

Lecture #6 of 14

(3: TThF, 5: MT*W*ThF, 4: MT*W*Th, 2: TW)

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Chemical Properties

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Chemical Properties

- Molecular nomenclature, Solutions, Balanced chemical reactions
- State functions, Standard states, Thermochemistry
- Non-ideal gases, Intermolecular forces, Physical properties, Phase changes, Colligative properties, Water activity
- Free energy, (X)Chemical potential, Entropic "force", Activity coefficients, Chemical equilibrium, van't Hoff equation, Le Chatelier's principle
- Schrödinger equation, Internal energy, Atomic orbitals, Hybridization
- Valence bond theory, Molecular orbital theory, Band diagrams
- Crystal field theory, Ligand field theory

(REVIEW... and then let's do all slides after this one again, just in case) 112 Schrödinger Equation

Elegant master equation that allows one to determine internal energies, E_n , of a system

$$\widehat{\boldsymbol{H}}\psi_n(x) = \mathbf{E}_n\psi_n(x)$$

... but this is not good enough for photochemists, or physicists, where <u>time-varying</u> oscillating electromagnetic fields often interact with matter... $E = hv = \hbar\omega$ (Planck)

$$\widehat{H}\Psi_n(x,t) = i\hbar \frac{\partial}{\partial t} \Psi_n(x,t)$$
 $E = mc^2 = pc$ (Einstein) $p = \frac{\hbar v}{c} = \frac{\hbar}{\lambda} = \hbar \overline{v} = \hbar k$ (de Broglie)

... so, how does one solve either of these Schrödinger equations?... We need to know \widehat{H} !

$$\widehat{H} = \widehat{T} + \widehat{V}$$

... um... well that didn't really help us at all... anyway, so instead, we need to know \hat{T} and \hat{V} ?

$$\widehat{T}(x) = KE = \frac{1}{2}m\widehat{v}^2 = \frac{\widehat{p}^2}{2m}... \text{ with } (\widehat{p}) \text{momentum} = (m) \text{ass x } (\widehat{v}) \text{elocity} = -i\hbar \frac{\partial}{\partial x}...$$

$$\hat{V}(x) = PE = 0...$$
 for particle-in-a-box...

$$\hat{V}(x) = \frac{1}{2}k_fx^2 = \frac{1}{2}m\omega^2x^2$$
... for harmonic oscillator, with k_f (force const), ω (angular freq)...

$$\hat{V}(r) = -\frac{q^2}{4\pi\varepsilon_0 r}$$
... for Hydrogen atom, with $F = qN_{\rm A}$ (Faraday const), ε_0 (vacuum permittivity)

... let's examine the hydrogen atom in more detail... because it seems like it may be fairly important to chemists ©

Solving the Schrödinger Equation

So, let's solve the Schrödinger equation using the Hamiltonian for the hydrogen(ic) atom...

$$\hat{H} = \hat{E}_{\text{K,electron}} + \hat{E}_{\text{K,nucleus}} + \hat{V}$$

$$= -\frac{\hbar^2}{2m_e} \nabla_e^2 - \frac{\hbar^2}{2m_N} \nabla_N^2 - \frac{Ze^2}{4\pi\epsilon_0 r}$$

... which can be written for the motion of the electron with respect to the nucleus using the reduced mass, μ , as...

$$-\frac{\hbar^{2}}{2\mu}\nabla^{2}\psi - \frac{Ze^{2}}{4\pi\varepsilon_{0}r}\psi = E\psi \qquad \frac{1}{\mu} = \frac{1}{m_{e}} + \frac{1}{m_{N}}$$

... the solutions to this equation are separable into radial (R) and angular (Y) components...

... and thus two equations ...
$$\Lambda^2 Y = -l(l+1)Y$$

$$\psi(r,\theta,\phi) = R(r)Y(\theta,\phi)$$

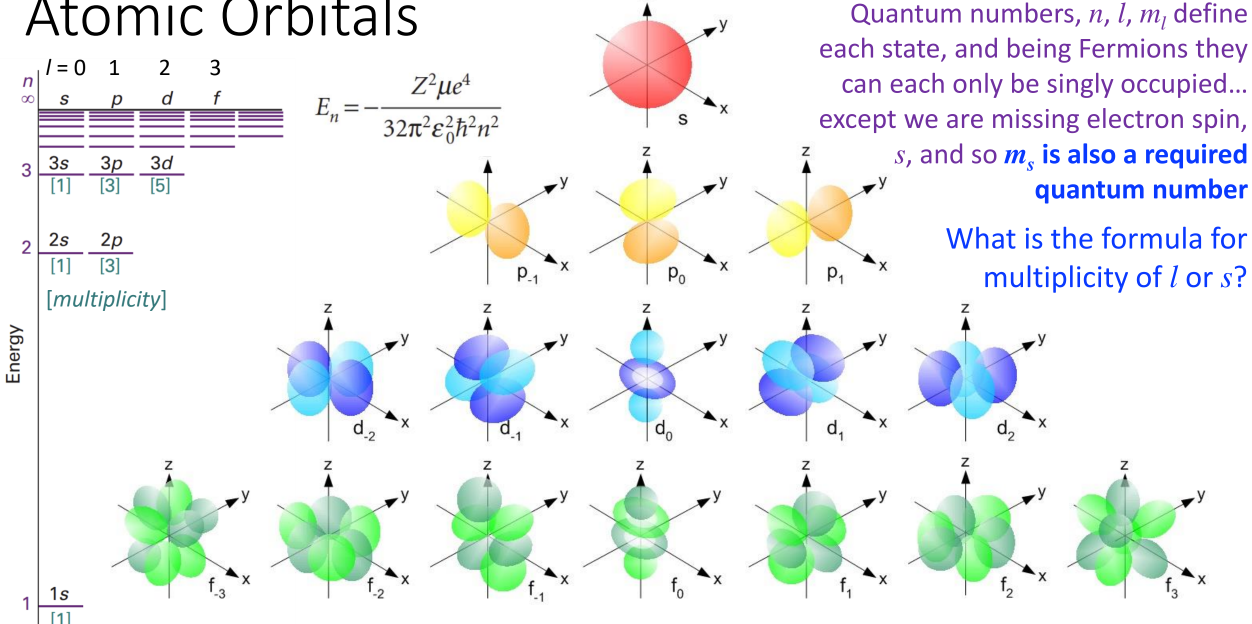
$$-\frac{\hbar^2}{2\mu}\frac{\mathrm{d}^2 u}{\mathrm{d}r^2} + V_{\text{eff}}u = Eu \qquad u = rR \qquad V_{\text{eff}} = -\frac{Ze^2}{4\pi\varepsilon_0 r} + \frac{l(l+1)\hbar^2}{2\mu r^2}$$

... solving the bottom radial equation gives energy eigenvalues, E_n , with $n \ge 1$...

... and solving the top spherical harmonics equation gives angular momentum eigenvalues,

$$l = [0, ..., n-2, n-1]$$
, with discrete projections on the z-axis, $m_l = [-l, -l+1, ..., 0, ..., l-1, l]$

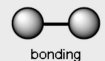
Atomic Orbitals

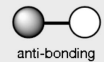


https://chembamm.files.wordpress.com/2015/09/single_electron_orbitals.jpg

Valence Bond Theory (and Orbital Hybridization)

The bonding nature of an orbital interaction is defined by the relative orbital phasing





The type of bonding is defined by the number of nodes parallel to the bond





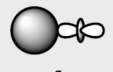






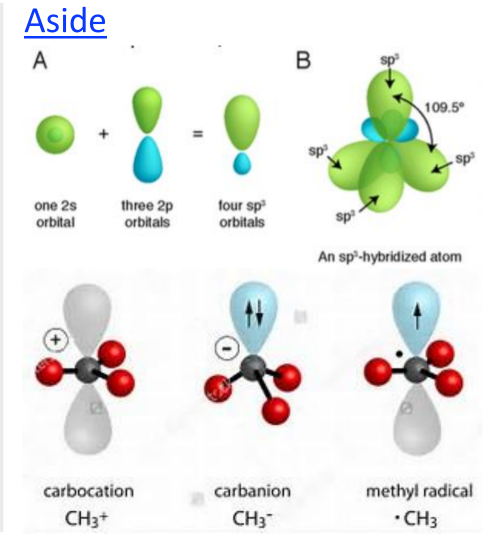


Interactions between different types of orbitals is okay as long as there is net overlap





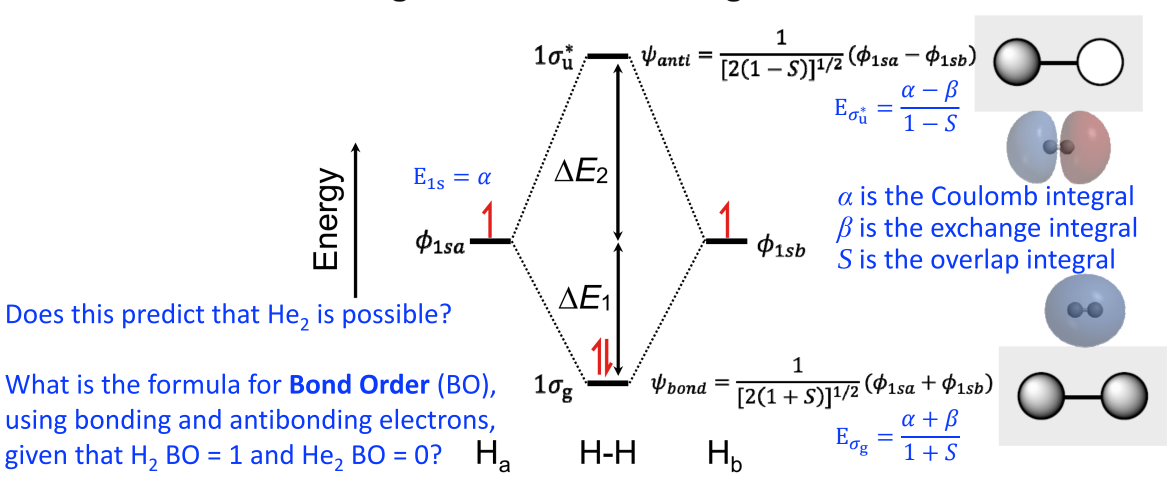




https://quizlet.com/190582013/1-6-sp3-hybrid-orbitals-and-the-structure-of-methane-flash-cards/https://socratic.org/questions/598bfedd7c01494e88568d61

Molecular Orbital Theory

H₂ molecule: two 1s atomic orbitals combine to make one bonding and one antibonding molecular orbital.



... just take Linear Combinations, ψ , of Atomic Orbitals, ϕ , (LCAO)

Molecular Orbital Theory

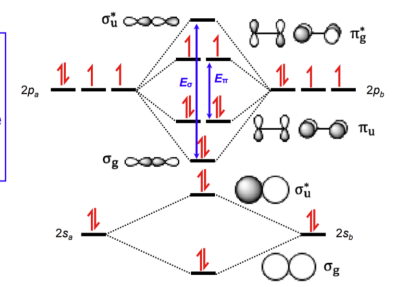
Homonuclear Diatomic Molecules

What happens when we move to more complicated systems? Consider O₂.

• The Lewis dot structure famously predicts the wrong electronic structure for O₂

• We can use LCAO-MO theory to get a better picture:

Notice that $E_{\sigma} > E_{\pi}$, because the σ bonds have more overlap than π bonds

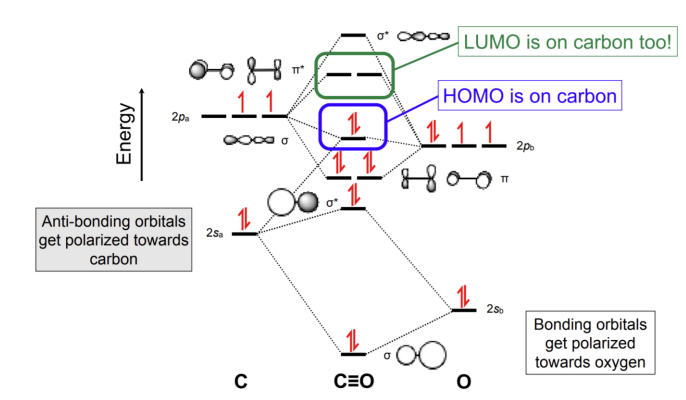


What is the Bond Order for O_2 ? What do the bonds look like?

What is the Bond Order for CO?

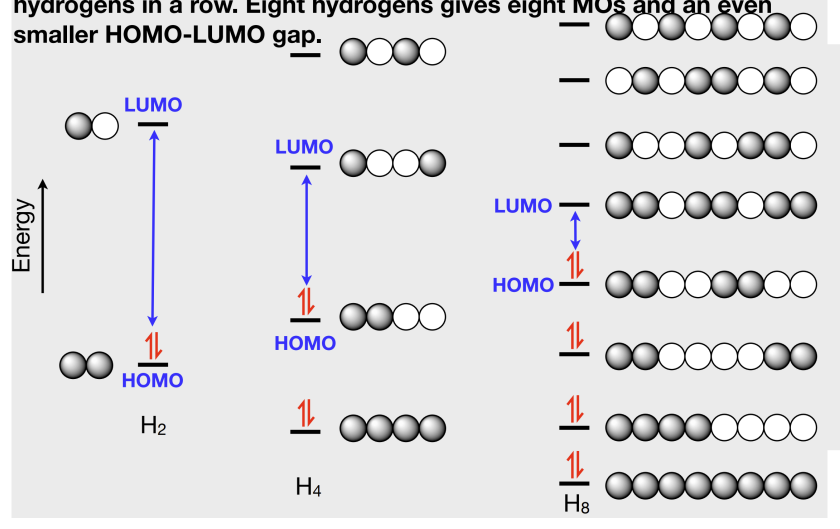
Heteronuclear Diatomic Molecules: CO

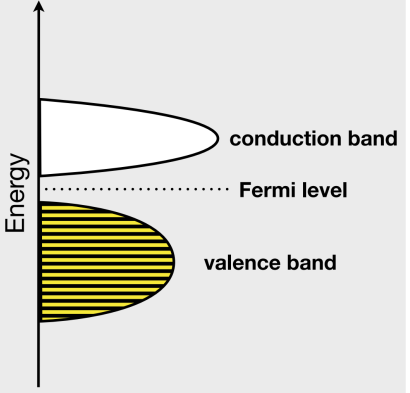
In molecules with more than one type of atom, MOs are formed from AOs that have different energies. Consider CO:



Band Diagrams

Look at what happens when we move from two hydrogens to four hydrogens in a row. Eight hydrogens gives eight MOs and an even





Bonds are delocalized over the entire "molecule" in this ideal representation

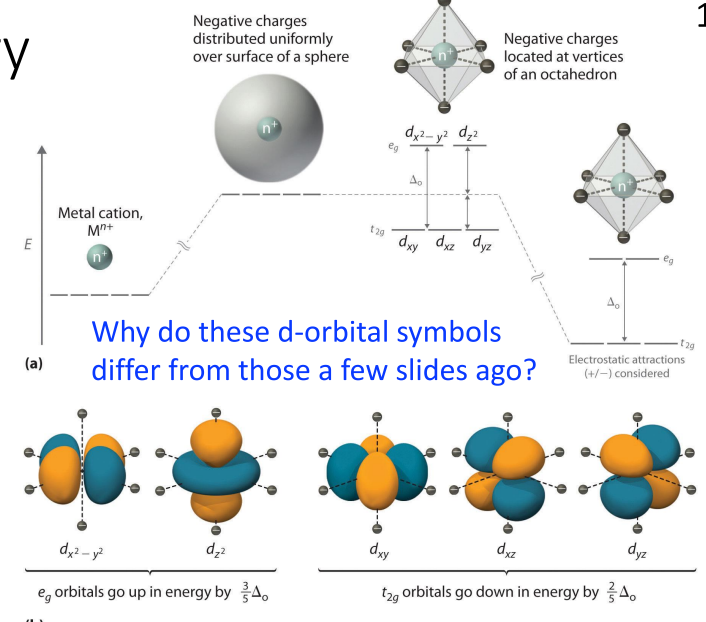
Crystal Field Theory

Crystal field stabilization energy (CFSE) = Δ_0 ...

... electron occupancy can either be **low spin** or **high spin** depending on the size of CFSE *vs.* the electron pairing energy...

... why do some orbitals go up in energy, and others down?

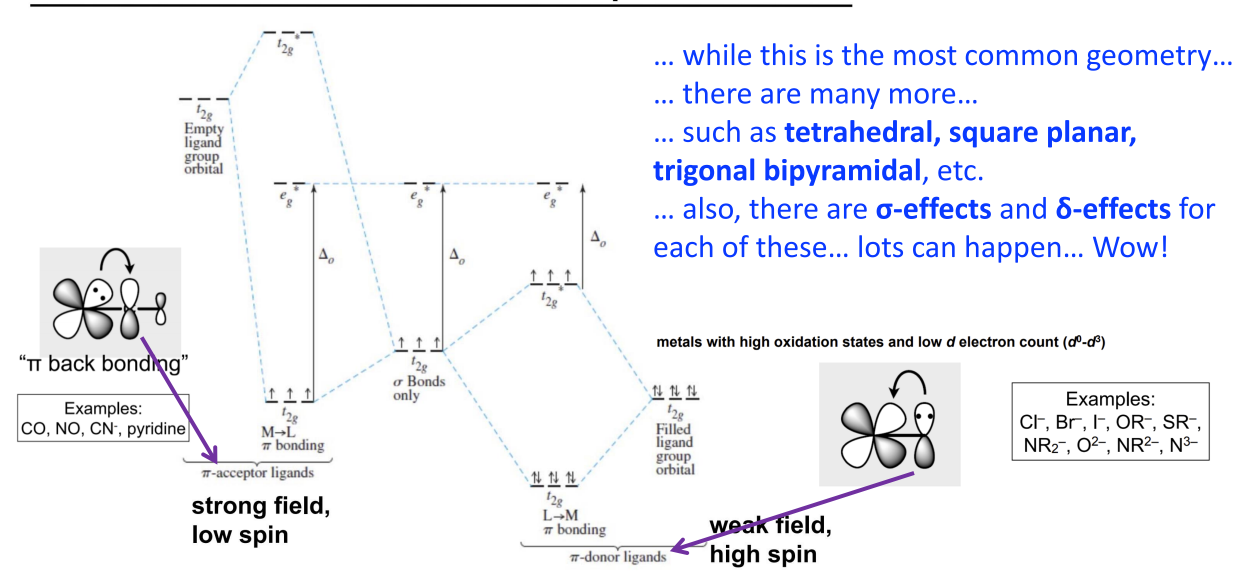
... moderately accurately predicts colors and magnetism of many molecules and materials



https://chem.libretexts.org/Courses/Douglas College/DC%3A Chem 2330 (O'Connor)/4%3A Crys tal Field Theory/4.3%3A High Spin and Low Spin Complexes

Ligand Field Theory

π-Effects in Octahedral Complexes



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