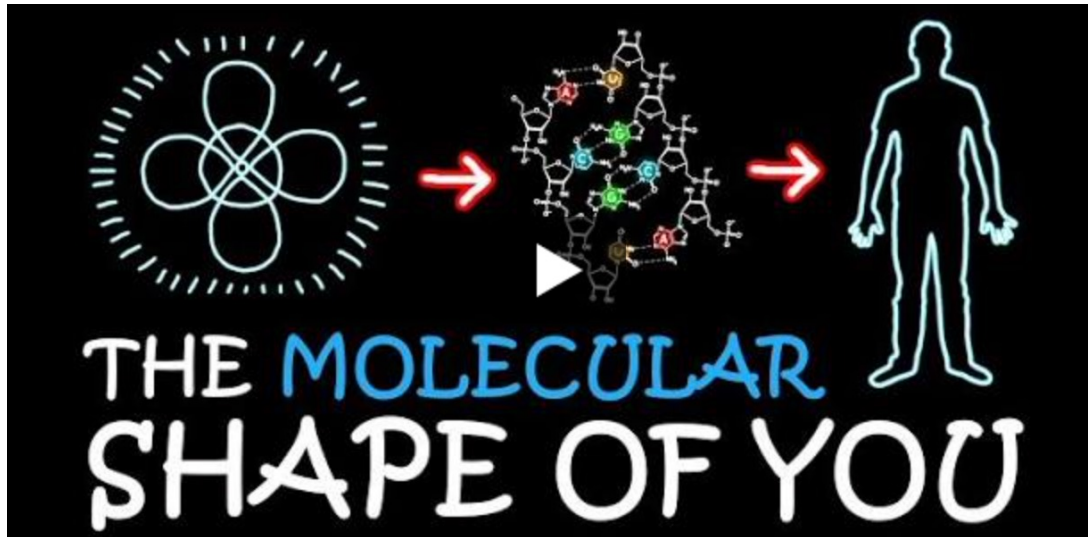


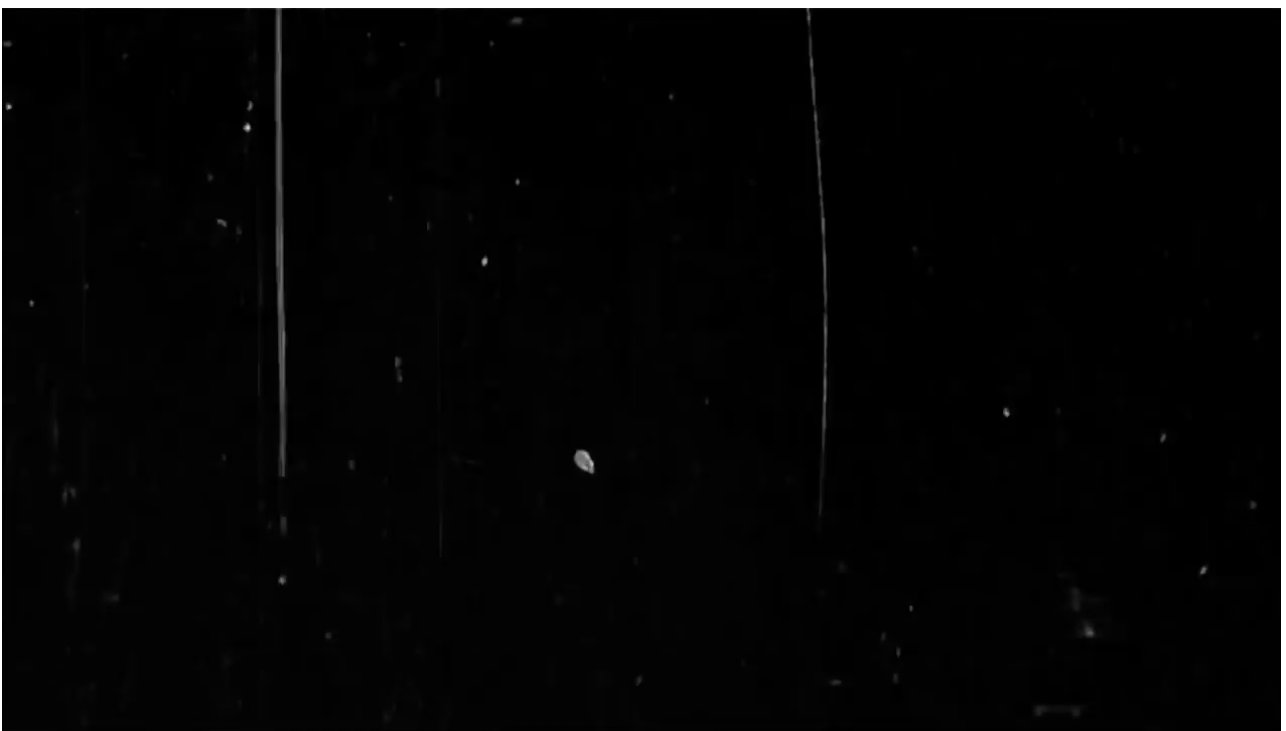
**MSE 423 Fall 2024 – Week 7**

# From atoms to molecules



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**MSE 423 Fall 2024 – Week 7**



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## Last week

- Two-electron atom and many-electron atom
- Energy of a collection of atoms
- Hartree ansatz and equations
- Spin-statistics connection
- Slater determinants and Hartree-Fock equations
- Orbital levels in multi-electron atoms
- Pauli exclusion principle
- Auf-bau principle and the periodic table
- Spin orbitals

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Exam: (double check in Dec/Jan):  
Monday 20 Jan 2025  
15h15 to 18h15 (AAC231)



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# Linear Combination of Atomic Orbitals

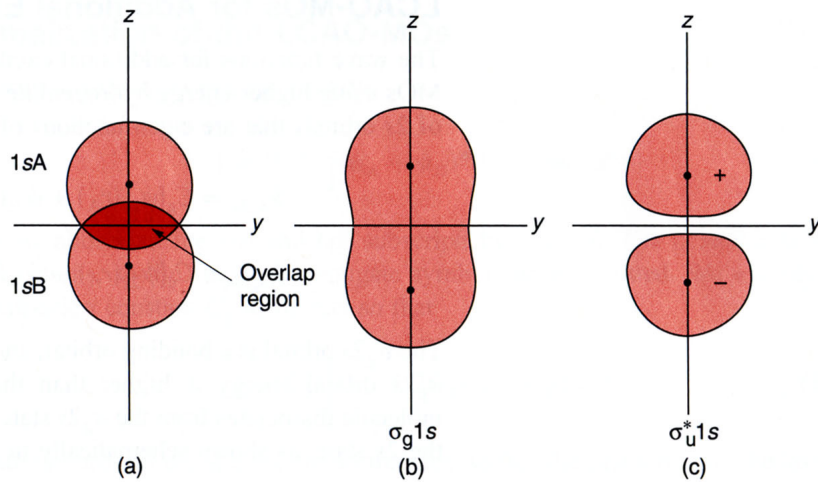
- Most common approach to find out the ground-state solution – it allows a meaningful definition of “hybridization”, “bonding” and “anti-bonding” orbitals.
- Also known as LCAO, LCAO-MO (for molecular orbitals), or tight-binding (for solids)
- Trial wavefunction is a linear combination of atomic orbitals – the variational parameters are the coefficients:

$$\Psi_{trial} = c_1 \Psi_{1s}(\vec{r} - \vec{R}_{H_1}) + c_2 \Psi_{1s}(\vec{r} - \vec{R}_{H_2})$$

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## How do we find the two coefficients?

# Bonding and Antibonding (I)



**Figure 18.7 The Orbital Region for the  $\sigma_g 1s$  and  $\sigma_u^* 1s$  LCAO Molecular Orbitals.** (a) The overlapping orbital regions of the  $1s_A$  and  $1s_B$  atomic orbitals. (b) The orbital region of the  $\sigma_g 1s$  LCAO-MO. (c) The orbital region of the  $\sigma_u^* 1s$  LCAO-MO. The orbital regions of the LCAO molecular orbitals have the same general features as the "exact" Born Oppenheimer orbitals whose orbital regions were depicted in Figure 18.4.

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## Formation of a Bonding Orbital

<http://winter.group.shef.ac.uk/orbitron/>



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## Formation of a Bonding Orbital



## Formation of an Antibonding Orbital

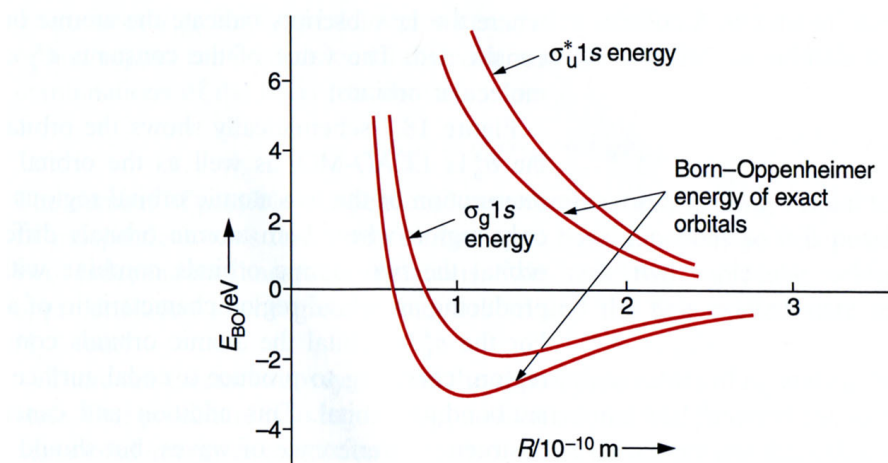


# Formation of an Antibonding Orbital



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## Bonding and Antibonding (II)



**Figure 18.8 The Orbital Energies for the  $\sigma_g 1s$  and  $\sigma_u^* 1s$  LCAO Molecular Orbitals.** This diagram shows qualitatively how the Born–Oppenheimer energies of the LCAO molecular orbitals compare with the Born–Oppenheimer energies of the “exact” orbitals. The approximate orbital energies must lie above the corresponding exact energies for all internuclear distances.

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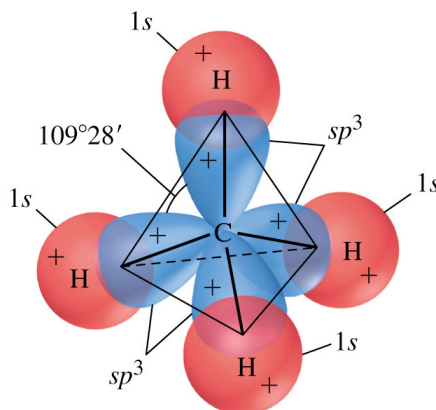
## More complex molecules: LCAO

$$\Psi_{trial} = \sum_{I,(nlm)} c_{(nlm)}^I \Psi_{(nlm)}^I(\vec{r} - \vec{R}_I)$$

$$E_{LCAO} = \min \frac{\langle \Psi_{trial} | \hat{H} | \Psi_{trial} \rangle}{\langle \Psi_{trial} | \Psi_{trial} \rangle}$$

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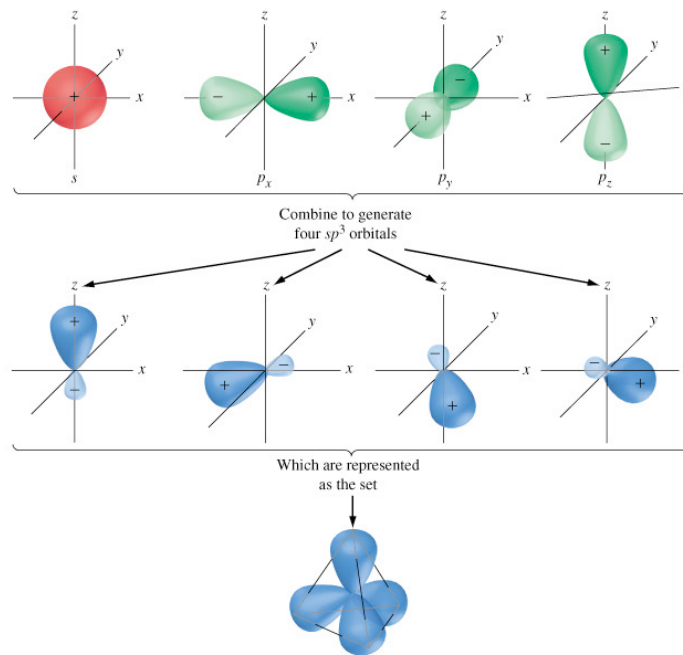
## More complex molecules: LCAO



$$\begin{aligned} \Psi_{trial} = & c_1 \Psi_{1s,H_a} + c_2 \Psi_{1s,H_b} + c_3 \Psi_{1s,H_c} + c_4 \Psi_{1s,H_d} \\ & + c_5 \Psi_{2s,C} + c_6 \Psi_{2p_x,C} + c_7 \Psi_{2p_y,C} + c_8 \Psi_{2p_z,C} \end{aligned}$$

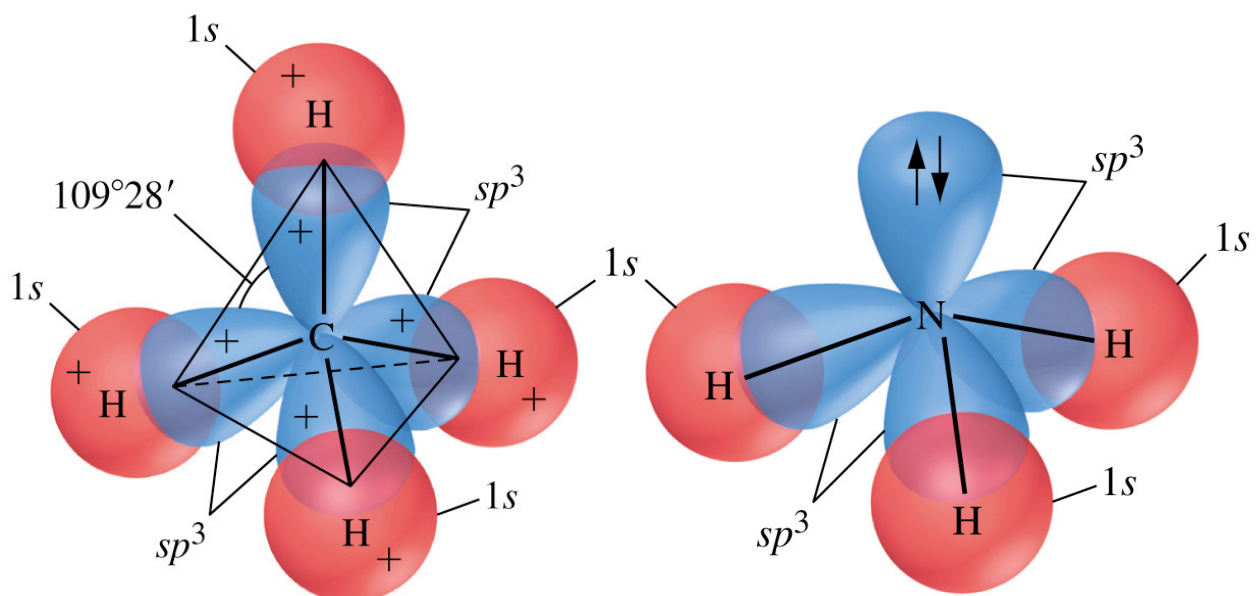
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# $sp^3$ hybridization



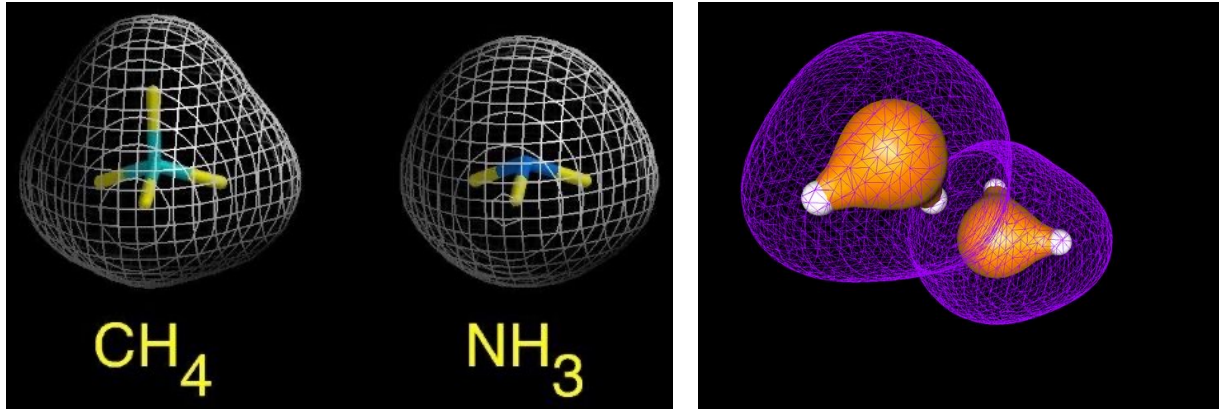
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# $sp^3$ hybridization



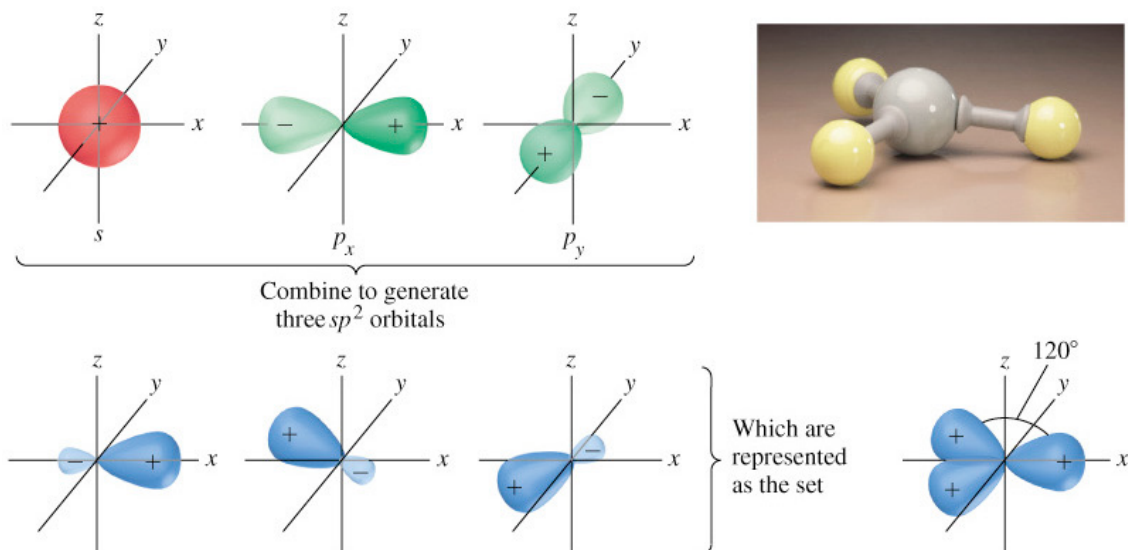
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# Great gases and liquids



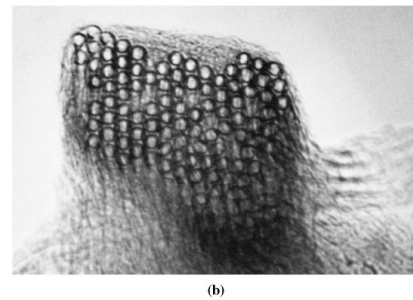
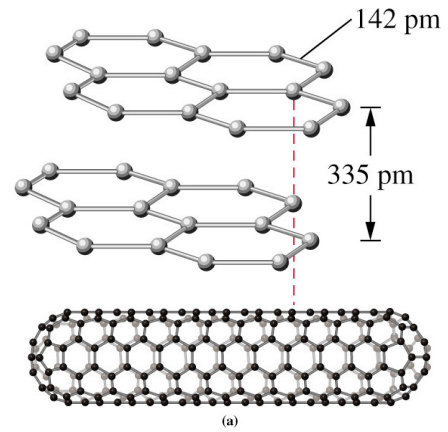
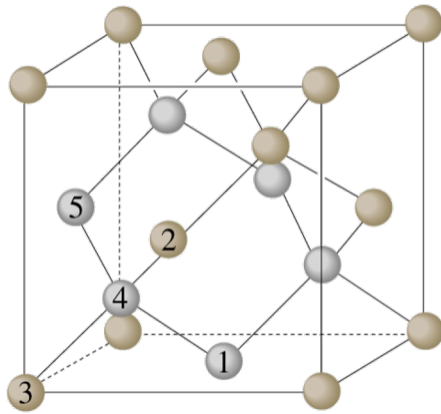
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## $sp^2$ hybridization



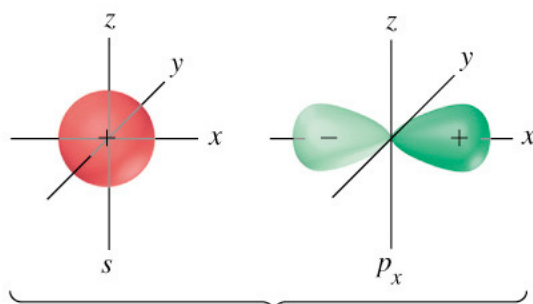
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# Carbon compounds

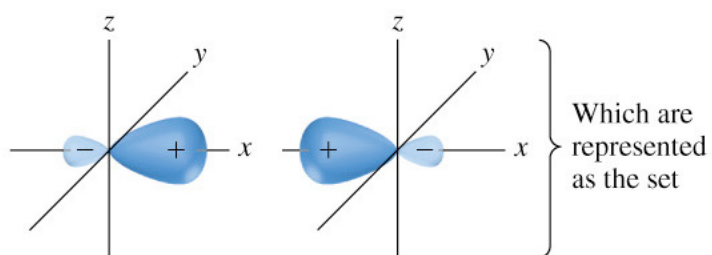


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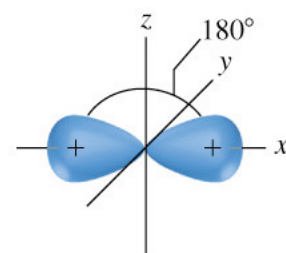
## sp hybridization



Combine to generate  
two *sp* orbitals

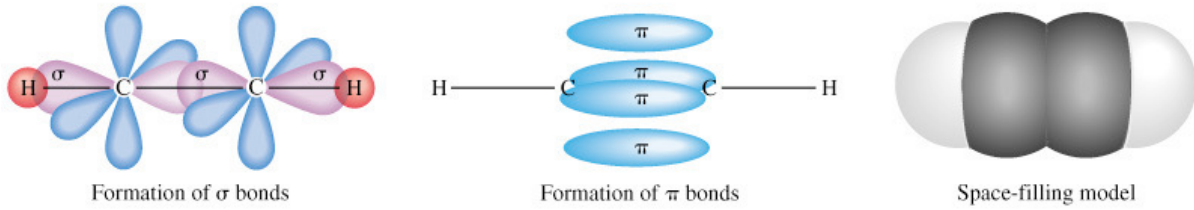


Which are  
represented  
as the set



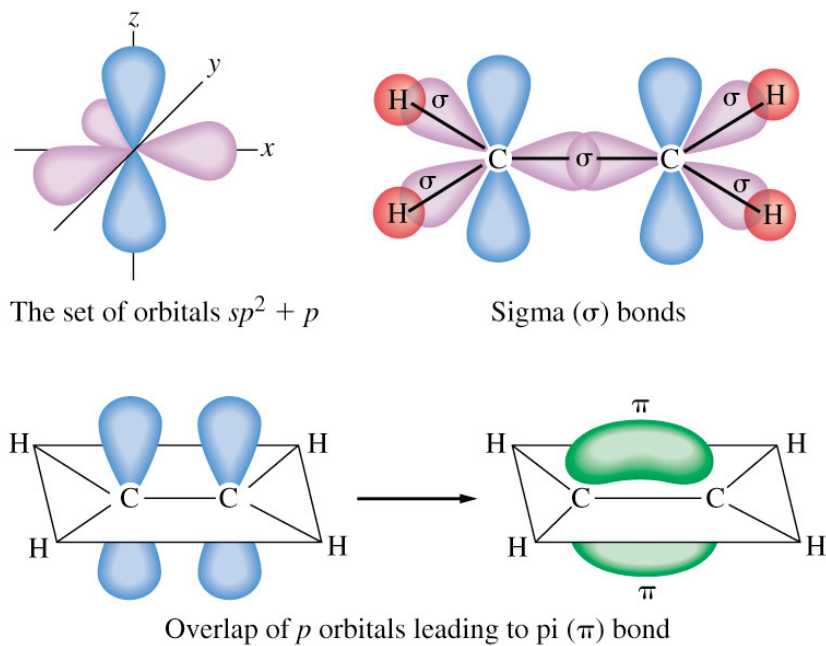
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# Ethyne (Acetylene)



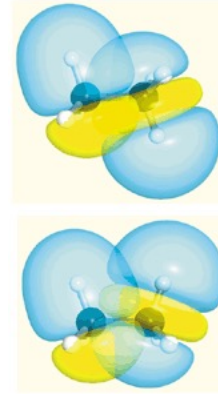
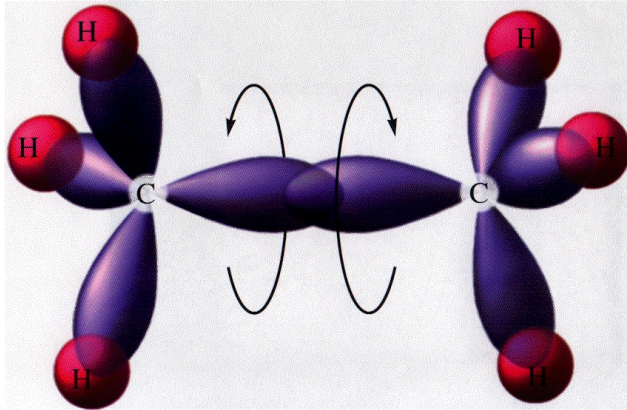
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# Ethene (Ethylene)



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# Ethane (saturated)



Goodman *et al.* Nature (2001)

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## Bond Lengths and Bond Energies

TABLE 11.2 Some Average Bond Lengths<sup>a</sup>

| Bond | Bond Length, pm | Bond | Bond Length, pm | Bond  | Bond Length, pm |
|------|-----------------|------|-----------------|-------|-----------------|
| H—H  | 74.14           | C—C  | 154             | N—N   | 145             |
| H—C  | 110             | C=C  | 134             | N=N   | 123             |
| H—N  | 100             | C≡C  | 120             | N≡N   | 109.8           |
| H—O  | 97              | C—N  | 147             | N—O   | 136             |
| H—S  | 132             | C=N  | 128             | N=O   | 120             |
| H—F  | 91.7            | C≡N  | 116             | O—O   | 145             |
| H—Cl | 127.4           | C—O  | 143             | O=O   | 121             |
| H—Br | 141.4           | C=O  | 120             | F—F   | 143             |
| H—I  | 160.9           | C—Cl | 178             | Cl—Cl | 199             |
|      |                 |      |                 | Br—Br | 228             |
|      |                 |      |                 | I—I   | 266             |

<sup>a</sup>Most values (C—H, N—H, C—H, ...) are averaged over a number of species containing the indicated bond and may vary by a few picometers. Where a diatomic molecule exists, the value given is the actual bond length in that molecule (H<sub>2</sub>, N<sub>2</sub>, HF, ...) and is known more precisely.

TABLE 11.3 Some Average Bond Energies<sup>a</sup>

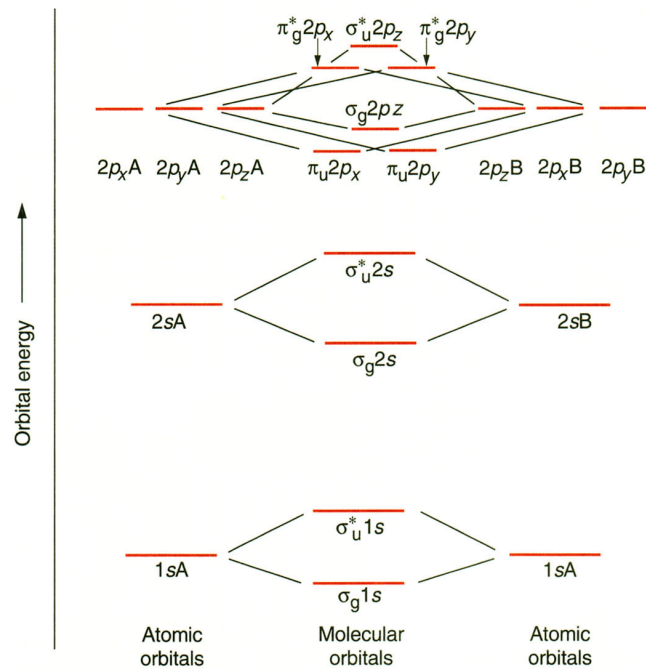
| Bond | Bond Energy, kJ/mol | Bond | Bond Energy, kJ/mol | Bond  | Bond Energy, kJ/mol |
|------|---------------------|------|---------------------|-------|---------------------|
| H—H  | 436                 | C—C  | 347                 | N—N   | 163                 |
| H—C  | 414                 | C=C  | 611                 | N=N   | 418                 |
| H—N  | 389                 | C≡C  | 837                 | N≡N   | 946                 |
| H—O  | 464                 | C—N  | 305                 | N—O   | 222                 |
| H—S  | 368                 | C=N  | 615                 | N=O   | 590                 |
| H—F  | 565                 | C≡N  | 891                 | O—O   | 142                 |
| H—Cl | 431                 | C—O  | 360                 | O=O   | 498                 |
| H—Br | 364                 | C=O  | 736 <sup>b</sup>    | F—F   | 159                 |
| H—I  | 297                 | C—Cl | 339                 | Cl—Cl | 243                 |
|      |                     |      |                     | Br—Br | 193                 |
|      |                     |      |                     | I—I   | 151                 |

<sup>a</sup>Although all data are listed with about the same precision (three significant figures), some values are actually known more precisely. Specifically, the values for the diatomic molecules: H<sub>2</sub>, HF, HCl, HBr, HI, N<sub>2</sub> (N≡N), O<sub>2</sub> (O=O), F<sub>2</sub>, Cl<sub>2</sub>, Br<sub>2</sub>, and I<sub>2</sub> are actually bond-dissociation energies, rather than average bond energies.

<sup>b</sup>The value for the C=O bonds in CO<sub>2</sub> is 799 kJ/mol.

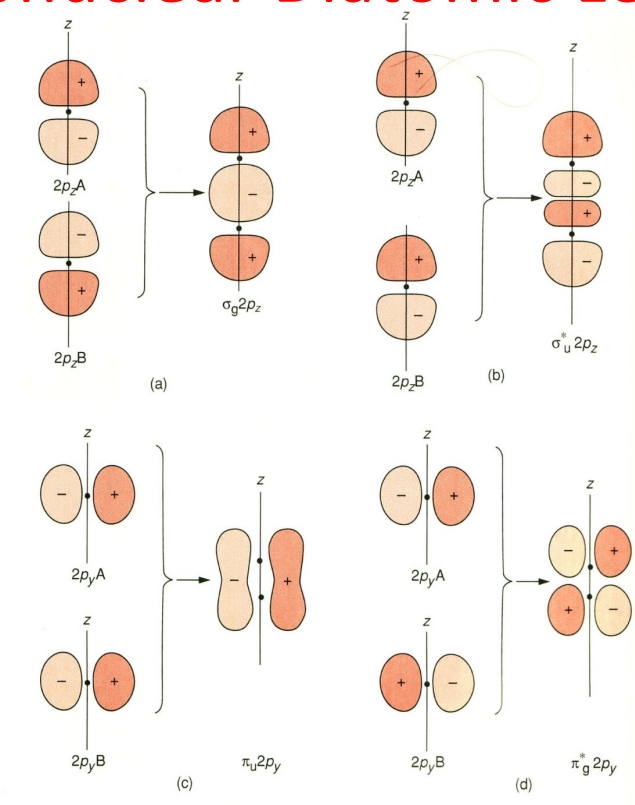
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# Homonuclear Diatomic Levels (I)

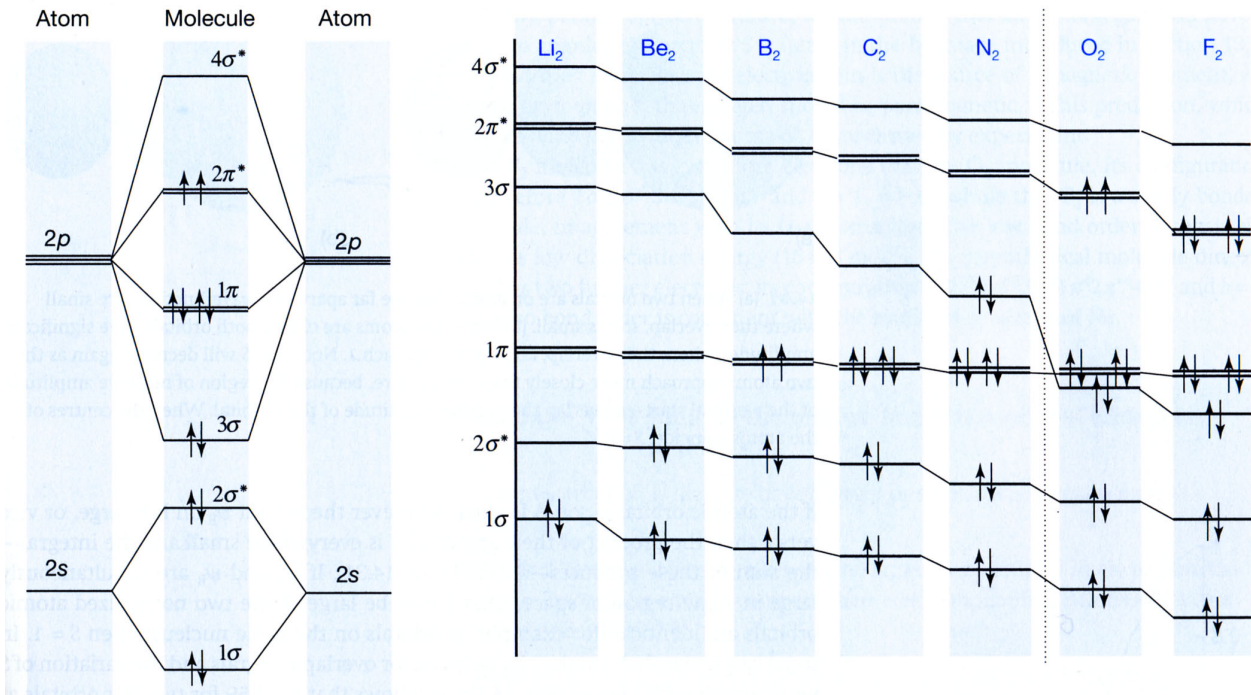


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# Homonuclear Diatomic Levels (II)



# Homonuclear Diatomic Levels (III)

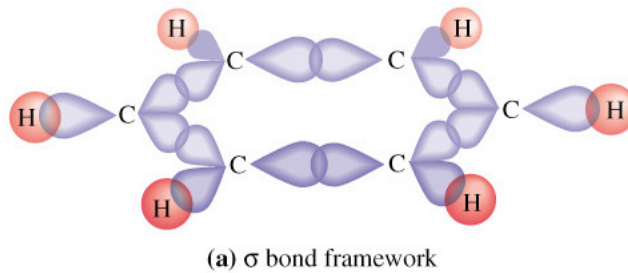
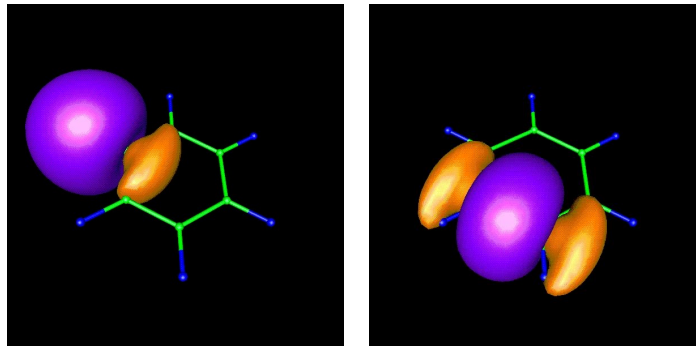


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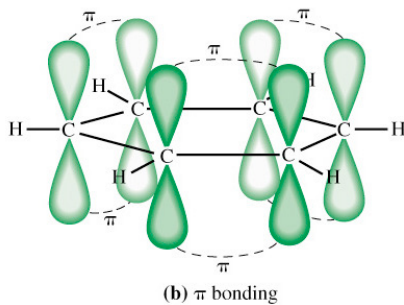
## Even simpler than LCAO: Empirical tight binding and Hückel approach

- TB: The matrix elements of the Hamiltonian are “universal empirical parameters”
- Hückel: Planar / quasi-planar systems with delocalized  $\pi$  bonding: two parameters
  - $\alpha$ : matrix element between same orbital
  - $\beta$ : matrix element between neighboring orbitals
  - Hamiltonian between further neighbors is 0

# Example: Benzene ( $C_6H_6$ )



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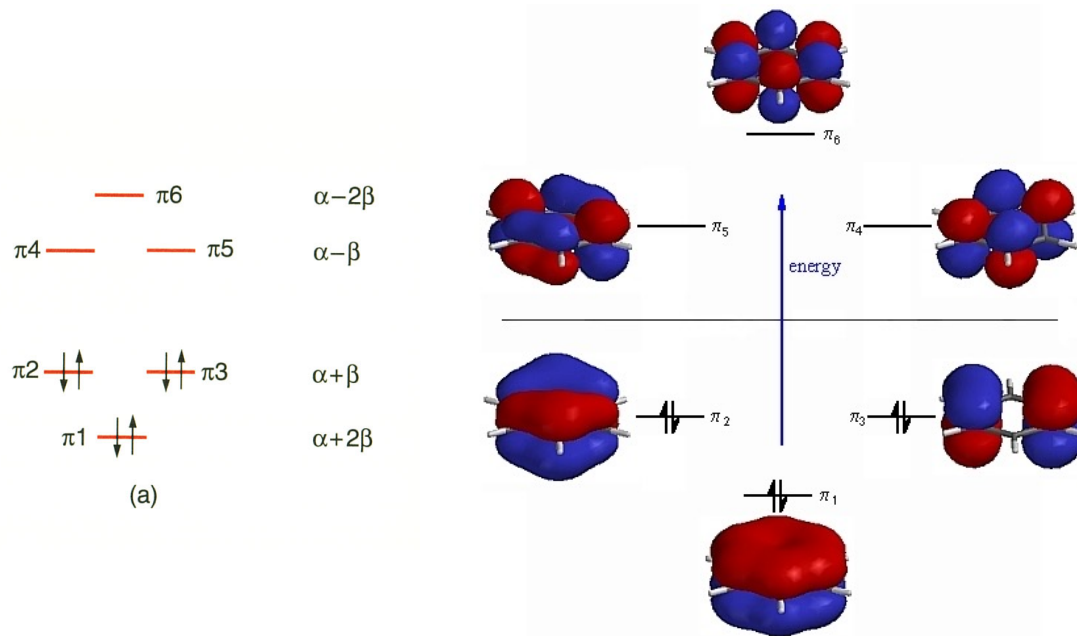


## Benzene – energy levels

$$\det \begin{pmatrix} \alpha - E & \beta & 0 & 0 & 0 & \beta \\ \beta & \alpha - E & \beta & 0 & 0 & 0 \\ 0 & \beta & \alpha - E & \beta & 0 & 0 \\ 0 & 0 & \beta & \alpha - E & \beta & 0 \\ 0 & 0 & 0 & \beta & \alpha - E & \beta \\ \beta & 0 & 0 & 0 & \beta & \alpha - E \end{pmatrix} = 0$$

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# Benzene – molecular orbitals



<http://www.chem.ucalgary.ca/SHMO/>