **Statistics**

This appendix will give a short introduction to many key aspects regarding the statistical terms used for calculating expectation values and its statistical error in the Variational Monte Carlo process. We will follow the work in [16] quite closely in order to get all the important points across.

A probability distribution function (PDF), p(x), is a function that describes the frequency of a stochastic value X to occur, that is

$$p(x) = \operatorname{Prob}(X = x). \tag{B.1}$$

This is for the discrete case.

For the continuous case, p(x) represents a probability distribution, where we must deal with probabilities of values being within some interval. That is, the probability of the stochastic variable X taking a value on the finite interval [a, b]is defined by an integral

$$Prob(a \le X \le b) = \int_{a}^{b} p(x)dx.$$
 (B.2)

All PDF's must take values that are both positive and less or equal to unity, the absolute maximum for any probability function to make sense. This means

$$0 \le p(x) \le 1,\tag{B.3}$$

In addition to this, both the discrete and continuous PDF's must be normalized to one as

$$\sum_{i} p(x_i) = 1 \tag{B.4}$$

$$\sum_{i} p(x_i) = 1$$

$$\int p(x)dx = 1.$$
(B.4)
(B.5)

#### Moments and expectation values

Since our wave functions are continuous, our PDF will be so as well, and the expectation value of a function f(x) is given by

$$\langle f \rangle \equiv f(x)p(x)dx.$$
 (B.6)

We also have the moments, as special case of expectation values, where the n-th moment is

$$\langle x^n \rangle \equiv \int x^n p(x) dx.$$
 (B.7)

The mean,  $\mu$ , is defined by the first moment,  $\langle x \rangle$ .

The *central moments* are defined by

$$\langle (x - \langle x \rangle)^n \rangle \equiv \int (x - \langle x \rangle)^n p(x) dx.$$
 (B.8)

The second central moment is called the *variance*,  $\sigma_X^2$  or Var(X), and can be written as

$$\sigma_X^2 = \text{Var}(X) = \langle (x - \langle x \rangle)^2 \rangle$$
$$= \langle x^2 \rangle - \langle x \rangle^2. \tag{B.9}$$

The standard deviation is then  $\sigma = \sqrt{\langle x^2 \rangle - \langle x \rangle^2}$ .

#### B.1 Correlated measurements

For now, we have been dealing with functions of one stochastic variable, so-called *univariate* PDF's. However, a PDF may as well consist of many variables. These PDF's are called *multivariate* PDF's. The variables themselves are independent, or *uncorrelated* is the multivariate PDF,  $P(x_1, x_2, ..., x_n)$  can be factorized as

$$P(x_1, x_2, ..., x_n) = \prod_{i=1}^{n} p_i(x_i),$$
(B.10)

where  $p_i(x_i)$  are the univariate PDF's.

The so-called *covariance* of two stochastic variables,  $X_i$  and  $X_i$ , is defined as

$$Cov(X_i, X_j) \equiv \langle (x_i - \langle x_i \rangle)(x_j - \langle x_j \rangle) \rangle = \langle x_i x_j \rangle - \langle x_i \rangle \langle x_j \rangle.$$
 (B.11)

We see that for j=i this reduces to the variance of  $X_i$ , that is

$$Cov(X_i, X_i) = Var(X_i) = \langle x_i^2 \rangle - \langle x_i \rangle^2.$$
 (B.12)

If the variables  $X_i$  and  $X_j$  are independent, or uncorrelated, the PDF's will factorize as in Eq. (B.10), and we will have  $\langle x_i x_j \rangle = \langle x_i \rangle \langle x_j \rangle$ . This will give  $\text{Cov}(X_i, X_j) = 0$ , when  $i \neq j$ .

Consider linear combinations of stochastic variables  $X_i$  and  $Y_j$  as

$$U = \sum_{i} a_i X_i, \tag{B.13}$$

and

$$V = \sum_{j} b_j Y_j, \tag{B.14}$$

where  $a_i$  and  $b_j$  are scalar coefficients. As given in [17], we will have

$$Cov(U, V) = \sum_{i,j} a_i b_j Cov(X_i, Y_j).$$
(B.15)

Since  $Var(X_i) = Cov(X_i, X_i)$ , the variance of U will be

$$Var(U) = \sum_{i,j} a_i a_j Cov(X_i, X_j),$$
(B.16)

which for uncorrelated variables will be reduced to

$$Var(U) = \sum_{i} a_i^2 Cov(X_i, X_i) = \sum_{i} a_i^2 Var(X_i),$$
 (B.17)

and

$$\operatorname{Var}\left(\sum_{i} a_{i} X_{i}\right) = \sum_{i} a_{i}^{2} \operatorname{Var}(X_{i}). \tag{B.18}$$

#### Approximations for finite data sets

Consider a computational experiment which produces a sequence of n stochastic values

$$\{x_1, x_2, \dots, x_n\} \tag{B.19}$$

which we will call the n measurements of our sample. We can hereby define the sample mean as

$$\bar{x}_n \equiv \frac{1}{n} \sum_{k=1}^n x_k,\tag{B.20}$$

with the sample variance and sample covariance being

$$Var(x) \equiv \frac{1}{n} \sum_{k=1}^{n} (x_k \bar{x}_n$$
 (B.21)

and

$$Cov(x) \equiv \frac{1}{n} \sum_{kl} (x_k - \bar{x}_n)(x_l - \bar{x}_n), \tag{B.22}$$

respectively. This sample covariance is a measure of the correlation between succeeding values in the sample. These values are not the same as the mean  $\mu_X$ , Var(X) and Cov(X) defined by the exact PDF,  $p_X(x)$ , but approximations to these quantities.

In the limit where  $n \to \infty$  it can be shown that the sample mean,  $\bar{x}_n$  approaches the true mean  $\mu_X$ , that is

$$\lim_{n \to \infty} \bar{x}_n = \mu_X,\tag{B.23}$$

and  $\bar{x}_n$  can be seen as an estimate of  $\mu_X$ . But how good is this approximation? In order to find this we also need the error estimate of our measurement. We can view the sample means themselves as measurements in a collection of sample means

(several experiments). To calculate the statistical error, we need the PDF of the sample means,  $p_{\bar{X}_n}(x)$ . The exact error will be given by the standard deviation of the sample means,  $\text{err}_X$ , also called sample error. However, we don't know the exact PDF, so we can only calulate an estimate of  $\text{err}_X$ .

The sample mean,  $\bar{x}_n$ , can be treated as a stochastic variable  $\bar{X}_n$ , because  $\bar{x}_n$  is a linear combination of stochastic variables  $X_i$  with 1/n as common coefficients. This means we can write

$$\bar{X}_n = \frac{1}{n} \sum_{i=1}^n X_i.$$
 (B.24)

We will now have  $p_{\bar{X}}(x)$  as the probability distributions of the sample means  $\bar{X}_n$ , but we cannot get a closed form expression for this. However, the *central limit theorem* (see [17]) opens for an approximation of  $p_{\bar{X}}(x)$  as  $n \to \infty$ . It states that

$$\lim_{n \to \infty} p_{\bar{X}}(x) = \left(\frac{n}{2\pi Var(X)}\right)^{1/2} e^{-\frac{n(x-\bar{x}_n)^2}{2Var(X)}},\tag{B.25}$$

which means that the PDF will approach a Gaussian distribution. This Gaussian distribution will have a mean and variance which equal to the true mean and variance,  $\mu_X$  and  $\sigma_X^2$ .

The error is given by

$$\operatorname{err}_{X}^{2} = \operatorname{Var}(\bar{X}_{n}) = \frac{1}{n^{2}} \sum_{ij} \operatorname{Cov}(X_{i}, X_{j}). \tag{B.26}$$

This exact error needs the true means  $\mu_X$  in order to be calculated. We will not be able to do this unless we have the exact PDFs of the variables  $X_i$ . However, we only have the measurements in a sample, and not the actual PDF.

Instead, we estimate the true means  $\mu_X$  by the sample means

$$\mu_{X_i} = \langle x_i \rangle \approx \frac{1}{n} \sum_{k=1}^n x_k = \langle x \rangle.$$
 (B.27)

The estimate of the covariance will then be

$$\operatorname{Cov}(X_{i}, X_{j}) = \langle (x_{i} - \langle x_{i} \rangle)(x_{j} - \langle x_{j} \rangle) \rangle \approx \langle (x_{i} - \bar{x})(x_{j} - \bar{x}) \rangle$$

$$\approx \frac{1}{n} \sum_{l}^{n} \left( \frac{1}{n} \sum_{k}^{n} (x_{k} - \bar{x}_{n})(x_{l} - \bar{x}_{n}) \right) = \frac{1}{n} \frac{1}{n} \sum_{kl} (x_{k} - \bar{x}_{n})(x_{l} - \bar{x}_{n})$$

$$= \frac{1}{n} \operatorname{Cov}(x). \tag{B.28}$$

The error estimate can be written using Eq. (B.28) as

$$\operatorname{err}_{X}^{2} = \frac{1}{n^{2}} \sum_{ij} \operatorname{Cov}(X_{i}, X_{j})$$

$$\approx \frac{1}{n^{2}} \sum_{ij} \frac{1}{n} \operatorname{Cov}(x) = \frac{1}{n^{2}} n^{2} \frac{1}{n} \operatorname{Cov}(x)$$

$$= \frac{1}{n} \operatorname{Cov}(x). \tag{B.29}$$

The same approximation goes for the variance, as

$$\operatorname{Var}(X_i) = = \langle (x_i - \langle x_i \rangle)^2 \rangle \approx \langle (x_i - \bar{x})^2 \rangle$$

$$= \frac{1}{n} \sum_{k=1}^n (x_k - \bar{x}_n)^2$$

$$= \operatorname{Var}(x), \tag{B.30}$$

which for uncorrelated stochastic variables will give an error estimation as

$$\operatorname{err}_{X}^{2} = \frac{1}{n^{2}} \sum_{ij} \operatorname{Cov}(X_{i}, X_{j})$$

$$= \frac{1}{n^{2}} \sum_{i} \operatorname{Var}(X_{i}) \approx \frac{1}{n^{2}} \sum_{i} \operatorname{Cov}(x)$$

$$= \frac{1}{n} \operatorname{Var}(x). \tag{B.31}$$

#### Autocorrelation function and correlation time

As given in [16], we can rewrite the error as

$$\operatorname{err}_{X}^{2} = \frac{1}{n} \operatorname{Cov}(x) = \frac{1}{n} \operatorname{Var}(x) + \frac{1}{n} \left( \operatorname{Cov}(x) - \operatorname{Var}(x) \right)$$
$$= \frac{1}{n^{2}} \sum_{k=1}^{n} (x_{k} - \bar{x}_{n})^{2} + \frac{2}{n^{2}} \sum_{k< l} (x_{k} - \bar{x}_{n})(x_{l} - \bar{x}_{n}).$$
(B.32)

We see the first term is just the variance, so the second term must be the error correction when the variables are correlated. By now introducing partial sums on the form

$$f_d = \frac{1}{n} \sum_{k=1}^{n-d} (x_k - \bar{x}_n)(x_{k+d} - \bar{x}_n),$$
 (B.33)

we can write the correlation term as

$$\frac{2}{n^2} \sum_{k < l} (x_k - \bar{x}_n)(x_l - \bar{x}_n) = \frac{2}{n} \sum_{d=1}^{n-1} f_d,$$
(B.34)

where now  $f_d$  is a measure of how correlated values separated by a distance d are. By dividing  $f_d$  by Var(x), we can define the *autocorrelation function*:

$$\kappa_d = \frac{f_d}{\text{Var}(x)}. ag{B.35}$$

We see that  $f_d$  equals the sample variance, Var(x), when d = 0, so the autocorrelation function has a value of 1 for d = 0.

We can now rewrite the sample error as

$$\operatorname{err}_{X}^{2} = \frac{1}{n} \operatorname{Var}(x) + \frac{2}{n} \operatorname{Var}(x) \sum_{d=1}^{n-1} \frac{f_{d}}{\operatorname{Var}(x)}$$

$$= \left(1 + 2 \sum_{d=1}^{n-1} \kappa_{d}\right) \frac{1}{n} \operatorname{Var}(x)$$

$$= \frac{\tau}{n} \operatorname{Var}(x), \tag{B.36}$$

where  $\tau$ ,

$$\tau = 1 + 2\sum_{d=1}^{n-1} \kappa_d, \tag{B.37}$$

is called the *autocorrelation time* and can be used to define the effective number of measurements,  $n_{eff} = n/\tau$ , for us to simply approximate the error by the sample variance.

# Appendix C

# DFP and energy minimization

Since we wish to minimize the energy using our Variational Monte Carlo machinery, an important part is to find the optimal parameters for the wave function. That is, the parameters for the wave function which minimizes the energy.

A rather brute force first approach is to manually vary the parameters and plot the results as in section 8.2. This will give a good estimate of what the optimal parameters are, but will in general not be precise enough for more quantitatively precise measurements.

The method we use is a so-called quasi-Newton method, the *Davidon-Fletcher-Powell*-algorithm (DFP) as given in [8]. This method builds on the conjugate gradient method (CGM) and the steepest descent method (both are described in [8]). These techniques however, are respectively too costly numerically, or too brute force for our liking.

Both CGM and DPF are based on approximating a function of variational parameters,  $f(\mathbf{x})$ , by a quadratic form, where now  $\mathbf{x}$  is the set of i variational parameters. This is done by writing the Taylor series of  $f(\mathbf{x})$  at some point  $\mathbf{P}$ , as

$$f(\mathbf{x}) = f(\mathbf{P}) + \sum_{i} \frac{\partial f}{\partial x_{i}} + \frac{1}{2} \frac{\partial^{2} f}{\partial x_{i} \partial x_{j}} \dots$$

$$\approx c - \mathbf{b} \cdot \mathbf{x} + \frac{1}{2} \mathbf{x} \cdot \mathbf{A} \cdot \mathbf{x}$$
(C.1)

where now

$$c \equiv f(\mathbf{P}) \qquad \mathbf{b} \equiv -\nabla f|_{\mathbf{P}} \qquad [\mathbf{A}]_{ij} \equiv \frac{\partial^2 f}{\partial x_i \partial x_j}|_{\mathbf{P}}.$$
 (C.2)

Matrix  $\mathbf{A}$  is called the *Hessian* matrix, the matrix with all second derivative. The gradient can easily be calculated as

$$\nabla f = \mathbf{A} \cdot \mathbf{x} - \mathbf{b}. \tag{C.3}$$

In the DFP method, we don't have to calculate the exact Hessian,  $\mathbf{A}$ , which takes a lot of computation time. The goal is then to iteratively construct a sequence of matrices,  $\mathbf{H}_i$ , such that

$$\lim_{i \to \infty} = \mathbf{A}^{-1}.\tag{C.4}$$

By its name, it should be clear that the Newton method for finding the zero of the gradient of function is similar to the quasi-Newton method. For the Newton method, we have to second order, near a current iteration point  $\mathbf{x}_i$ , the following

$$f(\mathbf{x}) = f(\mathbf{x}_i) + (\mathbf{x} - \mathbf{x}_i) \cdot \nabla f(\mathbf{x}_i) + \frac{1}{2} (\mathbf{x} - \mathbf{x}_i) \cdot \mathbf{A} \cdot (\mathbf{x} - \mathbf{x}_i)$$
 (C.5)

and

$$\nabla f(\mathbf{x}) = \nabla f(\mathbf{x}_i) + \mathbf{A} \cdot (\mathbf{x} - \mathbf{x}_i). \tag{C.6}$$

In order to find the next iteration point using Newton's method, we use that  $\nabla f(\mathbf{x}) = 0$ , giving

$$\mathbf{x} - \mathbf{x}_i = -\mathbf{A}^{-1} \cdot \nabla f(\mathbf{x}_i). \tag{C.7}$$

The quasi-Newton method is introduced since we do not compute the exact Hessian and its inverse, but an approximation  $\mathbf{H} \approx \mathbf{A}^{-1}$ . As explained in [8], this approach can often be better than using the true Hessian, as the matrix  $\mathbf{H}$  is constructed in such a way that it is always positive-definite and symmetric. The true Hessian is generally not always positive-definite when we are far from the minimum, which can lead to moving towards increasing values of f instead of decreasing values.

If we subtract Eq. (C.7) at  $\mathbf{x}_{i+1}$  from the same equation at  $\mathbf{x}_i$ , we get

$$\mathbf{x}_{i+1} - \mathbf{x}_i = \mathbf{A}^{-1} \cdot (\nabla f_{i+1}) - \nabla f_i), \tag{C.8}$$

where now  $f_j = f(\mathbf{x}_j)$ . At the point  $x_{i+1}$  we also have that  $\mathbf{H}_{i+1}$  is a good approximation for  $\mathbf{A}^{-1}$ , yielding

$$\mathbf{x}_{i+1} - \mathbf{x}_i = \mathbf{H}_{i+1} \cdot (\nabla f_{i+1}) - \nabla f_i). \tag{C.9}$$

The "outer" product of two vectors,  $\mathbf{u}$  and  $\mathbf{v}$ , is given as,  $\mathbf{u} \otimes \mathbf{v}$ . As opposed to the "inner" product of two vectors

$$\mathbf{u} \cdot \mathbf{v} = \mathbf{u}^T \mathbf{v} \tag{C.10}$$

which returns a scalar, the "outer" product,

$$\mathbf{u} \otimes \mathbf{v} = \mathbf{u}\mathbf{v}^T, \tag{C.11}$$

returns a matrix. Component  $(\mathbf{u} \otimes \mathbf{v})_{ij}$  is simply given as  $(\mathbf{u} \otimes \mathbf{v})_{ij} = u_i v_j$ . The DPF updating formula for  $\mathbf{H}_{i+1}$  is given in [8] as

$$\mathbf{H}_{i} = \mathbf{H}_{i+1} + \frac{(\mathbf{x}_{i+1} - \mathbf{x}_{i}) \otimes (\mathbf{x}_{i+1} - \mathbf{x}_{i})}{(\mathbf{x}_{i+1} - \mathbf{x}_{i}) \cdot (\nabla f_{i+1} - \nabla f_{i})} - \frac{[\mathbf{H}_{i} \cdot (\nabla f_{i+1} - \nabla f_{i})] \otimes [\mathbf{H}_{i} \cdot (\nabla f_{i+1} - \nabla f_{i})]}{(\nabla f_{i+1} - \nabla f_{i}) \cdot \mathbf{H}_{i} \cdot (\nabla f_{i+1} - \nabla f_{i})}.$$
(C.12)

We have used the function dfpmin.c from [8] which uses this quasi-Newton method for finding a minimum for  $f(\mathbf{x})$ , while finding an approximation for the inverse Hessian. This method takes as input a function that returns the mean energy, and a function that calculates the gradient of the energy.

In this thesis we have a function  $f = \bar{E}$ , the mean of the local energy operator,  $\hat{\mathbf{E}}_L$ , as given in 4.4.1. To use the DFP method we must have the gradient of f, that is the first derivatives of the energy expectation value with respect to the variational parameters,  $x_i$ . The first derivatives of the energy expectation value are given in [10] as

$$\frac{\partial \bar{E}}{\partial x_i} = \bar{E}_i = \left\langle \frac{\psi_i}{\psi} E_L + \frac{H\psi_i}{\psi} - 2\bar{E}\frac{\psi_i}{\psi} \right\rangle 
= 2\left\langle \frac{\psi_i}{\psi} (E_L - \bar{E}) \right\rangle = 2\left\langle \frac{\psi_i}{\psi} E_L \right\rangle - 2\left\langle \frac{\psi_i}{\psi} \right\rangle \bar{E}$$
(C.13)

where  $\psi_i = \partial \psi / \partial x_i$ . The expectation values in Eq. (C.13) cannot be calculated easily using closed form expressions, and must be evaluated numerically. In this work we have just used the simple *three-point estimation* of the first derivative

$$f'(x) \approx \frac{f(x+h) - f(x-h)}{2h},\tag{C.14}$$

with h = 0.001.

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