Lecture 6: Coupled Cluster Theory

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1 Quantum Chemistry Wavefunctions

In the last set of notes, we introduced the concept of a wave operator Ω , which transforms our reference determinant Φ into the exact wavefunction Ψ

$$|\Psi\rangle = \Omega |\Phi\rangle$$

Different valid choices of the wave operator Ω constitute different ansatz, which each arrive at valid form for the same exact wavefunction $|\Psi\rangle$. In perturbation theory (PT), our wave operator is of the form, $\Omega = P + \sum_{n=0}^{\infty} (\mathcal{R}_0 V')^n \mathcal{R}_0 V P$, so that our wavefunction is

$$|\Psi_{\mathrm{PT}}\rangle = |\Phi_0\rangle + \sum_{n=0}^{\infty} (\mathcal{R}_0 V')^n \mathcal{R}_0 V |\Phi\rangle$$

In configuration interaction (CI), our wave operator (though we did not call it that at the time) takes the form $\Omega = (1 + C_1 + C_2 + \cdots + C_n)$ so that

$$|\Psi_{\text{CI}}\rangle = (1 + C_1 + C_2 + \dots + C_n) |\Phi\rangle$$

In coupled cluster theory (CC), our wave operator, and therefore our wavefunction, is an exponential form,

$$|\Psi_{\rm CC}\rangle = \exp(T_1 + T_2 + \dots + T_n) |\Phi\rangle$$

Or simply,

$$|\Psi_{\rm CC}\rangle = \exp(T) |\Phi\rangle$$

Each of the wavefunctions, in their limit, approach the exact same wavefunction. However, perturbation theory is an odd duck, since it always has an infinite number of contributions to the wavefunction and energy, whereas the CC and CI wavefunctions for a given n-electron system terminate at n terms (you can't have a quadruple excitation in a 3-electron system).

2 Coupled Cluster Theory

Our coupled cluster wavefunction from before is

$$|\Psi_{\rm CC}\rangle = \exp(T) |\Phi\rangle$$

$$|\Psi_{\rm CC}\rangle = \exp(T_1 + T_2 + \dots + T_n) |\Phi\rangle$$

where T is the "cluster operator" which is a sum over k-electron excitation operators

$$T = T_1 + T_2 + T_3 + \cdots$$

$$T_k = \left(\frac{1}{k!}\right)^2 t_{a_1 \cdots a_k}^{i_1 \cdots i_k} \tilde{a}_{i_1 \cdots i_k}^{a_1 \cdots a_k}$$

where $t_{a_1 \cdots a_k}^{i_1 \cdots i_k}$ are the coupled cluster *amplitudes*. It is worth briefly pointing out some buried complexity here. Noting that the function e^T can be expanded according to it's Taylor series, the full form of the coupled cluster wavefunction looks like the following,

$$|\Psi_{\rm CC}\rangle = \left[1 + (T_1 + T_2 + T_3 + T_4 + \cdots) + \frac{1}{2}(T_1 + T_2 + T_3 + T_4 + \cdots)^2 + \frac{1}{6}(T_1 + T_2 + T_3 + T_4 + \cdots)^3 \cdots\right] |\Phi\rangle$$

Solving for the coupled cluster energy is done by doing the same thing we always do: project our reference determinant $\langle \Phi |$ on the left of both sides of the Schrödinger equation. Assuming intermediate normalization, we obtain

$$\langle \Phi | H_{\rm c} | \Psi_{\rm CC} \rangle = E_c \langle \Phi | \Psi_{\rm CC} \rangle$$

 $\langle \Phi | H_c \exp(T) | \Phi \rangle = E_c$

Equations for the cluster amplitudes $t_{ab\cdots}^{ij\cdots}$ can be found by left-projecting the Schrödinger equation by excited determinants

$$\langle \Phi_{ij...}^{ab...} | H_{c} \exp(T) | \Phi \rangle = E_{c} \langle \Phi_{ij...}^{ab...} | \exp(T) | \Phi \rangle$$

where an equation involving t_i^a would be obtained by projecting by $\langle \Phi_i^a |$, an equation for t_{ij}^{ab} by projecting by $\langle \Phi_{ij}^{ab} |$, and so on. To see this, one can take the right side of the above equation and expand the exponential $\exp(T) = 1 + T + \frac{T^2}{2!} + \cdots$, and only one term survives:

$$E_{\rm c} \langle \Phi_{ij...}^{ab...} | \exp(T) | \Phi \rangle = E_{\rm c} t_{ab...}^{ij...}$$

Plugging in the exponential expansion of $\exp(T)$ into the energy equation yields

$$\langle \Phi | H_{\rm c} | \Phi \rangle + \langle \Phi | H_{\rm c} T | \Phi \rangle + \langle \Phi | H_{\rm c} \frac{T^2}{2!} | \Phi \rangle = E_{\rm c}$$

These terms are the only ones that survive because higher powers of T contain no terms for which one can obtain complete contractions. An analgous equation exists for the amplitudes, but the powers of T which survive will be dictated by the type of excitation of the determinant in the bra $\langle \Phi_{ij}^{ab\cdots} |$, as this will influence the number of contractions one can make.

These equations is not easily solvable. The trick used in traditional coupled cluster theory to make these equations more workable is to multiply by $\exp(-T)$ on both sides, which effectively transforms our Hamiltonian $\bar{H}_c = \exp(-T)H \exp(T)$. We highlight the key results in blue.

$$\begin{split} \langle \Phi | H_{\rm c} | \Psi_{\rm CC} \rangle &= E_c \, \langle \Phi | \Psi_{\rm CC} \rangle \\ \langle \Phi | \exp(-T) H_{\rm c} \exp(T) | \Phi \rangle &= E \, \langle \Phi | \exp(-T) \exp(T) | \Phi \rangle \\ \langle \Phi | \exp(-T) H_{\rm c} \exp(T) | \Phi \rangle &= E_{\rm c} \end{split}$$

$$\begin{split} \langle \Phi_{ij}^{ab\cdots} | H_{\rm c} \exp(T) | \Phi \rangle &= E_{\rm c} \langle \Phi_{ij\cdots}^{ab\cdots} | \exp(T) | \Phi \rangle \\ \langle \Phi_{ij\cdots}^{ab\cdots} | \exp(-T) H_{\rm c} \exp(T) | \Phi \rangle &= E_{\rm c} \langle \Phi_{ij\cdots}^{ab\cdots} | \exp(-T) \exp(T) | \Phi \rangle \\ \langle \Phi_{ij\cdots}^{ab\cdots} | \exp(-T) H_{\rm c} \exp(T) | \Phi \rangle &= E_{\rm c} \langle \Phi_{ij\cdots}^{ab\cdots} | \Phi \rangle \\ \langle \Phi_{ij\cdots}^{ab\cdots} | \exp(-T) H_{\rm c} \exp(T) | \Phi \rangle &= 0 \end{split}$$

This transformation decouples the amplitude equation from the energy equation by setting the right side of the amplitude equation to zero. In the language of diagrams, this transformation removes "disconnected" diagrams, and $E_{c}t_{ab\cdots}^{ij\cdots}$ is one example. This transformation also can be simplified further via the *Hausdorff expansion*.

2.1 The Hausdorff Expansion

The Hausdorff expansion expresses our transformed Hamiltonian in terms of a sum over nested commutators. As we will see, upon simplification it produces an expansion in powers of T which naturally truncates, thus removing the infinite expansion due to the exponential $\exp(T)$. The Hausdorff expansion is,

$$\bar{H}_{c} = \exp(-T)H_{c}\exp(T) = H_{c} + [H_{c}, T] + \frac{1}{2!}[[H_{c}, T], T] + \frac{1}{3!}[[[H_{c}, T], T], T] + \frac{1}{4!}[[[H_{c}, T], T], T] + \cdots$$

This is simplified by evaluating the commutators using Wick's theorem. Generally, if two strings of operators Q and Q' are normal ordered, and one of them has an even operator count, then

$$[Q, Q'] = N[\overrightarrow{QQ'}] - N[\overrightarrow{Q'Q}]$$

In this specific case,

$$[H_{c},T] = H_{c}T - TH_{c} = N[H_{c}T] + N[\overline{H_{c}T}] - N[TH_{c}] - N[\overline{TH_{c}}]$$
$$[H_{c},T] = N[\overline{H_{c}T}] - N[\overline{TH_{c}}]$$

The second term above is actually 0, since T is composed entirely of quasi-particle creation operators, so any contraction of T with something on the right will always be zero (†'s on the right \implies 0). Therefore, we only have,

$$[H_{\rm c}, T] = N[\overrightarrow{H_{\rm c}T}]$$

This reasoning can be continued for nested commutations with T, and we find that the only terms which survive are ones in which every single cluster operator T is contracted with the Hamiltonian at least once.

$$[\cdot,T]^n = N[H_cTT\cdots T] = (H_cT^n)_C$$

This naturally truncates the Hausdorff expansion, since the only nonzero terms in the Hausdorff expansion are those in which the Hamiltonian has at least one contraction with every cluster operator on its right. In the language of diagrams, we call these terms *connected*, so we label them with a subscript capital C.

$$\bar{H}_{c} = (H_{c} \exp(T))_{C} = (H_{c} + H_{c}T + \frac{1}{2!}H_{c}T^{2} + \frac{1}{3!}H_{c}T^{3} + \frac{1}{4!}H_{c}T^{4})_{C}$$

This ends at the fourth power of T because H_c is a linear combination of one-electron and two-electron operators, and these can contract with at most two and four T's, respectively. To convince yourself of this, you can write out the full form of $T = T_1 + T_2 + T_3 + \cdots$, re-express each T_k according to their definition, take various powers of the expansion T^2, T^4, T^5 , and try contracting with the Hamiltonian. You will find that most terms in T^4 cannot be such that every T forms a contraction with H_c , and no terms in T^5 can have each cluster operator contract with H_c . There isn't enough "room" in H_c for that many contractions.

To summarize, we initially replaced $H_c \exp(T)$ in our energy and amplitude equations with $\exp(-T)H_c \exp(T)$. We then used the Hausdorff expansion, simplified, and found that it truncates at just connected terms, $\exp(-T)H_c \exp(T) = (H_c \exp(T))_C$. This gives us our final results for the coupled cluster energy and amplitude equations,

$$\langle \Phi | H_{c} \exp(T) | \Phi \rangle_{C} = E_{c}$$
$$\langle \Phi_{ij...}^{ab...} | H_{c} \exp(T) | \Phi \rangle_{C} = 0$$

Note that these are the same results as the equations previously derived (in blue), except we have used the result of the Hausdorff expansion to replace $\exp(-T)H_c\exp(T)$ with just connected terms $(H_c\exp(T))_C$. Practically, when one uses the equations above algebraically, one should look for complete contractions in which the Hamiltonian has at least one contraction with every cluster operator on its right.

2.2 Excitation Levels

There is an alternative to considering which terms form complete contractions when evaluating expectation values. The excitation level of a series of operators is equal to the net number of particles or quasi-particles it creates, divided by 2. The excitation levels of T_1 , T_2 , and T_3 are 1, 2, and 3, and \tilde{a}_{abcd}^{ijkl} , which would perhaps appear in a bra as $\langle \Phi_{ijkl}^{abcd}|$, would have an excitation level of -4. The total excitation level of a term in an expectation value must balance out to zero, otherwise the term is zero. Thus, this is another way of detecting which terms cannot form complete contractions. What is the excitation level of terms in our Hamiltonian H_c ? Excitation operators in the Hamiltonian are made up of general indices p, q, r, s (not necessarily occupied or virtual), so the excitation level of the Hamiltonian is variable. The one-electron part of the Hamiltonian can have an excitation level of 0, +1, or -1. The two-electron part of the Hamiltonian can have an excitation level of 0, +1, or -2. To see why, recall that the one-electron operator is $h_p^a \tilde{a}_p^a$, which can take on virtual and occupied indices in four ways: $h_a^b \tilde{a}_a^a$, $h_a^i \tilde{a}_a^a$, $h_i^i \tilde{a}_a^i$, $h_i^j \tilde{a}_j^i$. In the $h_a^i \tilde{a}_a^a$ case, it converts to two quasi-particle creation operators $h_a^b h_a^b$. According to our definition of excitation level, this has an excitation level of +1. In the case of $h_a^a \tilde{a}_a^i$, we have $h_a^a h_a^a$, two quasi-particle annihilation operators, so that the excitation level is -1. The other two cases have excitation levels of 0. Similar arguments can be made for the two-electron cases involving \tilde{a}_a^{pq} .

By analyzing excitation levels, we can decide if expectation values are 0 or not. Examples,

• The following expectation value is nonzero. Each T_1 has an excitation level of +1, \tilde{a}_{ab}^{ij} has an excitation level of -2, and the Hamiltonian excitation level can be chosen to be 0 so that 1+1-2+0=0.

$$\langle \Phi_{ij}^{ab} | H_{\rm c} T_1^2 | \Phi \rangle$$

• The following expectation value is 0. T_4 has an excitation level of +4, \tilde{a}_a^i has an excitation level of -1, and we can choose the Hamiltonian to have an excitation level of -2, at best. -1 + -2 + 4 = 1.

$$\langle \Phi_i^a | H_c T_4 | \Phi \rangle$$

One could come up with similar conclusions by looking at what complete contractions can be formed, but it is often much faster to count excitation levels.

In coupled cluster, we truncate our cluster operator at some level of excitation, and then perform the Hausdorff expansion to find which cluster operator terms to utilize in our expectation values. Analyzing what complete contractions survive (or counting excitation levels) can then be used to filter the relevant terms terms even further. Finally, you can evaluate the expectation values either algebraically or diagrammatically.

The utility of the Hausdorff expansion is the removal of terms which are "disconnected", (terms in the exponential expansion of T for which a stray cluster operator is not contracted with H_c at least once). One could in theory skip

2.3 The Amplitude Equations

The cleanest way to solve for the cluster amplitudes in our amplitude equation

$$\langle \Phi_{ij...}^{ab...} | H_{c} \exp(T) | \Phi \rangle_{C} = 0$$

requires an odd-looking partitioning of our Hamiltonian into two parts: the diagonal part of the Fock operator, and everything else.

$$H_0 = f_p^p \tilde{a}_p^p$$

$$V_{c} = H_{c} - H_{0} = f_{p}^{q} (1 - \delta_{p}^{q}) \tilde{a}_{q}^{p} + \frac{1}{4} \bar{g}_{pq}^{rs} \tilde{a}_{rs}^{pq}$$

 H_0 can be absorbed into an excited determinant to give back a term which depends on the orbital energies,

$$H_0 \Phi_{i_1 \cdots i_k}^{a_1 \cdots a_k} = \mathcal{E}_{i_1 \cdots i_k}^{a_1 \cdots a_k} \Phi_{i_1 \cdots i_k}^{a_1 \cdots a_k} \qquad \qquad \mathcal{E}_{q_1 \cdots q_k}^{p_1 \cdots p_k} \equiv \sum_{r=1}^k f_{p_r}^{p_r} - \sum_{r=1}^k f_{q_r}^{q_r}$$
 (1)

 $\mathcal{E}_{i_1\cdots i_k}^{a_1\cdots a_k}$ is essentially the change in the sum of the orbital energies from the reference state (Hartree-Fock) for the given excited determinant. A single \mathcal{E}_{ij}^{ab} is $\mathcal{E}_{ij}^{ab} = \epsilon_a + \epsilon_b - \epsilon_i - \epsilon_j$. Using the above equations in conjunction with our amplitude equations,

$$\langle \Phi_{ij\cdots}^{ab\cdots} | H_{c} \exp(T) | \Phi \rangle_{C} = 0$$

We plug in our partitioned Hamiltonian $H_c = H_0 + V_c$,

$$\langle \Phi_{ij...}^{ab...}|H_0 \exp(T)|\Phi\rangle_{\mathbf{C}} + \langle \Phi_{ij...}^{ab...}|V_{\mathbf{c}} \exp(T)|\Phi\rangle_{\mathbf{C}} = 0$$

The left term evalutes to

$$\langle \Phi^{ab\cdots}_{ij\cdots}|H_0 \exp(T)|\Phi\rangle_{\mathbf{C}} = \langle \Phi^{ab\cdots}_{ij\cdots}|H_0T|\Phi\rangle = \mathcal{E}^{ab\cdots}_{ij\cdots} \langle \Phi^{ab\cdots}_{ij\cdots}|T|\Phi\rangle = \mathcal{E}^{ab\cdots}_{ij\cdots}t^{ij\cdots}_{ab\cdots}$$

where we have used the fact that only T in $\exp(T) = 1 + T + \frac{1}{2}T^2 \cdots$ can create connected terms with H_0 , since it is a one-electron operator. Now our amplitude equation reads as

$$\langle \Phi_{ij\cdots}^{ab\cdots} | V_{c} \exp(T) | \Phi \rangle_{C} + \mathcal{E}_{ij\cdots}^{ab\cdots} t_{ab\cdots}^{ij\cdots} = 0$$

Since $\mathcal{E}^{ab\cdots}_{ij\cdots}$ is just a linear combination of orbital energies, we can absorb a minus sign by swapping upper and lower indices, $-\mathcal{E}^{ab\cdots}_{ij\cdots} = \mathcal{E}^{ij\cdots}_{ab\cdots}$, which gives us the following clean equation for solving for the amplitudes,

$$t_{ab\cdots}^{ij\cdots} = \left(\mathcal{E}_{ab\cdots}^{ij\cdots}\right)^{-1} \left\langle \Phi_{ij\cdots}^{ab\cdots} | V_{c} \exp(T) | \Phi \right\rangle_{C}$$

The amplitude equations must be solved iteratively, since $t_{ab\cdots}^{ij\cdots}$ is also embedded in $\exp(T)$.

2.4 The CCSDTQ amplitude equations

If we truncate the cluster operator $T = T_1 + T_2 + T_3 + T_4$, then $\exp(T) = \exp(T_1 + T_2 + T_3 + T_4)$. Not every operator in the exponential expansion of $\exp(T_1 + T_2 + T_3 + T_4)$ is relevant; disconnected terms are omitted, as well as terms for which the excitation level of the expectation value does not balance out to 0. After careful consideration, one can use the amplitude equation we derived above to obtain amplitude equations for CCSDTQ:

$$\begin{split} t_a^i = & (\mathcal{E}_a^i)^{-1} \left\langle \Phi_i^a \middle| V_{\text{c}} (1 + T_1 + T_2 + T_3 + \frac{1}{2} T_1^2 + T_1 T_2 + \frac{1}{3!} T_1^3) \middle| \Phi \right\rangle_{\text{C}} \\ t_{ab}^{ij} = & (\mathcal{E}_{ab}^{ij})^{-1} \left\langle \Phi_{ij}^{ab} \middle| V_{\text{c}} (1 + T_1 + T_2 + T_3 + T_4 + \frac{1}{2} T_1^2 + T_1 T_2 + T_1 T_3 + \frac{1}{2} T_2^2 + \frac{1}{3!} T_1^3 + \frac{1}{2} T_1^2 T_2 + \frac{1}{4!} T_1^4) \middle| \Phi \right\rangle_{\text{C}} \\ t_{abc}^{ijk} = & (\mathcal{E}_{abc}^{ijk})^{-1} \left\langle \Phi_{ijk}^{abc} \middle| V_{\text{c}} (T_2 + T_3 + T_4 + T_1 T_2 + T_1 T_3 + \frac{1}{2} T_2^2 + T_1 T_4 + T_2 T_3 + \frac{1}{2} T_1^2 T_2 + \frac{1}{2} T_1^2 T_3 + \frac{1}{2} T_1 T_2^2 + \frac{1}{3!} T_1^3 T_2 \right) \middle| \Phi \right\rangle_{\text{C}} \\ t_{abcd}^{ijkl} = & (\mathcal{E}_{abcd}^{ijkl})^{-1} \left\langle \Phi_{ijkl}^{abcd} \middle| V_{\text{c}} (T_3 + T_4 + T_1 T_3 + \frac{1}{2} T_2^2 + T_1 T_4 + T_2 T_3 + T_2 T_4 + \frac{1}{2} T_3^2 + \frac{1}{2} T_1^2 T_3 + \frac{1}{2} T_1 T_2^2 + \frac{1}{2} T_1^2 T_4 + T_1 T_2 T_3 + \frac{1}{3!} T_1^3 T_3 + \frac{1}{2!2!} T_1^2 T_2^2 \right) \middle| \Phi \right\rangle_{\text{C}} \end{split}$$

3 On the connection between Coupled Cluster and Configuration Interaction

Our CI wavefunction is

$$|\Psi_{\rm CI}\rangle = (1 + C_1 + C_2 + \cdots) |\Phi\rangle$$

Our CC wavefunction is

$$|\Psi_{\rm CC}\rangle = \exp(T_1 + T_2 + \cdots) |\Phi\rangle$$

$$|\Psi_{\rm CC}\rangle = \left[1 + (T_1 + T_2 + T_3 + T_4 + \cdots) + \frac{1}{2}(T_1 + T_2 + T_3 + T_4 + \cdots)^2 + \frac{1}{6}(T_1 + T_2 + T_3 + T_4 + \cdots)^3 \cdots\right] |\Phi\rangle$$

Noting that each T_k (C_k) is just made up of scalar cluster amplitudes $t_{ab\cdots}^{ij\cdots}$ (CI coefficients $c_{ab\cdots}^{ij\cdots}$) and a Phi-normal-ordered excitation operator, we can compare these wavefunction expansions term by term and find the following:

$$C_1 = T_1$$

$$C_2 = T_2 + \frac{1}{2}T_1^2$$

$$C_3 = T_3 + T_1T_2 + \frac{1}{6}T_1^3$$

$$C_4 = T_4 + \frac{1}{2}T_2^2 + T_1T_3 + \frac{1}{2}T_1^2T_2 + \frac{1}{24}T_1^4$$

CI does not discriminate between different kinds of n-tuple excitations, but in coupled cluster thoery, we can sub-classify various n-tuple excitations in different ways. Consider the quadruple excitation terms. The physical interpretation of T_4 as an excitation event on some quantum mechanical state is 4 electrons coming together at the same location and, upon interacting, they all have to simultaneously excite to a new state. $\frac{1}{2}T_2^2$, on the other hand, corresponds to an excitation event in which a pair of 2 electrons which are close enough to locally interact each excite to a new state, while simultaneously another distinct pair of electrons does the same. Which of these physical events $(T_4 \text{ or } \frac{1}{2}T_2^2)$ are more likely to occur in a molecule? Almost certainly the latter, since electrons tend to avoid each other, but they also tend to form electron pairs! We conclude that $\frac{1}{2}T_2^2$ is a more important contribution to our wavefunction than T_4 .

Now consider the CCSD wavefunction,

$$|\Psi_{\text{CCSD}}\rangle = \exp(T_1 + T_2) |\Phi\rangle$$

$$|\Psi_{\text{CCSD}}\rangle = (1 + T_1 + T_2 + \frac{1}{2}T_1^2 + T_1T_2 + \frac{1}{6}T_1^2 + \frac{1}{2}T_2^2 + \frac{1}{2}T_1^2T_2 + \frac{1}{24}T_1^4 + \dots) |\Phi\rangle$$

We obtain all single and double excitations according to CI, but we also obtain some triple excitations $(T_1T_2, \frac{1}{6}T_1^3)$ and some quadruple excitations $(\frac{1}{2}T_2^2 + \frac{1}{2}T_1^2T_2 + \frac{1}{24}T_1^4)$. This is why its called coupled cluster; higher order excitations (quadruple, $\frac{1}{2}T_2^2$) are included even in low-order truncation of the scheme (CCSD) via *coupling* two double excitations together. To put it another way, in CCSD, we obtain some information about quadruple excitations by considering two clusters which each doubly excite simultaneously. Not only that, but CCSD is giving us quadruple excitation of relatively high importance, according to our physical argument earlier.

It is clear that CCSD is more accurate than CISD. However, coupled cluster gets even better at higher orders. Looking at our expressions for C_1, C_2, C_3, C_4 above, we see that the majority of C_4 is made up of terms which occur at CCSDT; expanding the CCSDT exponential $\exp(T_1 + T_2 + T_3)$ will give you all of the terms in C_4 except T_4 (!!!). Thus, CCSDT is wayyyy more accurate than CISDT, and it's even almost as accurate correlation-wise as CISDTQ, since all that is missing is T_4 . Nevermind the fact that coupled cluster is size-consistent, which is probably making up for more accuracy than the relative importance of T_4 in some cases. In this view, it is even proper to say that CCSDT is more accurate than CISDTQ.

3.1 Deriving CC from CI

Starting from full-CI,

$$|\Psi\rangle = (1 + C_1 + C_2 + C_3 + \dots + C_n) |\Phi\rangle$$

 $|\Psi\rangle = (1 + C) |\Phi\rangle$

We can define the cluster operator $T = T_1 + T_2 + \cdots$ as $T = \ln(1+C)$ where $C = C_1 + C_2 + \cdots$. This way, $1 + C = e^T$, so it must be the case

$$|\Psi\rangle = e^T |\Phi\rangle$$

which is our coupled cluster wavefunction.

We can compare coupled cluster and CI operators using the Taylor expansion for $\ln(1+z)$

$$\ln(1+z) = \sum_{m=1}^{\infty} \frac{(-1)^{m+1} z^m}{m}$$

This implies that for T we have

$$T = \ln(1+C) = \sum_{m=1}^{\infty} \frac{(-1)^{m+1}C^m}{m}$$

This series always converges, since we always have a finite number of electrons in a system. Expanding the series we get,

$$T = C - \frac{C^2}{2} + \frac{C^3}{3} + \cdots$$

$$T = C_1 + C_2 + C_3 + \cdots - \frac{1}{2}(C_1 + C_2 + C_3 + \cdots)^2 + \frac{1}{3}(C_1 + C_2 + C_3 + \cdots)^3 + \cdots$$

4 On the connection between Coupled Cluster and Perturbation Theory

TODO; diagrams pre-requisite.