

Post-Hartree-Fock Methods

Methods use a Hartree-Fock calculation as starting point and try to improve the HF results by taking account of **electron correlation**:

- Configuration Interaction (CI)
- Many-Body Perturbation Theory (Moller-Plesset (MPn))
- Coupled Cluster (CC) Theory
(Chapter 7 of the script)

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Size Extensivity/Size Consistency

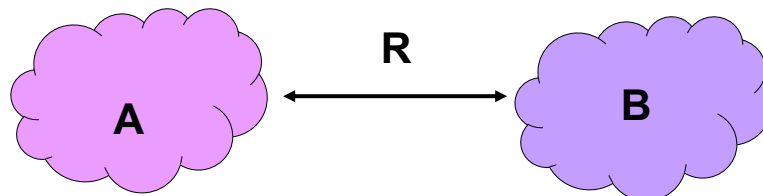
Extensive and intensive properties:

- *intensive* properties do not depend on system size (but only on the type of system) e.g. density, melting point etc..
- *extensive* properties depend on size (amount of matter) e.g. mass, volume etc..

Size Extensivity:

- correct linear scaling with number of electrons n (\Leftrightarrow error constant as a function of n)

Size Consistency:



$E(A+B) \rightarrow E(A)+E(B)$ for $R \rightarrow \infty$ (when interaction is nullified)
(*strict separability*)

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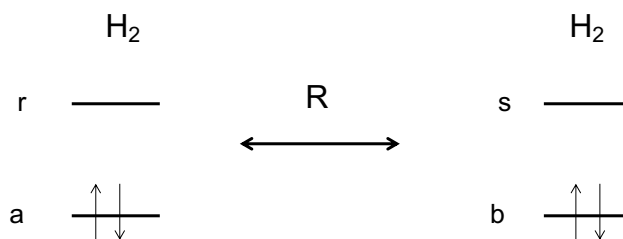
Quiz XV: Size-Consistency

- 1) Is Hartree-Fock size-consistent (RHF, UHF)?
- 2) Is full CI size-consistent? Truncated CI?
- 3) Is MPn size-consistent?

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Why is truncated CI not size-consistent?

Example: CISD of $(H_2)_2$ with minimal basis set



Determinants for full system:

gs: (aa'bb')

Singles: (arbb'), (aa'bs), (asbb'), (aa'br)

Doubles: (rr'bb'), (aa'ss'), (aa'rr'), (ss'bb'), (arbs)

For $R \rightarrow \infty$: **6 dets**

(aa'bb'), (arbb'), (aa'bs), (rr'bb'),
(aa'ss'), (arbs)

Single systems: 3 determinants each

gs: (aa')

Singles: (ar)

Doubles: (rr')

Total wavefunction product $3 \times 3 = 9$

Slater determinants (includes triply and quadruply excited configurations!)

(arss'), (rr'bs), (rr'ss')

Truncated CI is not size-consistent!

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Coupled Cluster (CC)

Yet another Ansatz for the Many-Body Wavefunction:

$$\boxed{|\Psi^{CC}\rangle = e^{\hat{T}} |\Phi_0\rangle}$$

where $|\Phi_0\rangle$ is a reference wavefunction. e.g. the Hartree-Fock determinant and \hat{T} is the cluster operator

$$\hat{T} = \hat{T}_1 + \hat{T}_2 + \dots$$

\hat{T}_1 is the one-particle excitation operator which generates singly excited configurations.
 \hat{T}_2 is the two-particle excitation operator which generates doubly excited Slater determinants.
 Etc..

In analogy to the terminology used in case of CI calculations, coupled cluster methods are labelled according to the order of excitations that is included e.g. **CCSD**, **CCSDT**, **CCSDTQ** etc.. In addition, acronyms of the type **CCSD(T)** refer to a coupled cluster calculation including singles and doubles and a perturbative treatment to include the effects of triple excitations.

Calculations of the coupled cluster type are nowadays one of the most popular approaches for very high accuracy calculations.

CCSD(T) is sometimes called the 'gold standard of quantum chemistry'.

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Another Notation: Second Quantization

Creation operator \hat{a}_i^\dagger creates an electron in orbital i
Annihilation operator \hat{a}_i removes an electron from orbital i

$$\hat{a}_i^\dagger |0\rangle = |\phi_i\rangle$$

$$\hat{a}_i |\phi_j\rangle = \delta_{ij} |0\rangle$$

Action is not commutative $\hat{a}_i^\dagger \hat{a}_i \neq \hat{a}_i \hat{a}_i^\dagger$ $[\hat{a}_i^\dagger, \hat{a}_i] = \hat{a}_i^\dagger \hat{a}_i - \hat{a}_i \hat{a}_i^\dagger \neq 0$
 For fermions: anticommutation $\{\hat{a}_i^\dagger, \hat{a}_i\} = \hat{a}_i^\dagger \hat{a}_i + \hat{a}_i \hat{a}_i^\dagger$

Canonical (normal) order: all creation operators left of annihilation operators

Cluster operators \hat{T}_n in second quantization:

$$\hat{T}_1 = \sum_a \sum_r t_a^r \hat{a}_r^\dagger \hat{a}_a$$

$$\hat{T}_2 = \sum_a \sum_{b \neq a} \sum_r \sum_{s \neq r} t_{ab}^{rs} \hat{a}_s^\dagger \hat{a}_r^\dagger \hat{a}_b \hat{a}_a$$

Coefficients = cluster amplitudes

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The Advantage of an Exponential Form

$$|\Psi^{CC}\rangle = e^{\hat{T}} |\Phi_0\rangle$$

Taylor expansion of $e^{\hat{T}}$:

$$e^{\hat{T}} = 1 + \hat{T} + \frac{1}{2!} \hat{T}^2 + \frac{1}{3!} \hat{T}^3 + \dots$$

Example: CCSD $\hat{T} = \hat{T}_1 + \hat{T}_2$

$$e^{(\hat{T}_1 + \hat{T}_2)} = 1 + \hat{T}_1 + \hat{T}_2 + \frac{1}{2!} (\hat{T}_1^2 + 2\hat{T}_1\hat{T}_2 + \hat{T}_2^2) + \dots$$

Includes given excitation level up to infinite order!

CC is size-consistent independent of reference wavefunction!

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Quiz XVI: Exponential Form

- 1) Is CISD or CCSD more accurate?

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Quiz XVI: Exponential Form

Given level of excitations can be realized in different ways: e.g. doubles

$$\hat{T}_1^2 = \sum_a \sum_r \sum_b \sum_s t_{ab}^{rs} \hat{a}_s^\dagger \hat{a}_r^\dagger \hat{a}_b \hat{a}_a$$

$$\hat{T}_2 = \sum_a \sum_{b \neq a} \sum_r \sum_{s \neq r} t_{ab}^{rs} \hat{a}_s^\dagger \hat{a}_r^\dagger \hat{a}_b \hat{a}_a$$

disconnected cluster amplitudes

Connected cluster amplitudes

Inclusion of disconnected clusters leads to size-consistency!

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Coupled Cluster Energy

$$|\Psi^{CC}\rangle = e^{\hat{T}} |\Phi_0\rangle$$

Insert into
Schrodinger Eqs

$$\hat{H} e^{\hat{T}} |\Phi_0\rangle = E^{CC} e^{\hat{T}} |\Phi_0\rangle$$

Multiply from left with
 $\langle \Phi_0 |$

$$\langle \Phi_0 | \hat{H} e^{\hat{T}} |\Phi_0\rangle = E^{CC} \langle \Phi_0 | e^{\hat{T}} |\Phi_0\rangle$$

$$\langle \Phi_0 | \hat{H} e^{\hat{T}} |\Phi_0\rangle = E^{CC} \langle \Phi_0 | (1 + \hat{T}_1 + \hat{T}_2 + \dots) |\Phi_0\rangle$$

$$\langle \Phi_0 | \hat{H} e^{\hat{T}} |\Phi_0\rangle = E^{CC}$$

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Coupled Cluster Energy: CCSD

e.g. for CCSD

$$E^{CCSD} = \langle \Phi_0 | \hat{H} \left(1 + \hat{T}_1 + \hat{T}_2 + \frac{1}{2} \hat{T}_1^2 + \hat{T}_1 \hat{T}_2 + \frac{1}{2} \hat{T}_2^2 + \dots \right) | \Phi_0 \rangle$$

$$E^{CCSD} = E_0 + \sum_a^{\text{occ}} \sum_r^{\text{virt}} t_a^r \langle \Phi_0 | \hat{H} | \Phi_a^r \rangle + \sum_{a,b>a}^{\text{occ}} \sum_{r,s>r}^{\text{virt}} (t_{ab}^{rs} + t_a^r t_b^s - t_a^s t_b^r) \langle \Phi_0 | \hat{H} | \Phi_{ab}^{rs} \rangle$$

$$E^{CCSD} = E_0 + \sum_{a,b>a}^{\text{occ}} \sum_{r,s>r}^{\text{virt}} (t_{ab}^{rs} + t_a^r t_b^s - t_a^s t_b^r) \langle \phi_a \phi_b | | \phi_r \phi_s \rangle$$

Still have to determine cluster amplitudes!

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Coupled Cluster Amplitudes t_μ

Variational approach: direct variation of the energy with respect to cluster amplitudes => intractable set of nonlinear equations

=> Projected coupled-cluster equations by left projecting each determinant μ

=> $\binom{M}{N}$ equations for energy and amplitudes

$$\begin{aligned} \langle \Phi_0 | \hat{H} e^{\hat{T}} | \Phi_0 \rangle &= E \\ \langle \mu | \hat{H} e^{\hat{T}} | \Phi_0 \rangle &= E \langle \mu | e^{\hat{T}} | \Phi_0 \rangle \end{aligned}$$

Have to be solved self-consistently!

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Convergence and Scaling

- The difference between the cc-pVDZ energies of H₂O from the full CI limit of various coupled-cluster methods (E_h). The HOH angle is 110.565° and $R_{ref} = 1.84345 a_0$.

method	$R = R_{ref}$	$R = 2R_{ref}$
RHF	0.217822	0.363954
CCSD	0.003744	0.022032
CCSDT	0.000493	-0.001405
CCSDTQ	0.000019	-0.000446
CCSDTQ5	0.000003	

- The series converges very fast at equilibrium and slower at stretched geometry where there a higher multi-reference character.

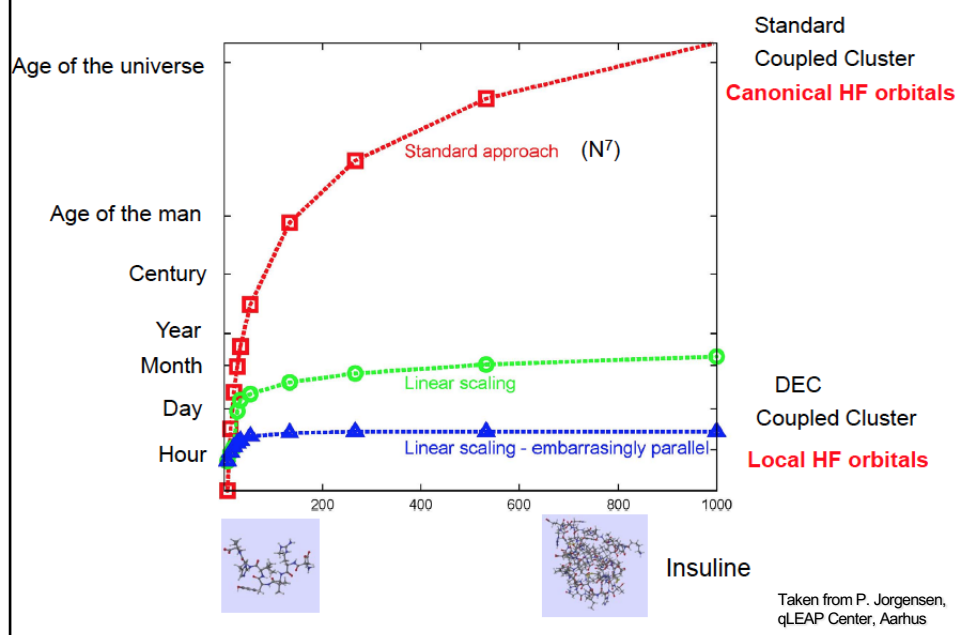
(from Wim Klopper and David P. Tew)

CCSD	N ⁶
CCSDT	N ⁸
CCSDTQ	N ¹⁰
CCSD(T)	N ⁷

Inclusion of triples via perturbation theory

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Scaling in Coupled Cluster theory CCSD(T)



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