

H_3 as an Example for Open Shell Systems: CIS with UHF and CUHF references

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1 Introduction

In computational chemistry, the time-independent Schrödinger equation plays a central role.

$$\hat{H}\Psi = E\Psi \quad (1)$$

All the information with regard to a certain molecule are linked to the wave function Ψ . This information must be gathered using quantum mechanical operators. In the Schrödinger equation we calculate this wave functions energy E using an operator known as the Hamiltonian. In its full form it can be seen in the equation below [1].

$$\hat{H} = -\sum_{i=1}^N \frac{1}{2} \nabla_i^2 - \sum_{A=1}^M \frac{1}{2M_A} \nabla_A^2 - \sum_{i=1}^N \sum_{A=1}^M \frac{Z_A}{r_{iA}} + \sum_{i=1}^N \sum_{j>i}^N \frac{1}{r_{ij}} + \sum_{A=1}^M \sum_{B>A}^M \frac{Z_A Z_B}{R_{AB}} \quad (2)$$

The Hamiltonian contains the kinetic energy of the N electrons i and the M nuclei A . It also accounts for the electrostatic potential between the nuclei and the electrons. Also note that this Hamiltonian was written down using atomic units. Unfortunately, the Schrödinger equation is not exactly solvable for systems that contain more than one electron, i.e. almost all chemically interesting systems. The reason for this is the repulsion that exists between two electrons. This term makes it impossible to solve the Schrödinger equation analytically for most systems. That is why we need to make approximations. In this paper, we will make use of the Hartree-Fock method combined with Single Excitation Configuration Interaction. In the Hartree Fock method, we use a single Slater determinant wave function that is an eigenfunction of the Fock operator [1].

$$\hat{f} = \hat{h}(1) + \sum_j^N \hat{J}_j - \hat{K}_j \quad (3)$$

The operators \hat{J}_j and \hat{K}_j are called the Coulomb and exchange operators respectively. The Coulomb operator accounts for interelectronic repulsion, while the exchange operator accounts for the stabilization that occurs between electrons of the same spin.

We can define an expectation value over the Hamiltonian and state that if we are in an energy minimum, a small variation in the wave function will not induce a change in energy.

$$\langle \psi + \delta\psi | \hat{H} | \psi + \delta\psi \rangle = E + \delta E + \delta E^* + \delta E^2 \quad (4)$$

In first order we say that if the energy of the wave function ψ is a minimum the wave function can be varied slightly without changing the expectation value over the Hamiltonian, so $\delta E = 0$. This approximation might take us a long way in the right direction, but does it take us all the way there? Here we need to introduce different methods within the Hartree-Fock theory. Restricted Hartree-Fock (RHF) forces the electrons into the same orbitals, thus creating doubly occupied orbitals. Even though this is a valid picture in organic chemistry, since we think about electrons in pairs, physically there is no obligation for the electrons to exist in these pairs. When we let go of the restriction, we enter the Unrestricted Hartree-Fock method (UHF). This is no longer confined to closed shell species since it has no problem with electrons that do not exist in pairs. However, we see that the wave function in this method does not contain the right symmetries. We can get to a lower energy since we give the wave function some extra freedom, yet the chemical intuition is tossed out of the window.

A molecule can have a lot of symmetry elements, simple examples could be rotational axes or mirror planes. This symmetry is reflected in that molecule's Hamiltonian and its wave function. Looking back at equation (4) and the constraint we applied, namely that δE should remain zero, we cannot state that our approximative Hartree-Fock wave function still abides by these symmetries. We can impose some additional constraints in order to get the right symmetries, but there is a cost involved. This leads us to Löwdin's symmetry dilemma [2]. On one side we can enforce the symmetry. This will lead us to a wave function that is chemically intuitive, but the energy we reach will be relatively high. We can also expect to keep the quantum numbers, such as S , associated with the wave function. On the other hand we can drop the symmetry. This will lead us to a lower energy wave function, but we will lose information about the spin quantum numbers.

We need some kind of theory to bring the best of both worlds together. On one side, we have the chemical intuition in the restricted wave functions, on the other the lower energy in the unrestricted wave functions. Roothaan's Restricted Open Shell theory (ROHF) [3] was an early attempt. As this approach is tedious and hard to implement, it is not very popular [4, 5]. One of the drawbacks in ROHF method is that the Fock operator is not unambiguously defined [4, 6]. UHF is rather easy to implement [4, 5], but as we stated above, there is a cost to this method. However, more recently Tsuchimochi and Scuseria presented their Constrained Unrestricted Hartree-Fock theory (CUHF) [4]. In CUHF we apply a constraint to the system that sets the spin contamination to zero (cfr. *infra*). This is because spin contaminated wave functions are no good starting point for post Hartree-Fock methods [4]. A lot has been said already about this method; Scuseria and Tsuchimochi themselves expanded on it in a follow-up paper [7]. Plakhutin and Davidson discussed some defects in the theory with respect to Koopmans theorem and ionisations [6]. In this work we will use the CUHF

wave function as a reference for Single Excitation Configuration Interaction (CIS). We will generate the single excitation energies for the CUHF wave function and compare them to the CIS excitation energies using ROHF and UHF references of the same system.

In this work we will start by introducing spin-contamination and the theory behind it. We will also give an overview of CUHF theory. Then we will discuss CIS and derive some important equations. Next we will look at spin-contamination in an actual system to stress the importance of uncontaminated spin. Then we will discuss the CIS results for CUHF, ROHF and UHF.

2 Theory

2.1 Spin-Contamination

In general one can define the expectation value of \hat{S}^2 like equation (5) [8].

$$\langle \hat{S}^2 \rangle = S_z^2 + S_z + q - \sum_{ij}^{pq} S_{ij}^2 \quad (5)$$

This equation is written down for a system with p α electrons with summation index i and q β electrons with summation index j . The S matrix contains the overlap between orbitals in the Molecular Orbital (MO) basis. S_z is the eigenvalue of the \hat{S}_z operator. When the overlap is diagonal in this MO basis, the last two terms vanish, and we are left with $S_z(S_z + 1)$. Spin-contamination is then defined as equation (6).

$$\delta_s = \langle S^2 \rangle - S_z(S_z + 1) \quad (6)$$

In UHF the overlap is not diagonal, so the overlap matrix between the α and β orbitals will not be diagonal. One can not expect the last two terms in equation (5) to vanish, which will result in δ_s not being equal to zero. This also means that the wave function is no longer an eigenfunction of \hat{S}^2 , as we have stated before. We lose a quantum number and with it the symmetry of the system. We can rewrite spin-contamination as equation (7) [9].

$$\delta_s = N_\beta - Tr(\gamma^\alpha \gamma^\beta) \quad (7)$$

In this equation, γ^α and γ^β are the one particle density matrices. In the Natural Orbital (NO) basis, these matrices are block diagonal [4], with blocks that can be seen in equations (8a) and (8b), where i corresponds to a certain closed shell orbital. When there are more α electrons than β electrons, the γ^α also contains a block that is a unity matrix with dimensions $n \times n$ where n is the amount of singly occupied orbitals. In γ^β this block is a zero matrix of the same dimensions. For clarity, we will always assume that the amount of α electrons is larger than the amount of β electrons. The results hold for any system where $N_\alpha \geq N_\beta$ [4].

$$\gamma_i^\alpha = \begin{bmatrix} n_i & m_i \\ m_i & 1 - n_i \end{bmatrix} \quad (8a)$$

$$\gamma_i^\beta = \begin{bmatrix} n_i & -m_i \\ -m_i & 1 - n_i \end{bmatrix} \quad (8b)$$

These matrices can now be written in their full form, as can be seen in equations (9a) and (9b)

$$\gamma^\alpha = \begin{bmatrix} \gamma_1^\alpha & \cdots & 0 & 0 & 0 \\ \vdots & \ddots & \vdots & \vdots & \vdots \\ 0 & \cdots & \gamma_{N_{cp}}^\alpha & 0 & 0 \\ 0 & \cdots & 0 & I_n & 0 \\ 0 & \cdots & 0 & 0 & 0 \end{bmatrix} \quad (9a)$$

$$\gamma^\beta = \begin{bmatrix} \gamma_1^\beta & \cdots & 0 & 0 & 0 \\ \vdots & \ddots & \vdots & \vdots & \vdots \\ 0 & \cdots & \gamma_{N_{cp}}^\beta & 0 & 0 \\ 0 & \cdots & 0 & 0 & 0 \\ 0 & \cdots & 0 & 0 & 0 \end{bmatrix} \quad (9b)$$

I_n is noted as the unity matrix in n dimensions, where n is the number of unpaired electrons. We have N_{cp} closed shell orbitals, each corresponding with a doubly occupied orbital. In our assumption that there are more α electrons than β electrons, N_{cp} is also equal to the amount of β electrons. In these equations n_i is the occupation of orbital i and $m_i = \sqrt{n_i - n_i^2}$ [4]. We can then define the spin density matrix $\mathbf{M} = (\gamma^\alpha - \gamma^\beta)/2$. The matrix subtraction leaves us with blocks in the \mathbf{M} matrix that correspond to equation (10).

$$M_i = \begin{bmatrix} 0 & m_i \\ m_i & 0 \end{bmatrix} \quad (10)$$

Of course, the open shell block will now consist of $\frac{1}{2}I_n$. Using this information we can rewrite equation (7) to equation (11) [4].

$$\delta_s = N_\beta - (N_\alpha + N_\beta)/2 + 2 * Tr(M^2) \quad (11)$$

We can indeed verify that this is true by working out the matrix product as was done in proof 2.1.

Proof 2.1. To a system with N_e electrons of which $2N_{cp}$ are in the closed shell and N_s are in the open shell applies:

$$\begin{aligned} Tr(\gamma^\alpha \gamma^\beta) &= \sum_i^{N_{cp}} (-4m_i + 1) \\ &= N_{cp} - \sum_i^{N_{cp}} 4m_i \\ &= \frac{N_e - N_s}{2} - \sum_i^{N_{cp}} 4m_i \\ &= \frac{N_e}{2} - 2Tr(M^2) \end{aligned}$$

□

In an additional step, we can use algebra to find equation (12). This equation can also be derived from proof 2.1.

$$\delta_s = 4 \sum_i^{N_{cp}} m_i^2 \quad (12)$$

Now if we want to make the spin contamination zero, we have to enforce double occupation or no occupation at all. We see here that in the closed shell itself, we can no longer have a difference between α and β orbitals. This will be the constraint that gives us the CUHF method. We could make the observation that in Roothaan's original paper, the start was very similar, as can be seen in equation (13) [3].

$$\Phi = (\phi_C, \phi_O) \quad (13)$$

Roothaan starts his exposition on open shell systems with this equation, representing the total wave function Φ as a combination of a closed shell system ϕ_C and an open shell system ϕ_O

2.2 Constrained Unrestricted Hartree-Fock

When talking about CUHF, there is more than one way to make sure that the spin-contamination remains zero. Scuseria and Tsuchimochi will enforce the constraint in the way we mentioned in the last section, where they allow only doubly occupied orbitals in the closed shell [4]. It can be shown that the normal UHF energy can be constructed by adding a correlation energy term to the closed shell energy. These terms can be seen in equations (14a) and (14b) [9].

$$E_{cs} = 2 \sum_{ij} h_{ij} P_{ij} + \sum_{ijkl} (2\langle ij|kl\rangle - \langle ij|lk\rangle) P_{ik} P_{jl} \quad (14a)$$

$$E_c = - \sum_{ijkl} \langle ij|lk\rangle M_{ik} M_{jl} \quad (14b)$$

Where the matrix \mathbf{M} is the spin density matrix (cfr. supra) and the \mathbf{P} matrix is the charge density matrix: $\mathbf{P} = (\gamma^\alpha + \gamma^\beta)/2$. If we take the derivative with respect to the density matrices we find equations (15a) and (15b).

$$\frac{\partial E_{cs}}{\partial \gamma_{ij}^\alpha} = \frac{\partial E_{cs}}{\partial \gamma_{ij}^\beta} = F_{ij}^{cs} \quad (15a)$$

$$-\frac{\partial E_c}{\partial \gamma_{ij}^\alpha} = \frac{\partial E_c}{\partial \gamma_{ij}^\beta} = \sum_{kl} \langle ik|lj\rangle M_{kl} = \Delta_{ij}^{UHF} \quad (15b)$$

So in fact, the UHF Fock operators can be written as a sum of this Δ^{UHF} and the closed shell Fock operator, as displayed in equations (16a) and (16b) [4].

$$F^\alpha = F_{cs} - \Delta^{UHF} \quad (16a)$$

$$F^\beta = F_{cs} + \Delta^{UHF} \quad (16b)$$

Now we need to expand this concept to CUHF. In general, we want to be able to write the CUHF Fock matrix \tilde{F} as the closed shell Fock matrix to which we add a certain Δ^{CUHF} . As we have already seen, the constraint is only applied to the virtual and closed shell orbitals. When we display the Fock matrix in MO basis it will exist in blocks, equation (17). The c, o and v stand for closed shell, open shell and virtual respectively.

$$\mathbf{F} = \begin{bmatrix} F_{cc} & F_{co} & F_{cv} \\ F_{oc} & F_{oo} & F_{ov} \\ F_{vc} & F_{vo} & F_{vv} \end{bmatrix} \quad (17)$$

We apply the constraint in the vc and cv blocks. We can do this by stating that $[\tilde{F}^\sigma, \gamma^\sigma] = 0$, where σ can equal either α or β . This basically means that the density matrices should share a complete set of eigenfunctions with the Fock matrices. This, in turn, means that the Fock matrices should be diagonal at self-consistent field (SCF) conditions in the NO basis. By subtracting the two conditions and algebraically working out the result we find that at SCF conditions Δ_{cv}^{CUHF} has to be equal to zero. This means that $\tilde{F}_{cv} = F_{cv}^{cs}$ [4]. For every other block, the CUHF Fock matrix equals the UHF Fock matrix, so the Δ^{CUHF} equals the Δ^{UHF} . When we enforce this constraint during every iteration, we will decrease the spin contamination. This does entail that we will need to transform the Δ^{UHF} to NO basis every time to enforce this constraint. This transformation will consist of two steps. First we need to transform the density matrix to an orthonormal basis, since AO basis is often not orthonormal. (Indeed, if it were, we would not expect any bonds to form.) We choose MO basis. The MO's are defined by the eigenvectors of the Fock operator, we will use the closed shell Fock operator. We start from the the Roothaan equation.

$$F_{cs}C = \epsilon SC \quad (18)$$

With a simple transformation, we can see that the Fock operator will become diagonal using the transformation in equation (19)

$$C^T F C = \epsilon C^T S C \quad (19)$$

This is true, since $C^T S C$ is the unity matrix if the C matrix holds the eigenvectors of the generalized eigen value problem. To transform to NO basis, we will also need the density matrix in the MO basis. This is not trivial, since the density matrix is usually defined by the coefficient matrices C. To transform the density matrix P , we will need to do a slightly different transformation. This transformation is given in equation (20) [10].

$$P^{MO} = C^{-1}P^{AO}(C^{-1})^T \quad (20)$$

We can then calculate the eigenvalues and subject the Δ^{UHF} to these transformations (21), we have easy access to the *cv* and *vc* blocks.

$$\Delta_{NO}^{UHF} = V^{-1}C^T \Delta_{AO}^{UHF} CV^{-1} \quad (21)$$

The V matrix contains the coefficients of the eigenvectors of the P matrix. Now we just switch the required blocks to zero, back transform and add the now Δ to the closed shell Fock matrix. To summarize this procedure we can look at the form of the Fock matrices in equations (22a) and (22b).

$$\tilde{F}^\alpha = F_{cs} - \Delta^{CUHF} \quad (22a)$$

$$\tilde{F}^\beta = F_{cs} + \Delta^{CUHF} \quad (22b)$$

Where Δ^{CUHF} is either zero or Δ^{UHF} depending on whether you are looking at the *cv* blocks or at the other blocks. Here we can also clearly see the difference with ROHF theory. In his original paper, Roothaan goes through great lengths to ensure that there is only one Fock matrix instead of separate ones for the α and β orbitals [3]. This makes it rather difficult to interpret results [4]. However, Scuseria and Tsuchimochi keep the separate matrices, like in a normal UHF approach. This means that we will end up with a slightly different picture from the ROHF picture, but the wave function and its energy remain the same [4].

2.3 Configuration interaction

The Hartree-Fock approximation gives us an initial guess for the energy and the wave function. However, it can be improved upon. The methods that do this are called post Hartree-Fock methods. The difference in energy between the normal Hartree-Fock method and the post Hartree-Fock method is called the correlation energy (23).

$$E_{corr} = \mathcal{E}_0 - E_0 \quad (23)$$

Here E_0 is the Hartree-Fock energy and \mathcal{E}_0 it the post Hartree-Fock energy. There are various approaches to post Hartree-Fock methods. We will discuss the Configuration Interaction

method(CI). If Φ_0 is the result of normal Hartree-Fock procedure, it is an approximation of the true wave function Γ . We know that we can make a better approximation using a linear combination of excited states that start from Φ_0 [1]. If Φ_0 is the Slater determinant that corresponds to the ground state of the system, we mix it with the excited states. Each determinant represents a certain configuration of the electrons. These configurations interact, as can be seen in the equation below.

$$|\Psi_0\rangle = c_0|\Phi_0\rangle + \sum_{ar} |\Phi_r^a\rangle + \sum_{a<b, r<s} c_{ab}^{rs} |\Phi_{ab}^{rs}\rangle + \dots \quad (24)$$

This wave function Ψ_0 is a better approximation to the real wave function Γ than Φ_0 . When we try to get the energy expectation value of this function, we can write it in matrix form, where each row and column correspond to a certain excited state. The full expression of this matrix is given below.

$$\mathbf{H} = \begin{bmatrix} \langle \Phi_0 | \hat{H} | \Phi_0 \rangle & \langle \Phi_0 | \hat{H} | \Phi_r^a \rangle & \dots & \langle \Phi_0 | \hat{H} | \Phi_{\{\chi_o\}}^{\{\chi_v\}} \rangle \\ \langle \Phi_r^a | \hat{H} | \Phi_0 \rangle & \langle \Phi_r^a | \hat{H} | \Phi_r^a \rangle & \dots & \langle \Phi_r^a | \hat{H} | \Phi_{\{\chi_o\}}^{\{\chi_v\}} \rangle \\ \vdots & \vdots & \ddots & \vdots \\ \langle \Phi_{\{\chi_o\}}^{\{\chi_v\}} | \hat{H} | \Phi_0 \rangle & \dots & \dots & \langle \Phi_{\{\chi_o\}}^{\{\chi_v\}} | \hat{H} | \Phi_{\{\chi_o\}}^{\{\chi_v\}} \rangle \end{bmatrix} \quad (25)$$

For a system with n occupied orbitals in the ground state, denoted with indices (r, s) and m unoccupied orbitals, denoted with indices (a, b) , the full CI matrix extends until every possible electron configuration has been accounted for. $\Phi_{\{\chi_o\}}^{\{\chi_v\}}$ is a state in which a set of m electrons is excited. In the ground state, these electrons occupy the set of orbitals χ_o , and they are excited to the set of virtual orbitals χ_v . We now have to determine what all the different matrix elements are. Since we are looking at CIS, we are only interested in the single excitations, so the matrix in equation (25) can be simplified quite a bit. All elements that contain a higher order excitation are set to zero. Then we already know that the expectation value of the Hartree-Fock wave function is the energy we can get from our normal Hartree-Fock calculation, so we can write this equation.

$$\langle \Phi_0 | \hat{H} | \Phi_0 \rangle = E_0 \quad (26)$$

When we look at single excitations, so at states Φ_r^a where the electron from orbital r has been excited to orbital a , we can write equation (27) [1] for the overlap between the ground state wave function and the excited states.

$$\langle \Phi_0 | \hat{H} | \Phi_r^a \rangle = \langle a | h | r \rangle + \sum_b \langle ab || rb \rangle = \langle \chi_a | \hat{f} | \chi_r \rangle = F_{ra} \quad (27)$$

Here $\langle ab || rb \rangle$ is shorthand notation for $\langle ab | rb \rangle - \langle ab | br \rangle$. The letter a represents the orbital χ_a . Now of course, by definition, at SCF conditions the off-diagonal elements of the Fock matrix are zero. This leads to Brillouin's theorem. It states that there is no overlap between single excited states and the ground state wave function when taking the expectation value over the Hamiltonian. This means that the ground state energy cannot be improved upon by doing a CIS calculation. Indeed, one can prove equation (28) [1]. The correlation energy is only defined by the overlap between second order excited states.

$$E_{corr} = \sum_{i < j, r < s} C_{ij}^{rs} \langle \Phi_0 | \hat{H} | \Phi_{ij}^{rs} \rangle \quad (28)$$

However, this does not mean that an excited state can not have a lower energy than the ground state. This can be understood by looking again at equation (4) and only considering columns and rows that contain the ground state or a single excitation. What we have now is in fact a block diagonal matrix. One block contains the ground state energy, so its dimensions are 1×1 . The other block contains all the excited state expectation values and overlap elements. It is a property of block diagonal matrices that their eigenvalues are the eigenvalues of the individual blocks. We can thus conclude that the ground state energy will be among the eigenvalues of this matrix. There is nothing that constrains the eigenvalues of the other block, so they might well be lower than the ground state energy. One can imagine a situation where the geometry of the molecule has not been optimized, in which it might be favorable to be in an excited state rather than in the ground state.

In CIS, we do not expect to lower the ground state energy. However, it can give us information about the excited states and the excitation energies that come with them. Now we are interested in the expectation values over the Hamiltonian of all the excited states, so we need an expression for $\langle \Phi_r^a | \hat{H} | \Phi_s^b \rangle$. There are four different situations that we need to discuss here [11]. The first scenario is an excitation where $a \neq b$ and $r \neq s$. The only term that will remain in that case is $\langle aj || ib \rangle$. Another case is when $r = s$ but $a \neq b$. Then the matrix element can be expressed using equation (29).

$$\langle \Phi_r^a | \hat{H} | \Phi_r^b \rangle = h_{ab} + \sum_{k \in r} \langle ak || bk \rangle = F_{ab} - \langle ar || br \rangle \quad (29)$$

Another case, where $r \neq s$ and $a = b$ we can write equation (30).

$$\langle \Phi_r^a | \hat{H} | \Phi_s^a \rangle = -h_{rs} - \sum_{k \in \{\Psi_0 + a\}} \langle ak || bk \rangle = -F_{rs} - \langle ra || sa \rangle \quad (30)$$

Finally, when both $a = b$ and $r = s$ we can find equation (31).

$$\begin{aligned} \langle \Phi_r^a | \hat{H} | \Phi_r^a \rangle = & \sum_{k \in \Psi_0} h_{kk} + \frac{1}{2} \sum_{k, l \in \Phi_0} \langle kl || kl \rangle - h_{rr} + h_{aa} \\ & - \sum_{k \in \Phi_0} \langle kr || kr \rangle + \sum_{k \in \Phi_0} \langle ka || ka \rangle - \langle ra || ra \rangle \end{aligned} \quad (31)$$

We can pull all these equations into one general equation (32).

$$\langle \Phi_r^a | \hat{H} | \Phi_s^b \rangle = E_0 \delta_{rs} \delta_{ab} + F_{ab} \delta_{rs} - F_{rs} \delta_{ab} + \langle as || jb \rangle \quad (32)$$

We notice that E_0 is added along the diagonal. Since we know that the trace of a matrix is equal to the sum of its eigenvalues, we can subtract E_0 from the diagonal and just add it to every eigenvalue at the end. At self-consistent field conditions the Fock operator equals $F_{ij} = \epsilon_i \delta_{ij}$, so equation (32) can be simplified to equation (33).

$$\langle \Phi_r^a | \hat{H} | \Phi_s^b \rangle = (\epsilon_a - \epsilon_r) \delta_{rs} \delta_{ab} + \langle as || jb \rangle \quad (33)$$

The ϵ_i values correspond to the orbital energies. If we now diagonalize the CIS matrix we end up with the excitation energies and contributions of the single excited states as eigenfunctions. However, since electrons in α orbitals can be excited to β orbitals, which is especially important in UHF, we have to do some extra steps. Indeed, when calculating the eigenfunctions of the Fock matrices we get separate coefficient matrices for α and β orbitals. We will need all these orbitals together to get the complete CIS picture. We will have to effectively double the dimensions of the problem. This becomes a bit clearer by looking at equation (32). In this equation for example a and b do not have the same spin. However, the expression F_{ab} implies that there is an overlap element in the Fock matrix between two orbitals that can have a different spin. We can see that the normal α and β Fock matrices will not be enough. We will need a Fock matrix that accounts for both α and β electrons, and has the respective orbitals as eigenfunctions. A block diagonal matrix like equation (34) is a logical choice. Its eigenvalues will be the combination of the sets of eigenvalues of the blocks, the same goes for its eigenvectors.

$$F = \begin{bmatrix} F^\alpha & 0 \\ 0 & F^\beta \end{bmatrix} \quad (34)$$

When looking at equation (32) we see that we also need the so-called two electron integrals. This is in fact a four dimensional matrix that holds the interelectronic repulsion in AO basis. We also need to double its dimensions in order to use it for CIS calculations. Since this is an $n \times n \times n \times n$ matrix, we will need to use the Kronecker product.

$$R' = I_2 \otimes (I_2 \otimes R)^T \quad (35)$$

The R' matrix now has double the dimensions of the original R matrix. R in this case is the two electron overlap matrix. This is necessary for the two electron integrals, since they only contain half the orbitals. These integrals are in the original basis, but we need to account for all possible excitations, meaning that we have to account for alpha and β orbitals at the same time. Since the two electron integrals are given in AO basis, they still need to be transformed to MO basis, since we want the Fock operators to be diagonal in equation (32). To do this transformation we will need the coefficient matrix that holds the eigenfunction of the matrix in equation (34).

3 Methodology

The RHF, UHF and CUHF algorithms were self-implemented. The following iterative algorithm was applied for (C)UHF:

1. Calculate the energy based on the current guess matrices (the core Hamiltonian for the first iteration). There is a different guess matrix for α and β electrons.
2. Calculate a new Fock matrix for both α and β electrons based on the eigenfunctions of the current guess matrices.
3. In CUHF we now manipulate the Fock matrices as described in the 2.2 section. In UHF this step is skipped
4. Set the new Fock matrices as new guess matrices.
5. Check for convergence. This was done using an energy criterion with a threshold of 10^{-12} Hartree.

The CIS algorithm was implemented using equation (33) to create the Hamiltonian matrix elements. Then a block was added containing only the expectation value of the reference. The matrix was then diagonalized. We used the packages NumPy [12], SciPy [13] and Psi4 [14]. NumPy and SciPy deliver tools for matrix manipulation and other linear algebra calculations that we had to do. Psi4 is an open source quantum chemistry package allows us to calculate certain properties of the system given a molecular geometry. These properties include nuclear repulsion, kinetic energy of the electrons among others. It also allows us to generate reference data for the molecular energy in RHF, UHF and CUHF. It also features a method for calculating ROHF energies. They also present the possibility to do configuration interaction for RHF and ROHF. The CIS reference data for UHF and CUHF were generated using the GQCP [15] library. We have also visualized some orbitals using Psi4's cubeprop function to generate a .cube file. This .cube file was then visualized using the IQmol software [16].

To demonstrate the impact of spin contamination and the importance of finding methods to overcome it, we will discuss the hydrogen gas dissociation problem [1]. Hydrogen gas is a molecule that has an RHF solution in its equilibrium geometry. However, when we stretch the bond the system evolves to a state of separate hydrogen atoms. For CIS H_3 will be used as a model system. Firstly because it is a simple system containing few electrons. This means that we do not need to track a lot of possible excitations, so the focus can shift towards the interpretation. For this same reason we will pick the STO-3G basis set. This set provides us with the minimal amount of basis functions needed to describe the system. This again makes interpretation the key point. We will also do a stretch of H_3 in the following way. Since the molecule is an equilateral triangle, we will stretch all the bonds equally. Then we will compare the CIS results for H_3 as generated by our implementation and the one used by Psi4 and GQCP. GQCP will generate the data for CUHF and UHF and Psi4 will generate the data for ROHF.

4 Results and discussion

4.1 H₂ Stretch

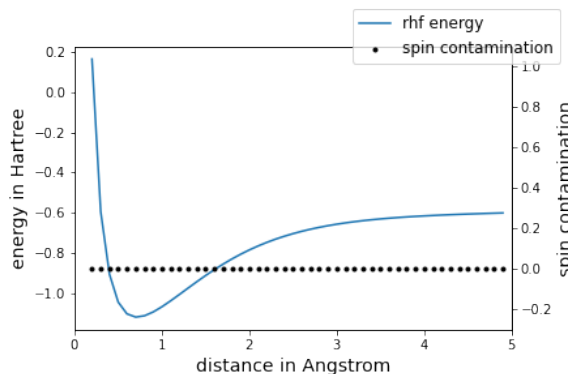


Figure 1: The RHF energy against the distance in an H₂ stretch. The zero level was chosen as the energy of two separate hydrogen atoms.

In figure 1 we have conducted a stretch of the H₂ molecule using an RHF calculation. We can see that the RHF energy does not behave as we would expect it to. Indeed, we expect the energy to evolve towards the energy of two separate hydrogen atoms, which clearly does not happen. This is because the physical picture we are seeing is intrinsically invalid. The RHF algorithm forces the electrons into the same orbital, the bonding orbital between the two nuclei. However, when the bond length increases, we expect the electrons to each be in their own respective orbitals around a single nucleus. At a distance of infinity, they should find themselves in a normal hydrogen s orbital. The RHF description is too limited and does not allow this. Worth noting is that the spin contamination remains zero all the way. This means that the wave function is an eigenfunction of the \hat{S}^2 [7].

We want to improve this image by using UHF. When we drop the constraint that the α and β orbitals must be identical we can indeed expect the energy to be equal to two separate hydrogen atoms. But since H₂ is a closed shell system we need to do a little manipulation before starting the algorithm. When we would do a core guess, so start from the molecular Hamiltonian as an approach for two Fock matrices, we would still find the restricted solution. To counteract this we will need to start from a mixed guess. For this we change the initial guess for the β electrons by mixing the Highest Occupied Molecular Orbital (HOMO) and the Lowest Unoccupied Molecular Orbital (LUMO) as a linear combination. This process is also known as orbital mixing. The result can be seen in figure 2.

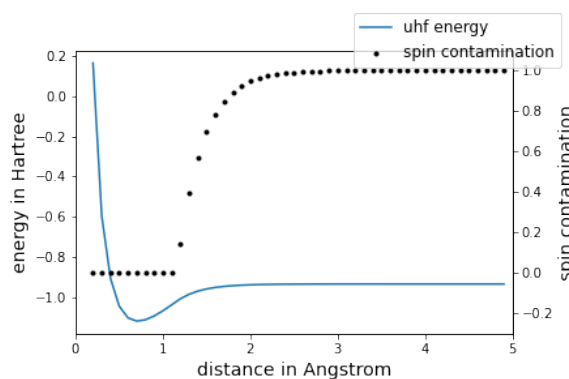


Figure 2: The UHF energy against the distance in an H_2 stretch.

This result is physically more accurate. We see that the system does evolve towards two separate hydrogen atoms. However, the spin contamination is rising, and after a certain point reaches the value one. This means that the wave function does no longer have the right symmetry. And as stated before, even if we cannot calculate the exact wave function we do know what symmetries it should have. Even though we might have a lower energy, the wave function is not as good as an approximation as it could be [17].

This is a prime example of the aforementioned symmetry dilemma by Löwdin. If we want the lowest possible energy, we lose the symmetry. If we want to keep the symmetry, we will not have the lowest energy. Now what would happen if we applied CUHF to the dissociation problem? This result has been plotted in figure 3.

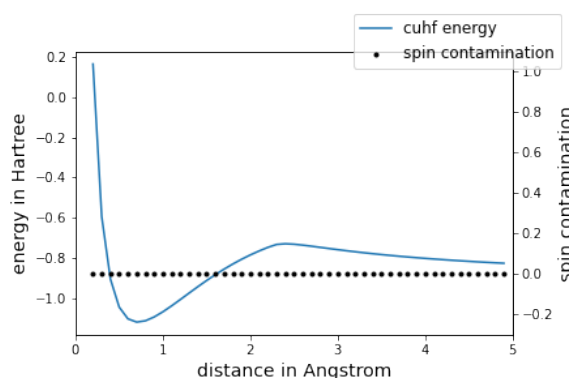


Figure 3: The CUHF energy against the distance in an H_2 stretch.

The CUHF picture is difficult to interpret. We see that the spin contamination remains zero, as can be expected. The energy seems to follow the RHF pattern (see also figure 4) until

a certain point, then it starts to exponentially decrease. Physically this seems like a logical behavior, as it is now again evolving to two separate hydrogen atoms. We can also notice that the spin contamination is zero everywhere, so the symmetry of the solution is also correct. For this plot a mixed guess was applied. In CUHF the mixed guess is done on the active space, which consists the orbitals to which the constraints are not applied, i.e. the open shell [7]. So when we would take the active space as all the electrons we reach the UHF solution. When we take only unpaired electrons as the active space expect to see the ROHF solution.

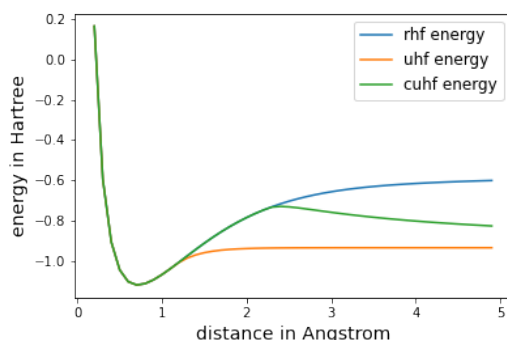


Figure 4: The stretch energies plotted together.

For comparison, we plotted all the stretches in one plot, figure 4. Here we can see that they all describe short bond lengths correctly, but as the internuclear distance increases, there are deviations visible. As one would expect, the CUHF energy is able to describe short bond lengths as correctly as both RHF and UHF. While RHF deviates from the true physical value at long bond distances, we see that CUHF does follow the expected limits (going to zero at infinity) while keeping the symmetry intact.

4.2 H_3 stretch

We will now discuss our model system, the trihydrogen molecule. This system is a ring of three hydrogen atoms, forming an equilateral triangle. Since UHF does not have a restricted solution, we do not need to do a mixed guess. This system is an open shell system, as stated before, with two α electrons and one β electron. It can not be described using RHF, since RHF needs closed-shell systems to work properly. For UHF, we have plotted the result in figure 5.

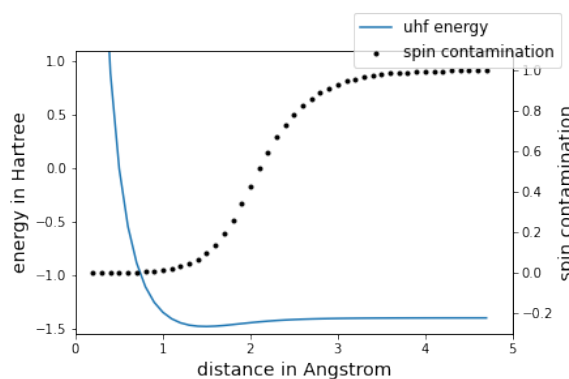


Figure 5: The stretch for H_3 using UHF.

We can see that the energy reaches the limiting case that we want, however the spin contamination is rising. In this figure, the last two data points were not plotted since convergence could not be reached. We can see that the molecule disintegrates into the three separate hydrogen atoms. After that, the energy reaches the level of three separate hydrogen atoms. We can also take a look at the CUHF result in figure 6.

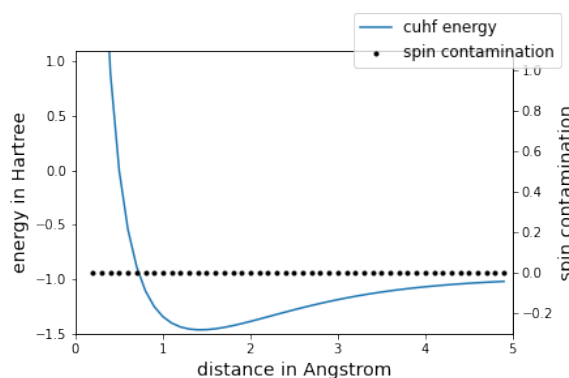


Figure 6: The stretch for H_3 using CUHF.

We can see that across the plotted data, the spin contamination remains zero. The curve is also smoother than the UHF curve. Where the UHF curve might suggest a disintegration, the CUHF curve rises above the level of three separate hydrogens. When we consider the constraint we have introduced this is hardly surprising. We have said that there are certain orbitals that have to be doubly occupied, as laid out in subsections 2.1 and 2.2. This is of course something we can not maintain at very long bondlengths. Physically we could state that if we were to pull the hydrogen atoms apart far enough, they would go their separate

ways, since this would be the more energetically stable solution, instead of trying to maintain the bonds across such large distances. It is then reasonable that the energy would rise as we continue to enforce this constraint.

4.3 CIS results

In H_3 we can consider nine excitations in total, so when we include the ground state, we should have ten states in total. We can take a closer look at figure 7.

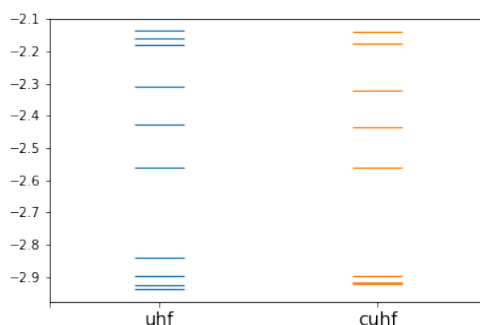


Figure 7: The CIS excitation energies for both UHF and CUHF as plotted by our implementation

When we look at the UHF energies, we have ten distinct values. In CUHF we have some degeneracies in the excitation energy levels. We can take a look at the table 1.

Table 1: The electronic excitation energies of H_3 in STO-3G basis. The energy is given in Hartree.

	UHF	CUHF
1	-2.938139	-2.922658
2	-2.923512	-2.915707
3	-2.896022	-2.915707
4	-2.840079	-2.895707
5	-2.560947	-2.560947
6	-2.428072	-2.436524
7	-2.309793	-2.323014
8	-2.179570	-2.176621
9	-2.160308	-2.176621
10	-2.137036	-2.139049

We do indeed notice that energies two and three are equal, as well as the energies eight and nine. Of course, it is logical to find a UHF result where all the expectation values are different. If all orbitals are different, We can expect all energy differences to be different as well. However, in CUHF we discover that there are some identical excitation energies. The reason is that the ground state and the mentioned excited state are degenerate. (cfr. infra) We can also note another odd feature in both energies. The lowest excitation does not equal the ground state. When we consult our data in table 2, we notice that the lowest excitation is lower in energy than the ground state.

Table 2: Electronic ground state energies for H_3 in the STO-3G basis. The energy is given in Hartree.

	UHF	CUHF
ground state energy	-2.923512	-2.915707

This data was verified using the GQCP library. This result is unexpected, since one does not expect an excitation to be lower in energy than the ground state. We could verify that this excitation is a triplet excitation, or an excitation in which an α electron is excited to a β orbital and vice versa. We could also determine the spin eigenvalue using the GQCP library as 0.5, which means that it still is a spin eigenstate. A possible explanation could be that the ground state of H_3 is degenerate, but in Hartree-Fock, we always take a single Slater determinant into account. In fact, the ground state contains two determinants. When we mix the determinants in CIS, the result is a decreased energy. We can see these determinants in figure 8. Furthermore, we can learn from this that for open shell species that have spin deficiencies, CIS is not the method of choice if we want to include the triplet excitations. When the triplet excitations are left out of the calculation, we find a result more in line with our expectations, in which the ground state is the lowest energy state. These results were displayed in table 3.

Table 3: The excitation energies when triplets are not considered. The energy is given in Hartree.

	UHF	CUHF
1	-2.923515	-2.915707
2	-2.840101	-2.895707
3	-2.428086	-2.436524
4	-2.160319	-2.176620
5	-2.137039	-2.139049

We can also notice that the degeneracies have also disappeared, meaning that one of the excited states in both degeneracies was a triplet excitation. We can explain this by looking at the specific excitations that contribute to these excited states. When we think of the H_3 molecule in STO-3G basis, there are three orbital pairs to consider. Let us take a look at the orbital diagram in figure 8.

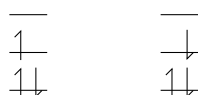


Figure 8: Energy diagrams of the ground state of H_3

We can see in this picture that it does not make a difference what spin the unpaired electron has towards the energy. So in fact we expect to see a degeneracy at the ground state energy level. Since this does involve an α to β excitation we do not expect to see it when leaving those out. The other degeneracy was seen at the energy level of -2.176620 Hartree. We can consult GQCP in order to find the exact excitation that is done. The configuration is given in figure 9.

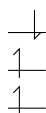


Figure 9: Energy diagram of the state at -2.176620 Hartree in CUHF

Based on this picture, we expect the same excitation energy if the α electron from ψ_1 would be excited. We can take a look at the orbitals in question. They are displayed in figure 10.



Figure 10: ϕ_1 for both α (left) and β electrons (right)

We detect some asymmetry in the orbitals. Normally we would expect H_3 to be a D_{3h} system. However, Psi4 has lowered the symmetry to a C_{2v} point group. The reason for this is that it does not contain all possible point groups. We will now compare the CIS data with CUHF reference to the CIS data with ROHF reference generated by Psi4. The Psi4 reference data for ROHF generates two states, one of which is the ground state. The energies are given in table 4. Psi4 does not consider triplet excitations by default, but we are still left with fewer excitations than one would expect.

Table 4: Electronic excitation energies of H_3 in ROHF according to psi4. The energy is given in Hartree.

	energy
1	-2.915707
2	-2.176621

We can see that the ground state energy is the same as CUHF method. However, the single excitation can also be seen returning to the CUHF excitations. Psi4 lists the excitation it considered as an excitation of the β electron into the open shell, so if we were to draw it up using an energy diagram, it would look like figure 11.



Figure 11: Energy diagram of the state at -2.176620 Hartree in ROHF

We can take a look at the orbital visualizations. These are displayed in next to each other in figures 12 through 15.

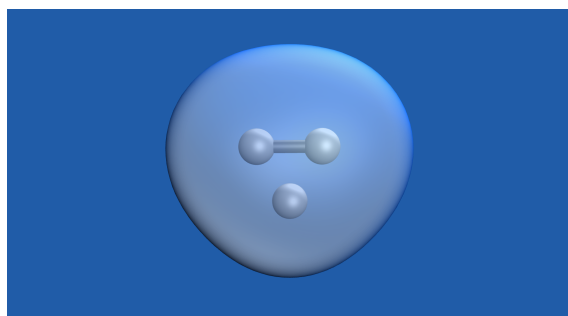


Figure 12: The lowest β MO in CUHF

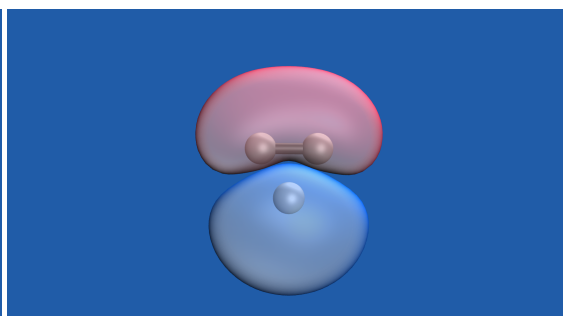


Figure 13: The highest β MO in CUHF

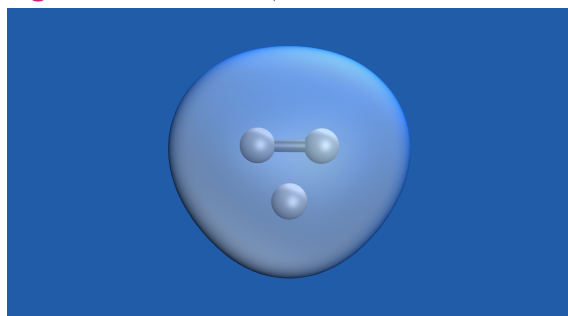


Figure 14: The lowest β MO in ROHF

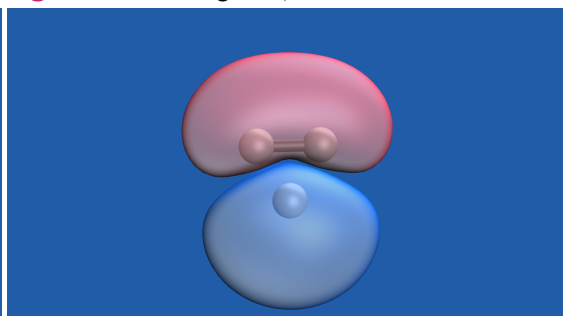


Figure 15: The second highest β MO in ROHF

All these orbitals are part of the same irreducible representation. Given that all the orbitals for ROHF shown above belong to the A_1 representation of the C_{2v} point group and that the

last orbital follows the B_2 representation, we will not see any excitations. We can verify that this is the case by looking at figure 16.

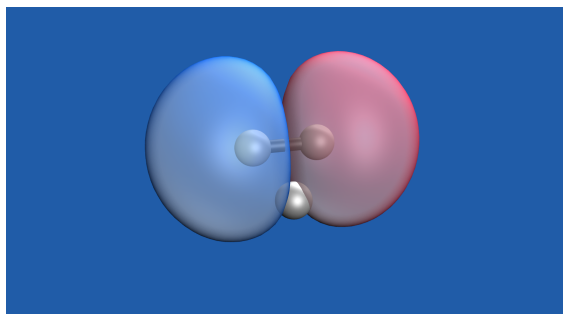


Figure 16: The highest β MO in H_3 using ROHF. It follows the B_2 representation

Interestingly enough this orbital is the same as the one orbital of CUHF we have not seen, the second highest β orbital. This is shown in figure 17.

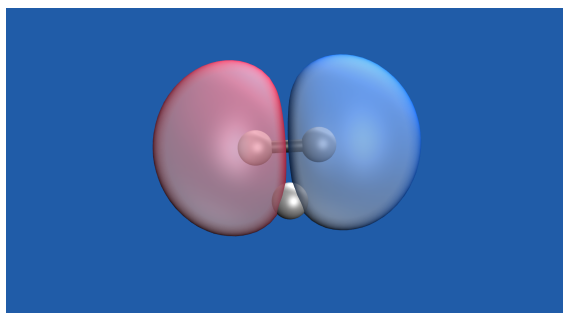


Figure 17: The second highest β MO in H_3 using CUHF.

Worth noting is that ROHF does not have a predefined set of operators. In this original work, Roothaan writes about coupling coefficients that vary between cases [3]. Plakhutin and Davidson list a table of possible values for these coefficients [6]. Depending on the coefficients used, we expect different orbital energies, so we can also expect different excitation energies. To conclude this section, we will look at the expectation values of the \hat{S}^2 operator of the excited states whose energies are listed in table 3. Plakhutin and Davidson mentioned that some excited states in CUHF are no longer eigenfunctions of \hat{S}^2 [6]. The main examples that are mentioned are the excitations of the open shell α electron to the α virtual orbital, and the excitation of the closed shell β electron to the open shell β orbital. In the 2.1 section, we saw that spin contamination could be defined like equation (6). For \hat{S}_z we expect the value 0.5 [18], so in a spin uncontaminated system, the expectation value over \hat{S}^2 should be 0.75. However, in table 5 we can clearly see that this is not always the case.

Table 5: The expectation value for the \hat{S}^2 operator in UHF and CUHF

	UHF	CUHF
1	0.75	0.75
2	0.759614	0.752480
3	1.098750	1.148304
4	1.739963	1.750000
5	1.401673	-1.34921

While the ground state may be uncontaminated, we see that the excited states are not. Even in CUHF, the excited state wave functions remain spin contaminated. As is clear from table 5, the constraint applied on the orbitals does not hold for the excited states.

5 Conclusions

In UHF the total spin symmetry of the Hamiltonian is broken, which results in spin contamination. We have seen that the CUHF method gives us an alternative that can deal with this spin contamination. We then explored the possibility of using CIS on this reference. Here we noticed that both CUHF and UHF feature excited states that are lower in energy than the ground state, when allowing for α -to- β and β -to- α excitations. We then concluded that CIS is not the ideal method for describing such systems if one wants to account for such excitations. This was based upon the fact that we found an excited state that was lower in energy than the ground state at equilibrium geometry. We have also seen that the singly excited states in CUHF are spin contaminated. If we used these states in a full CI wave function, we expect that wave function to be spin contaminated as well. An interesting next step would be to apply full CI to the CUHF wave function to see how the ground state energy might evolve if we do allow second order excitations. However, since spin contamination is not out of the picture for excited states, we would need to develop a way to account for this and maybe remove it entirely. Also, the study of other open shell systems could be done. H_3 is a system with one unpaired electron. O_2 has two unpaired electrons. It could be interesting to consider such a system, since we cannot assume that the same results will hold for such systems.

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