Quantitative Quantum Chemistry

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High-accuracy quantum-chemical calculations

- The last decade has seen a dramatic improvement in our ability to treat molecular electronic systems accurately
  - development of techniques for systematic convergence towards the exact solution
  - extensive benchmarking on small and light molecular systems
- In many cases, we can now confidently confirm or reject experimental observations
- Many black-box methods have been developed
  - well-defined levels of theory
  - relatively easy to use by the nonspecialist
- Still, the exact solution can be approached in infinitely many ways
  - at many (incomplete) levels of theory, agreement with experiment may be obtained
  - such agreement always arises by error cancellation
  - error cancellation is treacherous: the right answer for the wrong reason
- For a reliable application of quantum chemistry, we must be wary of the pitfalls of error cancellation
- We shall here review high-accuracy quantum chemistry, paying special attention to the problem of error cancellation
Overview

• The many-electron problem
  – Hartree–Fock theory
  – electron correlation and virtual excitations
  – coupled-cluster theory
  – convergence to the “exact” solution

• Orbital expansions and basis sets
  – electron cusp and the Coulomb hole
  – basis-set convergence
  – basis-set extrapolation
  – explicit correlation

• Practical calculations and error cancellation
  – atomization energies
  – reaction enthalpies
  – bond distances
  – vibrational frequencies
The many-body problem of quantum chemistry

- All information about the electronic system is contained in the wave function, obtained by solving the Schrödinger equation.

> “The underlying physical laws necessary for the mathematical theory of a large part of physics and the whole of chemistry are thus completely known, and the difficulty is only that the exact application of these laws leads to equations much too complicated to be soluble.”

P. A. M. Dirac, 1929

- We are thus forced to make approximations—that is, set up simplified descriptions that incorporate the most important features of the system.

- This should preferably be done in an orderly fashion, so that the exact solution can be approached in a systematic manner.

- In this manner, we establish hierarchies of approximations—that is, a system of ever more accurate and expensive computational models.
The Hartree–Fock approximation

- The Hartree–Fock model—the fundamental approximation of wave-function theory
  - each electron moves in the mean field of all other electrons
  - provides an uncorrelated description: average rather than instantaneous interactions
  - gives rise to the concept of molecular orbitals
  - typical errors: 0.5% in the energy; 1% in bond distances, 5%–10% in other properties
  - forms the basis for more accurate treatments

- The Hartree–Fock and exact wave functions in helium:
  - concentric Hartree–Fock contours, reflecting an uncorrelated description
  - in reality, the electrons see each other and the contours becomes distorted
Electron correlation and virtual excitations

- **electron correlation:**
  - to improve upon the Hartree–Fock model, we must take into account the instantaneous interactions among the electrons
  - in real space, the electrons are constantly being scattered by collisions
  - in the orbital picture, these collisions manifest themselves as excitations from occupied to virtual (unoccupied) spin orbitals

- **double excitations:**
  - the most important events are collisions between two electrons
  - in the orbital picture, such an event corresponds to an excitation from two occupied to two virtual spin orbitals, known as pair excitations or double excitations

- Consider the following double-excitation operator:
  \[
  \hat{X}_{ij}^{ab} = t_{ij}^{ab} a_i^\dagger a_j^\dagger a_a^\dagger a_a
  \]
  - the amplitude \( t_{ij}^{ab} \) represents the probability that the electrons in \( \phi_i \) and \( \phi_j \) will interact and be excited to \( \phi_a \) and \( \phi_b \)
  - by applying \( 1 + \hat{X}_{ij}^{ab} \) to the Hartree–Fock state, we obtain an improved, correlated description of the electrons:
    \[
    |\text{HF}\rangle \rightarrow (1 + \hat{X}_{ij}^{ab})|\text{HF}\rangle
    \]
Example: electron correlation in H$_2$

- Consider the effect of a double excitation in H$_2$:

\[ |1\sigma_g^2\rangle \rightarrow (1 + \hat{X}_{gg}^{uu})|1\sigma_g^2\rangle = |1\sigma_g^2\rangle - 0.11|1\sigma_u^2\rangle \]

- The one-electron density $\rho(z)$ is hardly affected:

- The two-electron density $\rho(z_1, z_2)$ changes dramatically:
Coupled-cluster theory

- In coupled-cluster (CC) theory, we generate the correlated state from the HF reference state by applying all possible excitation operators

\[ |CC\rangle = (1 + \hat{X}^a_i) \cdots (1 + \hat{X}^{ab}_{ij}) \cdots (1 + \hat{X}^{abc}_{ijk}) \cdots (1 + \hat{X}^{abcd}_{ijkl}) \cdots |HF\rangle \]

- with each excitation, there is an associated probability amplitude \( t_{abc} \cdots \)
- single excitations represent orbital readjustments rather than direct interactions
- double excitations are particularly important, arising from pair interactions
- higher excitations should become progressively less important

- This classification provides a hierarchy of ‘truncated’ CC wave functions:
  - CCS, CCSD, CCSDT, CCSDTQ, CCSDTQ5, \ldots
  - errors are typically reduced by a factor of three to four at each new level

- Lower-order excitations work in tandem to produce higher-order excited configurations

\[ \left(1 + \hat{X}^{ab}_{ij}\right) \left(1 + \hat{X}^{cd}_{kl}\right) |HF\rangle = |HF\rangle + \hat{X}^{ab}_{ij} |HF\rangle + \hat{X}^{cd}_{kl} |HF\rangle + \hat{X}^{ab}_{ij} \hat{X}^{cd}_{kl} |HF\rangle \]

- the important thing is to parameterize the excitations rather than the resulting states
For the virtual excitations, we must create flexible virtual orbital spaces

- we expand the MOs in one-electron Gaussian-type functions (GTOs):

\[ G_{ijk}(r_A, \alpha) = x_A^i y_A^j z_A^k \exp \left( -\alpha r_A^2 \right) \]

Basis sets of increasing size:

- minimal or single-zeta (SZ) basis sets:
  * one set of GTOs for each occupied atomic shell (2s1p)
  * gives a rudimentary description of electron structure (no virtual space)

- double-zeta (DZ) basis sets:
  * two sets of GTOs for each occupied atomic shell (3s2p1d)
  * sufficient for a qualitative description of the electrons (minimal virtual space)

- triple-zeta (TZ), quadruple-zeta (QZ), and larger basis sets:
  * needed for a quantitative description of the electrons (flexible virtual space)

The number of GTOs per atom increases rapidly:

<table>
<thead>
<tr>
<th></th>
<th>SZ</th>
<th>DZ</th>
<th>TZ</th>
<th>QZ</th>
<th>5Z</th>
<th>6Z</th>
<th>X</th>
</tr>
</thead>
<tbody>
<tr>
<td>cardinal number</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
<td>X</td>
</tr>
<tr>
<td>number of AOs</td>
<td>5</td>
<td>14</td>
<td>30</td>
<td>55</td>
<td>91</td>
<td>140</td>
<td>X</td>
</tr>
</tbody>
</table>

\[ \propto X^3 \]

- large basis sets are needed to generate a flexible virtual space!
The quality of nonrelativistic molecular electronic-structure calculations is determined by the description of
1. the $N$-electron space (wave-function model),
2. the one-electron space (basis set).

In each space, there is a hierarchy of levels of increasing complexity:

1. the $N$-electron hierarchy:
   coupled-cluster excitation levels
   
   HF, CCSD, CCSDT, CCSDTQ, ...

2. the one-electron hierarchy:
   correlation-consistent basis sets

   DZ, TZ, QZ, 5Z, 6Z, ...

The quality is systematically improved upon by going up in the hierarchies.
Atomization energies (kJ/mol)

1. **CCSD(T)**
   - DZ
   - TZ
   - QZ
   - 5Z
   - 6Z

2. **CCSD**
   - DZ
   - TZ
   - QZ
   - 5Z
   - 6Z

3. **MP2**
   - DZ
   - TZ
   - QZ
   - 5Z
   - 6Z

4. **HF**
   - DZ
   - TZ
   - QZ
   - 5Z
   - 6Z
Coupled-cluster (CC) vs. basis-set (AO) convergence

- Atomization energies of HF, N₂, F₂, and CO
  - contributions of each CC excitation level (left) and AO basis-set shell (right)

  ![Graphs showing CC and AO convergence](image)

- The excitation-level convergence is approximately linear (log-linear plot)
  - each new excitation level reduces the error by about an order of magnitude
- The basis-set convergence is much slower (log-log plot)
  - each shell contributes an energy proportional to $X^{-4}$ where $X$ is the cardinal number
- Convergence is smooth in both directions!
  - we shall consider the CC convergence first, the AO convergence next
### CC convergence: atomization energies (kJ/mol)

<table>
<thead>
<tr>
<th></th>
<th>RHF</th>
<th>SD</th>
<th>T</th>
<th>Q</th>
<th>rel.</th>
<th>vib.</th>
<th>total</th>
<th>experiment</th>
<th>error</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₂</td>
<td>531.1</td>
<td>218.3</td>
<td>9.5</td>
<td>0.4</td>
<td>−0.7</td>
<td>−43.2</td>
<td>715.4</td>
<td>714.8±1.8</td>
<td>0.6</td>
</tr>
<tr>
<td>H₂O</td>
<td>652.3</td>
<td>305.3</td>
<td>17.3</td>
<td>0.8</td>
<td>−2.1</td>
<td>−55.4</td>
<td>918.2</td>
<td>917.8±0.2</td>
<td>0.4</td>
</tr>
<tr>
<td>HF</td>
<td>405.7</td>
<td>178.2</td>
<td>9.1</td>
<td>0.6</td>
<td>−2.5</td>
<td>−24.5</td>
<td>566.7</td>
<td>566.2±0.7</td>
<td>0.5</td>
</tr>
<tr>
<td>N₂</td>
<td>482.9</td>
<td>426.0</td>
<td>42.4</td>
<td>3.9</td>
<td>−0.6</td>
<td>−14.1</td>
<td>940.6</td>
<td>941.6±0.2</td>
<td>−1.1</td>
</tr>
<tr>
<td>F₂</td>
<td>−155.3</td>
<td>283.3</td>
<td>31.6</td>
<td>3.3</td>
<td>−3.3</td>
<td>−5.5</td>
<td>154.1</td>
<td>154.6±0.6</td>
<td>−0.5</td>
</tr>
<tr>
<td>CO</td>
<td>730.1</td>
<td>322.2</td>
<td>32.1</td>
<td>2.3</td>
<td>−2.0</td>
<td>−12.9</td>
<td>1071.8</td>
<td>1071.8±0.5</td>
<td>−0.0</td>
</tr>
</tbody>
</table>

- agreement with experiment in all cases except for N₂
- all purely electronic contributions are positive (except Hartree–Fock for F₂)
  - very large correlation contributions
- relativistic corrections are small (≈ 0.5%) but needed for agreement with experiment
  - nearly cancel quadruples contributions ([error cancellation possible](#))
- vibrational corrections are substantial
  - similar to triples contributions but oppositely directed ([error cancellation](#))
CC convergence: bond distances (pm)

<table>
<thead>
<tr>
<th></th>
<th>RHF</th>
<th>SD</th>
<th>T</th>
<th>Q</th>
<th>5</th>
<th>rel.</th>
<th>theory</th>
<th>exp.</th>
<th>err.</th>
</tr>
</thead>
<tbody>
<tr>
<td>HF</td>
<td>89.70</td>
<td>1.67</td>
<td>0.29</td>
<td>0.02</td>
<td>0.00</td>
<td>0.01</td>
<td>91.69</td>
<td>91.69</td>
<td>0.00</td>
</tr>
<tr>
<td>N₂</td>
<td>106.54</td>
<td>2.40</td>
<td>0.67</td>
<td>0.14</td>
<td>0.03</td>
<td>0.00</td>
<td>109.78</td>
<td>109.77</td>
<td>0.01</td>
</tr>
<tr>
<td>F₂</td>
<td>132.64</td>
<td>6.04</td>
<td>2.02</td>
<td>0.44</td>
<td>0.03</td>
<td>0.05</td>
<td>141.22</td>
<td>141.27</td>
<td>−0.05</td>
</tr>
<tr>
<td>CO</td>
<td>110.18</td>
<td>1.87</td>
<td>0.75</td>
<td>0.04</td>
<td>0.00</td>
<td>0.00</td>
<td>112.84</td>
<td>112.84</td>
<td>0.00</td>
</tr>
</tbody>
</table>

- agreement with experiment to within 0.01 pm except for F₂
- Hartree–Fock theory underestimates bond distances by up to 8.6 pm (for F₂)
- all correlation contributions are positive
  - approximate linear convergence, slowest for F₂
  - triples contribute up to 2.0 pm, quadruples up to 0.4 pm, and quintuples 0.03 pm
- relativistic corrections are small except for F₂ (0.05 pm)
  - of the same magnitude and direction as the quintuples contributions
  - no sources of error cancellation (in a complete AO basis)
CC convergence: harmonic constants $\omega_e$ (cm$^{-1}$)

<table>
<thead>
<tr>
<th></th>
<th>RHF</th>
<th>SD</th>
<th>T</th>
<th>Q</th>
<th>5</th>
<th>rel.</th>
<th>theory</th>
<th>exp.</th>
<th>err.</th>
</tr>
</thead>
<tbody>
<tr>
<td>HF</td>
<td>4473.8</td>
<td>−277.4</td>
<td>−50.2</td>
<td>−4.1</td>
<td>−0.1</td>
<td>−3.5</td>
<td>4138.5</td>
<td>4138.3</td>
<td>0.2</td>
</tr>
<tr>
<td>N$_2$</td>
<td>2730.3</td>
<td>−275.8</td>
<td>−72.4</td>
<td>−18.8</td>
<td>−3.9</td>
<td>−1.4</td>
<td>2358.0</td>
<td>2358.6</td>
<td>−0.6</td>
</tr>
<tr>
<td>F$_2$</td>
<td>1266.9</td>
<td>−236.1</td>
<td>−95.3</td>
<td>−15.3</td>
<td>−0.8</td>
<td>−0.5</td>
<td>918.9</td>
<td>916.6</td>
<td>2.3</td>
</tr>
<tr>
<td>CO</td>
<td>2426.7</td>
<td>−177.4</td>
<td>−71.7</td>
<td>−7.2</td>
<td>0.0</td>
<td>−1.3</td>
<td>2169.1</td>
<td>2169.8</td>
<td>0.7</td>
</tr>
</tbody>
</table>

- agreement with experiment to within 1 cm$^{-1}$ except for F$_2$
- Hartree–Fock theory overestimates harmonic frequencies by up to 38% (in F$_2$).
- all correlation contributions are large and negative
  - triples contribute up to 95 cm$^{-1}$, quadruples 20 cm$^{-1}$, and quintuples 4 cm$^{-1}$
  - sextuples are sometimes needed for convergence to within 1 cm$^{-1}$
- relativistic corrections are of the order of 1 cm$^{-1}$
  - of the same magnitude and direction as the quadruples or quintuples
- no sources of error cancellation in a complete AO basis
Many-body perturbation theory: approximate coupled-cluster theory

- Coupled-cluster amplitudes may be estimated by perturbation theory
- Caveat: the resulting perturbation series is frequently divergent, even in simple cases
  - here are some examples for the HF molecule (10 electrons):

- However, to lowest order, perturbational corrections are very useful and popular
  - MP2 (approximate CCSD) and CCSD(T) (approximate CCSDT)
  - correlation effects are typically overestimated, leading to fortuitously good results
Basis-set convergence

- In all examples up to now, we have worked in a complete AO basis.
- However, the overall quality is determined by the description of one/Minus electron basis sets.

1. the wave-function model
2. the AO basis set

- We shall now consider basis-set convergence:
  - new sources of errors
  - new opportunities for error cancellation

- We begin by investigating convergence in the helium atom.
AO convergence: the helium atom

- The helium atom contains only two electrons
  - ideal system to study basis-set convergence
  - pair interactions dominate also molecules

- Historical interest: the first many-body system treated with quantum mechanics
  - experimental ionization potential of helium: 24.59 eV

- Unsöld 1927: 20.41 eV
  - first-order perturbation theory—not much better than Bohr theory

- Hylleraas 1928: 24.47 eV
  - expansion in antisymmetric orbital products
  - excruciatingly slow AO convergence

- Hylleraas 1929: 24.58 eV
  - introduced the interelectronic coordinate $r_{12}$ to arbitrary powers
  - the discrepancy of 0.01 eV due to relativistic corrections
  - full agreement between experiment and quantum mechanics

- Hylleraas discovered both the slow CI convergence and the efficacy of introducing $r_{12}$
  - the question of CI expansions vs. explicit correlation is still with us today
The local kinetic energy

- Consider the local energy of the helium atom

\[ E_{\text{loc}} = (H\Psi)/\Psi \quad \text{← constant for exact wave function} \]

- The electronic Hamiltonian has singularities at points of coalescence

\[ H = -\frac{1}{2} \nabla^2_1 - \frac{1}{2} \nabla^2_2 - \frac{2}{r_1} - \frac{2}{r_2} + \frac{1}{r_{12}} \]

  - the infinite potential terms must be canceled by infinite kinetic terms at coalescence

- Local kinetic energy in the helium atom
  - positive around the nucleus
  - negative around the second electron

- Negative kinetic energy counterintuitive
  - classical forbidden region
  - internal “tunneling”
  - w. f. decays towards the singularity
  - the Coulomb hole

- The difficulty Hylleraas ran into was the description of the Coulomb hole!
The Coulomb hole: the forbidden region

- Each electron is surrounded by a classically forbidden region: the Coulomb hole
  - without a good description of this region, our results will be inaccurate
The electron cusp and the Coulomb hole

- The behaviour of the wave function at coalescence was established by Slater (1928)
  - nuclear and electronic cusp conditions

\[
\left( \frac{\partial \Psi}{\partial r_i} \right)_{r_i=0, \text{ave}} = -Z \Psi (r_i = 0) \\
\left( \frac{\partial \Psi}{\partial r_{12}} \right)_{r_{12}=0, \text{ave}} = \frac{1}{2} \Psi (r_{12} = 0)
\]

- The electronic cusp condition implies the existence of a Coulomb hole
  - the helium atom, with one electron fixed at a distance of 0.5\(a_0\) from the nucleus
Basis-set convergence

- Convergence of the contributions to the atomization energy of CO (kJ/mol):

<table>
<thead>
<tr>
<th></th>
<th>$N_{bas}$</th>
<th>$\text{HF (}n^4\text{)}$</th>
<th>$\text{SD (}n^6\text{)}$</th>
<th>$\text{(T)(}n^7\text{)}$</th>
<th>CCSD(T)</th>
<th>error</th>
</tr>
</thead>
<tbody>
<tr>
<td>cc-pCVDZ</td>
<td>36</td>
<td>710.2</td>
<td>+</td>
<td>277.4</td>
<td>+</td>
<td>24.5</td>
</tr>
<tr>
<td>cc-pCVTZ</td>
<td>86</td>
<td>727.1</td>
<td>+</td>
<td>297.3</td>
<td>+</td>
<td>32.6</td>
</tr>
<tr>
<td>cc-pCVQZ</td>
<td>168</td>
<td>730.3</td>
<td>+</td>
<td>311.0</td>
<td>+</td>
<td>33.8</td>
</tr>
<tr>
<td>cc-pCV5Z</td>
<td>290</td>
<td>730.1</td>
<td>+</td>
<td>316.4</td>
<td>+</td>
<td>34.2</td>
</tr>
<tr>
<td>cc-pCV6Z</td>
<td>460</td>
<td>730.1</td>
<td>+</td>
<td>318.8</td>
<td>+</td>
<td>34.4</td>
</tr>
<tr>
<td>limit</td>
<td>$\infty$</td>
<td>730.1</td>
<td>+</td>
<td>322.1</td>
<td>+</td>
<td>34.6</td>
</tr>
</tbody>
</table>

- the doubles converge very slowly—chemical accuracy requires 460 AOs (6Z)!
- the Hartree–Fock and triples contributions are less of a problem.

- The slow convergence arises from a poor description of short-range (dynamical) correlation in the orbital approximation (since $r_{ij}$ is not present in the wave function):
The principal expansion and correlation-consistent basis sets

- The energy contribution from each AO in large CI calculations on helium:

  \[ \varepsilon_{nlm} \approx n^{-6} \]

- The principal expansion: include all AOs belonging to the same shell simultaneously, in order of increasing principal quantum number \( n \):

  \[ \varepsilon_n \approx n^2 n^{-6} = n^{-4} \]

- Practical realization: the correlation-consistent basis sets cc-pV\(X\)Z (Dunning, 1989)
- Energy-optimized AOs are added one shell at a time:

  \[
  \begin{array}{cccc|c}
  \text{SZ} & \text{cc-pVDZ} & \text{cc-pVTZ} & \text{cc-pVQZ} & \text{number of AOs} \\
  2s1p & +3s3p3d & +4s4p4d4f & +5s5p5d5f5g & \propto X^2 \\
  3s2p1d & 4s3p2d1f & 5s4p3d2f1g & \propto X^3 \\
  \end{array}
  \]

- The error in the energy is equal to the contributions from all omitted shells:

  \[ \Delta E_X \approx \sum_{n=X+1}^{\infty} n^{-4} \approx X^{-3} \]
Some observations

- The dependence of the error in the correlation energy on the number of AOs $N$:

$$\Delta \varepsilon_N \propto X^{-3}$$
$$N \propto X^3$$

$$\Rightarrow \Delta \varepsilon_N \propto N^{-1}$$

- The dependence of the error in the correlation energy on the CPU time:

$$\Delta \varepsilon_N \propto N^{-1}$$
$$T \propto N^4$$

$$\Rightarrow \Delta \varepsilon_n \propto T^{-1/4}$$

- Each new digit in the energy therefore costs 10000 times more CPU time!

1 minute $\rightarrow$ 1 week $\rightarrow$ 200 years

- The convergence is exceedingly slow!

- Clearly, a brute-force extension of the basis set until convergence (to some target accuracy) may not always be possible.

- Fortunately, the convergence is very smooth.

- Careful analysis of the convergence of the partial-wave expansion: Schwartz (1962)
Extrapolations

- From two separate calculations with basis sets $E_X$ and $E_Y$

\[
E_\infty = E_X + AX^{-3}
\]
\[
E_\infty = E_Y + AY^{-3}
\]

we eliminate $A$ to obtain the following two-point extrapolation formula:

\[
E_\infty = \frac{X^3 E_X - Y^3 E_Y}{X^3 - Y^3}
\]

- Mean absolute error in the electronic energy of CH$_2$, H$_2$O, HF, N$_2$, CO, Ne, and F$_2$:

<table>
<thead>
<tr>
<th>mE$_h$</th>
<th>DZ</th>
<th>TZ</th>
<th>QZ</th>
<th>5Z</th>
<th>6Z</th>
<th>R12</th>
</tr>
</thead>
<tbody>
<tr>
<td>plain</td>
<td>194.8</td>
<td>62.2</td>
<td>23.1</td>
<td>10.6</td>
<td>6.6</td>
<td>1.4</td>
</tr>
<tr>
<td>extr.</td>
<td>21.4</td>
<td>1.4</td>
<td>0.4</td>
<td>0.5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- For the error in the AE of CO relative to R12, we now obtain:

<table>
<thead>
<tr>
<th>kJ/mol</th>
<th>DZ</th>
<th>TZ</th>
<th>QZ</th>
<th>5Z</th>
<th>6Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>plain</td>
<td>$-73.5$</td>
<td>$-28.3$</td>
<td>$-11.4$</td>
<td>$-6.0$</td>
<td>$-3.5$</td>
</tr>
<tr>
<td>extr.</td>
<td>$-18.5$</td>
<td>$-0.7$</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
</tr>
</tbody>
</table>

- Chemical accuracy is now achieved with just 168 AOs (QZ), at a fraction of the cost.
Explicitly correlated methods

- To improve basis-set convergence, we may use *explicitly correlated wave functions* (Hylleraas, 1928).
- In such wave functions, the interelectronic distances $r_{ij}$ are used as variables.
- In its simplest manifestation, only terms linear in $r_{ij}$ occur: the R12 method

$$\Psi_{R12} = \sum_K C_K \Phi_K + C_R r_{12} \Phi_0$$

- Helium-atom convergence of
  - the standard CI expansion
  - the CI-R12 expansion
  - the Hylleraas expansion

  on a logarithmic energy scale:

- Here we investigate the convergence of traditional, orbital-based wave functions as parametrized in coupled-cluster theory.
- However, we often use R12 calculations as benchmarks.
Extrapolation vs. explicit correlation

- For some purposes, extrapolation is competitive with explicit correlation
  - logarithmic errors in plain and extrapolated energies relative to R12
Neon: explicit correlation and extrapolation

aug–cc–pCVXZ
Changes upon basis-set extension

- When the basis-set increases, the energy is lowered most for systems of low electronic energy—that is, differences increase upon basis-set extension.

- This observation helps predict changes in properties upon basis-set extension:
  1. AEs increase since molecular energies are lower than atomic energies.
  2. Exothermic reactions become more exothermic.
  3. Polarizabilities increase since the energy is lower in the field.
  4. Bond distances shorten since shorter bonds give lower electronic energy.

- Equipped with a knowledge of convergence rate and direction, we shall now consider a number of molecular properties:
  1. Atomization energies
  2. Reaction enthalpies
  3. Molecular bond distances
  4. Two-electron Darwin term

- We shall now examine the convergence more carefully, with emphasis on
  - the interplay between excitation-level and basis-set convergence
  - error cancellations
Atomization energies (AEs)

- Let us consider the situation for an important molecular property: AEs

\[ D_e = \sum_A E^A (2S+1 L) - E_{\text{mol}} (R_e) \]

- Statistics based 20 closed-shell organic molecules (kJ/mol)

- AEs increase with cardinal number
- AEs increase with excitation level in the coupled-cluster hierarchy:

\[ \text{HF} < \text{CCSD} < \text{CCSD(T)} < \text{MP2} \]

- MP2 overestimates the doubles contribution
  - benefits from error cancellation at the MP2/TZ level

- CCSD(T) performs excellently in large basis sets
  - DZ and TZ basis are inadequate for CCSD(T)
Atomization energies (kJ/mol)

-200 to 200 kJ/mol for various levels of theory (HF, CCSD, CCSD(T), CCSD(T) with different basis sets: DZ, TZ, QZ, 5Z, 6Z).
Comparison of CCSD(T) and experimental AEs

<table>
<thead>
<tr>
<th></th>
<th>cc-pCVQZ</th>
<th>cc-pCV(TQ)Z</th>
<th>cc-pCV6Z</th>
<th>exp</th>
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<tr>
<td>F₂</td>
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<td>−15.4</td>
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<tr>
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<tr>
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<tr>
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<td>H₂O</td>
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<td>−11.8</td>
<td>974.7</td>
<td>−0.5</td>
</tr>
<tr>
<td>CO</td>
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<td>1086.6</td>
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<td>2343.6</td>
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<td>2363.2</td>
<td>3.4</td>
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</table>

- Only O₃ and HOF have an error larger than 2.2 kJ/mol.
- Without extrapolation: cc-p(c)V6Z needed; with extrapolation: cc-pCV(TQ)Z sufficient
- Core correlation important: F₂ – 0.1 kJ/mol; CO – 4.8 kJ/mol; C₂H₂ – 10.8 kJ/mol
The (in)adequacy of CCSD(T)

<table>
<thead>
<tr>
<th></th>
<th>CCSD(T)</th>
<th>CCSDT</th>
<th>CCSDTQ</th>
<th>experiment</th>
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<tr>
<td></td>
<td>cc-pCV(56)Z</td>
<td>cc-pCV(Q5)Z</td>
<td>cc-pVTZ</td>
<td>De</td>
</tr>
<tr>
<td>CH₂</td>
<td>757.9</td>
<td>-0.9</td>
<td>758.9</td>
<td>0.1</td>
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<tr>
<td>H₂O</td>
<td>975.3</td>
<td>0.1</td>
<td>974.9</td>
<td>-0.3</td>
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<tr>
<td>HF</td>
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<td>-0.2</td>
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<tr>
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<tr>
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<tr>
<td>CO</td>
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<td>1084.4</td>
<td>-2.3</td>
</tr>
</tbody>
</table>

- The excellent performance of CCSD(T) for AEs relies on error cancellation:
  - relaxation of triples from CCSD(T) to CCSDT reduces the AEs;
  - inclusion of quadruples from CCSDT to CCSDTQ increases the AEs.

- The error incurred by treating the connected triples perturbatively is quite large (about 10% of the full triples contribution) but canceled by the neglect of quadruples.

- The rigorous calculation of AEs to chemical accuracy requires CCSDTQ/cc-pCV6Z!
Reaction enthalpies

- Closed-shell, isogyric exothermic reactions

\[ \Delta_r H_e^\circ \, (0K) = \sum_P E^P \, (R_e) - \sum_R E^R \, (R_e) \]

- Convergence faster than for AEs but still governed by short-range correlation:

<table>
<thead>
<tr>
<th></th>
<th>CCSD(T)</th>
<th>DZ</th>
<th>TZ</th>
<th>QZ</th>
<th>5Z</th>
<th>6Z</th>
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</thead>
<tbody>
<tr>
<td>REs plain</td>
<td>33.4</td>
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<td>0.4</td>
<td>-0.1</td>
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<tr>
<td>extr.</td>
<td>2.7</td>
<td>-1.1</td>
<td>-0.8</td>
<td>-0.3</td>
<td></td>
<td></td>
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<tr>
<td>AEs plain</td>
<td>-103.3</td>
<td>-34.9</td>
<td>-14.3</td>
<td>-7.4</td>
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<td></td>
</tr>
<tr>
<td>extr.</td>
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<td>-1.1</td>
<td>-0.6</td>
<td>-0.9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- Errors relative to experimental results corrected for vibrations and relativity:

**Mean absolute errors:**

- **CCSD(T):** 1.8 kJ/mol
- **CCSD:** 14.5 kJ/mol
- **MP2:** 13.5 kJ/mol
- **HF:** 43.0 kJ/mol
<table>
<thead>
<tr>
<th>Reaction</th>
<th>exp.</th>
<th>R12</th>
<th>G2</th>
<th>B3LYP</th>
<th>(56)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₂ + H₂ → CH₄</td>
<td>-544(2)</td>
<td>-542</td>
<td>-534</td>
<td>-543</td>
<td>-543</td>
</tr>
<tr>
<td>C₂H₂ + H₂ → C₂H₄</td>
<td>-203(2)</td>
<td>-204</td>
<td>-202</td>
<td>-208</td>
<td>-206</td>
</tr>
<tr>
<td>C₂H₂ + 3H₂ → 2CH₄</td>
<td>-446(2)</td>
<td>-447</td>
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<td>-450</td>
<td>-447</td>
</tr>
<tr>
<td>CO + H₂ → CH₂O</td>
<td>-21(1)</td>
<td>-22</td>
<td>-17</td>
<td>-34</td>
<td>-23</td>
</tr>
<tr>
<td>N₂ + 3H₂ → 2NH₂</td>
<td>-164(1)</td>
<td>-162</td>
<td>-147</td>
<td>-166</td>
<td>-165</td>
</tr>
<tr>
<td>F₂ + H₂ → 2HF</td>
<td>-563(1)</td>
<td>-562</td>
<td>-564</td>
<td>-540</td>
<td>-564</td>
</tr>
<tr>
<td>O₃ + 3H₂ → 3H₂O</td>
<td>-933(2)</td>
<td>-943</td>
<td>-912</td>
<td>-909</td>
<td>-946</td>
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<tr>
<td>CH₂O + 2H₂ → CH₄ + H₂O</td>
<td>-251(1)</td>
<td>-250</td>
<td>-235</td>
<td>-234</td>
<td>-250</td>
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<tr>
<td>H₂O₂ + H₂ → 2H₂O</td>
<td>-365(2)</td>
<td>-365</td>
<td>-360</td>
<td>-346</td>
<td>-362</td>
</tr>
<tr>
<td>CO + 3H₂ → CH₄ + H₂O</td>
<td>-272(1)</td>
<td>-272</td>
<td>-251</td>
<td>-268</td>
<td>-273</td>
</tr>
<tr>
<td>HCN + 3H₂ → CH₄ + NH₂</td>
<td>-320(3)</td>
<td>-320</td>
<td>-305</td>
<td>-320</td>
<td>-321</td>
</tr>
<tr>
<td>HNO + 2H₂ → H₂O + NH₂</td>
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<td>-445</td>
<td>-426</td>
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<td>-446</td>
</tr>
<tr>
<td>H₂O + F₂ → HOF + HF</td>
<td>-129(4)</td>
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<td>-123</td>
<td>-119</td>
<td>-118</td>
</tr>
<tr>
<td>CO₂ + 4H₂ → CH₄ + 2H₂O</td>
<td>-244(1)</td>
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<td>-216</td>
<td>-211</td>
<td>-244</td>
</tr>
<tr>
<td>2CH₂ → C₂H₄</td>
<td>-844(3)</td>
<td>-842</td>
<td>-829</td>
<td>-845</td>
<td>-845</td>
</tr>
</tbody>
</table>
Bond distances

- Statistics based on 28 bond distances at the all-electron cc-pVXZ level (pm):

  - bonds shorten with increasing basis: DZ > TZ > QZ
  - bonds lengthen with increasing excitations: HF < CCSD < MP2 < CCSD(T)
  - considerable scope for error cancellation: CISD/DZ, MP3/DZ
  - CCSD(T) mean errors: DZ: 1.68 pm; TZ: 0.01 pm; QZ: −0.12 pm
Bond distances $R_e$ of BH, CO, N$_2$, HF, and F$_2$ (cm$^{-1}$)

- SCF cc-pCVDZ: 2.5
- SCF cc-pCVTZ: 3.4
- SCF cc-pCVQZ: 3.5
- SCF cc-pCV5Z: 3.5
- CCSD/cc-pCVDZ: 1.2
- CCSD/cc-pCVTZ: 0.6
- CCSD/cc-pCVQZ: 0.9
- CCSD/cc-pCV5Z: 1.0
- MP2 cc-pCVDZ: 1.5
- MP2 cc-pCVTZ: 0.9
- MP2 cc-pCVQZ: 0.8
- MP2 cc-pCV5Z: 0.8
- CCSD(T) cc-pCVDZ: 2.1
- CCSD(T) cc-pCVTZ: 0.2
- CCSD(T) cc-pCVQZ: 0.1
- CCSD(T) cc-pCV5Z: 0.1
## Bond distances

<table>
<thead>
<tr>
<th></th>
<th>HF</th>
<th>MP2</th>
<th>CCSD</th>
<th>CCSD(T)</th>
<th>emp.</th>
<th>exp.</th>
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<tr>
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</table>
Bond distances: CCSD(T) error cancellation

- CCSD(T) distances compared with experiment:

<table>
<thead>
<tr>
<th>pm</th>
<th>DZ</th>
<th>TZ</th>
<th>QZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>∆</td>
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<td>0.01</td>
<td>−0.12</td>
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<tr>
<td></td>
<td>Δ</td>
<td>1.68</td>
<td>0.20</td>
</tr>
</tbody>
</table>

- The high accuracy of CCSD(T) arises partly because of error cancellation.

- Bond distances are further reduced by
  - basis-set extension QZ → 6Z: ≈ −0.10 pm
  - triples relaxation CCSD(T) → CCSDT: ≈ −0.04 pm

- Intrinsic error of the CCSDT model: ≈ −0.2 pm

- Connected quadruples increase bond lengths by about 0.1–0.2 pm
Error cancellation: some general considerations

- In an infinite basis, the CC hierarchy is related to experiment as follows:
  - \( \text{HF} < \text{CCSD} < \text{CCSDT} < \text{CCSDTQ} < \cdots < \text{FCI} = \exp \) (BDs and AEs)
  - no error cancellation may occur (ignoring relativity and adiabatic corrections)

- In a finite basis, results are shifted and cancellation occurs for BDs but not for AEs:
  - \( \text{HF} < \text{CCSD} \approx \exp < \text{CCSDT} < \text{CCSDTQ} < \cdots < \text{FCI} \) (BDs shifted up)
  - \( \text{HF} < \text{CCSD} < \text{CCSDT} < \text{CCSDTQ} < \cdots < \text{FCI} < \exp \) (AEs shifted down)

- Perturbative treatments of connected excitations tend to overestimate their effect:
  - \( \text{HF} < \text{CCSD} < \text{MP2} \) ← (over)compensates for lack of triples
  - \( \text{CCSD} < \text{CCSDT} < \text{CCSD(T)} \) ← compensates for lack of quadruples
  - a new source of error cancellation

- For BDs and AEs, we now observe
  - \( \text{HF} < \text{CCSD} < \text{MP2} \approx \text{CCSD(T)} \approx \exp \) (BDs)
  - \( \text{HF} < \text{CCSD} < \text{CCSD(T)} \approx \exp < \text{MP2} \) (AEs)
  - error cancellation may now occur also for AEs

- We shall see that vibrational frequencies behave in much the same manner as BDs since basis-set extension and increased excitation levels work in opposite directions
Vibrational frequencies of diatoms

- The frequency of a diatomic fundamental transition is given by

\[ \nu = \omega_e - 2\omega_ex_e \]

- the harmonic constant \( \omega_e \) requires 2nd derivatives of PES
- the anharmonic constant \( \omega_e x_e \) requires 4th derivatives of PES

- With the advent of CCSD(T) in the 1990s, it was soon realized that this model is capable of highly accurate vibrational constants (to within a few wavenumbers at the TZ level)

- However, with the development of codes capable of handling very large basis sets and high excitation levels, it has slowly transpired that things are perhaps not so simple

- It is easier to converge harmonic than anharmonic constants:

| \( |\Delta| \) (cm\(^{-1}\)) | RHF | MP2 | CCSD | CCSD(T) | judgment |
|----------------|-----|-----|------|--------|----------|
| \( \omega_e \)  | 287 | 71  | 72   | 10     | DIFFICULT! |
| \( \omega_e x_e \) | 4   | 3   | 1    | 0      | EASY!    |
Harmonic constants $\omega_e$ of BH, CO, N$_2$, HF, and F$_2$ (cm$^{-1}$)

<table>
<thead>
<tr>
<th>Method</th>
<th>Basis Set</th>
<th>BH</th>
<th>CO</th>
<th>N$_2$</th>
<th>HF</th>
<th>F$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCSD(T) cc-pCVDZ</td>
<td>-250</td>
<td>42</td>
<td>14</td>
<td>9</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>MP2 cc-pCVDZ</td>
<td>-250</td>
<td>68</td>
<td>81</td>
<td>73</td>
<td>71</td>
<td></td>
</tr>
<tr>
<td>SCF cc-pCVDZ</td>
<td>-250</td>
<td>269</td>
<td>288</td>
<td>287</td>
<td>287</td>
<td></td>
</tr>
</tbody>
</table>
Anharmonic constants $\omega_e x_e$ of BH, CO, N$_2$, HF, and F$_2$ (cm$^{-1}$)
Higher-order connected contributions to $\omega_e$ in $\text{N}_2$

- There are substantial higher-order corrections:

<table>
<thead>
<tr>
<th>Method</th>
<th>Contribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>HF</td>
<td>371.9</td>
</tr>
<tr>
<td>CCSD/FC</td>
<td>84.6</td>
</tr>
<tr>
<td>CCSD(T)/FC</td>
<td>4.1</td>
</tr>
<tr>
<td>CCSD(T)</td>
<td>13.8</td>
</tr>
<tr>
<td>CSDT</td>
<td>23.5</td>
</tr>
<tr>
<td>CCSDTQ</td>
<td>4.7</td>
</tr>
<tr>
<td>CSDTQ5</td>
<td>0.8</td>
</tr>
</tbody>
</table>

- Connected triples relaxation contributes 9.7 cm$^{-1}$ (total triples $-70.5$ cm$^{-1}$)
- Connected quadruples contribute $-18.8$ cm$^{-1}$
- Connected quintuples contribute $-3.9$ cm$^{-1}$
**Harmonic frequencies of HF, N$_2$, F$_2$, and CO**

- Harmonic frequencies were obtained in the following manner:

<table>
<thead>
<tr>
<th>Method</th>
<th>HF</th>
<th>N$_2$</th>
<th>F$_2$</th>
<th>CO</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCSD-R12</td>
<td>4191.0</td>
<td>2443.2</td>
<td>1026.5</td>
<td>2238.5</td>
</tr>
<tr>
<td>CCSD(T)–CCSD</td>
<td>−48.4</td>
<td>−80.6</td>
<td>−95.7</td>
<td>−71.5</td>
</tr>
<tr>
<td>aug-cc-pV6Z/FC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CCSDTQ–CCSD(T)</td>
<td>−4.5</td>
<td>−9.1</td>
<td>−12.2</td>
<td>−6.5</td>
</tr>
<tr>
<td>cc-pVTZ/FC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CCSDTQ5–CCSDTQ</td>
<td>−0.1</td>
<td>−3.9</td>
<td>−0.8</td>
<td>0.0</td>
</tr>
<tr>
<td>cc-pVDZ/FC</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CCSD(T) core correlation</td>
<td>4.0</td>
<td>9.8</td>
<td>1.6</td>
<td>9.9</td>
</tr>
<tr>
<td>aug-cc-pCV5Z</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>relativistic correction</td>
<td>−3.5</td>
<td>−1.4</td>
<td>−0.5</td>
<td>−1.3</td>
</tr>
<tr>
<td>adiabatic correction</td>
<td>0.4</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>theory</td>
<td>4138.9</td>
<td>2358.0</td>
<td>918.9</td>
<td>2169.1</td>
</tr>
<tr>
<td>experiment</td>
<td>4138.3</td>
<td>2358.6</td>
<td>916.6</td>
<td>2169.8</td>
</tr>
<tr>
<td>error</td>
<td>0.1</td>
<td>−0.6</td>
<td>2.3</td>
<td>0.7</td>
</tr>
</tbody>
</table>

- it does not seem possible yet to obtain harmonic frequencies to within 1 cm$^{-1}$
- error in F$_2$ may be explained by a too short bond distance (141.19 vs. 141.27 pm)
There are two distinct contributions to the vibrational frequency:
- a dominant positive nuclear contribution and a negative electronic contribution

Frequencies calculated at the **optimized geometry:**
- basis-set extension reduces the bond length
- the nuclear contribution increases more than the electronic decreases
- basis-set extension therefore increases the frequency

Frequencies calculated at the **experimental geometry:**
- the frequency is improved since the nuclear contribution is now exact
- basis-set extension reduces the frequency by reducing the electronic contribution
Harmonic frequencies at experimental geometry

- Frequencies are significantly improved when calculated at the experimental geometry.
  - equivalent to the addition of an empirical linear term to the force field

- Frequencies calculated at [optimized] and [experimental] geometries

- frequencies typically too large at optimized geometries (except in DZ basis)
- use of experimental geometries improves frequencies by extending bond lengths
- this is particularly efficacious at low levels of theory (HF, MP2, CCSD)
Conclusions

- Quantum chemistry provides a set of well-defined levels of approximation
  - excitation-level expansion
  - basis-set expansion
- The “exact” result can be approached in a systematic manner
- In practice, we have to be content with low levels of theory
  - low excitation levels, approximated by perturbation theory
  - small basis sets
- Sometimes complicated interplay between different approximations
  - useful and reliable (balanced) levels of theory have been implemented in codes
  - can be usefully applied without deep knowledge of quantum chemistry
- Still, some knowledge of these methods, their errors and the interplay of these errors, will make the application of these methods more reliable
- Discipline is always needed!