# Development of the Transcorrelated Full Configuration Interaction Quantum Monte Carlo Method

Von der Fakultät Chemie der Universität Stuttgart zur Erlangung der Würde eines Doktors der Naturwissenschaften (Dr. rer. nat.) genehmigte Abhandlung

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Tag der mündlichen Prüfung: .....



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It is important to realize that in physics today, we have no knowledge of what energy is.

— Richard Feynman, Feynman Lectures on Physics, Volume 1, Chapter 4

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# Erklärung über die Eigenständigkeit der Dissertation

Ich erkläre hiermit an Eides statt, dass ich die vorliegende Arbeit	
"Development of the Transcorrelated Full Configuration Interaction Quantum Monte Carlo Method "	$\iota m$
ohne Hilfe Dritter und ohne Benutzung anderer als der angegebenen Hilfsmittel an habe; die aus fremden Quellen direkt oder indirekt übernommenen Gedanken solche kenntlich gemacht. Die Arbeit wurde bisher in gleicher oder ähnlicher keiner anderen Prüfungsbehörde vorgelegt.	sind als

Datum, Ort

Unterschrift

# Acknowledgements

TODO: thank people

# Acronyms

<b>1RDM</b> 1-electron reduced density matrix	8, 49	
<b>2RDM</b> 2-electron reduced density matrix	10, 49	
AO atomic orbital	12	
API application programming interface	106	
BCH Baker-Campbell-Hausdorff	29	
<b>BOA</b> Born-Oppenheimer approximation	3, 4, 14, 74	
CABS complementary auxiliary basis set	27	
CAS complete active space	5	
CASPT2 complete active space perturbation theory	to second order 28	
<b>CASSCF</b> complete active space self-consistent field	16, 28	
CBS complete basis set	7, 8, 13–15, 21, 51, 65, 67, 74, 107	
CC coupled cluster	18, 19, 28, 50, 55	
$\operatorname{cc\text{-pV}} olimits_X \mathbf{Z}$ correlation-consistent polarised valence with)	th X-zeta quality (X=D, T, Q, 5, 6, 21	
CCSD coupled cluster with single and double excita	tions 19, 20	
CCSD(T) coupled cluster with single and double excitations and perturbative triples 20		
CCSDT coupled cluster with single, double and trip	ble excitations 19	
CI configuration interaction	13,15,16,18,19,24,43,46,49	
CSF configuration state function	42	
DC distinguishable cluster	20	
DCSD distinguishable cluster with single and double	e excitations 20	
DMC diffusion Monte Carlo	39–42, 53	

xii Acronyms

DMRG density matrix renormalization group	28, 50
ECG Explicitly Correlated Gaussian	24, 25
<b>EOM</b> equation of motion	20
FCI full configuration interaction	13, 15, 16, 19, 41–43
FCIQMC full configuration interaction quantum 46, 48–51, 53, 74, 75, 106	Monte Carlo 2, 14, 15, 28, 31, 41–43,
GTG Gaussian-type Geminal	24
GTO Gaussian-type orbital	8, 12
HF Hartree-Fock	11–13, 15–19, 25, 29, 31, 39, 51, 54, 75
MC Monte Carlo	32–35, 37, 38, 58
MCSCF multiconfigurational self-consistent field	15, 16, 42
MO molecular orbital	12
MP2 second-order Møller-Plesset perturbation th	neory 18, 26
MPI Message Passing Interface	106
MRCI multireference configuration interaction	28
QMC quantum Monte Carlo	2, 37, 41, 46, 52, 53
RHF restricted Hartree-Fock	12, 15, 74
RI resolution of identity	27, 31
<b>ROHF</b> restricted open-shell Hartree-Fock	12
SCF self consistent field	12
SD Slater determinant	7, 11, 13, 15, 16, 18, 25, 29, 42–44, 49
STO Slater-type orbital	8, 9, 22, 74
TC transcorrelation	2, 28, 29, 49–51, 104, 106
<b>UHF</b> unrestricted Hartree-Fock	12
VMC variational Monte Carlo	29, 37, 38, 41, 53, 54, 107
<b>xTC</b> transcorrelation via exclusion of explicit thr	ree-body components 68, 77, 105–107,

109

# Zusammenfassung

 $\operatorname{TODO}:$  Hier werde ich eine Zusammenfassung schreiben.

## **Abstract**

TODO: abstract

## Introduction

According to modern quantum theory, to fully describe a (nonrelativistic) system of  $N_A$  nuclei and  $N_e$  electrons, we must solve the time-dependent Schrödinger equation,

$$i\hbar \frac{\partial}{\partial t} \Psi(\mathbf{x}, t) = \hat{H} \Psi(\mathbf{x}, t)$$
 (1.1)

where  $\mathbf{x} = (\mathbf{r}, \sigma)$ ,  $\mathbf{r} \in \mathcal{R}^{3(N_e + N_A)}$  gives the spatial coordinates, and  $\sigma$  the corresponding spins. For nuclei with atomic numbers  $Z_I$  (positive charge  $Z_e$ ), the Hamiltonian operator  $\hat{H}$  may be written

$$\hat{H} = -\sum_{I=1}^{N_A} \frac{\hbar^2}{2m_I} \nabla_I^2 + \frac{1}{4\pi\epsilon_0} \frac{1}{2} \sum_{I,J=1;I\neq J}^{N_A} \frac{Z_I Z_J e^2}{|\mathbf{R}_I - \mathbf{R}_J|}$$

$$-\sum_{i=1}^{N_e} \frac{\hbar^2}{2m_e} \nabla_i^2 + \frac{1}{4\pi\epsilon_0} \frac{1}{2} \sum_{i,j=1;i\neq j}^{N_e} \frac{e^2}{|\mathbf{r}_i - \mathbf{r}_j|} - \sum_{i=1}^{N_e} \sum_{I=1}^{N_A} \frac{Z_I e^2}{|\mathbf{r}_i - \mathbf{R}_I|}$$
(1.2)

where  $N_A$  and  $N_e$  are the number of nuclei and electrons, respectively, and we have represented the nuclear coordinates by  $\mathbf{R}_I$  and the electron coordinates by  $\mathbf{r}_i$ .

Since quantum chemistry and condensed matter sciences are in general concerned with nonrelativistic processes involving electrons and nuclei, this might boldly be called the theory of everything.<sup>1</sup> Hence, we may be tempted to conclude this dissertation early, but in practice the evaluation of the Hamiltonian in equation 1.2 is impossible. First, there is no closed form solution of equation 1.1 with this Hamiltonian. Second, numerical evaluation of  $\hat{H}$  is far from trivial.

Let's consider a simple example, the Argon atom. Say we want to solve this partial differential equation on a grid. Let's choose a very coarse  $10 \times 10 \times 10$  grid. Then at each time step we need to store  $10^{3\times(18+18)}=10^{108}$  values, corresponding to all the particle positions and the grid. Considering there are "only"  $\sim 10^{80}$  atoms in the known universe,<sup>2</sup> this is completely unreasonable.

This system has only 18 electrons and 18 nuclei, a far cry from the  $10^{23}$  or higher number of electrons in a typical condensed matter system, for example. Moreover, we have not taken into account floating point precision, or that we would need to calculate

this for possibly many time steps. Clearly, drastic approximations and more sophisticated methods are required.

#### 1.1. Overview of the Thesis

This dissertation fits into the field of nonrelativistic electronic structure theory, the branch of quantum chemistry concerned with the description of electrons and their correlation inside molecules and materials. More specifically, this dissertation focuses on high-accuracy (but often high-cost) ab initio methodologies, especially full configuration interaction quantum Monte Carlo (FCIQMC) and transcorrelation (TC). As such, we will only be discussing small systems consisting of only a few atoms, as they are tractible with a full, all-electron treatment with these methods. In principle, these methods should be able to be embedded<sup>3,4</sup> into more large-scale calculations using multiscale techniques, but this is outside the scope of my work. Nevertheless, the work herein is focused on methodologies, and not on particular physical systems.

The outline of the dissertation is as thus:

- Chapter 1 (this chapter) provides a basic overview of electronic structure theory methods and some of its principal concepts. Sections 1.5 and 1.6 in particular are largely based on the appropriate chapters of reference 5.
- Chapter 2 reviews the current works in so-called "explicitly correlated" methods, notably the well-established R12/F12 and the recently-reinvigorated TC.
- Chapter 3 provides a basic introduction to quantum Monte Carlo (QMC) and how it relates to FCIQMC.
- Chapter 4 discusses optimization strategies of Jastrow factors in the context of TC.
- Chapter 5 discusses an extension of the methods in the previous chapters to ensure size consistency and success when targeting strongly multi-reference problems.
- Chapter A provides an overview of the software pytchint developed in the group for evaluation of TC integrals.
- Finally, chapter 7 provides a review and an outlook for the field.

## 1.2. Principal Approximations

As discussed, to make any progress in electronic structure theory, we must make use of approximations. Of course, we must always be cautious and suspicious when using these,

and make sure they are valid for the systems in question. Thankfully, over the course of the last century, a cornucopia of different approximations has been developed to tackle the Schrödinger question, some of which will be discussed in this section.

#### 1.2.1. The Born-Oppenheimer Approximation

The most important approximation used in electronic structure theory is the Born-Oppenheimer approximation (BOA).<sup>6</sup> The BOA relies on the fact that the nuclei are much heavier than the electrons, with the mass of a single proton being almost 2000 times the mass of an electron. As an intuitive picture, we may think of the nuclei as moving much slower than the electrons, which can adapt themselves to the instantaneous positions of the nuclei. In mathematical terms, this means we can take the total wave function to be a product of its nuclear and electronic components,

$$\Psi_{\text{total}} = \Psi_{\text{nuc}}\Psi_{\text{elec}}.\tag{1.3}$$

Notice that the first two terms of equation 1.2 are independent of the electronic coordinates and, ipso facto, have no effect on  $\Psi_{\text{elec}}$ . This leads to the *electronic Hamiltonian* under the BOA, which can be written as

$$\hat{H}_{\text{elec}} = -\sum_{i} \frac{1}{2} \nabla_{i}^{2} - \sum_{i,I} \frac{Z_{I}}{r_{iI}} + \sum_{i>j} \frac{1}{r_{ij}}, \tag{1.4}$$

where we have simplified notation by using miniscule roman letters for the electrons and capital roman letters for the nuclei, and  $r_{ij} = |\mathbf{r}_i - \mathbf{r}_j|$  and  $r_{iI} = |\mathbf{r}_i - \mathbf{R}_I|$ , as well as by using atomic units.

In the language of second quantisation, equation 1.4 can be written as

$$\hat{H}_{\text{elec}} = \sum_{pq} h_{pq} a_p^{\dagger} a_q + \frac{1}{2} \sum_{pqrs} V_{pqrs} a_p^{\dagger} a_r^{\dagger} a_s a_q, \tag{1.5}$$

where p, q, r, s are general spin-orbital indices, and  $a_p$  ( $a_p^{\dagger}$ ) is the annihilation (creation) operator for an electron in spin-orbital p. These must obey the anti-commutation relations

$$\{a_p, a_q^{\dagger}\} := a_p a_q^{\dagger} + a_p^{\dagger} a_q = \delta_{pq} \tag{1.6}$$

$$\{a_p, a_q\} = \{a_p^{\dagger}, a_q^{\dagger}\} = 0$$
 (1.7)

so that the electrons satisfy the Pauli exclusion principle.

In equation 1.5,  $h_{pq}$  is a one-body integral,

$$h_{pq} = \int d^3r \; \phi_p^*(\mathbf{r}) \left( -\frac{1}{2} \nabla^2 - \sum_I \frac{Z_I}{|\mathbf{R}_I - \mathbf{r}|} \right) \phi_q(\mathbf{r})$$
 (1.8)

and  $V_{pqrs}$  is a two-body integral,

$$V_{pqrs} = \int d^3r \int d^3r' \ \phi_p^*(\mathbf{r}) \phi_r^*(\mathbf{r}') \frac{1}{|\mathbf{r} - \mathbf{r}'|} \phi_q(\mathbf{r}) \phi_s(\mathbf{r}'), \tag{1.9}$$

with a spatial-orbital basis  $\{\phi_i(\mathbf{r})\}$ .

For our purposes, we neglect the first term of equation 1.2, and treat the second term as approximately constant, so  $\hat{H}_{\text{nuc}} = \sum_{IJ} Z_I Z_J r_{IJ}^{-1}$ . Thus, we are left with the problem of solving the electronic structure problem, which is the subject of this dissertation. Solving the electronic problem for different nuclear positions gives us the potential energy surface.

Note that while chemists and physicists typically talk about the positions of nuclei in a molecule or solid as if they are fixed in place, as will be done in this dissertation, this is really a colloquialism. If the nuclei had an exact position and zero kinetic energy, the BOA would be in direct contradiction of the Heisenberg uncertainty principle. Instead, on the timescale of the electrons, due to the much higher mass of nuclei, in the BOA we treat the nuclei as approximately localised in a state in which their motion is much slower than that of the electrons (but, importantly, not zero). This keeps the approximation from being in conflict with the fundamental postulates of quantum theory.

The BOA is an immensely practical tool as it substantially simplifies our equations, and in many applications it is an excellent approximation. It will be a fundamental assumption throughout the rest of this dissertation, though it need not always be valid in all of quantum chemistry.

While we have done a lot to drastically reduce the complexity of equation 1.2, equation 1.5 is still intractible for large system sizes, scaling combinatorially with the size of the Hilbert space, as a function of the system size  $N_e$  (henceforth N), and the basis set size M. Hence, in addition to using a smaller basis, we still need extra approximations and sophisticated methodologies.

#### 1.2.2. Core Orbitals

In the ground state of a single atom, electrons first occupy the lowest orbitals. The first occupied shells are typically where the electrons are the most tightly bound.\*

Since they tend to be further from the nucleus in an atom, the valence (non-core) orbitals are typically the most affected by the introduction of additional atoms in the system. It is therefore often the valence orbitals that are most important in chemical systems. For this reason, we sometimes "freeze" the core orbitals, meaning they remain doubly occupied by electrons, and only correlate the valence orbitals. It is also common to refer to core electrons and valence electrons, meaning those electrons occupying the core and valence orbital space, respectively (though this is technically imprecise language, as electrons are indistinguishable).

<sup>\*</sup>This neglects symmetries: energy may be higher due to the angular momentum while staying spatially closer to the nucleus.

Similarly, we may also delete virtual orbitals, typically those of high energy, to further reduce the size of the problem. The remaining space is known as an active space, and is the basic idea of the complete active space (CAS) methods.

#### 1.2.3. Model Hamiltonians

In 1929, the surrealist artist René Magritte displayed a now-famous painting of a pipe with the caption *Ceci n'est pas une pipe* (French for "This is not a pipe"). It was meant to depict the idea that the painting itself is in a way treacherous: it may appear to be a pipe, but you cannot stuff it or smoke from it, as it is a representation of a pipe.

In a similar way, physicists and chemists often use *model Hamiltonians*, which do away with aspects of equation 1.2 that are not expected to be relevant to the problem at hand, resulting in new Hamiltonians that may be considered a representation of equation 1.2, much like Magritte's painting.\* Compared to *ab initio* Hamiltonians like equation 1.5, model Hamiltonians are generally much simpler, but depend on parameters whose values we may not necessarily know a priori.

The most famous model Hamiltonian is the Hubbard model,<sup>7</sup> most typically used to describe electrons in a periodic lattice, and can be written as

$$\hat{H}_{\text{Hubb}} = -\sum_{\langle m,n \rangle} \sum_{\sigma} t_{mn} a_{n\sigma}^{\dagger} a_{m\sigma} + \sum_{m} U_{m} \hat{n}_{m\uparrow} \hat{n}_{m\downarrow}, \qquad (1.10)$$

where  $\langle m, n \rangle$  denotes nearest neighbour sites,  $t_{mn}$  (often taken to be constant, t) is a parameter called the hopping amplitude, and  $U_m$  (often taken to be constant, U) is a parameter called the on-site repulsion. Here we have used a spatial orbital basis labels  $\{m, n\}$  and spin labels  $\sigma \in \{\uparrow, \downarrow\}$ .  $\hat{n}_{m\sigma} := a_{m\sigma}^{\dagger} a_{m\sigma}$  is the number operator.

Despite its apparent simplicity, the Hubbard model is a rich model, and has been used to describe a variety of phenomena, from metal-insulator transitions to high-temperature superconductivity. Moreover, it is not always so simple to solve, and has been the subject of much research.<sup>8,9</sup>

Let's consider an especially simple version of the model, with only two sites, and open boundary conditions:

$$\hat{H}_{\text{Hubb}} = -t \sum_{\sigma} (a_{1\sigma}^{\dagger} a_{2\sigma} + a_{2\sigma}^{\dagger} a_{1\sigma}) + U(\hat{n}_{1\uparrow} \hat{n}_{1\downarrow} + \hat{n}_{2\uparrow} \hat{n}_{2\downarrow}). \tag{1.11}$$

We could use this as an approximate description of the  $H_2$  molecule, where the electrons are restricted to only the lowest 1s orbitals. Compared to equation 1.2, this is a tremendously easy problem now.

<sup>\*</sup>We might further argue that all of science can be described this way, as scientific models are always "mere" representations of Nature, and are not Nature itself.

Of course, when solving a problem using a model Hamiltonian, we must be careful in choosing the correct one. In particular, the exact solution to the Hubbard model in one dimension<sup>8</sup> does not appear to be in agreement with the *ab initio* solution of a one-dimensional chain of hydrogen atoms.<sup>10,11</sup>

#### 1.3. The Variational Principle

One way of finding approximations to the lowest energy eigenstate of a quantum problem like that of equation 1.5 is the variational method. Several methods discussed in this dissertation build on the variational method, and the variational method itself is based on the variational principle, which states<sup>12</sup> for any "trial wave function"  $|\tilde{\Psi}\rangle$  and Hermitian operator (such as the electronic Hamiltonian)  $\hat{H}$ ,

$$\frac{\left\langle \tilde{\Psi} \middle| \hat{H} \middle| \tilde{\Psi} \right\rangle}{\left\langle \tilde{\Psi} \middle| \tilde{\Psi} \right\rangle} \ge E_0 := \frac{\left\langle \Psi \middle| \hat{H} \middle| \Psi \right\rangle}{\left\langle \Psi \middle| \Psi \right\rangle} \tag{1.12}$$

where  $E_0$  is the lowest eigenvalue (e.g. the ground state energy). That is, the energy of  $|\tilde{\Psi}\rangle$  is an upper bound of the exact energy of the system.

To prove this, consider an expansion of  $\left|\tilde{\Psi}\right\rangle$  terms of the exact eigenkets  $|k\rangle$  of  $\hat{H}$ , so that  $\hat{H}\left|k\right\rangle = E_{k}\left|k\right\rangle$  and  $\left|\tilde{\Psi}\right\rangle = \sum_{k} c_{k}\left|k\right\rangle$ . Then

$$\frac{\left\langle \tilde{\Psi} \middle| \hat{H} \middle| \tilde{\Psi} \right\rangle}{\left\langle \tilde{\Psi} \middle| \tilde{\Psi} \right\rangle} = \frac{\sum_{kk'} c_k^* c_{k'} \left\langle k \middle| \hat{H} \middle| k' \right\rangle}{\sum_{kk'} \left\langle k \middle| k' \right\rangle c_k^* c_{k'}}$$

$$= \frac{\sum_{k} |c_k|^2 (E_k - E_0)}{\sum_{k} |c_k|^2} + E_0$$

$$\geq E_0$$
(1.13)

where, in going from the first to the second line, we used the orthonormality of the eigenbasis  $\{|k\rangle\}$  and that the left- and right-eigenvectors are the same. That is, we used the fact the  $\hat{H}$  is a Hermitian operator. This is an important observation for future discussions, such as in section 2.5.

## 1.4. The Schrödinger Equation as a Matrix Problem

Implicit in the operator algebra of the fermionic creation and annihilation operators  $a^{\dagger}$  and a in equation 1.5 is the Pauli exclusion principle: we cannot have two or more fermions occupying identical states. In the context of electronic structure theory and this dissertation, the fermions and states in question are electrons and orbitals, respectively.

This handles the antisymmetry requirement of the fermionic wave function,  $\hat{P}_{ij}^{\sigma}\Psi = -\Psi$ , where  $\hat{P}_{ij}^{\sigma}$  permutes the *i*th and *j*th electrons with spin  $\sigma$ .

Our wave function can hence be expanded by a linear combination of Slater determinants (SDs),

$$\Psi = \sum_{i} c_{i} |D_{i}\rangle, \quad \text{where } |D_{i}\rangle = \frac{1}{\sqrt{N!}} \begin{vmatrix} \chi_{1}(\boldsymbol{x}_{1}) & \chi_{1}(\boldsymbol{x}_{2}) & \cdots & \chi_{1}(\boldsymbol{x}_{N}) \\ \chi_{2}(\boldsymbol{x}_{1}) & \chi_{2}(\boldsymbol{x}_{2}) & \cdots & \chi_{2}(\boldsymbol{x}_{N}) \\ \vdots & \vdots & \ddots & \vdots \\ \chi_{N}(\boldsymbol{x}_{1}) & \chi_{N}(\boldsymbol{x}_{2}) & \cdots & \chi_{N}(\boldsymbol{x}_{N}) \end{vmatrix}$$
(1.14)

for a spin-orbital basis  $\{\chi_p(\mathbf{x}_q)\}$ . A spin-orbital is typically written as a product of a spatial part and a spin part, e.g.  $\chi(\mathbf{x}) = \phi(\mathbf{r})\omega(\sigma)$ .

Given such a finite spin-orbital basis of size 2M (M spatial orbitals with 2 different spins each),\* the SD expansion is also finite. Therefore, we may write  $\hat{H}$  as a square matrix and equation 1.5 as an eigenvalue equation,

$$\hat{H}\Psi = E\Psi. \tag{1.15}$$

Note, however, that in order to make this a finite-dimensional matrix eigenvalue problem, we needed to use a finite spin-orbital basis. Thus, E in equation 1.15 is not the true ground state energy of the system, but indeed only an approximation within that basis set. This is the case even if we were to solve equation 1.15 exactly. In order to get the true ground state energy, we need to reach the complete basis set (CBS), i.e.  $M \to \infty$ .

With equation 1.15, we have transformed the hopelessly intractible partial differential equation 1.2 into a finite algebraic eigenvalue problem, which is much better suited for solving on a computer.

If we have  $N_{\uparrow}$  spin-up electrons and  $N_{\downarrow}$  spin-down electrons, we then have

$$|\mathcal{H}| = \binom{M}{N_{\uparrow}} \binom{M}{N_{\downarrow}} \tag{1.16}$$

where  $|\mathcal{H}|$  denotes the size of the Hilbert space, and we have assumed that the number of spin-up orbitals M is equal to the number of spin-down orbitals. Here,  $\binom{n}{r} = \frac{n!}{r!(n-r)!}$  is a binomial coefficient.

The combinatorial scaling is still unfavourable. For a closed-shell system with 20 electrons in 20 orbitals,  $|\mathcal{H}| = 3.41 \times 10^{10}$ . Assuming we wish to have double precision,<sup>†</sup> to store the whole matrix we would need about  $10^{23}$  bytes, or 100 zettabytes. So, while

<sup>\*</sup>We assume here the same number of spin- $\uparrow$  and spin- $\downarrow$  orbitals for simplicity, though it need not always be the case.

 $<sup>^{\</sup>dagger}$ A double precision floating-point number has 1 bit for the sign, 11 bits for the exponent, and 52 bits for digits in the number (known as the mantissa or significand). This resolves to 64 bits or 8 bytes.

we've substantially reduced the complexity of the problem, it is still extremely demanding. More approximations are therefore needed.

#### 1.4.1. Orbitals

Orbitals (and spin-orbitals), a term already used in this text, describe the spatial distribution (and in the case of spin-orbitals, also the spin) of a single electron. We assume that spatial orbitals form an orthonormal set, and that they form a complete space for an arbitrary wave function. However, in theory to have a CBS would mean needing an infinite number of basis functions. In practice, we must of course have a finite basis, and so this naturally leads to a hierarchy of basis sets, where the typically larger basis sets are both more expensive and more accurate.

Many different basis set families exist, but among the most popular are those due to Dunning and coworkers.<sup>14</sup> Additionally, many different basis set extrapolation techniques have also been developed.<sup>15–22</sup>

Strictly speaking, any kind of set of functions can be used as a basis set, as long as they are a complete set. However, in practice we may broadly categorise them into two groups: Slater-type orbitals (STOs) and Gaussian-type orbitals (GTOs).

The exact solution for the Hydrogen atom is the Slater-type function  $\frac{1}{\sqrt{\pi}}e^{-r}$ , and similarly we know that molecular orbitals also decay as  $\sim e^{-\zeta r}$ .

However, these STOs are difficult to work with, as integrals of the form

$$\int d^3 r_1 \int d^3 r_2 \, \phi_p^*(\boldsymbol{r}_1) \phi_r^*(\boldsymbol{r}_2) \frac{1}{r_{12}} \phi_q(\boldsymbol{r}_1) \phi_s(\boldsymbol{r}_2)$$

$$\tag{1.17}$$

are routinely needed.

Therefore, GTOs have been developed,<sup>23</sup> which have the form  $e^{-\alpha r^2}$ . GTOs are much easier to handle, as the product of a gaussian is also a gaussian, and the integral of a gaussian is exactly known, so equations like 1.17 have known solutions. With this comes a tradeoff, since we know STOs more faithfully capture the form of molecular orbitals, but GTOs are much better suited for computations. For this, the most common basis sets typically use a linear combination (or "contraction") of gaussians fitted to a STO. An illustration is shown in figure 1.1.

It must be noted, however, that STOs and GTOs have different asymptotic behaviour for  $r \to \infty$  and  $r \to 0$ . The latter in particular will be discussed in section 2.1.

#### 1.4.2. Density Matrices

The 1-electron reduced density matrix (1RDM) is defined by its matrix elements

$$\gamma_{pq} = \langle \Psi | \, a_q^{\dagger} a_p \, | \Psi \rangle \,, \tag{1.18}$$

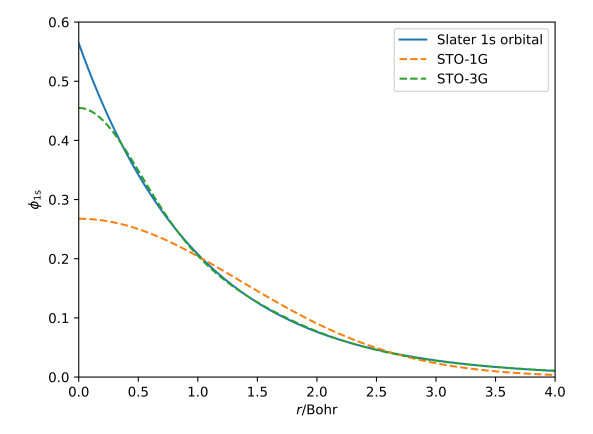


Figure 1.1. Illustration of an s-type orbital and the fitted gaussians (STO-xG means x gaussians are used to approximate the STO). Note that for much of the curve, there is good agreement, but not for the short- and long-range. The short-range behaviour in particular leads to complications, and is the focus of the methods discussed in chapter 2. The long-range behaviour is typically less of a problem, since the integrals we are interested in typically decay with r. Fitting parameters for the gaussians were taken from reference 24.

or written in a spin-orbital basis:

$$\gamma(\boldsymbol{x}_1, \boldsymbol{x}_2) = \sum_{pq} \phi_q(\boldsymbol{x}_2)^* \phi_p(\boldsymbol{x}_1) \gamma_{pq}$$
(1.19)

Similarly, the 2-electron reduced density matrix (2RDM) is defined as

$$\Gamma_{pqrs} = \langle \Psi | a_s^{\dagger} a_r^{\dagger} a_q a_p | \Psi \rangle. \tag{1.20}$$

Any one- and two-electron Hermitian operator  $\Omega$  can be written (in a spin-orbital basis) as

$$\langle \Psi | \Omega | \Psi \rangle = \sum_{pq} \gamma_{pq} \Omega_{pq} + \sum_{pqrs} \Gamma_{pqrs} \Omega_{rpsq}. \tag{1.21}$$

#### 1.4.3. Electron Correlation

Two variables are independent (or uncorrelated) if the joint probability distribution is the product of their expected values, i.e.

$$P(x_1, x_2) = P(x_1)P(x_2). (1.22)$$

By Bayes' theorem, <sup>25,26</sup> this can be rewritten in terms of the conditional probability,

$$P(x_1|x_2) = \frac{P(x_1, x_2)}{P(x_2)} = \frac{P(x_1)P(x_2)}{P(x_2)} = P(x_1).$$
(1.23)

The variables are said to be correlated if the above is not true.

In the case of electronic structure, the variables in question are  $\mathbf{x} := (\mathbf{r}, \sigma)$ , the spatial coordinates and spin of the electrons. Therefore, when we speak of electron correlation, we are referring to these relations. Furthermore, as electrons are indistinguishable, for every pair  $(\mathbf{x}_1, \mathbf{x}_2)$ ,

$$P(\boldsymbol{x}_1) = P(\boldsymbol{x}_2) = \frac{1}{N}\rho(\boldsymbol{x}) \tag{1.24}$$

$$P(x_1, x_2) = \frac{1}{N(N-1)} \rho(x_1, x_2)$$
 (1.25)

where  $\rho(\mathbf{x})$  is the electron density and  $\rho(\mathbf{x}_1, \mathbf{x}_2)$  is the probability of finding electrons at  $(\mathbf{x}_1, \mathbf{x}_2)$  simultaneously, known as the pair density,

$$\rho(\boldsymbol{x}) = N \int d^3x_2 \cdots d^3x_N \Psi^*(\boldsymbol{x}, \boldsymbol{x}_2, \boldsymbol{x}_3, \cdots, \boldsymbol{x}_N) \Psi(\boldsymbol{x}, \boldsymbol{x}_2, \boldsymbol{x}_3, \cdots, \boldsymbol{x}_N)$$
(1.26)

$$\rho(\boldsymbol{x}_1, \boldsymbol{x}_2) = N(N-1) \int d^3 x_2 \cdots d^3 x_N \Psi^*(\boldsymbol{x}_1, \boldsymbol{x}_2, \boldsymbol{x}_3, \cdots, \boldsymbol{x}_N) \Psi(\boldsymbol{x}_1, \boldsymbol{x}_2, \boldsymbol{x}_3, \cdots, \boldsymbol{x}_N).$$
(1.27)

There are two key sources of electron correlation: Fermi correlation, which arises from the fact that electrons obey Fermi statistics, i.e. the wave function is antisymmetric with respect to exchange of  $x_1$  and  $x_2$ , and Coulomb correlation, which arises from the fact that electrons repel each other.

#### 1.5. The Hartree-Fock Method

The typical starting point to most *ab initio* electronic structure methods is the Hartree-Fock (HF) method.<sup>27–29</sup> The key approximation in HF is that we treat the exact N-body wave function solution as approximately a single SD.\* Then, we variationally (see section 1.3) optimise the energy of the system with respect to parameters pertaining to the spin-orbitals. We will refer to this SD as  $|\Phi_0\rangle$ .

Then, for some arbitrary initial configuration  $|0\rangle$ ,

$$|\Phi_0\rangle = e^{-\kappa} |0\rangle. \tag{1.28}$$

Here,

$$\kappa = \sum_{pq} \kappa_{pq} a_p^{\dagger} a_q \tag{1.29}$$

is an anti-Hermitian operator (i.e.  $\kappa_{pq}^* = -\kappa_{qp}$ ) whose matrix elements  $\kappa_{pq}$  are the orbital rotation coefficients and will be the variational parameters. The anti-Hermiticity of  $\kappa$  ensures that the orbital rotation  $e^{-\kappa}$  is unitary.

With  $\langle \Phi_0 | \Phi_0 \rangle = 1$ , we want

$$E_{\rm HF} = \min_{\kappa_{pq}} \langle \Phi_0 | \hat{H} | \Phi_0 \rangle. \tag{1.30}$$

This is a nonlinear equation, and hence the parameters  $\kappa_{pq}$  must be determined iteratively. Since  $e^{-\kappa}$  is unitary, the orthonormality of the spin-orbitals is preserved,  $\langle \phi_p | \phi_q \rangle = \delta_{pq}$ .

The HF method can also be understood as a mean-field theory. That is, we treat the N-body problem as N one-body problems, where a single electron is in an effective "averaged" potential from the other N-1 electrons. This effective one-body Hamiltonian is known as the Fock operator,

$$f = \sum_{pq} f_{pq} a_p^{\dagger} a_q. \tag{1.31}$$

The Fock operator f may be written as

$$f = h + V_{\text{eff}} \tag{1.32}$$

where the Fock potential's matrix elements are  $V_{\text{eff},pq} = \sum_{i \in \text{occ}} (V_{pqii} - V_{piiq})$ .

<sup>\*</sup>More precisely, the HF approximation is to treat the wave function as a single configuration. In the context of this dissertation, that configuration will always be a SD.

The Hartree-Fock orbitals, which ultimately dictate the parameters for the minimisation procedure, are determined by diagonalising the Fock matrix,

$$f_{pq} = \epsilon_p \delta_{pq} \tag{1.33}$$

where the eigenvalues  $\epsilon_p$  are known as the orbital energies.

Since f itself depends on the orbitals, the solution must be determined self-consistently. Hence HF is often referred to as a self consistent field (SCF) method.

Often, the orbitals for the  $\uparrow$  and  $\downarrow$  electrons are restricted to be the same. This is known as the restricted Hartree-Fock (RHF) method. Otherwise, we might instead use the unrestricted Hartree-Fock (UHF) method, where they are treated independently. However, since this breaks spin symmetry, we might have spin contamination. Restricted open-shell Hartree-Fock (ROHF) is a variant that can treat open-shell systems which RHF otherwise would not be able to treat while remaining an eigenstate of the  $S^2$  (total spin) operator.

#### 1.5.1. Correlation Energy

The correlation energy (for a given basis set) is defined as the difference between the exact energy and the HF energy,

$$E_{\rm corr} = E_{\rm exact} - E_{\rm HF}. \tag{1.34}$$

It should be noted, however, that despite this name the HF method captures Pauli exchange, and hence does actually correlate electrons. Nevertheless, we typically neglect this when we speak of "correlation energy".

Similarly, the Coulomb hole is defined as the remaining part of the wave function that is not captured by the HF method,

$$\Psi_{\text{hole}} = \Psi_{\text{exact}} - \Psi_{\text{HF}}.\tag{1.35}$$

#### 1.5.2. The Roothaan-Hall Equations

The Roothaan-Hall equations are a matrix representation of the HF approximation.<sup>30,31</sup> Given a basis set, typically GTOs, the Roothaan-Hall equations are given by the generalised eigenvalue problem,

$$FC = SC\epsilon \tag{1.36}$$

where C is a matrix of coefficients, F is the Fock matrix (which depends on C), S is the overlap matrix (which reduces to the identity matrix for orthonormalised bases), and  $\epsilon$  is a diagonal matrix of orbital energies. The Roothaan-Hall equations are obtained by expanding the unknown molecular orbitals (MOs) in a basis of known functions, typically atomic orbitals (AOs).

Since this representation is in matrix form, instead of in terms of derivatives and integrals, it is more amenable to conventional computational techniques.

#### 1.6. Post-Hartree-Fock Methods

While the HF method is convenient and can be used efficiently to solve for many electrons, it is oftentimes not sufficient to describe the electronic structure of a molecule. To account for the remaining correlation, numerous post-Hartree-Fock methods (that is, methods run after HF) have been formulated, some of which are described in this section.

Correlation energy is typically categorised as either dynamical or static, although their effects are not mutually exclusive and so sometimes the distinction can be hazy.

Dynamical correlation is remaining correlation that arises due to the instantaneous repulsion from the motion (dynamics) of the electrons.

Static correlation, on the other hand, is related to the degeneracy or near-degeneracy of configurations. In general, to accurately describe a system with strong static correlation, a so-called multi-reference method is needed. As will be seen in chapter 5, static correlation plays a big role for molecules at dissociation.

We therefore have two "axes" on which to approach the exact solution to equation 1.5:

- The basis set chosen, where going to a larger basis set (in a systematic way) improves the result, e.g. cc-pVDZ to cc-pVTZ to cc-pVQZ, until eventually reaching the CBS limit.
- The "hierarchy of theories" where the more accurate the method, the closer to the full configuration interaction (FCI) limit (discussed below). Many post-Hartree-Fock methods can be systematically improved, for example by considering additional excitations in a coupled cluster method.

#### 1.6.1. Configuration Interaction

Configuration interaction (CI) is perhaps the most conceptually straightforward post-Hartree-Fock method. Instead of using a single SD, we approximate the wave function as a linear combination of SDs. That is,

$$|\Phi_{\rm CI}\rangle = \sum_{p} c_p |D_p\rangle \tag{1.37}$$

where  $|D_p\rangle$  is a SD. This may alternatively be written\*

$$|\Phi_{\text{CI}}\rangle = (\mathbb{1} + \sum_{ia} c_i^a \hat{a}_a^{\dagger} \hat{a}_i + \sum_{ijab} c_{ij}^{ab} a_a^{\dagger} a_b^{\dagger} a_i a_j + \dots) |\Phi_0\rangle$$

$$(1.38)$$

<sup>\*</sup>We adopt the common notation convention where i, j, k, ... denote occupied orbitals, a, b, c, ... virtual (or unoccupied) orbitals, and p, q, r, ... unspecified.

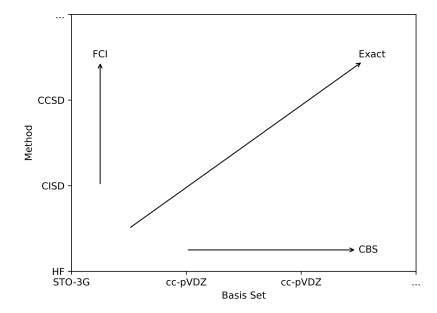
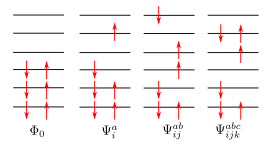


Figure 1.2. A sketch for the hierarchy of theories and basis sets. There are two axes on which to systematically improve until reaching the exact solution to the time-independent Schrödinger equation under the BOA: increasing basis set size (x axis) and increasing the level of theory (y axis). For most of this dissertation, we assume to be at or close to the highest level of theory (the FCI limit), as we mostly employ FCIQMC. We therefore focus on methods of rapidly converging to the CBS limit without the inherently expensive process of adding more basis functions.

where the number of terms in the sum depends on how many excitations we consider, i.e. how far we truncate the CI expansion. The energy and wave function are then found as the lowest eigenvalue and corresponding eigenvector, respectively, of the Hamiltonian written in the basis of these SDs. This can be done, for example, by exact diagonalisation. If all possible excitations are considered, then we have reached the CBS limit, and the solution is exact within that basis set.



**Figure 1.3.** Illustration of a RHF reference and examples of single-, double-, and triple-excitations. The method CISDT, for example, would consider all determinants of the forms illustrated.

There are some important limitations to this method, however. Perhaps most notable is that the CI method is not size consistent. For two noninteracting systems A and B, we expect the composite system A + B to have energy equal to the sum of that for A and B, i.e.  $E_{A+B} = E_A + E_B$ . However, within a truncated CI method, this is not the case\*, due to the additive ansatz 1.38.

Additionally, if we wish to go to the FCI limit and ensure size consistency and accurate results, we once again have bad scaling, as discussed in section 1.4. However, there exist sparse diagonalisation routines that need not store the entire matrix H, such as the Davidson,  $^{32}$  Lanczos,  $^{33}$  and Arnoldi $^{34}$  (for non-Hermitian H) methods, which target the ground state and need to store only a fraction of the matrix. Still, these scale with the size of the eigenvector, which is still prohibitively expensive for large systems. A more practical approach to reaching FCI accuracy will be discussed later in section 3.4, where we discuss the FCIQMC algorithm.

#### 1.6.2. Multi-Configurational Self-Consistent Field

As a generalisation of the HF method, multiconfigurational self-consistent field (MCSCF) is particularly well-suited for problems with static correlation.<sup>5,35,36</sup> As in HF, we minimise the electronic energy with respect to variational parameters; however, now we also simultaneously minimise with respect to expansion coefficients for a set of configurations

<sup>\*</sup>Of course, in FCI we necessarily have size consistency, since we have the exact solution for the basis set.

(e.g. SDs). i.e. consider the MCSCF wave function

$$|\kappa, \mathbf{c}\rangle = e^{-\kappa} \sum_{i} c_i |D_i\rangle,$$
 (1.39)

where  $\kappa$  is familiar from HF, and the  $c_i$  coefficients familiar from CI. In MCSCF, we optimise

$$E_{\text{MCSCF}} = \frac{\langle \kappa, \boldsymbol{c} | H | \kappa, \boldsymbol{c} \rangle}{\langle \kappa, \boldsymbol{c} | \kappa, \boldsymbol{c} \rangle}.$$
 (1.40)

Like in HF, the variational parameters appear nonlinearly and must be optimised iteratively. Determining an appropriate set  $\{|D_i\rangle\}$  tends to be challenging, and even for small systems generating an MCSCF wave function can prove intractible.

#### Complete Active Space Self-Consistent Field

One "flavour" of MCSCF that has proven particularly successful is complete active space self-consistent field (CASSCF).<sup>37–40</sup> In CASSCF, instead of inspecting individual configurations, we consider a set of configurations that satisfy a set of criteria. In particular, we partition the orbitals into three sets: the core, active, and virtual (unoccupied) regions. The core orbitals, as alluded to in section 1.2.2, are approximated to be doubly occupied. The virtual orbitals are approximated to always be unoccupied. The active orbitals are the remaining orbitals, which can have occupations between 0 and 2.

The MCSCF expansion is found by considering all possible excitations of the active electrons in the active space. Notice that in the limit of an empty active space, we recover the HF method, and in the limit of an active space containing all orbitals, we recover the FCI method.

#### 1.6.3. Perturbation Theory

Perturbation theory is a set of approximate mathematical methods for solving problems involving small disturbances (perturbations) to a problem with a known solution (the unperturbed problem). If these perturbations are not too large, then the solution of the perturbed problem is close to that of the unperturbed problem, and can be expressed as the solution of the unperturbed problem plus some corrections. Perturbation theory fails, however, if the perturbation is large.

#### Rayleigh-Schrödinger

Perturbation theory applied to time-independent problems is sometimes referred to as Rayleigh-Schrödinger perturbation theory. 12,41,42 Consider a Hamiltonian,

$$H_{\rm PT} = H_0 + \lambda H',\tag{1.41}$$

where  $H_0$ , referred to as the unperturbed Hamiltonian,  $\lambda$  is an arbitrary (real) parameter controlling the strength of the perturbation, and H' is the perturbation. We assume that we know the exact solution to  $H_0$ , such that we have all eigenstates  $\{|\Psi_n^{(0)}\rangle\}$  and their corresponding eigenvalues  $\{E_n^{(0)}\}$ .

To obtain the solution to the true Hamiltonian H, we expand in terms of the perturbation  $\lambda$ ,

$$|\Psi_n\rangle = \sum_{k=0}^{\infty} \lambda^k \left| \Psi_n^{(k)} \right\rangle \tag{1.42}$$

and

$$E_n = \sum_{k=0}^{\infty} \lambda^k E_n^{(k)}.$$
 (1.43)

Since in this work we are generally interested in the ground state, we are typically targeting n = 0.

By inserting these expansions into the time-independent Schrödinger equation, we equate terms of the same order in  $\lambda$ , which leads to the corrections to the energy and wave function. We may then truncate the expansion to some order. This allows for a systematically improvable result.

#### Møller-Plesset

Møller-Plesset perturbation theory $^{43}$  is a special case of Rayleigh-Schrödinger perturbation theory, and is the variant most commonly seen in quantum chemistry.

In Møller-Plesset perturbation theory, the unperturbed Hamiltonian is chosen to be the Fock operator 1.31. The perturbed Hamiltonian is known as the fluctuation potential and is the difference between the true Coulomb interaction and the effective one-electron potential discussed in section 1.5.

That is,

$$H_0 = f, \quad H' = \sum_{i < j} r_{ij}^{-1} - V_{\text{eff}} = H - H_0.$$
 (1.44)

If we apply Rayleigh-Schrödinger perturbation theory, we find the zeroth order wave function to be the HF wave function,

$$\left|\Psi_{0}^{(0)}\right\rangle = \left|\Phi_{0}\right\rangle, \quad f\left|\Phi_{0}\right\rangle = \underbrace{\sum_{i}}_{E_{0}^{(0)}} \epsilon_{i} \left|\Phi_{0}\right\rangle$$
 (1.45)

where we have also identified the zeroth-order energy as the sum of orbital energies  $E^{(0)} = \sum_{i} \epsilon_{i}$ . With a bit more busywork, we get

$$E^{(1)} = \left\langle \Psi_0^{(0)} \middle| H' \middle| \Psi_0^{(0)} \right\rangle = \left\langle \Phi_0 \middle| H' \middle| \Phi_0 \right\rangle \tag{1.46}$$

$$E^{(2)} = \sum_{n>0} \frac{|\langle \Psi_n^{(0)} | H' | \Psi_0^{(0)} \rangle|^2}{E_0^{(0)} - E_n^{(0)}} = -\sum_{i>j,a>b} \frac{|\langle \Phi_{ij}^{ab} | H' | \Phi_0 \rangle|^2}{\epsilon_a + \epsilon_b - \epsilon_i - \epsilon_j},$$
(1.47)

where  $|\Phi_{ij}^{ab}\rangle = a_a^{\dagger} a_b^{\dagger} a_i a_j |\Phi_0\rangle$  is a doubly-excited state with respect to the HF wave function, and  $\epsilon_a + \epsilon_b - \epsilon_i - \epsilon_j = E_0^{(0)} - E_n^{(0)}$  is the energy difference between two eigenstates of the Fock operator.

Some features worth noting about Møller-Plesset perturbation theory:

- The sum of zeroth- and first-order energies is the HF energy:  $E_{\rm HF}=E_0^{(0)}+E_0^{(1)}$ .
- By the variational principle,  $E_0^{(0)} < E_n^{(0)}$  for all n > 0 (except for degenerate ground states), so  $E_0^{(2)} < 0$ , i.e. the energy always decreases.
- If the ground state is degenerate, the term diverges. It might be possible to lift the degeneracy by a change of basis, however.
- If the HF solution is already a good approximation for the system, then Møller-Plesset perturbation theory can provide surprisingly good (and size consistent) results. Hence, this method is typically only applicable to single-reference problems (i.e. those without strong static correlation).

While higher-order approaches exist and see use, the most popular is second-order Møller-Plesset perturbation theory (MP2), due to its excellent compromise between cost and accuracy. That is, for only a little extra work after a successful HF, applying MP2 can improve our results considerably.

#### 1.6.4. Coupled Cluster Theory

The lack of size consistency in CI theory arises from the linear ansatz in section 1.6.1. By instead using an exponential ansatz, we arrive at one of the most successful theories in electronic theory, coupled cluster (CC). 44–47 While the methods presented here are inherently single-reference and build on a single SD, multi-reference generalisations to CC theory exist and are the study of active research. 48–53 Therefore, the presented "flavours" of CC are generally expected to fail for systems with strong static correlation.

#### Standard Coupled Cluster

To ensure a multiplicatively-separable wave function, we use a multiplicative ansatz,

$$|\Psi_{\rm CC}\rangle = e^T |\Phi_0\rangle, \qquad (1.48)$$

where  $|\Phi_0\rangle$  is the reference (typically HF) wave function, and T is the cluster operator. The cluster operator is defined  $T=T_1+T_2+T_3+\cdots$ , where  $T_n$  is the n-body cluster operator, made up of all possible n-body excitations, for example,  $T_1=\sum_{i,a}t_{ia}a_a^{\dagger}a_i$  for the one-body cluster operator, and  $T_2=\sum_{i< j,a< b}t_{ij}^{ab}a_a^{\dagger}a_b^{\dagger}a_ja_i$  for the two-body cluster operator.

It is worth noting that CC and CI are identical, differing only in their parametrisation, when neither are truncated. They both provide FCI-level accuracy.

By inserting equation 1.48 into the time-independent Schrödinger equation and premultiplying by  $e^{-T}$ , we obtain

$$\langle \Phi_0 | e^{-T} H e^T | \Phi_0 \rangle = E \tag{1.49}$$

$$\left\langle \Phi_{ij...}^{ab...} \right| e^{-T} H e^{T} \left| \Phi_{0} \right\rangle = 0$$
 (1.50)

where  $\left|\Phi_{ij...}^{ab...}\right\rangle$  is a *n*-body excitation with respect to the reference wave function.

Note, however, that these equations do not rely on the Variational Principle. Instead of minimising a functional, we solve equations 1.50 for the amplitudes, which give the energy. While we still cannot reasonably always include all excitations in the cluster operator T, due to the multiplicative ansatz, even if we truncate the cluster operator, we get size-consistent results. In particular, any truncated CC wave function ansatz will contain contributions from all determinants in Fock space. The simplest CC truncation commonly used is the coupled cluster with single and double excitations (CCSD) method. In a sense, CCSD and any other truncated CC method is an approximation for FCI where the coefficients are not approximated to be zero (like in CI) but instead the higher-order excitations terms are generated from the lower-order excitation terms in a size-consistent fashion.

CC theory owes much of its success to its high accuracy at a reasonable cost. CCSD and coupled cluster with single, double and triple excitations (CCSDT) in particular scale as  $\mathcal{O}(M^6)$  and  $\mathcal{O}(M^8)$  respectively, with M the number of spatial orbitals. Every additional set of excitations incurs an additional factor of  $\mathcal{O}(M^2)$  scaling.

#### Distinguishable Cluster Theory

The more recently-developed distinguishable cluster with single and double excitations (DCSD) method's central philosophy is removing irrelevant or nonphysical terms while maintaining many of CC's desireable properties, such as size consistency.<sup>54</sup>

This method was shown to give quantitatively better results than standard CC for a variety of systems, and qualitatively correct results for systems where CC can fail spectacularly, such as the N<sub>2</sub> binding curve. Since its first publication, distinguishable cluster (DC) has been extended to explicitly-correlated F12 theory,<sup>55</sup> equation of motion (EOM),<sup>56</sup> tailored coupled cluster,<sup>57</sup> and more.

#### Combining Coupled Cluster with Perturbation Theory

It is possible to avoid the computationally-expensive scaling of CC somewhat, by treating some terms perturbatively. The most famous of these is the coupled cluster with single and double excitations and perturbative triples (CCSD(T)) method,<sup>58</sup> in which the single-and double-excitations are treated fully, whereas the triple-excitations are treated as perturbations, as in section 1.6.3.

CCSD(T) scales as  $\mathcal{O}(M^7)$  and for the price provides extremely accurate results. The perturbative correction is a step that is applied only once at the end of a CCSD calculation. This is why sometimes this method is referred to as the "gold standard" of quantum chemistry.

## **Explicitly Correlated Methods**

With the advent of modern computers combined with a vast array of sophisticated algorithms from which to choose, *ab initio* quantum chemistry has become a tremendously powerful tool, going beyond the study of small atoms, to molecules and solids, and are among the most effective and systematically improvable techniques to date. Nevertheless, convergence to the CBS limit is notoriously slow.

In particular, consider the popular basis set family developed by Dunning and coworkers, correlation-consistent polarised valence with X-zeta quality (X=D, T, Q, 5, 6, ...) (cc-pVXZ).<sup>14,21,59-61</sup> The size of these basis sets scale as  $M \in \mathcal{O}(X^3)$ , and since for standard post-Hartree-Fock discussed in chapter 1.6 we require four-index integrals, our computation time will scale at best as  $t \in \mathcal{O}(X^{12})$ .<sup>62</sup>

Meanwhile, the CBS correlation error scales as  $\epsilon \in \mathcal{O}(X^{-3})^{16,18}$  or  $\epsilon \in \mathcal{O}(M^{-1})$ ,  $^{63}$  resulting in time  $t \in \mathcal{O}(M^{-\frac{1}{4}})$ . Thus, the methods discussed so far come with the painful cost of requiring very large basis sets to approach high-accuracy results.

Explicitly correlated methods are a class of electronic structure methods specifically designed to address this unfavourable scaling by explicitly including the interelectronic distance  $r_{12}$ , and is the subject of this chapter. As the R12/F12 family of methods is the most mature of the explicitly correlated electronic structure methods, many reviews focusing on this topic already exist. This chapter in particular is in large part based on references [62, 64–66].

## 2.1. The Cusp Conditions

Consider two charged point particles in a system described by the Hamiltonian of equation (1.5). By the Schrödinger equation, the local energy

$$E_L := \frac{H\Psi}{\Psi} \tag{2.1}$$

must be constant in the exact solution. However, when these two particles coalesce, i.e.  $r_{12} \to 0$ , the Coulomb potential,  $r^{-1}$ , diverges. Thus, for the local energy to be constant, we must have that near coalescence points, the kinetic energy exactly cancels the Coulomb energy. A more formal treatment of this argument leads to the (antiparallel)

electron-electron Kato cusp condition,<sup>67</sup>

$$\left. \frac{\widetilde{\partial \Psi}}{\partial r_{12}} \right|_{r_{12} \to 0} = \frac{1}{2} \Psi(r_{12} = 0)$$
 (2.2)

where the tilde represents spherical averaging.

This cusp condition was also later generalised.<sup>68,69</sup> Early literature on the subject suggested that the success of explicitly correlated methods were due to the superior description of short-range correlation effects, and in particular in their much more faithful capturing of the cusp conditions like equation 2.1.<sup>70–73</sup> However, further study found that the correlation error from a bad description of the wave function around a small sphere centred on the cusp is actually negligible.<sup>62,74–76</sup> Instead, the success of explicitly correlated methods is actually due to the superior description of the overall shape and size of the Coulomb hole, which has a radius on the order of the atomic radius.

To understand why gaussian-type basis sets fail so spectacularly at capturing the cusp behaviour, it is instructive to consider a simpler example, like that of approximating |x| by its Fourier decomposition. Such an illustration is found in figure 2.1. It is also worth noting that STOs do not suffer as badly from this limitation,<sup>77</sup> but as discussed in chapter 1.4.1, they are unsuccessful due to their lack of practicality.\*

## 2.2. Hylleraas Methods

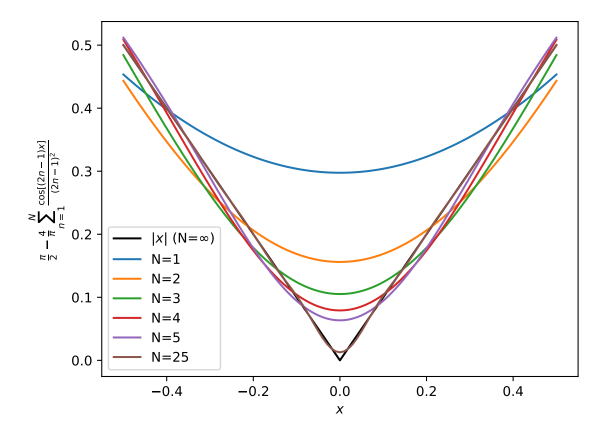
Almost 30 years prior to Kato's landmark paper describing cusps in the analytical form for the wave function, there was already work being done to understand the significance of including  $r_{12}$  terms in the wave function. Instead of being motivated by rigorous mathematics like Kato's, Slater was motivated by studies in the He atom. In particular, he tried to construct a wave function which faithfully represents both the core region as well as the Rydberg limit (i.e. a highly excited atom where the electron is very far from the nucleus).<sup>77–80</sup> This led him to suggest multiplying the wave function by a factor

$$e^{-2(r_1+r_2)+r_{12}/2},$$
 (2.3)

which can easily be shown to satisfy the cusp equation (2.2).

However, the first successful explicitly correlated electronic structure calculation is typically attributed to Hylleraas,<sup>65</sup> where he aimed to improve convergence of orbital expansions for helium.<sup>81,82</sup> In this method, the coordinates  $s := r_1 + r_2$ ,  $t := r_1 - r_2$  and

<sup>\*</sup>Note that STOs do not suffer from the electron-nuclear cusp, which is in any case not the focus of many explicitly correlated methods, such as F12, as discussed later in this chapter. Even with STOs, we have electron-electron cusps.



**Figure 2.1.** A toy example of the Coulomb cusp. Here, the Fourier expansion  $x \approx \frac{\pi}{2} - \frac{4}{\pi} \sum_{n=1}^{N} \frac{\cos[(2n-1)x]}{(2n-1)^2}$  is plotted for a few values of N, including the exact solution. As can be seen, even for many terms, the Fourier expansion is a poor descripter in the cusp region. Indeed, the only way to describe it exactly is with an infinite number of terms.

 $u := r_{12}$  are used to construct the wave function

$$\Psi_N(s, t, u) = e^{-\alpha s} \sum_{k}^{N} c_k s^{l_k} t^{2m_k} u^{n_k}.$$
 (2.4)

In particular, using only three terms (N=3), and variationally optimising for the parameters  $c_1, c_2, c_3$ , Hylleraas was able to reach within 1.3 millihartree from the exact result.

Since then, there was rapid development on this approach and combining it with CI (which came to be known as the CI-Hyl methods).<sup>83–91</sup> In CI-Hyl methods, the wave function is expanded as in CI,

$$\Psi = \sum_{k} c_k \Phi_k \tag{2.5}$$

where

$$\Phi_k = \mathcal{A}r_{ij}^{\nu_k} \prod_i \chi_{k_i}(\boldsymbol{x}_i) \tag{2.6}$$

where  $\chi_k$  is a spin-orbital basis and  $\mathcal{A}$  is the antisymmetriser operator.

However, CI-Hyl methods were to eventually fall out of favour. This is because the expansions involve exceedingly difficult integrals involving many electrons and over products of correlation factors. This significantly restricts the tractibility and scalability of the method, and it has since largely gone unused.

## 2.3. Explicitly Correlated Gaussians

Boys<sup>92</sup> and Singer<sup>93</sup> independently introduced gaussian basis functions with explicit correlation for calculations on molecules.<sup>94</sup> These methods are referred to as Explicitly Correlated Gaussians (ECGs), or in the case of functions of two electrons, Gaussian-type Geminals (GTGs). A spherical GTG may be written (with nuclar coordinates  $\mathbf{R}_1$  and  $\mathbf{R}_2$ ) as

$$g(\mathbf{r}_1, \mathbf{r}_2) = \exp(-\zeta_1 |\mathbf{r}_1 - \mathbf{R}_1|^2 - \zeta_2 |\mathbf{r}_2 - \mathbf{R}_2|^2 - \gamma |\mathbf{r}_1 - \mathbf{r}_2|^2).$$
(2.7)

This can be interpreted as two s-type orbitals and an interelectronic correlation factor  $e^{-\gamma r_{12}^2}$ . While they do not have the correct cusp behaviour, a linear combination of GTGs approximately capture the electron-electron cusp. This is similar to how gaussian basis functions do not individually capture the electron-nuclear cusp, but a linear combination of them approximately does.

One major strength of this approach is that all integrals have closed-form algebraic expressions, <sup>95</sup> and avoids nonlinear optimisation. <sup>96,97</sup> ECG methods have been extended to post-Hartre-Fock methods, such as MP2<sup>98,99</sup>, and methods to avoid its many difficult integrals have also been developed. <sup>100–104</sup>

2.4. F12 Methods 25

While not as popular as F12 methods (see section 2.4), ECGs have been used for highly accurate variational calculations, <sup>105</sup> as well as for applications outside of standard electronic structure theory, such as bosons, <sup>106</sup> positronium (a bound state of an electron and a positron), <sup>107</sup> and non-Born-Oppenheimer systems. <sup>108</sup>

#### 2.4. F12 Methods

The most influential class of explicitly correlated methods to date are the F12 methods. The core principle of these methods is to augment the wave function from a conventional (typically SD) basis with an explicitly-correlated correction, called the F12 (or R12) correction. The original formulation parametrised a two-electron system (such as helium) wave function as

$$|\Psi\rangle = (1 + tQ_{12}F_{12}(r_{12}))|\Psi_0\rangle + \sum_p t_p |\Psi_p\rangle$$
 (2.8)

where  $\Psi_0$  is a reference wave function (such as HF),  $|\Psi_p\rangle$  are excited-state SDs,  $t, t_p$  are parameters ("amplitudes") to be optimised, and

$$Q_{12} := \sum_{\alpha\beta} |\alpha\beta\rangle \langle \alpha\beta| \tag{2.9}$$

is referred to as a "strong orthogonality projector", which ensures the  $F_{12}(r_{12})$  term is orthogonal to the reference and singly-excited determinants. Here  $\alpha, \beta$  refer to virtual orbitals in the formally complete basis. The fact that, due to  $Q_{12}$ , the  $F_{12}(r_{12})$  term commutes with the standard excitation operators aids in including this explicit correlation into conventional electron correlation methods. This geminal term is added a correction to the standard wave function.

In the original formulation,  $F_{12}(r_{12}) = r_{12}$  (hence the name "R12" or "F12" for the more general methodology). However, another popular choice is a Slater-type geminal<sup>110</sup> fitted to a linear combination of gaussians, <sup>111</sup>

$$F_{12}(r_{12}) = -\gamma^{-1} e^{-\gamma r_{12}} = -\gamma^{-1} + r_{12} - \frac{1}{2} \gamma r_{12}^2 + \dots \approx \sum_{i} c_i e^{-\alpha_i r_{12}^2}.$$
 (2.10)

The choice of the length scale  $\gamma$  can either be optimised or (more typically) kept fix, although formally it is orbital dependent and a poor choice may result in a loss of accuracy.<sup>112</sup> Other choices for the correlation factor also exist.<sup>113</sup>

#### 2.4.1. MP2-F12

Following the discussion in reference 66, and adopting their notation, we consider the MP2-F12 method as an illustrative example.

An alternative derivation of the MP2 equations from section 1.6.3 is to minimise the Hylleraas functional  $^{114,115}$ 

$$E^{(2)} = \min_{\Psi} \left\langle \Psi^{(1)} \middle| (H_0 - E^{(0)}) \middle| \Psi^{(1)} \right\rangle + 2 \left\langle \Psi^{(1)} \middle| H \middle| \Psi^{(0)} \right\rangle \tag{2.11}$$

where  $H_0$  is the unperturbed Hamiltonians, and the (i) superscript denotes the order of the correction, as introduced in section 1.6.3.

The earliest generalisation of equation 2.8 was an ansatz for MP2, 116-120

$$\left|\Psi^{(1)}\right\rangle = \frac{1}{2} \sum_{ij} \left( \sum_{ab} t_{ij}^{ab} \left| \Psi_{ij}^{ab} \right\rangle + \sum_{kl} t_{ij}^{kl} \sum_{\alpha\beta} \left\langle \alpha\beta \right| Q_{12} F_{12}(r_{12}) \left| kl \right\rangle \left| \Psi_{ij}^{\alpha\beta} \right\rangle \right), \qquad (2.12)$$

$$Q_{12} = (1 - o_1)(1 - o_2) - v_1 v_2 (2.13)$$

where we define the one-electron projection operators

$$o_m = \sum_{i} |\phi_i(\mathbf{r}_m)\rangle \langle \phi_i(\mathbf{r}_m)| \qquad (2.14)$$

and

$$v_m = \sum_{n} |\phi_a(\mathbf{r}_m)\rangle \langle \phi_a(\mathbf{r}_m)| \qquad (2.15)$$

such that  $o_m$  and  $v_m$  project onto the occupied and virtual orbitals, respectively, with

$$\langle \phi_i(\mathbf{r}_m) | \Omega | jk \rangle = \int d^3 r_m \ \phi_i^*(\mathbf{r}_m) \Omega \phi_j(\mathbf{r}_1) \phi_k(\mathbf{r}_2)$$
 (2.16)

for operator  $\Omega$ . The first term in brackets in equation 2.12 are from the conventional MP2 wave function correction, and the extra term is a contraction of a formally-infinite set of double excitations, representing the F12 correction (again, orthogonal to the conventional wave function thanks to  $Q_{12}$ ).

It was also shown that solving for the amplitudes  $t_{ij}^{kl}$  can be avoided,  $t_{ij}^{21,111,121-123}$  and can instead be derived from the cusp conditions to give

$$t_{ij}^{kl} = \frac{3}{8}\delta_{ik}\delta_{jl} + \frac{1}{8}\delta_{il}\delta_{jk}.$$
 (2.17)

This also tends to be more accurate, as it avoids the geminal basis set superposition error.  $^{122,124}$ 

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#### 2.4.2. Many-Electron Integrals

The Hylleraas functional, equation 2.11, will contain terms such as

$$\sum_{\alpha\beta} \langle \Phi_0 | H | \Psi_{ij}^{\alpha\beta} \rangle \langle \alpha\beta | Q_{12} F_{12}(r_{12}) | kl \rangle = 2 \langle ij | r_{12}^{-1} Q_{12} F_{12}(r_{12}) | kl \rangle - \langle jl | r_{12}^{-1} Q_{12} F_{12}(r_{12}) | kl \rangle.$$
(2.18)

Plugging in the form for  $Q_{12}$  from equation 2.13, we get integrals of the form

$$\langle ij| r_{12}^{-1} o_1 F_{12}(r_{12}) |kl\rangle$$
. (2.19)

For  $F_{12}(r_{12}) = r_{12}$ , this is equal to

$$\sum_{o} \langle ijo| r_{12}^{-1} r_{23} | olk \rangle, \qquad (2.20)$$

which is a three-electron integral. Here, o is an occupied orbital. Clearly, with more operators multiplied together in the kernel of these integrals, we might expect even higher-order integrals. This may seem distrastrous, as this would be a massive bottleneck.

However, it was found that by insertion of the resolution of identity (RI), <sup>116,117,125</sup> these integrals can be reduced to a linear combination of two-electron integrals. In particular, the strong projection operator becomes

$$Q_{12} = 1 - \sum_{o} \sum_{\alpha} (|o\alpha\rangle \langle o\alpha| + |\alpha o\rangle \langle \alpha o|) + \sum_{o,o'} |oo'\rangle \langle oo'| - \sum_{a,b} |ab\rangle \langle ab|, \qquad (2.21)$$

and the discussed three-body integral becomes

$$\sum_{o} \langle ijo| \, r_{12}^{-1} r_{23} \, | olk \rangle = \delta_{ik} \delta_{kl} 
- \sum_{o} \sum_{\alpha} \left( \langle ij| \, r_{12}^{-1} \, | o\alpha \rangle \, \langle o\alpha | \, r_{12} \, | kl \rangle + \langle ij| \, r_{12}^{-1} \, | \alpha o \rangle \, \langle \alpha o | \, r_{12} \, | kl \rangle \right) 
+ \sum_{o,o'} \langle ij| \, r_{12}^{-1} \, \left| oo' \right\rangle \langle oo' | \, r_{12} \, | kl \rangle - \sum_{a,b} \langle ij| \, r_{12}^{-1} \, | ab \rangle \, \langle ab | \, r_{12} \, | kl \rangle .$$
(2.22)

In the RI approximation, the summation over the additional orbitals  $\alpha$  is in principle infinite. In practice, of course, these must be finite. Initially, the basis in the RI expansion was set equal to the orbital basis (dubbed the "standard approximation").  $^{116-118,125-130}$  However, normally larger bases are used, such as in the complementary auxiliary basis set (CABS) approach, where the RI basis is equal to the orbital basis, plus some "auxiliary" basis functions that are orthogonal to the orbital basis.  $^{123,129}$  Many more important advances were since made, such as more efficient RI methods and ways of generating intermediate values have also been developed.  $^{110,111,121,123,131-147}$ 

Moreover, it has been found that the original  $F_{12}(r_{12}) = r_{12}$  form requires larger auxiliary basis functions, since its long-range behaviour is unphysical. <sup>110,121</sup> Instead, functions such as that in equation 2.10 is used.

#### 2.4.3. Higher-Order F12 Theories

F12 theory has been extended to many other theories besides MP2, such as CC, CASSCF, multireference configuration interaction (MRCI), complete active space perturbation theory to second order (CASPT2), density matrix renormalization group (DMRG), and FCIQMC.<sup>66,138,148–163</sup> The key change compared to conventional methods such as those in chapter 1 is in the excitation operators, particularly the double excitations  $E_{ij}^{ab}$ . In particular, for a multireference method with reference determinants  $|\Phi_I\rangle$ , the wave function may be parametrised as

$$|\Psi\rangle = |\Psi_{\text{conv}}\rangle + \frac{1}{2} \sum_{I} \sum_{ijkl} t_{klI}^{ij} \sum_{\alpha\beta} \langle \alpha\beta | Q_{12}^{I} F_{12}(r_{12}) | kl \rangle E_{ij}^{\alpha\beta} |\Phi_{I}\rangle, \qquad (2.23)$$

where  $|\Psi_{conv}\rangle$  is the conventional wave function. In this way, we extend the excitations to include geminal terms in the (formally complete) auxiliary basis.

These F12 methodologies have been particularly successful for large systems, proving only marginally more expensive than the conventional methods in some cases. 124,164

Since F12 methods work directly on the wave function, it is also natural to extend to excited states, particularly for multireference methods. <sup>152,161,162,165,166</sup> However, since the geminal terms include only occupied orbitals, there is an inherent bias to the ground state. <sup>152,153</sup> Several means of ameliorating this have been proposed, such as combining F12 with response theory, <sup>153</sup> or by including extending the geminal basis to include virtual orbitals. <sup>154</sup>

## 2.5. The Transcorrelated Method

One of the main focuses of this dissertation is TC. Hirschfelder was the first to propose a similarity-transformed Hamiltonian method in the 1960s, <sup>167</sup> which was further developed by Jankowski, <sup>168,169</sup> and later by Boys and Handy. <sup>170–173</sup>

The core concept is to similarity-transform the Hamiltonian using some invertible operator P, such that

$$\hat{H}_{TC} = P^{-1}\hat{H}P. \tag{2.24}$$

For any eigenvalue and eigenfunction pair  $(E_i, \Psi)$  of  $\hat{H}$ , the transformed Hamiltonian  $\hat{H}_{TC}$  has eigenfunction  $\Phi := P\Psi$  with the same eigenvalue  $E_i$ . This is called isospectrality, and it allows us to solve  $\hat{H}_{TC}$  to get the spectrum of  $\hat{H}$ .

However, note that there is no requirement that  $\hat{H}_{TC}$  is Hermitian, and in fact it typically is not. Recall from section 1.3 that the variational principle relies on the

Hermiticity of the operator. It is therefore nontrivial to make TC variational, and not all conventional methods can be used to treat  $\hat{H}_{TC}$  (although, as discussed in section 3.4, projector methods such as FCIQMC and CC may be used).

#### 2.5.1. The Method of Boys and Handy

One of the earliest practical approaches, on which most modern variations build, is that of Boys and Handy. <sup>170–173</sup> In this methodology, we start with a Jastrow ansatz <sup>174</sup>

$$\Psi = e^J \Phi. \tag{2.25}$$

In this way, TC shares similarities with variational Monte Carlo (VMC), to be discussed in section 3.2. Inserting 2.25 into the Schrödinger equation, we get

$$\hat{H}e^{J}\Phi = Ee^{J}\Phi \implies \underbrace{e^{-J}\hat{H}e^{J}}_{\hat{H}_{TC}}\Phi = E\Phi,$$
 (2.26)

i.e. similarity-transforming  $\hat{H}$  with the operator  $P \equiv e^{J}$ .

In the original formulation,  $\Phi$  was chosen to be a single SD (HF), and J was chosen such that

$$J = \sum_{i < i} u(\mathbf{r}_i, \mathbf{r}_j) \tag{2.27}$$

where  $u(\mathbf{r}_i, \mathbf{r}_j) = u(\mathbf{r}_j, \mathbf{r}_i)$  is a symmetric two-electron correlation function.

The Baker-Campbell-Hausdorff (BCH) expansion of  $\hat{H}_{TC}$  truncates exactly to second order, i.e.

$$\hat{H}_{TC} := e^{-J} \hat{H} e^{J} = \hat{H} + [\hat{H}, J] + \frac{1}{2} [[\hat{H}, J], J].$$
 (2.28)

This may be rewritten

$$\hat{H}_{TC} = \hat{H} - \sum_{i < j} \hat{K}(\boldsymbol{r}_i, \boldsymbol{r}_j) - \sum_{i < j < k} \hat{L}(\boldsymbol{r}_i, \boldsymbol{r}_j, \boldsymbol{r}_k), \qquad (2.29)$$

where, using the shorthand  $u_{ij} := u(\mathbf{r}_i, \mathbf{r}_j)$ ,

$$\hat{K}(\boldsymbol{r}_{i},\boldsymbol{r}_{j}) = \frac{1}{2} \left( \nabla_{i}^{2} u_{ij} + \nabla_{j}^{2} u_{ij} + (\nabla_{i} u_{ij})^{2} + (\nabla_{j} u_{ij})^{2} \right) + (\nabla_{i} u_{ij} \cdot \nabla_{i}) + (\nabla_{j} u_{ij} \cdot \nabla_{j}),$$
(2.30)

and

$$\hat{L}(\mathbf{r}_i, \mathbf{r}_i, \mathbf{r}_k) = \nabla_i u_{ii} \cdot \nabla_i u_{ik} + \nabla_i u_{ii} \cdot \nabla_i u_{ik} + \nabla_k u_{ki} \cdot \nabla_k u_{ki}. \tag{2.31}$$

In second quantisation, the TC Hamiltonian can then be written

$$\hat{H}_{TC} = \sum_{pq} h_q^p a_p^{\dagger} a_q + \frac{1}{2} \sum_{pqrs} \left( V_{rs}^{pq} - K_{rs}^{pq} \right) a_p^{\dagger} a_q^{\dagger} a_s a_r - \frac{1}{6} \sum_{pqrstu} L_{stu}^{pqr} a_p^{\dagger} a_q^{\dagger} a_r^{\dagger} a_u a_t a_s, \quad (2.32)$$

where

$$\begin{split} h_q^p &= \langle p|\,h\,|q\rangle\,,\\ V_{rs}^{pq} &= \langle pq|\,r_{12}^{-1}\,|rs\rangle\,,\\ K_{rs}^{pq} &= \langle pq|\,\hat{K}\,|rs\rangle\,,\\ L_{stu}^{pqr} &= \langle pqr|\,\hat{L}\,|stu\rangle\,. \end{split} \tag{2.33}$$

Particularly noteworthy are the three-electron integrals from  $\hat{L}$  and the non-Hermitian terms from  $\hat{K}$ .

Boys and Handy's original choice for the form of u was

$$u(\mathbf{r}_{i}, \mathbf{r}_{j}) = \sum_{l,m,n}^{l+m+n \leq S} c_{lmn} \bar{r}_{ij}^{l} (\bar{r}_{Ii}^{m} \bar{r}_{Jj}^{n} + \bar{r}_{Ij}^{m} \bar{r}_{Ji}^{n})$$
(2.34)

where S is some integer  $(S = 6 \text{ in reference } 170)^{175}$  and

$$\bar{r} = \frac{r}{1+r}. (2.35)$$

In this form, electron-electron (e-e), electron-nucleus (e-n), electron-electron-nucleus (e-e-n) and electron-electron-nucleus-nucleus (e-e-n-n) terms appear when m = n = 0 and l > 0, m, n > 0 and l = 0, one of m, n > 0 and l > 0, or m, n, l > 0 respectively.

In the original formulation,  $^{170}$  the following equations are approximately solved via quadrature for the orbital coefficients  $a_i$  and the coefficients  $c_{lmn}$  in u.

$$\langle \Phi | (\hat{H}_{TC} - E) | \Phi \rangle = 0 \tag{2.36}$$

$$\langle \partial \Phi / \partial a_i | (\hat{H}_{TC} - E) | \Phi \rangle = 0$$
 (2.37)

$$\langle \Phi | (\hat{H}_{TC} - \hat{H}_{TC}^{\dagger}) | \Phi \rangle = 0,$$
 (2.38)

and the energy is calculated via projection,

$$\langle \Phi | \hat{H}_{\text{TC}} | \Phi \rangle$$
. (2.39)

Equation 2.38 in particular is to make the TC Hamiltonian as close to Hermitian as possible for orbital optimisation.

This formulation was further refined by Handy.<sup>176</sup> We notice that since  $\hat{H}_{TC}$  is non-Hermitian, we cannot directly minimise the energy. Instead, he minimised the transcorre-

lated variance,

$$\sigma^2 = \langle [(\hat{H}_{TC} - E)\Phi]^2 \rangle. \tag{2.40}$$

Furthermore, equation 2.34 is not the only possible choice for the Jastrow factor, and in particular Handy chose a form with simpler analytical integrals.

#### 2.5.2. Modern Resurgence

For several decades TC received little attention, though some more developments were made, such as calculations using a fixed Jastrow factor,  $^{177}$  more efficient integration,  $^{178,179}$  multi-reference  $\Phi$ ,  $^{180}$  among others.  $^{181-183}$  More recently, there has been a renewal of interest in TC,  $^{175,181-223}$  notably the demonstrations that the TC Hamiltonian may be treated by various electronic structure methods, to be discussed more in section  $3.4.7.^{175,185-188,196,203-205,213,214,217}$  TC has since been applied to a variety of systems, and in the context of quantum computing.  $^{193,201,207,208,215}$  After an overview of stochastic methods, the rest of this dissertation will be focused on discussing recent developments in TC.

In typical modern TC workflows,  $\Phi$  is initially kept fixed (e.g. to the HF determinant, though it need not be this), and then the parameters of the Jastrow factor, J, are optimised. The integrals for the similarity-transformed Hamiltonian  $\hat{H}_{TC}$  are then calculated, and finally an electronic structure method such as FCIQMC is applied to  $\hat{H}_{TC}$ , solving for E and in effect "recalculating"  $\Phi$ , which was previously kept fixed. The choice of J has ranged from especially simple 188,194,216 to much more sophisticated. 175,224,225

TC is still behind F12 in terms of its performance to cost ratio, but F12 is ahead several decades of development. However, TC shows promise thanks in part to the high degree of flexibility owing to the Jastrow factor. Moreover, the additive ansatz of F12 leads to many-electron integrals, whereas the multiplicative ansatz of TC leads to at most three-electron integrals.\* This is ameliorated in F12 via the use of RI approximations and in TC via other approximations, like discussed in section 4.9.1.

<sup>\*</sup>This assumes the Jastrow factor may be written  $J = \sum_i u_i(\mathbf{r}_i, \mathbf{r}_j)$ .

# Monte Carlo Methods

Monte Carlo (MC) methods are a class of numerical methods that use random sampling to numerically solve problems. It has found applications in an impressive range of fields, from physics to finance.<sup>226–231</sup> It is particularly useful for problems with high dimensionality, where deterministic methods are often impractical. In quantum chemistry and physics, since a 'dimension' can refer to any degree of freedom, high-dimensional problems are commonplace, and so MC methods are a natural choice.

While the name *Monte Carlo* was coined by Stanislav Ulam, after the famous casino in Monaco,  $^{232,233}$  the foundational concept was already developed in the 18th century by the French mathematician Georges-Louis Leclerc, Comte de Buffon. As one of the earliest example applications, in the Buffon needle problem, one can randomly toss needles onto a lined sheet of paper and determine  $\pi$ .  $^{234-236}$ 

Monte Carlo methods is a broad term, and as such it is not possible to give a comprehensive overview in a single chapter, and there exist many reviews and textbooks on MC and related topics.<sup>237–242</sup> Here, we will focus on only a few concepts particularly relevant for this dissertation, largely following reference 230 and the relevant chapters of reference 231.

#### 3.1. Classical Monte Carlo Methods

We start our discussion with classical MC methods. While there are numerous possible applications, notably in molecular dynamics,  $^{237}$  here we restrict ourselves to the topic of Monte Carlo integration. In particular, we consider the classical textbook problem of calculating the value of  $\pi$ , then we provide a more rigorous framework.

#### 3.1.1. A Very Bad Game of Darts

If we imagine throwing darts at a dartboard randomly, we can approximate  $\pi$ . If the radius of the circle is r, then its area is  $\pi r^2$ . The area of the square circumscribing the circle is  $4r^2$ . Therefore, the ratio of the area of the circle to the area of the square is  $\pi/4$ . If we randomly sample a point in the square ("throw a dart"), the probability that the point is inside the circle is proportional to its area. Since we sample inside the square,

the probability of landing inside the circle is

$$P(\text{inside circle}) = \frac{\pi r^2}{4r^2} = \frac{\pi}{4}.$$
 (3.1)

Therefore, if we sample a large number of points, the ratio of the number of darts that land inside the unit circle to the total number of darts, we can approximate the probability distribution P and thus get an estimate for  $\pi$  as

$$\pi \approx 4 \frac{N_{\rm in}}{N_{\rm out}}.$$
 (3.2)

This is illustrated in figure 3.1, and captures the core philosophy of MC methods.

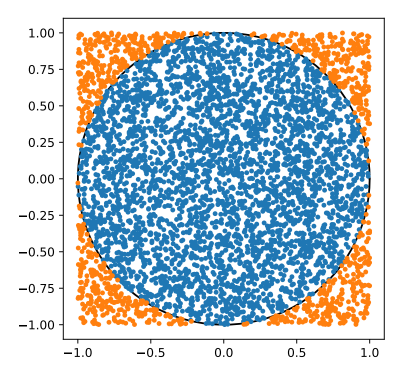


Figure 3.1. Our "game of darts". Points inside the unit circle are coloured blue whereas points outside are orange. Using stochastic sampling, this naive approach uses 5000 randomly generated coordinates  $(x,y) \in [-1,1] \times [-1,1]$  to approximate  $\pi \approx 4N_{\rm in}/N_{\rm out} \approx 3.1464$ .

Controlling the stochastic error is critical in MC methods. To estimate the reliability of our estimate, we can determine the standard deviation of the estimate. Our sampling scheme is a binomial distribution, with  $p = \pi/4$ . Hence,

$$\sigma_{\pi}^2 = \text{Var}(4P(\text{inside circle})) = \frac{16p(1-p)}{N}$$
 (3.3)

or

$$\sigma_{\pi} = 4\sqrt{\frac{\frac{\pi}{4}(1 - \frac{\pi}{4})}{N}} \propto \frac{1}{\sqrt{N}}.$$
(3.4)

#### 3.1.2. A More Mathematical Description

As we have expressed the problem of the previous section in terms of areas, it is clear that it can also be formulated in terms of integrals. For this particular problem, we have:

$$\pi = \int_{-1}^{1} dx \int_{-1}^{1} dy \ \Theta(1 - x^2 - y^2), \tag{3.5}$$

where  $\Theta$  is the Heaviside step function. More generally, consider the integral of some smooth function f over  $[a, b] \subseteq \mathbb{R},^*$ 

$$I = \int_{a}^{b} \mathrm{d}x \ f(x). \tag{3.6}$$

Standard finite element methods for solving integrals of this type typically involving dividing the integration domain into N subintervals of length h and determining the weights  $w_i$  from e.g. a polynomial approximation. That is,

$$I \approx \sum_{i=1}^{N} w_i f(x_i). \tag{3.7}$$

The error  $\sigma$  in these sorts of methods is typically  $\sigma \propto h^{-k} \propto N^{-k}$ , where  $k \in \mathbb{Z}_{>0}$ . For a multi-dimensional integral,  $\sigma \propto N^{-k/d}$ , where d is the number of dimensions.<sup>13</sup>

In MC methods,  $\forall i$  take  $w_i = 1$  and  $x_i \in [a, b]$  randomly sampled. That is,

$$I \approx \sum_{i=1}^{N} f(x_i). \tag{3.8}$$

If, for example, we choose N to be randomly sampled, then the variance is

$$\sigma^2 = \left\langle \left( \frac{b-a}{N} \sum_{i=1}^N f_i \right)^2 \right\rangle - \left( \left\langle \frac{b-a}{N} \sum_{i=1}^N f_i \right\rangle \right)^2 \tag{3.9}$$

$$=\frac{(b-a)^2}{N}(\bar{f}^2-\bar{f}^2) \tag{3.10}$$

where  $f_i := f(x_i)$  and  $x_i$  is a random number drawn, the angular brackets denote an average over all possible realisations, and the overbar represents an average of the function over the domain ([a, b] in this discussion). i.e. the error in this method is proportional to the variance of f. Perhaps more interestingly,  $\sigma \propto N^{-1/2}$ , in line with the central limit

<sup>\*</sup>We present the one-dimensional case for simplicity. The generalization to higher dimensions is straightforward.

theorem.<sup>243</sup> Comparing this error with standard quadrature, we see that MC integration is more efficient than an order-k algorithm when d > 2k. That is, although this particular MC example is naive, using simply a uniform distribution, it is still more efficient than standard methodologies for very large dimensions.

There exist several methods to reduce errors in MC methods.<sup>244</sup> Among the most important ones is importance sampling.<sup>245</sup> In the prevous example, it is clear that if significant contributions to the integral come from a small region of the domain, only a few points would be sampled by the MC algorithm there when using a uniform distribution. This would lead to large statistical errors. Mathematically, this is from the large variance of the function. In importance sampling, we sample from a distribution p which has roughly the same shape as f such that f/p is roughly constant over the integration domain. Of course, being a probability distribution, we require

$$p(x) > 0 \quad \forall x \tag{3.11}$$

and

$$\int \mathrm{d}x \ p(x) = 1. \tag{3.12}$$

The integral is

$$I = \int \mathrm{d}x \ f(x) = \int \mathrm{d}x \ \frac{f(x)}{p(x)} p(x). \tag{3.13}$$

Then, if we sample points according to p, we have

$$I \approx \frac{1}{N} \sum_{i=1}^{N} \frac{f(x_i)}{p(x_i)},\tag{3.14}$$

where the naive MC method is recovered when p(x) = 1/(b-a), i.e. the uniform distribution.

In this case, the variance in the result is

$$\sigma^2 = \left\langle \left( \frac{1}{N} \sum_{i=1}^N \frac{f(x_i)}{p(x_i)} \right)^2 \right\rangle - \left( \left\langle \frac{1}{N} \sum_{i=1}^N \frac{f(x_i)}{p(x_i)} \right\rangle \right)^2. \tag{3.15}$$

From this, we can see that if f/p is constant, the error vanishes, so it is important to choose a good p. If p is chosen poorly, the variance can be worsened, so this is a delicate problem.

In practice it is often difficult to estimate f, and a choice for p is problem-specific. However, when we have some approximation for its overall shape, importance sampling can be a powerful tool. Alternatively, another method known as adaptive Monte Carlo<sup>246</sup> seeks the most significant regions of f by random sampling, so that no a priori knowledge of the functional form is required.

#### 3.1.3. Metropolis-Hastings Algorithm

A particularly successful method for generating random samples  $x_i$  from a probability distribution  $\pi(x)$ , where direct sampling of  $\pi$  is difficult, is the Metropolis-Hastings algorithm.<sup>233,247</sup> Often,  $\pi(x)$  is known only up to a normalisation constant,  $\pi(x) = \pi(x)/C$  where  $C = \int dx \, \pi(x)$  is intractible.

The Metropolis-Hastings algorithm is based on Markov chains,<sup>248</sup> and we construct such a Markov chain so that its stationary distribution is  $\pi(x)$ , which is to say if the Markov chain starts at  $\pi(x)$  for step t, it is still  $\pi(x)$  for step t+1.

The probability of having a sequence of states  $x_1, x_2, \ldots, x_N$  is

$$p(x_1, x_2, \dots, x_N) = p(x_1)p(x_2|x_1)p(x_3|x_2) \cdots p(x_N|x_{N-1}).$$
(3.16)

where p(x|x') is the transition probability from  $x' \to x$ . Then, the probability at step t to be in state x,  $\pi_t(x)$  is given by

$$\pi_{t+1}(x) = \sum_{x' \in \Omega} \pi_t(x') p(x|x')$$
(3.17)

where  $\Omega$  is the set of all possible states. This is called the master equation. In the stationary state,  $\pi_t(x) = \pi(x)$ , so

$$\pi(x) = \sum_{x' \in \Omega} \pi(x') p(x|x'). \tag{3.18}$$

Finding the general solution to this problem is nontrivial. However, a sufficient (but not necessary) condition is called detailed balance:

$$\pi(x)p(x|x') = \pi(x')p(x'|x). \tag{3.19}$$

This ensures that the probability of going from x to x' is the same as the probability of going from x' to x, which implies that the probability is stationary.

In order to actually construct the algorithm, we must introduce the trial step probability  $\omega(x|x')$ , and the acceptance probability A(x|x'). Then,

$$p(x|x') = \omega(x|x')A(x|x'). \tag{3.20}$$

 $\omega(x|x'), A(x|x') \in [0,1]$  for each pair x, x', and  $\sum_{x'} \omega(x|x') = 1$ . Furthermore, the original formulation of the algorithm required  $\omega(x|x') = \omega(x'|x)$ , which leads to (plugging into the master equation)

$$\frac{A(x|x')}{A(x'|x)} = \frac{\pi(x')}{\pi(x)}. (3.21)$$

The algorithm proceeds in two stages: we propose a step  $x' \to x$ , and then accept or reject it. The probability of accepting is

$$A(x|x') = \min\left(1, \frac{\pi(x')}{\pi(x)}\right). \tag{3.22}$$

If we accept, we set the new state to x, otherwise we stay at x'.

In practice, on a computer, accepting is done by generating a uniform random number  $r \in [0,1)$  and accepting if  $r < \frac{\pi(x')}{\pi(x)}$ , and otherwise rejecting. Furthermore, we don't just have a single Markov chain, but a collection of so-called "walkers", that each perform their own Markov chain (for parallelisation). The integrand is then sampled at each position where the walkers reach.

One final note, is that since the current state of a Markov chain is dependent on the previous state, the Markov chain is not independent of itself. This is referred to as autocorrelation. One method to reduce this autocorrelation and give essentially independent samples is known as blocking.<sup>249</sup>

## 3.2. Variational (Quantum) Monte Carlo

As our first foray into QMC, we consider VMC. In section 1.3 we introduced the Variational Principle by parametrising the wave function and then finding the minimum of the expectation value of the energy occurring in the parametrisation space. If we have a large number of electrons and/or a large number of parameters, then the integrals involved in the evaluation of the energy will necessarily be high-dimensional. This is where MC comes in. For basic trial wave functions and small atoms like Hydrogen or Helium, direct integration may be possible, but as discussed in the previous section, this quickly becomes impossible for larger systems.

For a trial wave function  $\Psi$  (we omit the tilde from section 1.3), the expectation value of the energy is

$$\langle E \rangle = \frac{\langle \Psi | H | \Psi \rangle}{\langle \Psi | \Psi \rangle} \tag{3.23}$$

$$= \frac{\int \mathrm{d}^{3N} r \ \Psi^* H \Psi}{\int \mathrm{d}^{3N} r \ |\Psi|^2} \tag{3.24}$$

$$= \frac{\int \mathrm{d}^{3N} r \ \Psi^* \Psi \frac{H\Psi}{\Psi}}{\int \mathrm{d}^{3N} r \ |\Psi|^2} \tag{3.25}$$

$$= \frac{\int \mathrm{d}^{3N} r |\Psi|^2 E_L(\mathbf{r})}{\int \mathrm{d}^{3N} r |\Psi|^2},\tag{3.26}$$

where we have introduced the local energy, defined as

$$E_L(\mathbf{r}) := \frac{H\Psi}{\Psi}.\tag{3.27}$$

This rewriting of the integral is particularly suitable for evaluation using MC. Then we may vary the parameters and stop according to some minimisation algorithm. Notice that the more strongly  $\Psi$  resembles the exact wave function, the less strongly  $E_L$  varies with r. In particular, if  $\Psi$  is equal to an exact eigenstate, then  $E_L$  is constant. Therefore, an alternative objective function to the energy expectation of equation 3.23 is the variance.  $^{250-252}$ 

Defining

$$p(\mathbf{r}) = \frac{|\Psi|^2}{\int \mathrm{d}^{3N} r \ |\Psi|^2},\tag{3.28}$$

we may write equation 3.23 as

$$\langle E \rangle = \int d^{3N} r \ p(\mathbf{r}) E_L(\mathbf{r}).$$
 (3.29)

This form of the energy is amenable to the Metropolis-Hastings approach outlined in section 3.1.3. Recall also that p need not be normalised when using the Metropolis-Hastings algorithm for sampling.

In principle, the algorithm is doable with just a single walker, but in practice we reduce the statistical error by using many. The algorithm then becomes

```
Initialise N walkers in random positions. until convergence criterion met* for each walker at position r sample the local energy E_L at r propose a new position r' with probably p=|\Psi(r')/\Psi(r)|^2 if r' is accepted set r to r'
```

The energy is then calculated as the expectation value of the local energy, averaged over the samples generated in this procedure. Steps at the beginning (before equilibrium) are discarded in a process called equilibration, <sup>253</sup> and a blocking procedure should be employed. The decision to stop is generally based on compute time and/or precision required.

#### 3.2.1. Trial Wave Functions

While VMC is a powerful tool, the quality of the solution is constrained by the quality of the trial wave function. Moreover, the evaluation of the trial wave function is expensive, and we therefore want a form that is easy to evaluate.

<sup>\*</sup>The convergence criterion in VMC is typically just to sample a predefind number of configurations.

The form of the trial wave function is typically chosen based on a Jastrow factor, as already introduced in section 2.5. Generally, <sup>174</sup>

$$\Psi_{\text{trial}} = e^J \Phi. \tag{3.30}$$

For computational efficiency, the Jastrow factor typically only retains one- and two-body terms,  $^{230}$ 

$$J = \sum_{i} \chi(\boldsymbol{x}_i) - \frac{1}{2} \sum_{i \neq j} u(\boldsymbol{x}_i, \boldsymbol{x}_j), \tag{3.31}$$

where  $\chi$  describes electron-nuclear correlation and u describes two-electron correlation (including e.g. electron-electron-nuclear correlation). There exist a number of more specific forms for J,  $^{225}$  though one unifying principle is for them to adhere to expected short-range (cusp conditions) and long-range (1/r) behaviour.

 $\Phi$  is typically chosen to be a single Slater determinant. <sup>230,254–257</sup> In particular, by choosing the HF determinant, typically only a small portion of the energy (that is, the correlation energy) is left for the Jastrow factor to describe, and typically even simple Jastrow factors can do this. <sup>230</sup> However, sometimes this is not enough, for example, when the HF determinant is not sufficient to describe all symmetries of the true wave function. <sup>258–260</sup>

#### 3.3. Diffusion Monte Carlo

Diffusion Monte Carlo (DMC) starts by performing a Wick rotation (that is, substitute  $t \to i\tau$ ) on the time-dependent Schrödinger equation, equation 1.1. We then get the imaginary-time Schrödinger equation

$$\frac{\partial}{\partial \tau} \Psi(\mathbf{r}, \tau) = -\hat{H} \Psi(\mathbf{r}, \tau). \tag{3.32}$$

This equation is a diffusion (or heat) equation, and  $\Psi$  may be interpreted as the density distribution for a large number of independent particles\* (walkers).<sup>261</sup> More specifically, the kinetic energy term, being second order, describes the movement of the walkers whereas the potential energy term describes generation or annihilation of walkers. The solution of this equation is

$$\Psi(\mathbf{r},\tau) = e^{-(\hat{H}-S)\tau}\Psi(\mathbf{r},0), \tag{3.33}$$

where we have introduced the energy shift S which is adapted throughout the simulation every n steps to keep the number of walkers constant, and to estimate the ground state energy.<sup>262</sup> In particular, if we expand  $\Psi(\mathbf{r},0)$  in terms of the eigenfunctions of  $\hat{H}$ ,  $\{\psi_i\}$ ,

<sup>\*</sup>Note that the particles, henceforth referred to as walkers, are not the particles being described by the Hamiltonian. One walker represents a configuration of all the particles described by the Hamiltonian.

labeled in such a way that  $i > j \implies E_i > E_j$ , we get

$$\Psi(\mathbf{r},\tau) = \sum_{i} a_{i} e^{-(\hat{E}_{i} - S)\tau} \psi(\mathbf{r}). \tag{3.34}$$

Here,  $a_i$  is the coefficient of the *i*th eigenfunction, and  $\hat{E}_i$  is the corresponding eigenvalue. After the excited-state solutions have decayed, the only possible value of S that stabilises the simulation is  $S = E_0$ . It is also clear from equation 3.34 that when  $S = E_0$ , all excited states decay exponentially to zero, and we are left with the ground state for large  $\tau$ .

We may also express  $\Psi$  in terms of its Green's function, in order to get a form that expresses a small time step,

$$\Psi(\mathbf{r}, \tau + \Delta \tau) = \int d^{3N} r' G(\mathbf{r}, \mathbf{r}'; \Delta \tau) \Psi(\mathbf{r}, \tau), \qquad (3.35)$$

where the Green's function

$$G(\mathbf{r}, \mathbf{r}'; \Delta \tau) = \langle r | e^{-\Delta \tau (H-S)} | r' \rangle$$
(3.36)

also obeys the imaginary-time Schrödinger equation, but with the initial condition

$$G(\mathbf{r}, \mathbf{r}'; 0) = \delta(\mathbf{r} - \mathbf{r}'). \tag{3.37}$$

For small time steps, G may be approximated by the Lie-Trotter-Suzuki decomposition, <sup>263–265</sup> i.e. for operators A,  $A_k$ ,

$$A = \sum_{k} A_{k} \implies e^{-\alpha A} \approx \left(\prod_{k} e^{-\alpha A_{k}/M}\right)^{M}$$
 (3.38)

for some value  $\alpha$  and for large M. From this, we can get the Green's function,

$$G(\mathbf{r}, \mathbf{r}'; \Delta \tau) = e^{-\Delta \tau (V(r) - S)} \frac{1}{\sqrt{2\pi \Delta \tau}} e^{-(r - r')^2/(2\pi \Delta \tau)} + \mathcal{O}(\Delta \tau^2), \tag{3.39}$$

which describes our time evolution.

In DMC, walkers diffuse through configuration space in two stages: diffusion and branching. In the diffusion step, walkers are moved according to the kinetic terms in the Green's function. In the branching step, unfavourable walkers are removed and favourable ones produce new walkers.

For the energy shift estimate, we wish to choose a value such that the overall population does not change too much. If the walkers proliferate too much, then the computational cost becomes infeasible, whereas if too many walkers die, we are left with poor statistics. One way to choose S is by keeping track of the population of walkers  $N_w$  for a target

populaton  $N_w^{\text{target}}$  and adjust,

$$S = S_{\text{old}} + \alpha \ln \left( \frac{N_w^{\text{target}}}{N_w} \right) \tag{3.40}$$

for small  $\alpha$ .

More concretely, in pseudocode, the basic DMC algorithm is:

```
Initialise N walkers in random positions. until convergence criterion met for each walker at position r shift position r to r' according to \hookrightarrow transition probability G evaluate q = \mathrm{e}^{-\Delta \tau(V(r') - S)} walker survives with probability q if q > 1 create 1 - q new walkers \hookrightarrow (with stochastic rounding) update S.
```

Notice that in contrast to VMC, DMC does not rely on a trial wave function. There is also a discretisation error incurred by the operator decomposition, though this can be handled via an application of the Metropolis-Hastings alogirthm.<sup>231</sup> However, in practice a trial wave function is actually necessary for fermions in order to improve the statistics. The walker distribution is positive, but for fermions this is not the case due to their exchange statistics. Since in electronic structure, we focus on fermions (electrons), this is an important feature. In order to still practically use DMC,\* an approximation called the fixed node approximation is made, where the nodal structure (i.e. the roots of  $\Psi$ ) of the distribution is determined by a trial function. Thus, the accuracy of DMC is still limited by the quality of the trial wave function (or more precisely, by its nodal surface). In a Slater-Jastrow-type trial wave function like equation 3.30, only the Slater-component will affect the nodal surface (and hence the accuracy). Thus, the choice of orbitals as well as the number of determinants can be important considerations.

# 3.4. Full Configuration Interaction Quantum Monte Carlo

The main post-Hartree-Fock method used throughout this dissertation is the FCIQMC algorithm.<sup>266</sup> For this, there exist a few different implementations, <sup>267–270</sup> but we focus on NECI.<sup>267</sup> As the name implies, FCIQMC combines concepts from FCI (discussed in section 1.6.1) and QMC concepts already discussed in this chapter. In particular, it implements FCI, retaining many of its desirable properties such as size consistency and accuracy while

<sup>\*</sup>Other simple solutions such as assigning a sign to each walker leads to large errors.

being able to handle much larger systems thanks to its stochastic framework (however, the scaling remains exponential, like FCI). Like FCI, it is non-perturbative, and so can handle multi-reference problems, and unlike other QMC methods, it does not rely on a trial wave function. It can also be used as a solver in place of FCI, like in a MCSCF calculation.<sup>271,272</sup>

#### 3.4.1. Main Concepts

Like DMC, FCIQMC is based on the imaginary-time Schrödinger equation, equation 3.32, repeated here:

$$\frac{\partial}{\partial \tau} \Psi(\mathbf{r}, \tau) = -\hat{H} \Psi(\mathbf{r}, \tau). \tag{3.41}$$

The solution to this equation is

$$\Psi(\mathbf{r},\tau) = e^{-\tau \hat{H}} \Psi(\mathbf{r},0). \tag{3.42}$$

Expanding  $e^{-\tau \hat{H}}$  to first order in small imaginary time step  $\Delta \tau$  gives us the FCIQMC projector

$$P = 1 - \Delta \tau \hat{H} \to 1 - \Delta \tau (\hat{H} - S), \tag{3.43}$$

where we have introduced an arbitrary shift S in analogy to DMC. Notice, in particular, that if  $S = E_0$  the ground state energy, all other states decay away. Hence, repeated application of P while adjusting S to get as close to  $E_0$  as possible will give us the ground state, as long as our initial state has nonzero overlap with the ground state.

Like FCI, the wave function is expanded in some basis. Throughout this dissertation, SDs are used, though configuration state functions (CSFs) are also possible.<sup>273</sup> The basis will be kept constant, and the coefficients may change with each time step,

$$|\Psi(\tau)\rangle = \sum_{i} c_i(\tau) |D_i\rangle.$$
 (3.44)

Like in DMC, we adjust S to approximate the ground state energy  $E_0$ , as  $S = E_0$  would project out all states other than the ground state, as long as  $\langle \Psi(\tau=0)|\Psi_0\rangle \neq 0$  where  $|\Psi_0\rangle$  is the ground state. Similarly, S affects the walker population. In particular, for walker population  $N_w$ , if  $S > E_0$  then the population grows, and if  $S < E_0$  it decays. If we have a target population  $N_w^{\text{target}}$  then we keep  $S > E_0$  (typically, just by setting S = 0) and "turn on" the shift once  $N_w^{\text{target}}$  is reached. However, unlike DMC, FCIQMC walkers can have either negative or positive weight. These weights correspond to the  $c_i$ , and the walkers "diffuse" across SD (or CSF) space. The walker population is reported in terms of absolute values, i.e.

$$N_w = \sum_{i \in \{SDs\}} |N_i|, \tag{3.45}$$

where we sum over the walkers across all SDs. Once in variable-shift mode, in practice S is updated every m cycles:

$$S(\tau + m\Delta\tau) = S(\tau) - \frac{\zeta}{m\Delta\tau} \ln\left(\frac{N_w(\tau + m\Delta\tau)}{N_w(\tau)}\right),\tag{3.46}$$

where  $\zeta$  is a damping parameter.

The key advantage of FCIQMC over ordinary FCI is no longer needing to store the full CI vector, making use of sparsity. Even for relatively large and strongly-correlated systems, only a small subspace of the entire Hilbert space needs to be sampled in FCIQMC, whereas in FCI, the full CI vector must be stored (or if done via dense diagonalisation, the entire H matrix, which is unrealistic for even small systems).

#### 3.4.2. Basic Algorithm

FCIQMC is typically run in three steps: (a) **spawning**, (b) **death/cloning**, and (c) **annihlation**. Each step can be recognised in the application of the projector operator of equation 3.43 onto some state  $|\Psi\rangle = \sum_i c_i |D_i\rangle$ ,

$$P|\Psi\rangle = |\Psi\rangle - \sum_{i} \sum_{j \neq i} \Delta \tau H_{ji} c_i |D_j\rangle - \sum_{i} (\Delta \tau (H_{ii} - S)) c_i |D_i\rangle, \tag{3.47}$$

where the right-hand side of the equation is the new wave function,  $|\Psi(\tau + \Delta \tau)\rangle$ . Or, in terms of the coefficients (relabelling  $i \leftrightarrow j$  in the (a) terms),

$$c_i(\tau + \Delta \tau) = c_i(\tau) - \Delta \tau (H_{ii} - S)c_i(\tau) - \Delta \tau \sum_{j \neq i} H_{ij}c_j(\tau).$$
 (3.48)

In practice, we start the algorithm with a population of walkers on a reference determinant (typically, but not always, the Hartree-Fock determinant), and calculate the correlation energy via FCIQMC. Instead of H in the working equations we would use the matrix

$$K_{ij} = H_{ij} - E_{\text{ref}} \delta_{ij} \tag{3.49}$$

where  $E_{\text{ref}} := \langle D_{\text{ref}} | H | D_{\text{ref}} \rangle$  is the energy of the reference determinant. However, FCIQMC is capable of adapting to a new reference determinant whenever another determinant exceeds the population of the reference determinant, and is not single-reference like canonical coupled cluster, for example. This shift is primarily a convenience.

The basic FCIQMC algorithm can be summarised by the following pseudocode (with further details below). Note that there are many extensions to the basic algorithm, beyond what is discussed in this dissertation.

```
Initialise N_{\mathrm{start}} walkers on the reference determinant. until stop criterion met for each occupied determinant D_i select a random determinant based on p_{\mathrm{gen}} spawn p_s(j|i) walkers on D_j for each occupied determinant D_i remove p_d (death/cloning) walkers from D_i for each occupied determinant D_i annihilate opposite signed walkers calculate the projected energy if \sum |N_i| \geq N_w update S.
```

#### **Spawning**

The first step, spawning, is realised by the application of the off-diagonal matrix elements. In this step, each walker on SD  $D_i$  may spawn a new set of walkers to connected\* SD  $D_j$ . As we are sampling the sum  $\Delta \tau \sum_{j\neq i} H_{ij}c_j(\tau)$ , as in the Metropolis-Hastings algorithm, it is beneficial (in terms of efficiency) to sample more where  $H_{ij}$  is larger. That is, the probability of spawning on  $D_j$  from  $D_i$ , denoted  $p_{\text{gen}}(j|i)$  (the generation probability) should be proportional to  $H_{ij}$ . Finally, the spawning probability is

$$p_s(j|i) = \Delta \tau \frac{|H_{ij}|}{p_{\text{gen}}(j|i)}.$$
(3.50)

Thus, at each FCIQMC iteration, every determinant  $D_i$  spawns  $N_i p_s(j|i)$  new walkers on a determinant  $D_j$  selected by an excitation generation algorithm. There,  $N_i$  is the (signed) number of walkers on  $D_i$ . If  $H_{ij} < 0$  then the child walkers are the same sign as the parent; otherwise, they have opposite sign. The original formulation of the algorithm used integer number of walkers, and spawned a walker with probability  $p_s(j|i)$  per walker on  $N_i$ , but it is more efficient to simply use noninteger values and spawn  $p_s(j|i)$  walkers.

#### Death/Cloning

The death/cloning step is characterised by the diagonal matrix elements. In this step,  $p_d$  walkers are removed from the simulation<sup>†</sup> where

$$p_d = \Delta \tau (H_{ii} - S). \tag{3.51}$$

<sup>\*</sup> $D_i$  and  $D_j$  are said to be connected if  $\langle D_i | H | D_j \rangle \neq 0$ .

<sup>&</sup>lt;sup>†</sup>The original formulation removed (or cloned) walkers with a probability of  $p_d$ .

Notice that if  $p_d$  is the same sign as the walker, then the number of walkers on  $D_i$  increases (hence "cloning", as opposed to "death"). "Cloning" is especially common in the growth stage of the population, where the population is growing exponentially towards the target  $N_w$ .

#### **Annihilation**

After spawning and death/cloning, we may have walkers of opposite sign on the same determinant. This is dealt with in the "annihilation" step of the algorithm, realised by combining the terms in equation 3.47. This step, while simple in principle, amounts to cancelling opposite walkers. Determinants without walkers left on them are then removed as well, to keep the storage of  $|\Psi\rangle$  sparse. However, so far the steps have all been embarrassingly parallel, but in this step we need to communicate in order to locate determinants on the same determinant, so this step is critical for parallelisation. Nevertheless, efficient implementations such as in NECI is able to scale efficiently to over 40000 cores.<sup>277</sup>

#### 3.4.3. Energy Estimators

The shift S is updated to approximate  $E_0$ , so we already have an estimator for the energy. However, since S is used to also control the population, this estimator can be noisy.<sup>266</sup> A more commonly used energy estimator is the projected energy,

$$E_{\text{proj}} = \frac{\langle D_{\text{ref}} | \hat{H} | \Psi \rangle}{\langle D_{\text{ref}} | \Psi \rangle}, \tag{3.52}$$

where we project onto the reference (most occupied) determinant.\*

In principle, any wave function could be used in place of the trial determinant in equation 3.52 (so long as it has nonzero overlap with  $|\Psi\rangle$ ). The choice of the reference determinant is for ease of computation, and because the larger the overlap with  $|\Psi\rangle$ , the smaller the error in the stochastic sampling. Since the reference determinant (by definition) has the largest number of walkers on it, it is the sensible choice for a single-determinant projected energy.

Another energy estimator is the "trial energy",  $^{279,280}$  which is just the projected energy on a multi-determinant wave function instead of  $|D_{\text{ref}}\rangle$ . In this method, after some time in variable shift mode, a "trial space" is constructed based on the top  $N_T$  determinants. The Hamiltonian spanned within the trial space is diagonalised exactly (sparsely or densely), and the eigenvector  $|\Psi_{\text{trial}}\rangle$  is stored. Then, the trial projected energy can be calculated as

$$E_{\text{trial}} = \frac{\langle \Psi_{\text{trial}} | \hat{H} | \Psi \rangle}{\langle \Psi_{\text{trial}} | \Psi \rangle}.$$
 (3.53)

<sup>\*</sup>While it may be tempting to instead use the energy estimator  $\langle \Psi | \hat{H} | \Psi \rangle / \langle \Psi | \Psi \rangle$ , this comes with a prohibitive cost, and has a bias. <sup>266,278</sup>

Note that while this is expected to reduce the error in the energy estimator, it also increases the cost by a factor of  $N_T$ .

An example of these energy estimators is shown in figure 3.2. As in other QMC methods, these energy estimators have autocorrelation, so in reporting the values, a blocking analysis is required.<sup>249</sup>

#### 3.4.4. Annihilation Plateaus

FCIQMC has its own kind of sign problem in the form of resolving the correct relative sign structure of the sampled states in the CI vector.<sup>281</sup> This manifests in the form of a so-called "annihilation plateau", illustrated in figure 3.3.

In this stage of the simulation, walkers are being spawned with incoherent signs, causing a massive amount of walker annihilation. This causes the population to stay relatively constant, until the sign structure is finally resolved and the population continues to grow again. The height of the plateau (i.e. the constant number of walkers where the simulation stagnates) quantifies the difficulty of the sign problem. In particular, for certain large systems, it can be prohibitively expensive to overcome. If we are below the annihilation plateau, then the original formulation of FCIQMC is not reliable, as the correct sign structure is not yet resolved.

#### 3.4.5. The Initiator Approximation

Since incoherent walker proliferation is a major source of the annihilation plateau, a sensible strategy to overcome this is by forcing growth to be coherent. This is achieved by the initiator approximation.<sup>282</sup> An FCIQMC calculation with the initiator approximation is sometimes referred to as i-FCIQMC; however, since the approximation is so useful and ubiquituous, we simply referred to it as FCIQMC. Indeed, in this dissertation all FCIQMC calculations used the initiator approximation, unless otherwise specified.

The initiator approximation allows only those determinants (dubbed "initiators") with at least  $N_{\text{thresh}}$  walkers to spawn new walkers on unoccupied determinants. If the population is greater than this threshold, the assumption is that the (relative) sign of that state is correct, and hence the spawning event should produce the correct sign.

Since the approximation truncates spawning, the growth phase of the simulation is slower. However, the annihilation plateau disappears. Note that as  $N_w \to \infty$ , the initiator approximation becomes exact, as all sampled states are initiators. While there is a small bias introduced by the initiator approximation, the efficiency is greatly improved and the range of potential applications is greatly increased.

#### 3.4.6. Reduced Density Matrix Sampling

As described in section 1.4.2, reduced density matrices are a useful tool for understanding the properties of a system. These can be sampled in FCIQMC by considering the RDMs

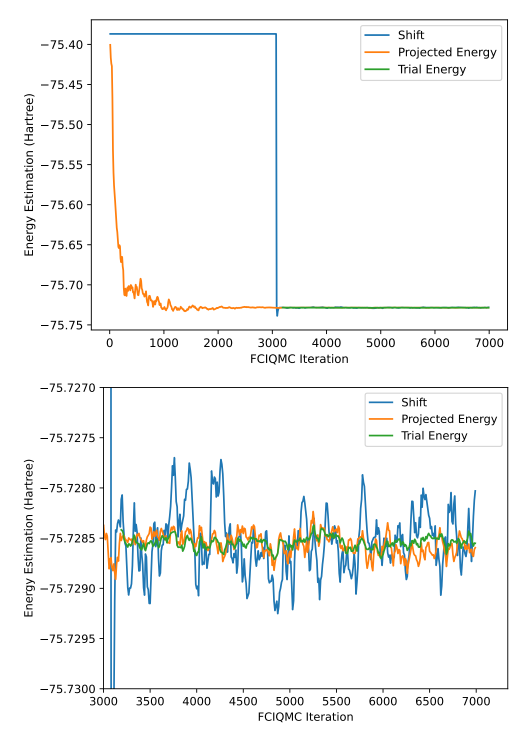


Figure 3.2. An example of the energy estimators used in an FCIQMC simulation. The shift is the most noisy, whereas the trial energy is the least noisy, but only available in variable shift mode and carries the highest cost. In this case, the trial energy is not a substantial improvement from the reference-projected energy. This calculation was done on the  $C_2$  molecule with the cc-pVDZ basis, at equilibrium geometry, 1.2425 Å.

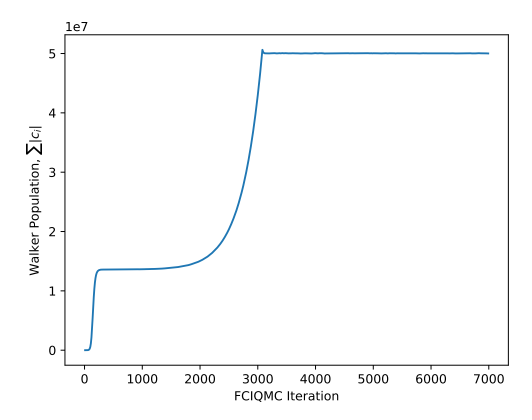


Figure 3.3. An example of an annihilation plateau in the context of a FCIQMC simulation. Despite not yet reaching the target population, the total population is roughly constant before increasing again. This stage is referred to as the annihilation plateau, and the higher the plateau, the more difficult the sign problem is. This calculation was done on the  $C_2$  molecule with the cc-pVDZ basis, at equilibrium geometry, 1.2425 Å. In this particular case, the plateau appears around  $1.36 \times 10^7$  walkers, and the target number is  $5 \times 10^7$ .

in the same basis.<sup>283</sup> The 1RDM is given by

$$\gamma_{pq} = \sum_{ij} c_i c_j \langle D_i | a_p^{\dagger} a_q | D_j \rangle \tag{3.54}$$

and the 2RDM by

$$\Gamma_{pqrs} = \sum_{ijkl} c_i c_j \langle D_i | a_p^{\dagger} a_q^{\dagger} a_s a_r | D_j \rangle.$$
(3.55)

In FCIQMC, we have the CI coefficients  $c_i$  and  $c_j$  by the walker distributions. However, using the same simulation for both  $c_i$  and  $c_j$  introduces a bias, hence to stochastically sample RDMs, we use an independent simultaneous simulation (referred to as a replica). One simulation samples  $c_i$  whereas the other samples  $c_j$ . Since the statistical errors are independent, the overall error is reduced. Unfortunately this requires twice the computing power, but thankfully since the calculations are independent, they are trivially parallelisable.

Note also that by inserting the Hamiltonian (energy) operator into equation 1.21 and using the replica trick to obtain the RDMs, we can also obtain yet another energy measure in FCIQMC.

#### 3.4.7. Combining TC with Modern Electronic Structure

The FCIQMC algorithm may also be combined with the TC method described in section  $2.5.^{205}$  Consider solving for the eigenstates  $\Phi$  of the TC Hamiltonian  $\hat{H}_{\text{TC}}$  using the imaginary-time Schrödinger equation,

$$-\frac{\partial}{\partial \tau} \Phi = (\hat{H}_{TC} - S)\Phi. \tag{3.56}$$

Since  $\hat{H}_{TC}$  and  $\hat{H}$  are isospectral, we have stationary states for the same values as the non-TC method. That is, we can control the walker population by setting the shift to the ground state energy,  $S = E_0$ .

The state  $\Phi$  is described by a linear combination of SDs, as in non-TC-FCIQMC,

$$|\Phi\rangle = \sum_{i} c_i |D_i\rangle. \tag{3.57}$$

However, since  $\hat{H}_{TC}$  is non-Hermitian, the coefficients  $\tilde{c}_i$  of the left-eigenvector with the same energy are not necessarily the same,

$$\langle \Phi | = \sum_{i} \tilde{c}_{i} \langle D_{i} |, \qquad (3.58)$$

and we must be careful when considering matrix connections  $\langle D_i | \hat{H}_{TC} | D_j \rangle$  and  $\langle D_j | \hat{H}_{TC} | D_i \rangle$ . In particular, the probability of spawning a walker from  $|D_i\rangle$  to  $|D_j\rangle$  may not be the same as the probability of spawning from  $|D_j\rangle$  to  $|D_i\rangle$ . Furthermore, when considering replicas such as in RDM sampling, the replica needs to be of the adjoint operator,  $\hat{H}_{\text{TC}}^{\dagger}$ , in order to target  $\tilde{c}_i$ . This is accomplished by a simple transform of the Jastrow factor,  $J \to -J$ , though this may cause practical complications. Otherwise, FCIQMC may be extended to TC by simply applying the method directly to  $\hat{H}_{\text{TC}}$ . An appropriate transformation from  $\hat{H}$  is therefore necessary beforehand.

Similar arguments may be made for applying TC to CC methods<sup>213,214</sup> and DMRG,<sup>189</sup> and TC variants of these methods have already been developed and successfully applied. In addition to TC-FCIQMC, this also continues to be an active area of research.

# Optimising Jastrow Factors for the Transcorrelated Method

This chapter is based in large part on the following paper, and most of the following discussion can already be found there:

Haupt, J. P.; Hosseini, S. M.; López Ríos, P.; Dobrautz, W.; Cohen, A.; Alavi, A. "Optimizing Jastrow Factors for the Transcorrelated Method". The Journal of Chemical Physics **2023**, *158*, 224105

Images have been reused from this paper (with permission).

#### 4.1. Introduction

In this chapter, we investigate the use of flexible Jastrow factors and a novel optimisation strategy for use in TC as introduced in section 2.5. As a brief recapitulation, the TC method amounts to a similarity transformation of the Hamiltonian  $\hat{H}$ ,  $\hat{H}_{TC} = \mathrm{e}^{-J}\hat{H}\mathrm{e}^{J}$ . However, as this is a non-unitary transformation, methods used to solve  $\hat{H}_{TC}$  are in general not variational, and hence we are not guaranteed to converge to the CBS limit from above. It is therefore important to choose J wisely, as otherwise the method may be highly non-variational, and we may suffer from poor error cancellation.

As an illustration of the method, we compute the all-electron atomisation energies for the challenging first-row molecules  $C_2$ , CN,  $N_2$  and  $O_2$  and find that TC-FCIQMC (that is, FCIQMC performed on a transcorrelated Hamiltonian) yields chemically accurate results using only a cc-pVTZ basis, which requires a much larger cc-pV5Z basis for non-TC.

# 4.2. Computational Details

We compute the ground-state energies of the all-electron C, N, and O atoms, as well as that for the  $C_2$ , CN,  $N_2$  and  $O_2$  molecules at their equilibrium geometries,  $^{285-287}$  listed in table 4.1. TC- and non-TC-FCIQMC calculations used HF orbitals (restricted open-shell in the case of open-shell systems) expanded in the standard cc-pVXZ family of basis sets.  $^{288}$ 

The quality of the energy differences is assessed using the atomisation energies of these molecules. In order to determine if our methodology yields chemically-accurate, i.e. within

wing reference 200.			
	System	State	$r_{\rm eq}  (\rm \AA)$
	$C_2$	$1\Sigma_q^+$	1.2425
	CN	$^2\Sigma^{g}_+$	1.1718
	$N_2$	$^{1}\Sigma_{q}^{+}$	1.0977
	$O_2$	$3\sum_{a}$	1.2075

**Table 4.1.** Electronic ground states and equilibrium bond lengths used for the molecules considered in this work, following reference 285.

an error of 1 kcal/mol  $\approx$  1.6 mHa, we also keep each individual error to be well within this threshold. We expect a total bias in our resulting relative energies of not more than 0.5 mHa.

For all our calculations, we generate our orbitals and integration grids using pyscf, <sup>289</sup> optimise the Jastrow factors using the CASINO continuum QMC package, <sup>263</sup> compute TC matrix elements using the tchint library, for which more details are presented in Appendix A, and perform (TC-)FCIQMC calculations using the NECI package. <sup>267</sup> FCIQMC energies reported are the standard HF-projected energies.

(TC-)FCIQMC values presented here were produced using a walker-number extrapolation scheme presented in another dissertation.<sup>290</sup>

## 4.3. Jastrow Factor

In continuum quantum Monte Carlo methods, the Jastrow factor for a molecule is typically expressed as the sum of electron-electron, electron-nucleus, and electron-electron-nucleus terms,\*

$$J = \sum_{i < j}^{N} v(r_{ij}) + \sum_{i}^{N} \sum_{I}^{N_A} \chi(r_{iI}) + \sum_{i < j}^{N} \sum_{I}^{N_A} f(r_{ij}, r_{iI}, r_{jI}), \tag{4.1}$$

where  $N_A$  is the number of nuclei, N the number of electrons, and each of u,  $\chi$ , and f are expressed as natural power expansions.<sup>224</sup> That is,

$$v(r_{ij}) = t(r_{ij}, L_v) \sum_{k} a_k r_{ij}^k,$$
 (4.2)

$$\chi(r_{iI}) = t(r_{iI}, L_{\chi}) \sum_{k} b_k r_{iI}^k,$$
(4.3)

$$f(r_{ij}, r_i, r_j) = t(r_{iI}, L_f)t(r_{jI}, L_f) \sum_{k,l,m} c_{klm} r_{ij}^k r_{iI}^l r_{jI}^m,$$
(4.4)

<sup>\*</sup>Of course, these are not all the possible terms. We may, for example, also choose to include electron-nucleus-nucleus terms.

4.3. Jastrow Factor 53

where  $\{a_k\}$ ,  $\{b_k\}$ , and  $\{c_{klm}\}$  are linear parameters,  $L_v$ ,  $L_\chi$ , and  $L_f$  are cut-off lengths,  $t(r,L) = (1-r/L)^3\Theta(r-L)$  is a cut-off function, and  $\Theta(r-L)$  is the Heaviside step function.

As described in chapter 2, accurately describing the (electron-electron and electron-nucleus) Kato cusp conditions<sup>67</sup> substationally improves the accuracy of our method. Also, as described in chapter 3, VMC and DMC methods sample electronic configurations  $\{R\}$  from a probability distribution based on an analytical trial wave function  $\tilde{\Psi}_{\rm T}(R)$  to produce a variational estimate of the total energy as an average of the local energy,  $E_{\rm L}(R) = \tilde{\Psi}_{\rm T}^{-1}(R)\hat{H}(R)\tilde{\Psi}_{\rm T}(R)$  over the sampled configurations. In the case of VMC, accurate description of the electron-electron and electron-nucleus Kato cusp conditions suppresses extreme outliers in the local energy sampling, allowing meaningful wave function parameters.

The most obvious way to enforce the electron-electron and electron-nucleus cusp conditions is by enforcing them in the form of the Jastrow factor through the relevant terms, namely v (equation 4.2) for the electron-electron cusp, and  $\chi$  (equation 4.3) for the electron-nucleus cusp. However, in the context of continuum QMC, it has been found to be better<sup>224,253,291</sup> to enforce the electron-nucleus cusp by modifying the l=0 (s-type) component of the cuspless molecular orbitals,  $\phi(r)$ , such that they exhibit a cusp.

Since we are interested in performing a post-Hartree-Fock calculation on  $\hat{H}_{TC}$ , such as FCIQMC, it is preferable to use unmodified molecular orbitals from standard basis sets during the optimisation process. If we optimise the Jastrow factor in VMC in the presence of cusp-corrected orbitals and then use them in TC-FCIQMC without the cusp-corrected orbitals, the Jastrow factor would be sub-optimal for the Hamiltonian, by construction.

Instead, we recast the cusp-correction scheme of Ref. 291 as an electron-nucleus Jastrow factor term, called  $\Lambda$ , to be added (rather than replacing) the  $\chi$  term of equation 4.3. We construct this term as

$$\Lambda(r) = \left[ \ln \tilde{\phi}(r) - \ln \phi(r) \right] \Theta(r - r_c), \tag{4.5}$$

where, adopting the notation of Ref. 291,  $r_c$  is a cutoff radius,  $\phi(r)$  is the s-type component of the target orbital, and  $\tilde{\phi}(r)$  is its cusp-corrected counterpart,

$$\tilde{\phi}(r) = e^{\sum_{l=0}^{4} \alpha_l r^l} + C \quad , \quad r < r_c.$$
 (4.6)

Here,  $\{\alpha_l\}$  are parameters determining the shape of the corrected orbital and the shift C is only set to a non-zero value in the presence of nodes of  $\phi(r)$  near the nucleus. More precisely, the shift C is chosen such that  $\tilde{\phi}(r_c) - C$  is of one sign within the radius  $r_c$ . This is necessary since we wish to impose an exponential correction, which is necessarily of one sign.

Following Ref. 291, we impose the cusp condition at  $r = r_c$ , as well as twice continuous differentiability at  $r = r_c$ . This leaves only  $\alpha_0$  and  $r_c$  as free parameters from equations

4.5 and 4.6.  $r_c$  is chosen to be small but within the same sign, as described above, while  $\alpha_0$  is determined by enforcing smoothness for the so-call "effective one-electron local energy",

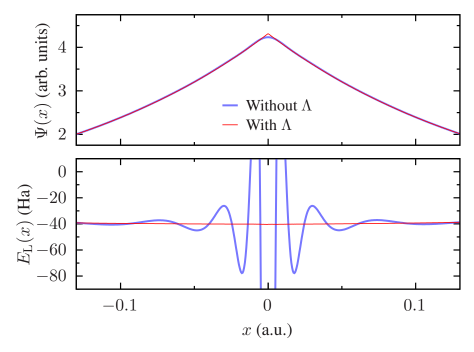
$$E_L^s(r) := \tilde{\phi}(r)^{-1} \left[ -\frac{1}{2} \nabla^2 - \frac{Z_{\text{eff}}}{r} \right] \tilde{\phi}(r). \tag{4.7}$$

Here, the effective nuclear charge  $Z_{\text{eff}}$ ,

$$Z_{\text{eff}} = Z \left( 1 + \frac{\eta(0)}{\tilde{\phi}(0)} \right) \tag{4.8}$$

ensures that  $E_L^s$  is finite at the origin, and is derived from the cusp condition.  $\eta$  is the rest of the orbital, leftover from removing the s-type component.

Figure 4.1 illustrates the effect of using a  $\Lambda$  term in practice.



**Figure 4.1.** HF wave function value and local energy as a function of the x coordinate of an electron in a carbon atom as it crosses the nucleus at x = 0, both with and without the  $\Lambda$  cusp-correcting Jastrow factor term. This is in the cc-pVDZ basis.

For the calculations in this chapter, we use a total of 44 optimisable Jastrow factor parameters for the atoms and homonuclear dimers, and 88 parameters for CN. We keep the  $L_v$ ,  $L_\chi$  and  $L_f$  cutoff lengths fixed at 4.5, 4, and 4 Bohr, for simplicity.

# 4.4. Optimisation Strategy

We optimise J using VMC. VMC provides a variational framework in which parameters  $\alpha$  present in a trial wave function  $\Psi_{\rm T}$  can be optimised. In this chapter,  $|\Psi_{\rm T}\rangle = {\rm e}^{J(\alpha)} |D_{\rm HF}\rangle$ .

In continuum QMC, wave function optimisation is usually carried out using a correlated-sampling approach in which a set of  $n_{\rm opt}$  electronic real-space configurations  $\{R_i\}_{i=1}^{n_{\rm opt}}$  distributed according to the initial wave function squared,  $|\Psi_{\rm T}(R;\alpha_0)|^2$  is generated, and then a target function is minimised by varying  $\alpha$  at fixed  $\{R_i\}$ .

The variational energy estimate for this trial wave function may be written

$$E_{\text{VMC}} = \frac{\langle \Psi_{\text{T}} | \hat{H} | \Psi_{\text{T}} \rangle}{\langle \Psi_{\text{T}} | \Psi_{\text{T}} \rangle}, \tag{4.9}$$

which may be used as a target function, as presented in section 3.2.

Another popular target function is the "variance of the VMC energy," 252,292

$$\sigma_{\text{VMC}}^2 = \frac{\langle \Psi_{\text{T}} | (\hat{H} - E_{\text{VMC}})^2 | \Psi_{\text{T}} \rangle}{\langle \Psi_{\text{T}} | \Psi_{\text{T}} \rangle}, \tag{4.10}$$

which reaches its minimum of zero when the trial wave function is an eigenstate of the Hamiltonian. In practice, minimising  $\sigma_{\text{VMC}}^2$  yields variational energies, but is affected by large fluctuations, as shown in this chapter.

In continuum QMC methods, modifications have been devised to circumvent this problem, such as weight limiting, unreweighted variance minimization, or the minimization of other measures of spread such as the median absolute deviation from the median energy. $^{253}$ 

The computational cost of optimizing Jastrow factors within VMC scales as a small power of system size, typically estimated to be  $\mathcal{O}(N^3)$ .

#### 4.4.1. Variance of the Reference Energy Minimisation

In the context of TC, the reference energy

$$E_{\text{ref}} = \langle D_{\text{HF}} | e^{-J} \hat{H} e^{J} | D_{\text{HF}} \rangle \tag{4.11}$$

is of particular significance since it represents the starting point of a TC-FCIQMC calculation (i.e. the walker distributions at imaginary time  $\tau=0$  has this energy). It is also the zeroth-order contribution to the TC-CC energy.

We refer to its associated variance,

$$\sigma_{\text{ref}}^2 = \langle D_{\text{HF}} | e^{-J} (\hat{H}^{\dagger} - E_{\text{ref}}) (\hat{H} - E_{\text{ref}}) e^{J} | D_{\text{HF}} \rangle$$

$$(4.12)$$

as the "variance of the reference energy," which is easily evaluated for a finite VMC sample of size  $n_{\text{opt}}$  as the sample variance of the Slater-Jastrow energy over the HF distribution,

$$S_{\text{ref}}^2 = \frac{1}{n_{\text{opt}} - 1} \sum_{n=1}^{n_{\text{opt}}} \left| \frac{\hat{H}(\boldsymbol{R}_n) \Psi_{\text{SJ}}(\boldsymbol{R}_n)}{\Psi_{\text{SJ}}(\boldsymbol{R}_n)} - \bar{E}_{\text{ref}} \right|^2, \tag{4.13}$$

which tends to  $\sigma_{\text{ref}}^2$  as  $n_{\text{opt}} \to \infty$ , where  $\Psi_{\text{SJ}} := e^J D_{\text{HF}}$  is the Slater-Jastrow wave function,  $\{\boldsymbol{R}_n\}_{n=1}^{n_{\text{opt}}}$  are electronic configurations distributed according to  $D_{\text{HF}}^2$ , and the VMC estimate of the reference energy is

$$\bar{E}_{\text{ref}} = \frac{1}{n_{\text{opt}}} \sum_{n=1}^{n_{\text{opt}}} \frac{\hat{H}(\boldsymbol{R}_n) \Psi_{\text{SJ}}(\boldsymbol{R}_n)}{\Psi_{\text{SJ}}(\boldsymbol{R}_n)}.$$
(4.14)

The variance of the reference energy has been used as a target function for optimising Jastrow factors for the TC method before, albeit in different theoretical frameworks.<sup>223,293</sup>

To understand the physical significance of the variance of the reference energy, note that equation 4.12 may be rewritten as

$$\sigma_{\text{ref}}^2 = \sum_{I \neq \text{HF}} \langle D_I | \hat{H}_{\text{TC}} | D_{\text{HF}} \rangle, \qquad (4.15)$$

where I runs over a complete basis set.\*

As evident by equation 4.15, minimising  $\sigma_{\text{ref}}^2$  essentially amounts to minimising the coupling of the reference determinant with the remainder of the space, which in the context of FCIQMC translates to a reduced spawning rate from the reference determinant to its connected excited-state determinants, thereby increasing the amplitude of the reference determinant in the resulting CI vector.

Note also that if the Slater-Jastrow wave function were an exact eigenstate of  $\hat{H}$ , then a TC-FCIQMC simulation starting from the HF determinant would immediately converge to a strictly single-determinant solution. Although this ideal scenario cannot be achieved in practice, it nevertheless illustrates the potential benefits of obtaining a relatively single-reference CI solution by minimising this target function. We expect that this increased single-reference character will also benefit other approaches, particularly those based on single references, such as TC-CC.

We therefore investigate the performance of minimising the variance of the reference as an alternative to minimising the variance of the VMC energy. In figure 4.2, we compare the VMC energy and variance obtained by variance minimisation methods along with energy-minimised<sup>294–296</sup> results for reference, for the systems considerd in this chapter, using  $n_{\rm opt} = 10^5$  VMC configurations.

Minimizing the variance of the VMC energy produces lower average values of  $\sigma_{\rm VMC}^2$ , as one would expect, but also erratic VMC energies with very large standard deviations (up to  $\sim 50$  mHa in our tests). Minimizing the variance of the reference energy, on the other hand, produces values of  $\sigma_{\rm VMC}^2$  which are only slightly higher on average than those obtained from minimizing the variance of the VMC energy (1–5% in our tests), while producing more stable VMC energies with much smaller standard deviations (up to

<sup>\*</sup>Here we have a complete basis set because we are optimising with continuum Monte Carlo. If we were to optimise this by directly calculating the matrix elements in equation 4.15, then the sum must be truncated. This is the subject of ongoing work.

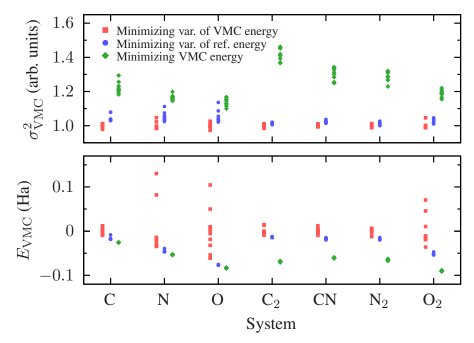


Figure 4.2. Variance of the VMC energy (top) and VMC energy (bottom) of the systems considered in this chapter using the cc-pVTZ basis and Jastrow factors obtained by minimising the variance of the VMC energy (red squares), the variance of the reference energy (blue circles), or the VMC energy (green diamonds) in each of ten independent optimisation runs with  $n_{\rm opt}=10^5$  VMC configurations. To ease comparisons, variances have been rescaled and energies shifted by their average values from minimising the variance of the VMC energy (i.e. the red squares average to a variance of 1 and an energy of 0 in the plot). The subpar ability of VMC energy variance minimisation to yield consistent VMC energies is evident in the bottom panel, suggesting to use the variance of the reference.

 $\sim 3$  mHa in our tests). We therefore do not use "regular" variance minimization since it introduces large stochastic noise, making it unsuitable for optimizing Jastrow factors, and from this point on we use the term "variance minimization" to refer to the minimization of the variance of the reference energy.

## 4.4.2. Choosing an Appropriate Sample Size

While expectation values relevant for most continuum QMC calculations converge using relatively few VMC configurations, it has also been known<sup>297</sup> that in order to converge other quantities, far more configurations are needed. That is,  $n_{\rm opt}$  is larger. In the spirit of MC, we may estimate the convergence of the expectation value for some quantity by performing multiple optimisation runs with different random number seeds but otherwise the same inputs. This would give us a standard deviation.

The value we want to converge in this case is not the VMC energy, nor the reference energy, but instead the TC-FCI energy. In practice, we use the uncertainty of the VMC estimate of the reference energy  $\bar{E}_{\rm ref}$  as a proxy for the standard deviation of the TC-FCIQMC energy. This is justified because:

- The standard deviation of the TC-FCIQMC energy is not larger than the standard deviation of the reference energy, as illustrated in figure 4.3. It is usually significantly smaller thanks to the ability of TC-FCIQMC to conpensate for the presence of a bias in  $E_{\rm ref}$  via the correlation energy.
- The standard deviation of the reference energy is not larger than the statistical uncertainty of the VMC estimate of the reference energy obtained with  $n_{\rm opt}$  configurations. It is usually significantly smaller due to the use of variance reduction techniques in QMC.

For the atoms and molecules considered in this chapter, we use  $n_{\rm opt} = 2 \times 10^7$  to yield TC-FCIQMC energies with standard deviations of less than 0.1 mHa.

#### 4.4.3. Energy Minimisation

The obvious alternative to variance minimisation is minimising the VMC energy, <sup>294–296</sup> which, as already demonstrated in figure 4.2, results in lower VMC energies but higher VMC variances. Energy minimisation yields wave functions which minimise the statistical fluctuations of the local energy in DMC calculations <sup>298</sup> and is the typical choice for continuum QMC. However, for our purposes, it is unclear whether the resulting wave functions provide a better description of the system than those produced by variance minimisation.

In figure 4.4, we compare the convergence with basis-set size of TC-FCIQMC total energies of the C, N, and O atoms using energy- and variance-minimised Jastrow factors.

4.5. Grid Sizes 59

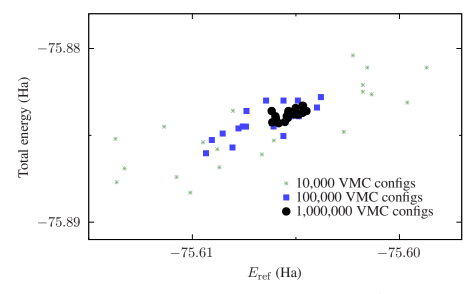


Figure 4.3. TC-FCIQMC energy of the  $C_2$  molecule using  $10^6$  walkers with the cc-pVDZ basis as a function of the reference energy for multiple independent Jastrow factor parameter sets obtained by variance minimisation using three different VMC sample sizes. The horizonal spread is about 1.8 times larger than the vertical spread, in line with the expectation that the standard deviation of the TC-FCIQMC energy is smaller than that of the reference energy.

Variance minimisation appears to produce wave functions which converge quickly and largely variationally to the basis set limit, which energy-minimised wave functions tend to yield non-variational TC-FCIQMC energies which converge more slowly to the basis set limit.

In figure 4.5 we plot the atomisation energies of the dimers as a function of reciprocal basis-set size, again demonstrating variance-minimised Jastrow factors exhibit favourable convergence properties.

Considering this evidence, we choose to use variance minimisation to optimise the Jastrow factors for subsequent post-HF calculations.

#### 4.5. Grid Sizes

How matrix elements are evaluated is presented in Appendix A. As mentioned there, we use Treutler-Ahlrichs integration grids,  $^{299,300}$  which are atom-centred grids constructed as the combination of a radial grid running up to the Bragg radius and a Lebedev angular grid. These are obtained using pyscf, which provides an integer parameter  $l_{\rm grid}$  to control the grid density. Here we test grid errors by evaluating TC-FCIQMC energies at  $l_{\rm grid}=1$ –5 and defining the integration error as the difference of each of these results with the value obtained by linear extrapolation to the  $1/n_{\rm grid} \rightarrow 0$  limit.

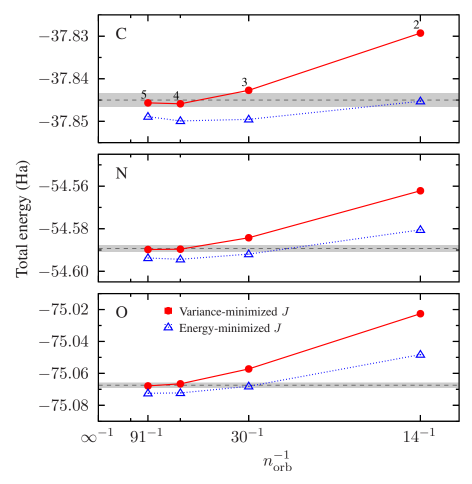


Figure 4.4. Total energy of the C, N, and O atoms as a function of the reciprocal number of molecular orbitals in the cc-pVXZ basis set family. The non-variational behaviour of up to about 5 mHa is evident for the energy-minimised Jastrow factors, for which convergence to the exact energy as a function of basis-set size is slow. The shaded areas represent  $\pm 1$  kcal/mol (so-call "chemical accuracy") around the exact non-relativistic total energy from reference 286. Points in the top panel are annotated with the basis set cardinal number X.

4.5. Grid Sizes 61

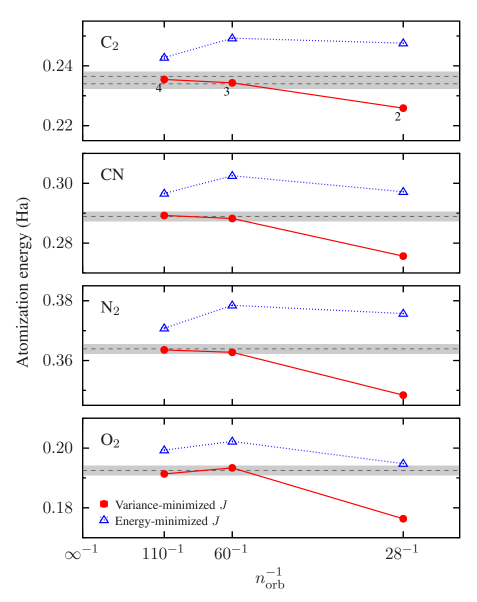
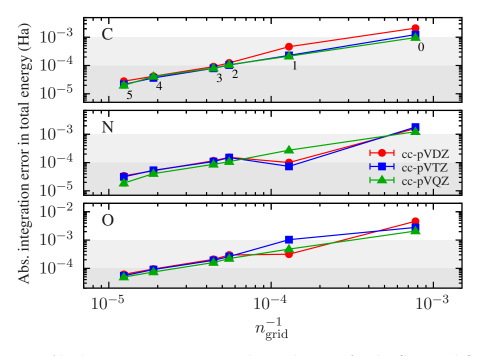


Figure 4.5. Atomisation energy of the  $C_2$ , CN,  $N_2$ , and  $O_2$  molecules as a function of the reciprocal of the number of molecular orbitals using the cc-pVXZ family of basis sets and Jastrow factors obtained by variance and energy minimisation. The shaded areas represent  $\pm 1$  kcal/mol around the theoretical estimate of the exact non-relativistic atomisation energies from references [285, 286]

.

In figure 4.6 we plot the absolute integration error in the total energy of the atoms as a function of  $1/n_{grid}$  for cc-pVDZ, cc-pVTZ and cc-pVQZ basis sets. We find that the basis set size has little to no effect on the integration error.



**Figure 4.6.** Absolute integration errors in the total energy for the C, N, and O atoms as a function of the reciprocal of grid size using basis sets in the cc-pVXZ family. The gray areas correspond to integration errors of less than 1 and 0.1 mHa. Points in the top panel are annotated with the value of **pyscf**'s grid density parameter.

In figure 4.7 we plot the absolute integration error in the total energies of the molecules, the atoms that conform them, and atomisation energies using the cc-pVDZ basis. We find integration-error cancellation in energy differences, with atomisation energy for all four molecules reaching 0.1 mHa for  $l_{\rm grid}=2$ . This represents a 60-point radial grid combined with a 302-point angular grid, for a total of 18120 grid points per atom. We use this throughout this dissertation.

### 4.6. Compactification of the CI Vector

The more compact the CI wave function is, the easier it is for FCIQMC to sample the wave function accurately and the smaller the initiator error becomes. The TC method has already been shown to make CI wave functions more compact for the two-dimensional Hubbard model.<sup>301</sup> Let  $\{c_I\}$  be the  $L^2$ -normalized coefficients of the CI wave function such that  $\sum_I c_I^2 = 1$ . The quantity

$$\xi = \frac{c_{\rm HF}^{\rm (TC)} - c_{\rm HF}^{\rm (non-TC)}}{1 - c_{\rm HF}^{\rm (non-TC)}}$$
(4.16)

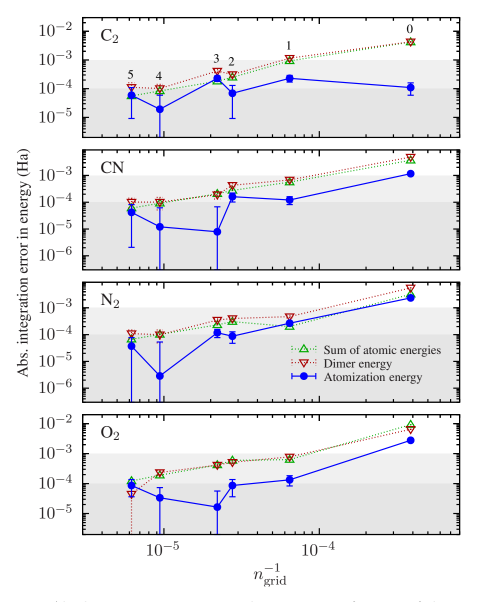


Figure 4.7. Absolute integration error in the energy as a function of the reciprocal number of grid points used for the evaluation of TC integrals. The shaded areas correspond to integration errors of less than 1 and 0.1 mHa. Points in the top panel are annotated with the value of pyscf's grid density parameter. These results demonstrate that  $l_{\rm grid}=2$  is sufficient to achieve sub-mHa accuracy in total energies and sub-0.1-mHa accuracy in relative energies.

is then a measure of the enhancement in the compactness of the wave function, going from 0 for no enhancement to 1 if the TC wave function becomes exactly single-determinantal.

From reference 301, TC yields a maximum of  $\xi = 0.64$  for the 18-site 2D Hubbard model. We find that the values of  $\xi$  for atomic and molecular systems are not dissimilar from this, as shown in table 4.2.

	С	N	О	$C_2$	CN	$N_2$	$O_2$
cc-pVDZ	0.46	0.63	0.71	0.14	0.23	0.38	0.53
cc- $pVTZ$	0.45	0.61	0.69	0.15	0.24	0.40	0.55
$\operatorname{cc-pVQZ}$	0.44	0.60	0.69	0.15	0.24	0.41	0.57

**Table 4.2.** Enhancement of the compactness of the CI wave function,  $\xi$  in Eq. 4.16, between our non-TC and TC-FCIQMC calculations.

#### 4.7. Neglecting Three-Body Excitations

As described in section 3.4, sampling in FCIQMC involves spawning walkers from occupied determinants onto connected determinants. The  $\hat{L}$  matrix, that is the three-body operator arising from the similarity transformation in TC, results in TC also connecting determinants by three-body excitations. This represents a huge increase in the connectivity of the Hilbert space compared to non-TC-FCIQMC.

These three-body excitations are represented by  $L_{ijk}^{abc}$  where each index is distinct. These have been found to typically be small.<sup>192</sup> Moreoever, simultaneous interaction of three electrons is unlikely due to Pauli repulsion (as necessarily at least two of the electrons will be of the same spin). Therefore, we neglect these contributions. The only distinct six-index integrals that we must keep are those occupied in the HF determinant, as these matrix elements are necessary to evaluate the projected energy.

Neglecting pure three-body excitations reduces the amount of storage needed to hold  $\hat{L}$  from  $\mathcal{O}(M^6)$  to  $\mathcal{O}(M^5 + N^3M^3)$ , where M is the number of orbitals and N is the number of electrons. Reduction factors obtains for the molecules studied in this chapter are reported in table 4.3.

	С	N	О	$C_2$	CN	$N_2$	$O_2$
cc-pVDZ	1.23	1.17	1.17	1.87	1.78	1.78	1.58
cc- $pVTZ$	2.04	2.02	1.93	3.72	3.66	3.66	3.46
cc- $pVQZ$	3.31	3.44	3.13	6.60	6.54	6.57	6.41

**Table 4.3.**  $\hat{L}$  matrix storage reduction factor from neglecting pure three-body excitations, computed as the number of non-zero matrix elements in a the full  $\hat{L}$  matrix divided by the number of non-zero matrix elements with repeated indices or three or more indices corresponding to orbitals occupied in the HF determinant.

Since two-body excitations are typically more expensive to attempt in practice compared to triple excitations, neglecting the latter actually increases the cost per step of the calculation. However, neglecting pure three-body excitations allows the TC-FCIQMC time step to be larger, thereby resulting in reduced serial correlation in the statistics, which enables reaching the target accuracy in fewer time steps, so one can expect a net cost reduction thanks to this approximation. Walltime reduction factors for the molecules studied in this chapter are reported in table 4.4.

	С	N	О	$C_2$	CN	$N_2$	$O_2$
cc-pVDZ	0.9	1.0	1.1	1.7	1.2	1.5	1.6
cc- $pVTZ$	1.0	1.0	0.8	2.4	1.0	1.8	2.0
$\operatorname{cc-pVQZ}$	1.5	1.5	1.0	3.1	0.9	1.9	2.3

**Table 4.4.** Reduction factor in the walltime required to advance one unit of imaginary time at fixed population from neglecting pure three-body excitations in the TC-FCIQMC calculation.

The effect of neglecting these pure three-body excitations on atomisation energy is also small, as shown in table 4.5. We find that this approximation results in errors of the order of  $\sim 0.3$  mHa at the cc-pVTZ level, which is a relatively small bias considering the substantial storage and cost benefits of the approximation.

	L		,	0
	$C_2$	(CN	$N_2$	$O_2$
cc-pVDZ	-0.62(2)	-0.46(0)	-0.56(2)	-0.55(2)
cc- $pVTZ$	-0.36(5)	-0.30(2)	-0.32(5)	-0.20(3)
cc- $pVQZ$	-0.45(6)	0.21(2)	-0.32(7)	-0.27(5)

**Table 4.5.** Error in the atomization energy of the molecules considered in this work incurred by neglecting pure three-body excitations from the FCIQMC dynamics, in mHa.

#### 4.8. Atomisation Energies

In this section, we compare results of TC-FCIQMC with individually variance-optimised Jastrow factors as a function of basis-set size with the corresponding non-TC results and with benchmark CBS values from references [285–287].

Table 4.6 shows a list of total energies obtained for each system and basis set. We find the TC total energies to be remarkably accurate already at the cc-pVQZ basis set level, differing by less than 2 mHa per atom from benchmark CBS values, while the non-TC total energies still miss the benchmarks by 25–30 mHa per atom with the cc-pV5Z basis set, and 20 mHa per atom at the cc-pV6Z level. The TC total energies exhibit slightly non-variational convergence, with the atomic energies reaching values 0.5 mHa below the benchmark before increasing again towards it for larger basis-set sizes. While

		С	N	O
	cc-pVDZ	-37.7619	-54.4801	-74.9117
$\Sigma$	cc-pVTZ	-37.7900	-54.5252	-74.9853
non-T	cc-pVQZ	-37.8126	-54.5535	-75.0236
noı	cc-pV5Z	-37.8199	-54.5627	-75.0369
	cc-pV6Z	-37.8263	-54.5697	-75.0447
	cc-pVDZ	-37.8293	-54.5622	-75.0226
$\circ$	cc-pVTZ	-37.8427	-54.5842	-75.0572
Ĭ	cc-pVQZ	-37.8459	-54.5896	-75.0665
	cc-pV5Z	-37.8457	-54.5898	-75.0678
]	Ref. 286	-37.8450	-54.5893	-75.0674
]	Ref. 287	-37.8450	-54.5893	-75.0674

		$C_2$	CN	$N_2$	$O_2$
7	cc-pVDZ	-75.7320	-92.4970	-109.2809	-149.9915
)L·	cc-pVTZ	-75.8094	-92.5954	-109.4014	-150.1554
non-	cc-pVQZ	-75.8578	-92.6517	-109.4653	-150.2362
п	cc-pV5Z	-75.8752	-92.6717	-109.4881	-150.2655
	cc-pVDZ	-75.8844	-92.6671	-109.4727	-150.2216
$\circ$	cc-pVTZ	-75.9197	-92.7152	-109.5312	-150.3078
Ľ	cc-pVQZ	-75.9272	-92.7247	-109.5428	-150.3244
]	Ref. 285	-75.9240	-92.7232	-109.5425	-150.3273
]	Ref. 286	-75.9265		-109.5427	-150.3274
]	Ref. 287		-92.7229	-109.5425	-150.3275

**Table 4.6.** Total energies in Ha obtained for atoms (top) and molecules (bottom) considered in this work, along with benchmark non-relativistic results. Statistical uncertainties from Monte Carlo sampling are smaller than 0.0001 Ha.

non-variationality is undesirable in a method, the amount by which the TC results dip below the CBS limit is sufficiently small for this not to be an issue in practice.

From a chemical perspective, relative energies are more important than totally energies. The atomisation energies of the  $C_2$ , CN,  $N_2$ , and  $O_2$  molecules obtained from the total energies in table 4.6 are given in table 4.7 and plotted in figure 4.8.

		$C_2$	CN	$N_2$	$O_2$
	cc-pVDZ	208.2	255.0	320.7	168.0
Ĥ	cc-pVTZ	229.4	280.1	351.0	184.9
non-	cc-pVQZ	232.6	285.6	358.4	189.0
n	cc-pV5Z	235.5	289.2	362.6	191.7
	cc-pVDZ	225.9	275.6	348.4	176.4
$\Gamma$ C	cc-pVTZ	234.3	288.2	362.7	193.4
	cc-pVQZ	235.5	289.3	363.6	191.4
	Ref. 285	234.0	288.9	363.9	192.5
	Ref. 286	236.5		364.1	192.6
	Ref. 287		288.6	363.9	192.7

**Table 4.7.** Atomisation energies in mHa obtained for the molecules considered in this chapter, along with benchmark non-relativistic results. Statistical uncertainties arising from Monte Carlo sampling are smaller than 0.1 mHa in all cases.

As with the total energies, we find that the TC atomisation energies converge much faster with respect to basis-set size compared to the non-TC calculations. In fact, the TC results are already chemically accurate at the cc-pVTZ basis-set level, matching accuracy of cc-pV5Z non-TC results. The application of the TC method to quantum chemical methods in general could be presumed to be problematic because any theoretical guarantee of cancellation of errors in energy differences disappears with the introduction of separately-optimized Jastrow factors for each system. However, the fact that in our results the relative energy converges at smaller basis-set sizes than the total energy implies that substantial error cancellation is at play in practice.

#### 4.9. Conclusion and Outlook

In this chapter, we presented a new method to optimise flexible Jastrow factors of a form common in continuum QMC methods for applications to TC. We tested these within the TC-FCIQMC framework. Minimising the variance of the reference energy is shown to be especially good for the TC method, since it maximises the single-reference character of the CI wave function.

These results show that this workflow can provide remarkably accurate total and relative energies, with relative energies at cc-pVTZ rivalling non-TC results at cc-pV5Z. These results also provide better energy estimates compared to other promising methods, such as neural-network based trial wave functions. $^{302,303}$ 

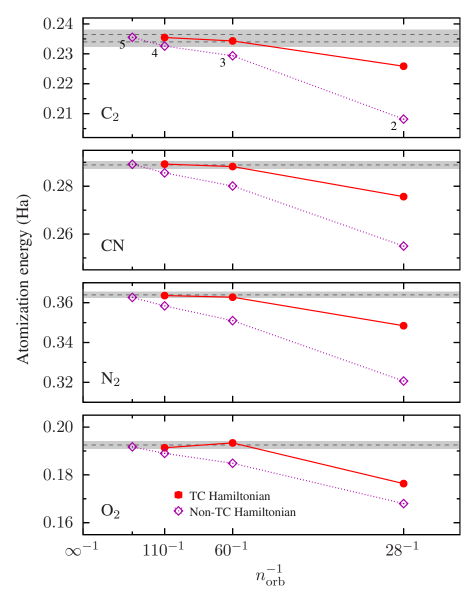


Figure 4.8. Atomisation energy of the  $C_2$ , CN,  $N_2$ , and  $O_2$  molecules obtained with FCIQMC and TC-FCIQMC as a function of the reciprocal of the number of molecular orbitals using the cc-pVXZ family of basis sets. Points in the top panel are annotated with the basis set cardinal number X. The shaded areas represent  $\pm 1$  kcal/mol around the theoretical estimate of the non-relativistic atomisation energy of reference 285; the distinct estimate of reference 286 is also shown for  $C_2$ .

Future work on this topic have focused on technical advancements and efficient approximations for dealing with the three-body integrals.

#### 4.9.1. The xTC Approximation

One particularly significant refinement of this method is the so-called transcorrelation via exclusion of explicit three-body components (xTC) approximation, <sup>190</sup> wherein upon

neglecting explicit three-body contributions, the remaining three-body terms are "folded" into the remaining lower-order terms.

The Hamiltonian is normal-ordered with respect to  $\Phi$  (where  $\Psi = e^{J}\Phi$ , with  $\Phi$  being the HF determinant in this chapter)<sup>304,305</sup>

$$H_N = \hat{H}_{TC} - \langle \Phi | \hat{H}_{TC} | \Phi \rangle = F_N + V_N + L_N \tag{4.17}$$

where the one-, two-, and three-body operators are (using Einstein summation)

$$F_N = [h_q^p + (U_{qs}^{pr} - U_{sq}^{pr})\gamma_s^r$$
(4.18)

$$-\frac{1}{2}(L_{qsu}^{prt} - L_{qus}^{prt} - L_{usq}^{prt})\gamma_{us}^{rt}]\tilde{a}_{q}^{p}, \tag{4.19}$$

$$V_N = \frac{1}{2} [U_{qs}^{pr} - (L_{qsu}^{prt} - L_{qus}^{prt} - L_{usq}^{prt})] \tilde{a}_{qs}^{pr}, \tag{4.20}$$

$$L_N = \frac{1}{6} L_{qsu}^{prt} \tilde{a}_{qsu}^{prt} \tag{4.21}$$

and U=V-K,  $\tilde{a}_{p...}^{q...}$  are the normal-ordered excitation operators, and  $\gamma_{p...}^{q...}=\langle\Phi|\,a_{p...}^{q...}\,|\Phi\rangle$  are the density matrices.

Ignoring the three-body terms  $L_N$  leads to an improved scaling with respect to basis set size while directly contracting intermediates to the lower-body corrections obviates the necessity to calculate L and results in excellent energy agreement. Once the integrals are calculated,  $\hat{H}_{TC}$  has, like  $\hat{H}$ , at most two-body interactions and therefore needs storage for four indices (albeit we no longer have Hermiticity). This means that calculations done on these integrals (in principle) are not much more expensive than the conventional non-TC calculations.

# The Transcorrelated Method for Multi-Reference Problems

This chapter is based in large part on the following upcoming publication:
Haupt, J. P.; Ríos, P. L.; Christlmaier, E. M. C.; Kats, D.; Alavi, A. "The Transcorrelated Method for Strongly Multireference Problems". **2024**, To be submitted Images have been reused from this paper (with permission).

#### 5.1. Introduction

In this chapter, we apply the new framework for the transcorrated method described in chapter 4 to problems of multireference character and find these methods may yield non-physical results. We propose an updated workflow wherein we use conventional post-Hartree-Fock methods as input to Jastrow optimisation for TC-FCIQMC. Results suggest size-consistent results and rapid basis set convergence compared to conventional methods, with the binding curve of  $N_2$  at aug-cc-pVTZ being within chemical accuracy of experiment.

#### 5.2. Motivation

Consider the methodology outlined in chapter 4. If we again use the same Jastrow factors, TODO: write Jastrow factor down again with the same objective function, TODO: write variance of reference with a single(!) reference

$$TODO$$
 (5.1)

and we use the workflow to calculate points along the binding curve of  $N_2$ , we find a non-physical "bump", as shown in figure 5.1.

Figure 5.1. TODO: figure showing a plot of the binding curve(s) with a single reference!

TODO: problem description, including the "dip"

#### 5.3. Size Consistency

#### 5.3.1. Jastrow Factor Forms

#### 5.3.2. Multireference Ansatzes

TODO: Present: "circular" approach, CASCI, CASSCF

#### 5.4. Transcorrelated Trial Wavefunction

#### 5.5. Results

#### 5.5.1. Binding Curves

TODO: mention that we fix the binding curves TODO: also include a table about just "how size-consistent" the results actually are.

#### 5.5.2. Excitation Energies

TODO: mention can target the exact state(s) in question

#### 5.6. Conclusion and Outlook

TODO: make sure to mention the cutoff analysis and constructing Jastrow factors from atomic Jastrow factors. Mention ECPs, since it's already been submitted... TODO: TC-CAS... TODO: self-consistent TC (i.e. feed back into itself)

## (Quasi-)Universal Jastrow Factors

The contents of this chapter are planned to be expanded for a future publication. Some images may be repeated there.

#### 6.1. Introduction

TODO: describe motivation (ease of use, scalability, generalisability)

#### 6.2. Theory

TODO: describe how to construct them, the different forms (Fournais, r12 only, atomic, quasi-atomic)

#### 6.3. Results

TODO: N2 binding curve, some atomisation energies (I guess C2, N2 and CN are enough)

#### 6.4. Conclusion and Outlook

TODO: mention the shortcoming of not being able to target specific states like the previous methodology (except for quasi-atomic) TODO: mention the fournais/universal form does not faithfully capture long-range correlation TODO: mention possibility of combining methods, also using the orbital cusp correction for the Fournais factor TODO:

TODO: Also mention (maybe by then you even have data for) the Fournais Jastrow factor

# Summary and Outlook

TODO: ... TODO: brief review TODO: outlook: ECPs (if not already published), efficient deterministic optimisation, better understanding of the form of the Jastrow factor and other options, pytchint optimisation, larger molecules, solids, embedding, CUDA

1.1.	Illustration of an $s$ -type orbital and the fitted gaussians (STO- $x$ G means $x$ gaussians are used to approximate the STO). Note that for much of the curve, there is good agreement, but not for the short- and long-range. The short-range behaviour in particular leads to complications, and is the focus	
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# The PyTCHInt Library

This appendix is based on the following software, to be released:

Haupt, J. P.; López Ríos, P.; Christlmaier, E. M. C.; Filip, M.-A.; Simula, K.; Hauskrecht, J.; Liao, K.; Dobrautz, W.; Guther, K.; Cohen, A. J.; Alavi, A. *Pytchint: Transcorrelated Hamiltonian Integrals Library*, 2024

The basis for how the library works is partially discussed in the following paper and its supplementary material:

Cohen, A. J.; Luo, H.; Guther, K.; Dobrautz, W.; Tew, D. P.; Alavi, A. "Similarity Transformation of the Electronic Schrödinger Equation via Jastrow Factorization". Journal of Chemical Physics **2019**, *151*, 061101

### A.1. Introduction

All transcorrelated matrix element calculations presented in this dissertation have been performed using the group's tchint library or its Python extension, pytchint. This library works by either producing transformed integral files, dubbed FCIDUMP files for four-index integrals, or TCDUMP files for the TC six-index integrals. Note that since the TC transformation is non-Hermitian, we do not have the same symmetries in these files as we do with conventional methods, or by interfacing with another program (such as NECI) or the Python interpreter (in the case of pytchint).

As described in section 2.5,

$$\hat{H}_{TC} = \sum_{pq} h_q^p a_p^{\dagger} a_q + \frac{1}{2} \sum_{pqrs} \left( V_{rs}^{pq} - K_{rs}^{pq} \right) a_p^{\dagger} a_q^{\dagger} a_s a_r - \frac{1}{6} \sum_{pqrstu} L_{stu}^{pqr} a_p^{\dagger} a_q^{\dagger} a_r^{\dagger} a_u a_t a_s, \quad (A.1)$$

where

$$\begin{split} h_q^p &= \langle p|\,h\,|q\rangle\,,\\ V_{rs}^{pq} &= \langle pq|\,r_{12}^{-1}\,|rs\rangle\,,\\ K_{rs}^{pq} &= \langle pq|\,\hat{K}\,|rs\rangle\,,\\ L_{stu}^{pqr} &= \langle pqr|\,\hat{L}\,|stu\rangle\,. \end{split} \tag{A.2}$$

 $h_q^p$  and  $V_{rs}^{pq}$  are the one- and two-body terms from the electronic Schrödinger equation, familiar from conventional methods. Therefore, the key quantities to be evaluated by

tchint are the non-Hermitian two-body integrals  $K_{rs}^{pq}$  and the Hermitian three-body integrals  $L_{stu}^{pqr}$ .

Once these integrals are evaluated, as long as we take care about using the correct bra and ket (a detail not important for Hermitian problems), many different methodologies can be applied with these integrals, such as the frozen-core approximation,  $^{309-311}$  or  $^{xTC}$ .

### A.2. Matrix Element Evaluation

The matrix elements not present in conventional methods are  $K_{rs}^{pq}$  and  $L_{stu}^{pqr}$ . For these, we integrate on a grid, typically Treutler-Ahlrichs integration grids,  $^{299,300}$  which are atom-centred grids commonly used in density functional theory. These grids are obtained via pyscf.

The two-electron matrix elements we need to evaluate are:

$$K_{rs}^{pq(1)} = \langle pq | \nabla_1 u(\mathbf{r}_1, \mathbf{r}_2) \cdot \nabla_1 | rs \rangle \tag{A.3}$$

$$K_{rs}^{pq(2)} = \langle pq | \nabla_1^2 u(\boldsymbol{r}_1, \boldsymbol{r}_2) | rs \rangle \tag{A.4}$$

$$K_{rs}^{pq(3)} = \langle pq | (\nabla_1 u(\mathbf{r}_1, \mathbf{r}_2))^2 | rs \rangle. \tag{A.5}$$

Discretising on a grid of  $N_{grid}$  points with weights w, we have

$$K_{rs}^{pq(1)} = \sum_{mn}^{N_{\text{grid}}} \phi_p(\boldsymbol{r}_m) \nabla_{\boldsymbol{r}_m} \phi_r(\boldsymbol{r}_m) \cdot \nabla_{\boldsymbol{r}_m} u(\boldsymbol{r}_m, \boldsymbol{r}_n) \phi_q(\boldsymbol{r}_n) \phi_s(\boldsymbol{r}_n) w(\boldsymbol{r}_m) w(\boldsymbol{r}_n).$$
(A.6)

Naive integration of this value yields  $\mathcal{O}(N_{\mathrm{grid}}^2 M^4)$  performance (where M is the number of basis functions). However, we may improve this by first integrating over one coordinate and storing the intermediate value. Moreover, for  $K_{rs}^{pq(2)}$ , it is more efficient to integrate by parts. Therefore, for the full K matrix, we calculate the intermediate value

$$X_s^q(\mathbf{r}_2) = \int d^3r_1 \, \nabla_1 u(\mathbf{r}_1, \mathbf{r}_2) \cdot [\phi_p(\mathbf{r}_1) \nabla_1 \phi_r(\mathbf{r}_1) - \phi_r(\mathbf{r}_1) \nabla_1 \phi_p(\mathbf{r}_1)]$$
(A.7)

$$+ \int d^3r_1 \, \phi_p(\mathbf{r}_1) (\nabla u(\mathbf{r}_1, \mathbf{r}_2))^2 \phi_r(\mathbf{r}_1). \tag{A.8}$$

We can then obtain the K matrix by

$$K_{rs}^{pq} = \int d^3r_2 \ \phi_q(\mathbf{r}_2) X_s^q(\mathbf{r}_2) \phi_s(\mathbf{r}_2)$$
(A.9)

for a cost of  $\mathcal{O}(N_{\mathrm{grid}}^2 M^2 + N_{\mathrm{grid}} M^4)$ .

The three-body matrix,

$$L_{stu}^{pqr} = \langle pqr | \nabla_1 u(\boldsymbol{r}_1, \boldsymbol{r}_2) \cdot \nabla u_1(\boldsymbol{r}_1, \boldsymbol{r}_3) | stu \rangle$$
(A.10)

may similarly be resolved by calculating the intermediate value

$$\mathbf{Y}_{qt}(\mathbf{r}_1) = \int d^3 r_2 \, \phi_q(\mathbf{r}_2) \nabla_1 u(\mathbf{r}_1, \mathbf{r}_2) \phi_t(\mathbf{r}_2)$$
(A.11)

to give

$$L_{stu}^{pqr} = \int d^3r_1 \, \phi_p(\mathbf{r}_1) \mathbf{Y}_{qt}(\mathbf{r}_1) \cdot \mathbf{Y}_{ru}(\mathbf{r}_1) \phi_s(\mathbf{r}_1). \tag{A.12}$$

These integrations are simply parallelisable, and tchint takes advantage of this by leveraging distributed memory with the Message Passing Interface (MPI) $^{312-315}$  standard and the OpenMP $^{316}$  application programming interface (API). We also make liberal use of BLAS $^{317}$  routines for further performance gains. Moreover, we take advantage of the fact that L is Hermitian, substantially reducing storage requirements. We may also store L sparsely since many elements are small, or use the xTC approximation to bypass storing L altogether and instead calculating modified four-index integrals and outputting a non-Hermitian FCIDUMP.

#### A.3. Interface

tchint is predominantly written in Fortran, but most of the important interfacing subroutines (such as returning given matrix elements) have C bindings. This allows tchint to be easily interfaced with other libraries, notably for TC-FCIQMC in NECI<sup>267</sup> and M7.<sup>270</sup> By using this interface, these programs may be used to perform transcorrelated calculations, with all the features present in tchint.

tchint has also been given a Python interface called pytchint by making use of the Cython language extension. <sup>318–320</sup> Cython features C-like performance with easy integration in Python to allow for interactive computing and interfacing with the vast scientific package ecosystem available in Python. Python is typically also easier for rapid prototyping, a crucial feature in the world of scientific research, while Cython provides efficiency and a comprehensive profiling toolset.

With pytchint, a user might interactively work with transcorrelated integrals by first calculating them,

```
import pytchint
options = pytchint.options.TcOptions(yaml_input="tchint.yml")
options.eval_mode = "xtc"
solver = pytchint.solver.Solver(options=options)
solver.run()
```

and then e.g. query individual matrix elements calculated via Slater-Condon rules with solver.sltcnd or return the variance of the reference with solver.refconn.

#### A.3.1. Deterministic Optimisation

One of the major motivations for developing the Python interface is removing VMC from the Jastrow optimisation pipeline, as this can be a particularly expensive step, especially for larger systems. Thanks to the Python interface, we can interact directly with Python's large ecosystem of optimisation packages, such as SciPy.<sup>321</sup> This allows for rapid prototyping and ease of development.

For each optimisation step, we pass the Jastrow parameters in Python, then calculate the relevant integrals for the variance of the reference (see chapter 4) by passing them to Fortran, and update the parameters according to an optimisation algorithm by passing the parameters back to Python. All that is needed to be added are gradients for the variance of the reference with respect to Jastrow factor parameters. As the Jastrow factors are linear in their parameters, this is a tractible problem.

However, as calculations are done entirely in pytchint, we are restricted to a basis set and do not enjoy the CBS optimisation as we do in continuum VMC. As an example, we consider calculations with the Li atom. Consider, moreover, the simple parameter-free electron-electron-cusp-correcting Jastrow factor

$$J_0 = \frac{1}{2}\bar{r}_{ij} \tag{A.13}$$

with the additional terms for the electron-nucleus cusp described in chapter 4. Here,  $\bar{r}_{ij} = 1 - \exp(-\alpha r_{ij})/\alpha$  with  $\alpha = 2.0$ . Consider two possible additional terms, each with one variational parameter,

$$J_{ee} = c_{ee}\bar{r}_{ij}^2 \tag{A.14}$$

$$J_{en} = c_{en}\bar{r}_i^2 \tag{A.15}$$

where for the electron-nucleus term  $\bar{r}_i = 1 - \exp(-\beta r_{ij})/\beta$  with  $\beta = 2.0$ .

Reference energies and variance of the reference values are presented in figure A.1. As illustrated, the minimum may shift based on basis set, so a careful selection is necessary for obtaining the optimal Jastrow factor.

Preliminary results are promising, and this may prove an interesting alternative to VMC, or a way of "correcting" smaller VMC calculations, to improve efficiency as well as reproducibility.

# A.4. Parallel Efficiency

As a benchmark, we calculate  $N_2$  with the aug-cc-pVQZ basis (14 electrons in 160 orbitals) with the xTC approximation. Elapsed time is plotted as a function of the number of compute cores used in figure A.2, with favourable scaling. This illustrations the parallel performance of the pytchint library.

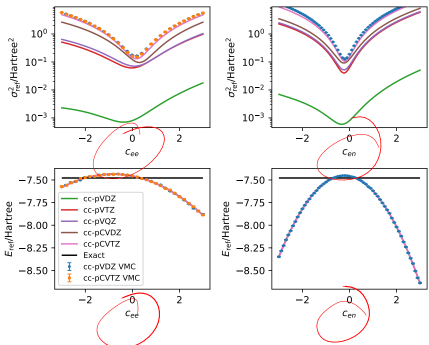


Figure A.1. Variance of the reference and reference energy values (top and bottom panels, respectively) with a single-parameter Jastrow factor, for the Li atom with the cc-pVXZ basis sets, with X = D,T,Q, as well as basis sets core-valence correlation at the double- and triple-zeta level. The left panel introduces an electron-electron term  $J_{ee}$  whereas the right panel introduces an electron-nucleus term  $J_{en}$ . The location of the minimum may be different compared to that obtained by continuum VMC, resulting in a suboptimal Jastrow factor. Data courtesy of Maria-Andreea Filip, who is leading this investigation. Results are preliminary and are intended to be presented in a future publication.

Moreover, thanks to the use of Cython, performance when directly running Fortran when compared to running via Python is roughly identical (468 seconds or 465 seconds with 128 cores for Fortran and Python, respectively).

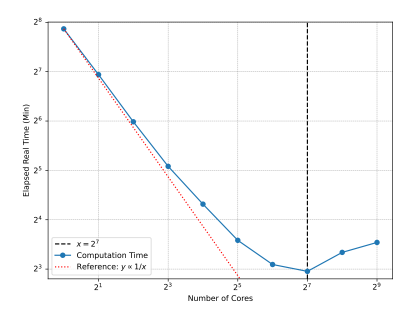


Figure A.2. Elapsed real time (or walltime) of calculating all-electron transcorrelated integrals under the xTC approximation for N<sub>2</sub> with the aug-cc-pVQZ basis. Note the logarithmic axes. Performance improves drastically when given additional cores, highlighting the parallel capabilities of the pytchint library. The dotted red line indicates ideal scaling. The dotted vertical line at 2<sup>7</sup> cores indicates additional nodes. Introducing more nodes, walltime slightly increases, indicating the additional MPI overhead of internode communication is not sufficiently compensated for by the additional cores for this size problem. However, it is worth noting that due to the distributed memory model, these calculations offer similar time performance but the memory load per node is reduced by roughly a factor equal to the number of nodes. Calculations were performed on AMD EPYC 9554 64-Core processors, and each node used 128 cores.

#### A.5. Conclusion and Outlook

Our in-house transcorrelated integral evaluation library is already flexible and performant, and is already used for most TC-related publications from the group. For wider adoption, we plan to improve documentation and ease of use through the Python interface, and have a form of Jastrow optimisation or (quasi-)universal Jastrow form such as those described in chapter 6. This way, pytchint is a standalone library and would not rely on external software.

Additional features we intend to investigate in the near future are: parallism with graphics processing units (as our integrals are easily parallelisable), which is already being

investigated in the context of TC, further support for periodic solids, and implementing more of the code in high-performance Python to allow easier use of additional features such as automatic differentiation  $^{322}$  with JAX,  $^{323}$  which would allow support for more complex Jastrow factors without the need for manually calculating analytic gradients.

# Curriculum vitae

#### Education

2020 - 2025 PhD student, Theoretical Chemistry

Max Planck Institute for Solid State Research, Stuttgart, Germany **Thesis:** Development of the Transcorrelated Full Configuration

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Computer Science

University of British Columbia, Vancouver, Canada

**Thesis:** Effects of Anisotropic Long-Range Hopping on Anderson

Localization

Advisor: Prof Roman Krems

#### **Publications**

TODO: Not really sure if there is a point to including a publications list when so few are actually published...

#### First Author

TODO: publications, I guess: optimization paper, binding curve paper, pytchint paper (likely with a "to be submitted"), universal jastrows is probably a long way away

- Haupt, J. P.; Hosseini, S. M.; López Ríos, P.; Dobrautz, W.; Cohen, A.; Alavi, A. "Optimizing Jastrow Factors for the Transcorrelated Method". The Journal of Chemical Physics 2023, 158, 224105
- (2) Haupt, J. P.; Ríos, P. L.; Christlmaier, E. M. C.; Kats, D.; Alavi, A. "The Transcorrelated Method for Strongly Multireference Problems". **2024**, To be submitted
- (3) Haupt, J. P. et al. Pytchint: Transcorrelated Hamiltonian Integrals Library, 2024

#### Coauthor

TODO: publications, I guess: ECPs, 2nd row atoms, deterministic optimisation, spin-dependent Jastrows TODO: Pablo's surname doesn't appear correctly in some of these

- (4) Filip, M.-A.; Ríos, P. L.; Haupt, J. P.; Christlmaier, E. M. C.; Kats, D.; Alavi, A. "Transcorrelated Methods for Second Row Elements". 2024
- (5) Simula, K.; López Ríos, P.; Christlmaier, E. M. C.; Haupt, J. P.; Alavi, A. "Effective Core Potentials in the Transcorrelated Method". 2024