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## Lecture Presentation

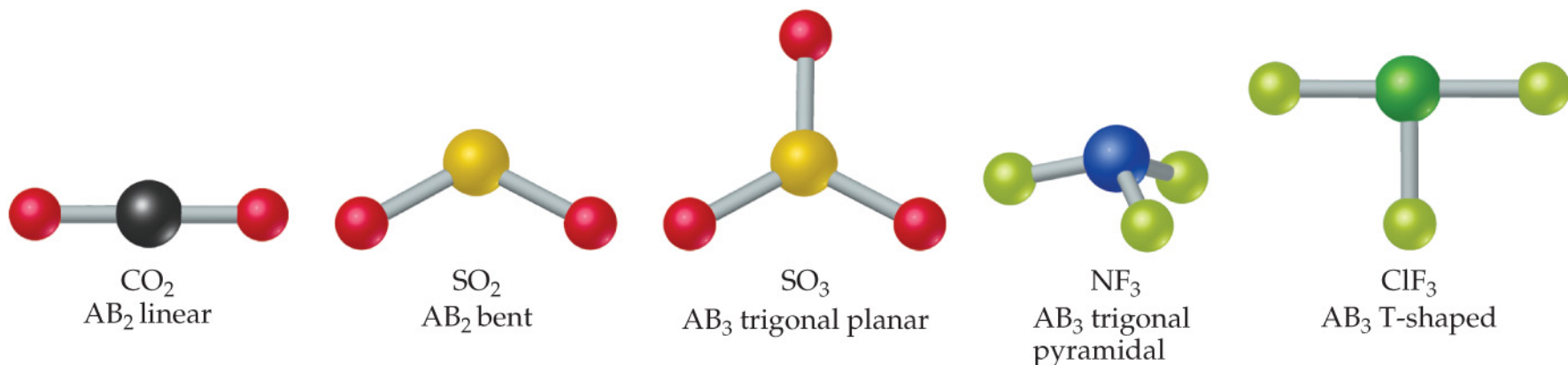
# Chapter 9

# Molecular Geometries and Bonding Theories

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St. Charles Community College  
Cottleville, MO

# Molecular Shapes

- The shape of a molecule plays an important role in its reactivity.
- By noting the number of bonding and nonbonding electron pairs, we can easily predict the shape of the molecule.



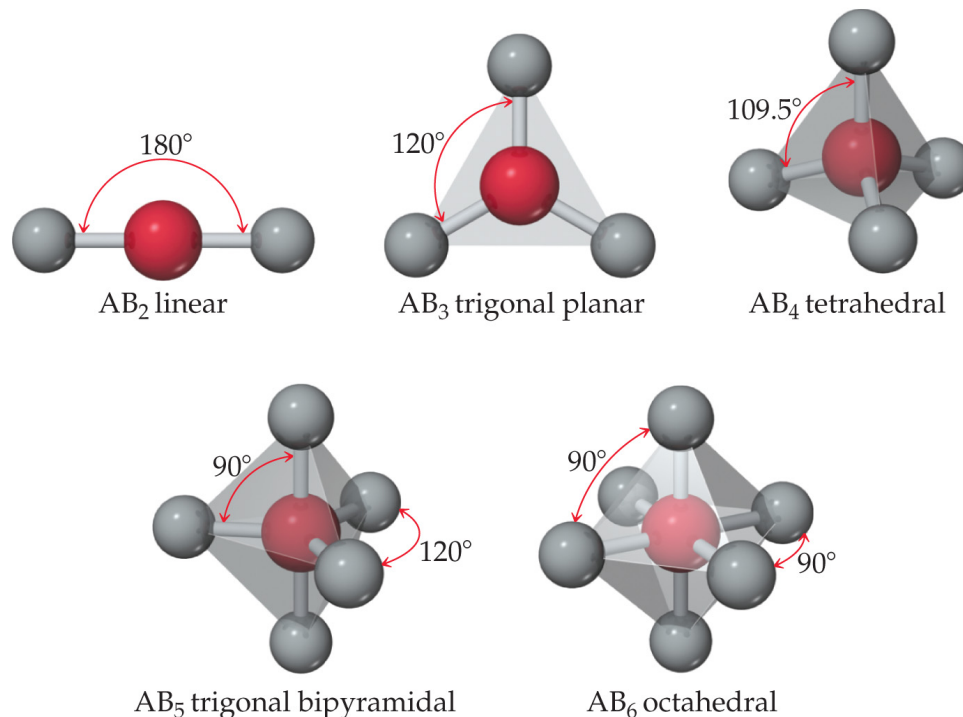
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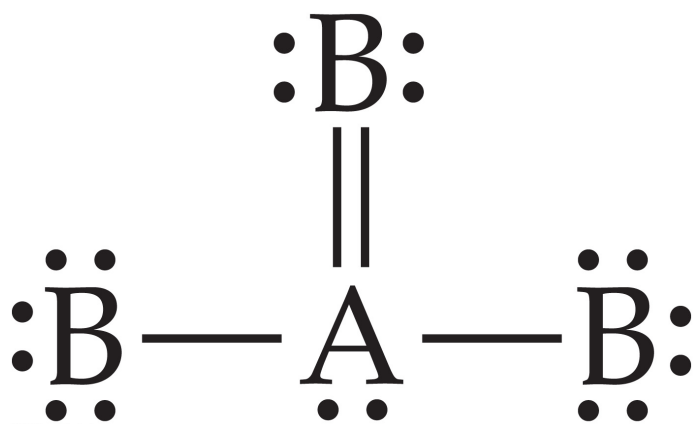
# What Determines the Shape of a Molecule?

- Simply put, electron pairs, whether they be bonding or nonbonding, repel each other.
- By assuming the electron pairs are placed as far as possible from each other, we can predict the shape of the molecule.



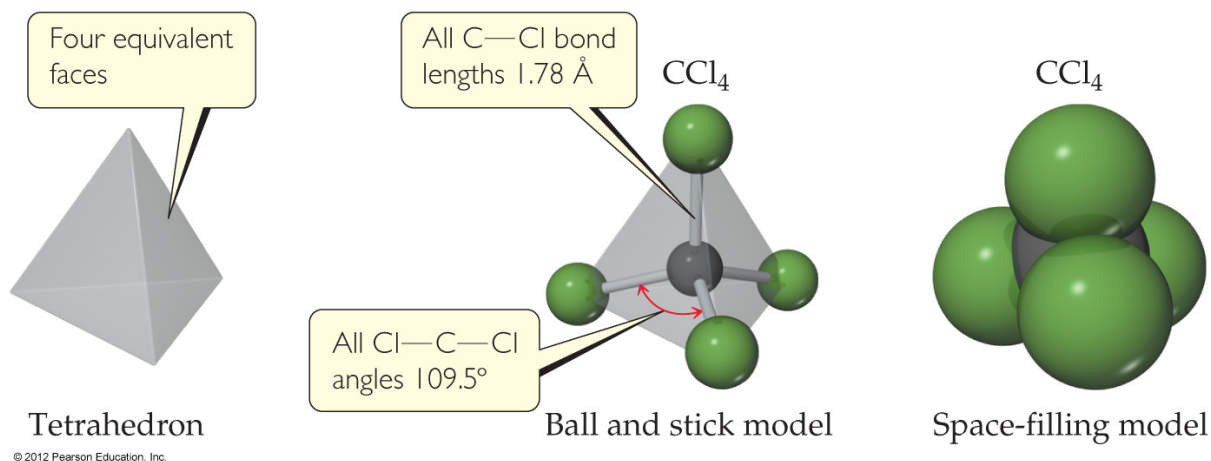
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# Electron Domains



- The central atom in this molecule, A, has four electron domains.
- We can refer to the electron pairs as **electron domains**.
- In a double or triple bond, all electrons shared between those two atoms are on the same side of the central atom; therefore, they count as one electron domain.

# Valence-Shell Electron-Pair Repulsion Theory (VSEPR)



*“The best arrangement of a given number of electron domains is the one that minimizes the repulsions among them.”*

# Electron-Domain Geometries

TABLE 9.4 • Geometric Arrangements Characteristic of Hybrid Orbital Sets

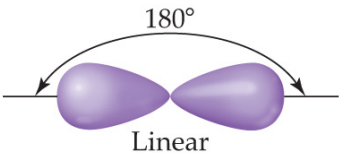
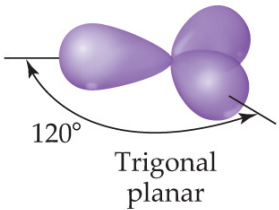
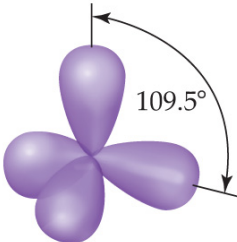
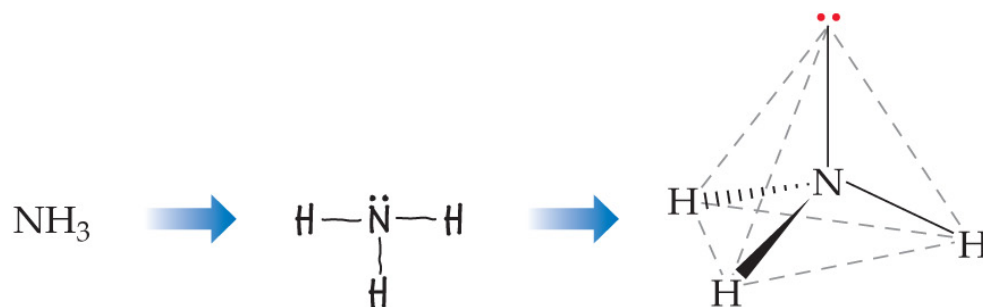
Atomic Orbital Set	Hybrid Orbital Set	Geometry	Examples
$s, p$	Two $sp$	 <p>Linear</p>	$\text{BeF}_2, \text{HgCl}_2$
$s, p, p$	Three $sp^2$	 <p>Trigonal planar</p>	$\text{BF}_3, \text{SO}_3$
$s, p, p, p$	Four $sp^3$	 <p>Tetrahedral</p>	$\text{CH}_4, \text{NH}_3, \text{H}_2\text{O}, \text{NH}_4^+$

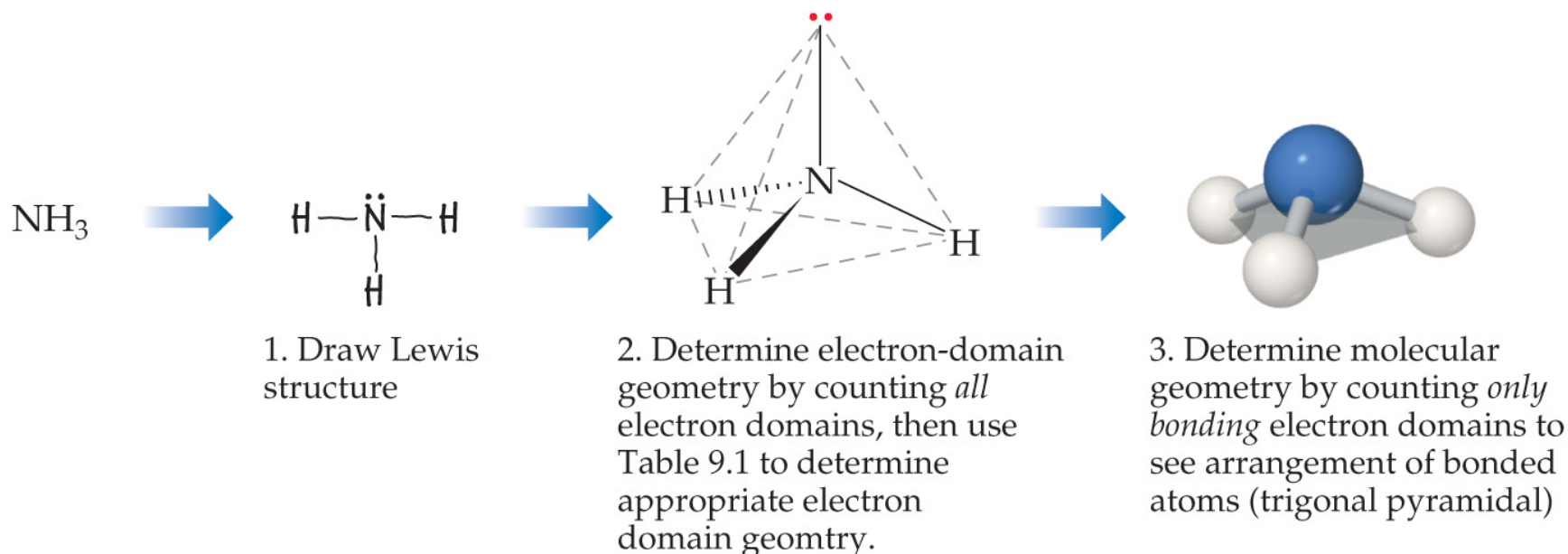
Table 9.1 contains the electron-domain geometries for two through six electron domains around a central atom.

# Electron-Domain Geometries

- All one must do is count the number of electron domains in the Lewis structure.
- The geometry will be that which corresponds to the number of electron domains.



# Molecular Geometries



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- The electron-domain geometry is often *not* the shape of the molecule, however.
- The molecular geometry is that defined by the positions of *only* the atoms in the molecules, not the nonbonding pairs.

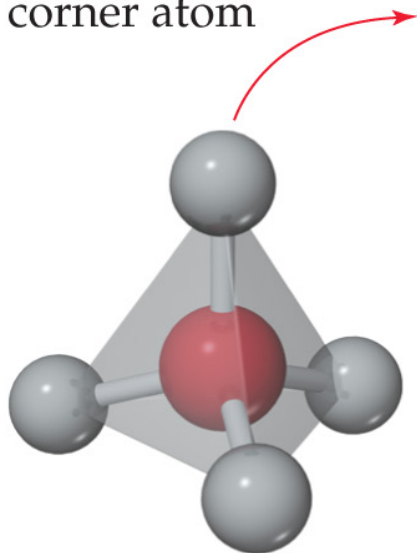
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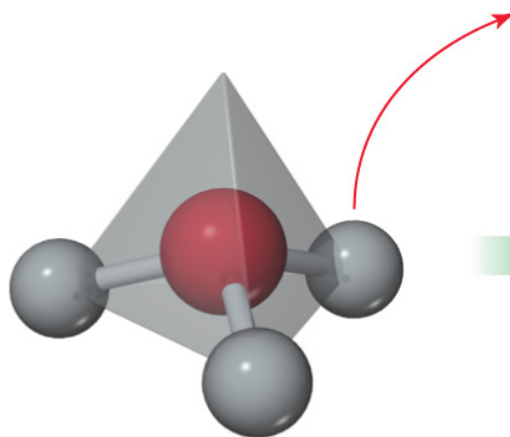
# Molecular Geometries

Removal of one  
corner atom

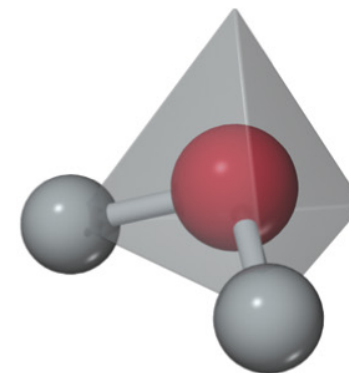


Tetrahedral

Removal of a  
second  
corner atom



Trigonal pyramidal



Bent

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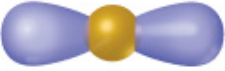

Within each electron domain, then, there might be more than one molecular geometry.

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# Linear Electron Domain

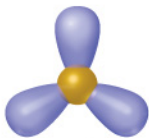
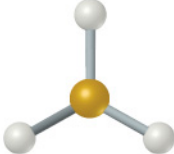
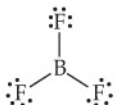
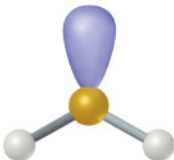
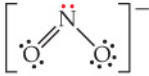
TABLE 9.2 • Electron-Domain and Molecular Geometries for Two, Three, and Four Electron Domains around a Central Atom

Number of Electron Domains	Electron-Domain Geometry	Bonding Domains	Nonbonding Domains	Molecular Geometry	Example
2	 Linear	2	0	 Linear	$\ddot{\text{O}}=\text{C}=\ddot{\text{O}}$

- In the linear domain, there is only one molecular geometry: linear.
- NOTE: If there are only two atoms in the molecule, the molecule will be linear no matter what the electron domain is.

# Trigonal Planar Electron Domain

TABLE 9.2 • Electron-Domain and Molecular Geometries for Two, Three, and Four Electron Domains around a Central Atom

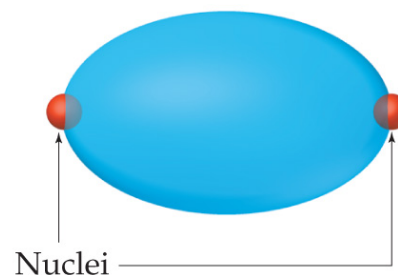
Number of Electron Domains	Electron-Domain Geometry	Bonding Domains	Nonbonding Domains	Molecular Geometry	Example
3	 Trigonal planar	3	0	 Trigonal planar	
		2	1	 Bent	

- There are two molecular geometries:
  - Trigonal planar, if all the electron domains are bonding,
  - Bent, if one of the domains is a nonbonding pair.

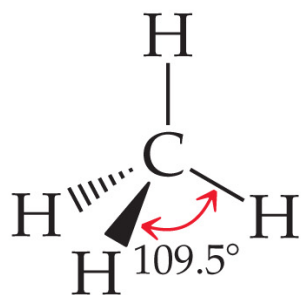
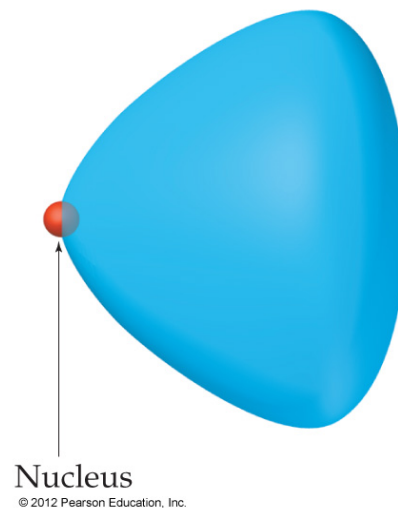
# Nonbonding Pairs and Bond Angle

- Nonbonding pairs are physically larger than bonding pairs.
- Therefore, their repulsions are greater; this tends to decrease bond angles in a molecule.

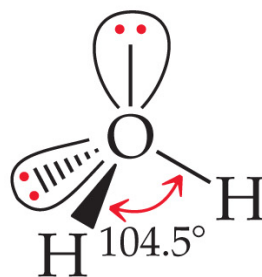
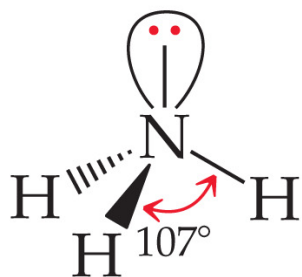
Bonding electron pair



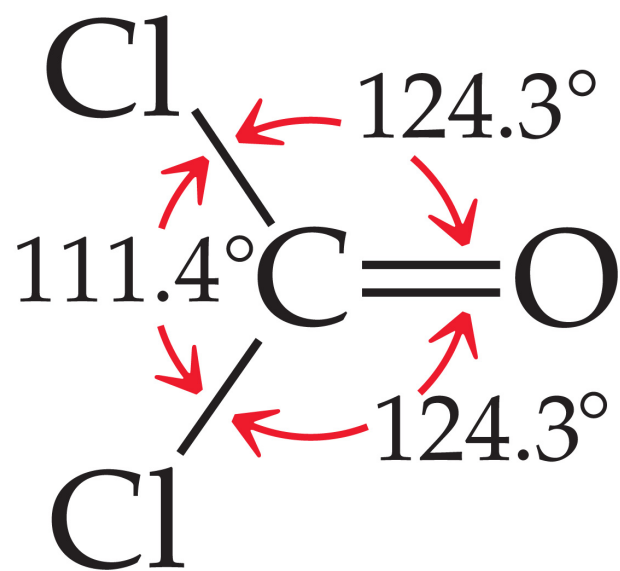
Nonbonding pair



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# Multiple Bonds and Bond Angles

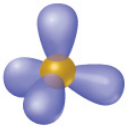
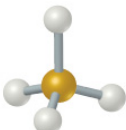

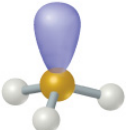

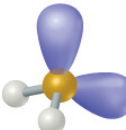



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- Double and triple bonds place greater electron density on one side of the central atom than do single bonds.
- Therefore, they also affect bond angles.

# Tetrahedral Electron Domain

TABLE 9.2 • Electron-Domain and Molecular Geometries for Two, Three, and Four Electron Domains around a Central Atom

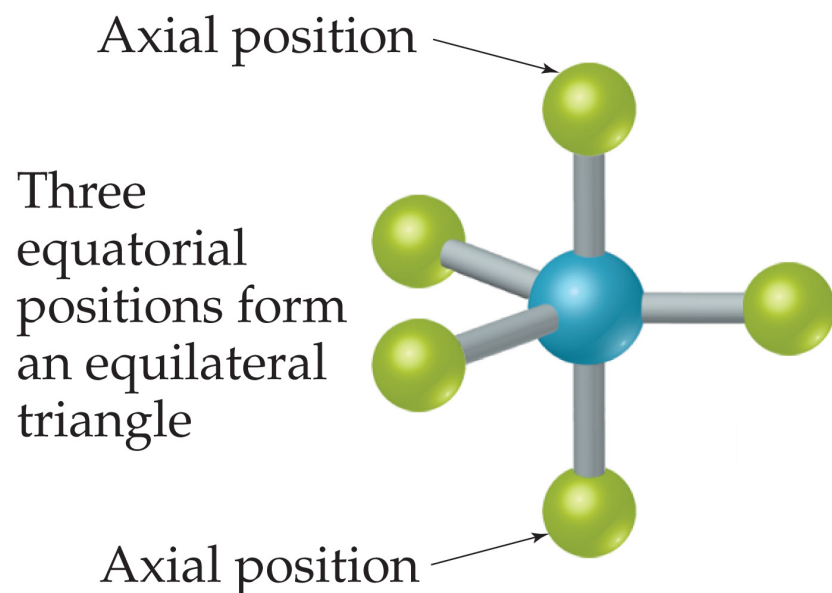
Number of Electron Domains	Electron-Domain Geometry	Bonding Domains	Nonbonding Domains	Molecular Geometry	Example
4	 Tetrahedral	4	0	 Tetrahedral	
		3	1	 Trigonal pyramidal	
		2	2	 Bent	

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- There are three molecular geometries:
  - Tetrahedral, if all are bonding pairs,
  - Trigonal pyramidal, if one is a nonbonding pair,
  - Bent, if there are two nonbonding pairs.

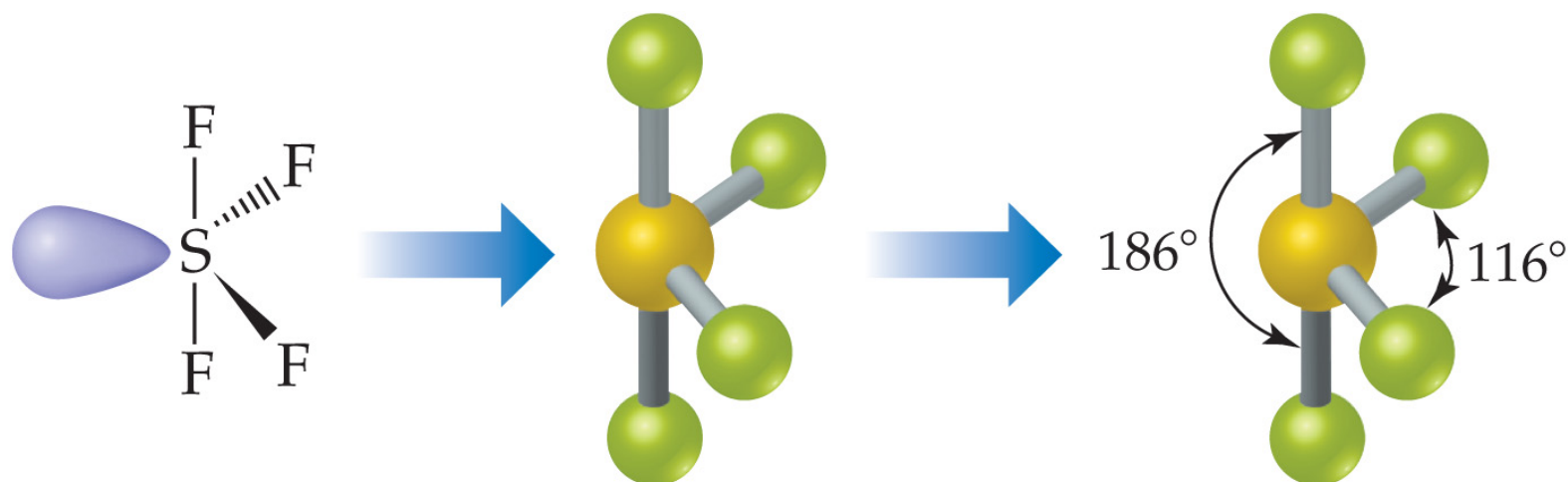
Molecular Geometries and Bonding

# Trigonal Bipyramidal Electron Domain



- There are two distinct positions in this geometry:
  - Axial
  - Equatorial

# Trigonal Bipyramidal Electron Domain



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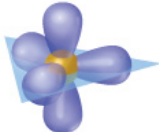
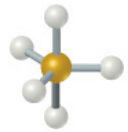
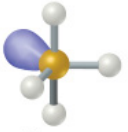
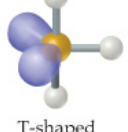
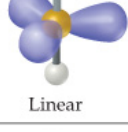
Lower-energy conformations result from having nonbonding electron pairs in equatorial, rather than axial, positions in this geometry.



# Trigonal Bipyramidal Electron Domain


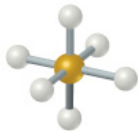
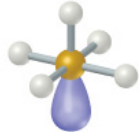
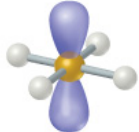
- There are four distinct molecular geometries in this domain:
  - Trigonal bipyramidal
  - Seesaw
  - T-shaped
  - Linear

TABLE 9.3 • Electron-Domain and Molecular Geometries for Five and Six Electron Domains around a Central Atom

Number of Electron Domains	Electron-Domain Geometry	Bonding Domains	Nonbonding Domains	Molecular Geometry	Example
5	 Trigonal bipyramidal	5	0	 Trigonal bipyramidal	PCl <sub>5</sub>
		4	1	 Seesaw	SF <sub>4</sub>
		3	2	 T-shaped	ClF <sub>3</sub>
		2	3	 Linear	XeF <sub>2</sub>

# Octahedral Electron Domain

TABLE 9.3 • Electron-Domain and Molecular Geometries for Five and Six Electron Domains around a Central Atom

Number of Electron Domains	Electron-Domain Geometry	Bonding Domains	Nonbonding Domains	Molecular Geometry	Example
6	 Octahedral	6	0	 Octahedral	SF <sub>6</sub>
		5	1	 Square pyramidal	BrF <sub>5</sub>
		4	2	 Square planar	XeF <sub>4</sub>

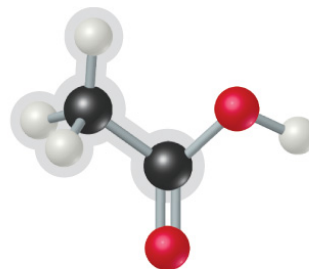
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- All positions are equivalent in the octahedral domain.
- There are three molecular geometries:
  - Octahedral
  - Square pyramidal
  - Square planar

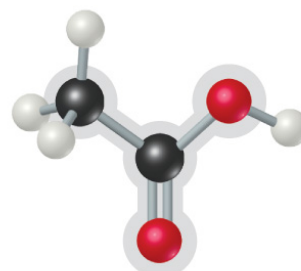
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# Larger Molecules

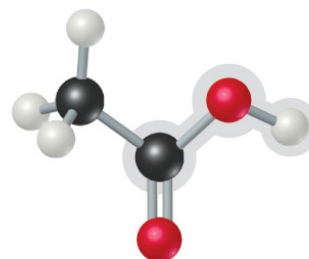
In larger molecules, it makes more sense to talk about the geometry about a particular atom rather than the geometry of the molecule as a whole.



Electron-domain geometry tetrahedral,  
molecular geometry tetrahedral



Electron-domain geometry trigonal planar,  
molecular geometry trigonal planar



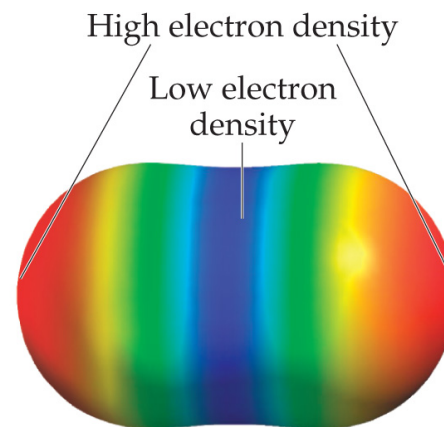
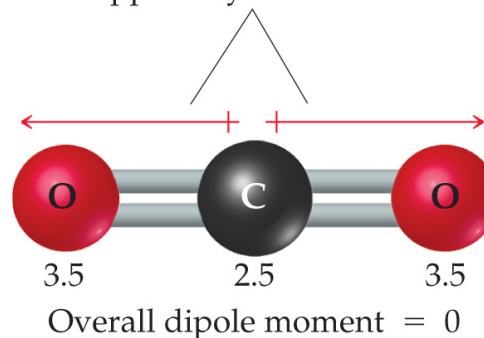
Electron-domain geometry tetrahedral,  
molecular geometry bent

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# Polarity

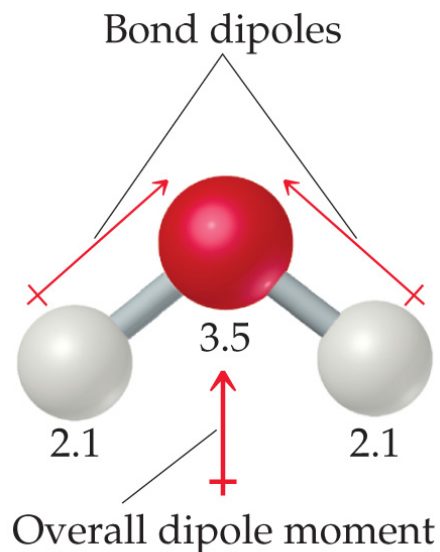
- In Chapter 8, we discussed bond dipoles.
- But just because a molecule possesses polar bonds does not mean the molecule *as a whole* will be polar.

Equal and oppositely directed bond dipoles

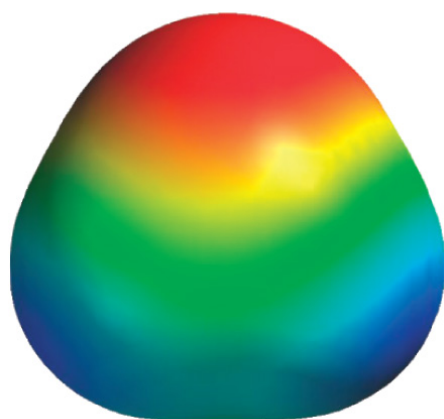


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# Polarity



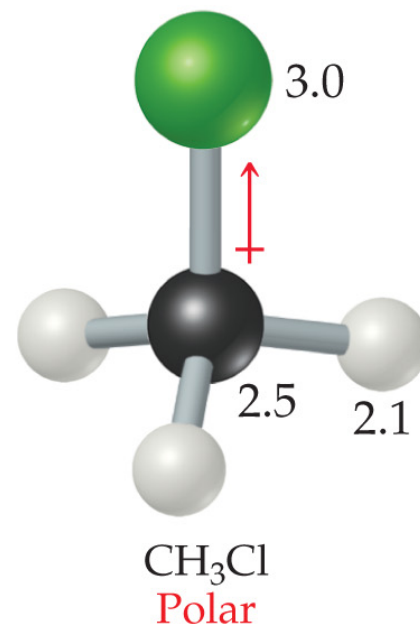
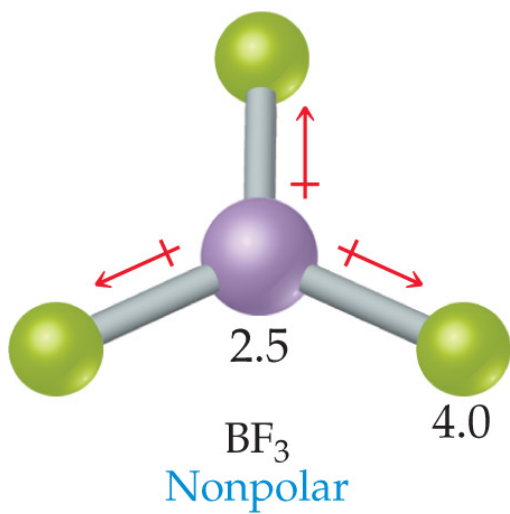
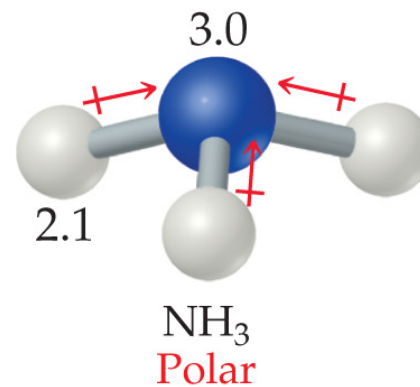
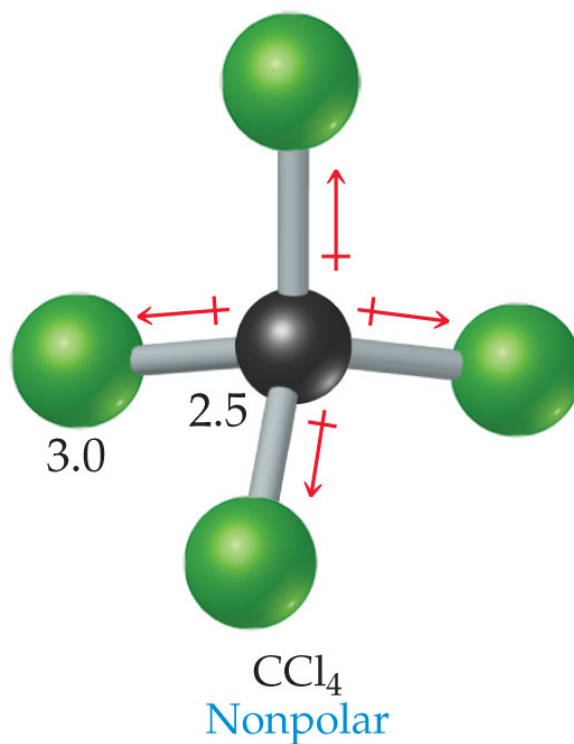
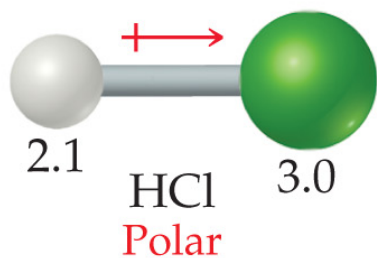
By adding the individual bond dipoles, one can determine the overall dipole moment for the molecule.



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# Polarity

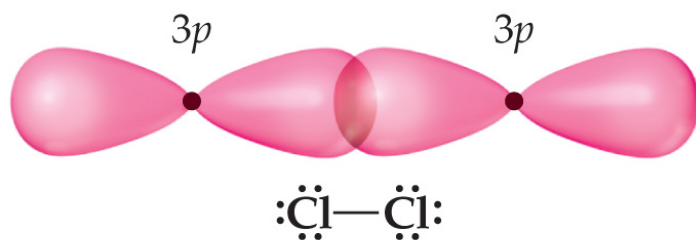
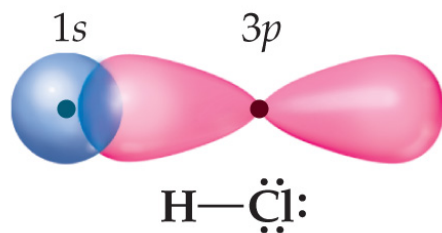
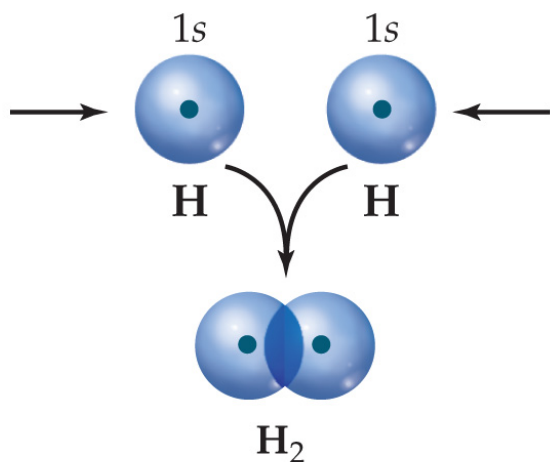


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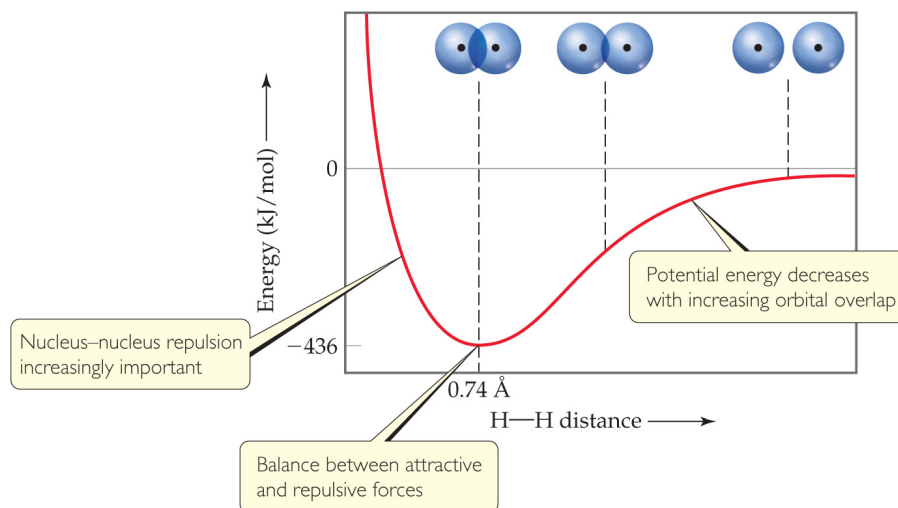
# Overlap and Bonding



- We think of covalent bonds forming through the sharing of electrons by adjacent atoms.
- In such an approach this can only occur when orbitals on the two atoms overlap.

# Overlap and Bonding

- Increased overlap brings the electrons and nuclei closer together while simultaneously decreasing electron–electron repulsion.
- However, if atoms get too close, the internuclear repulsion greatly raises the energy.

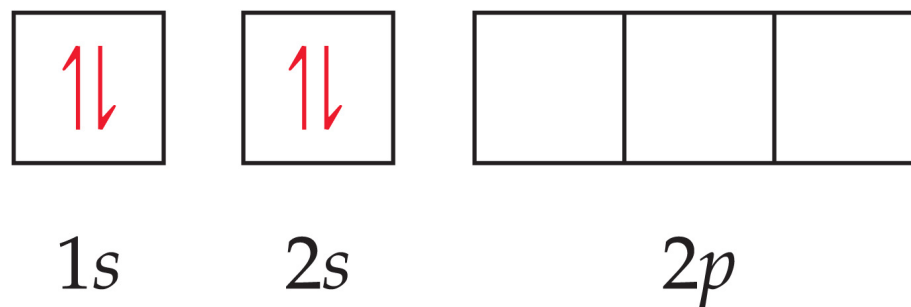


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# Hybrid Orbitals

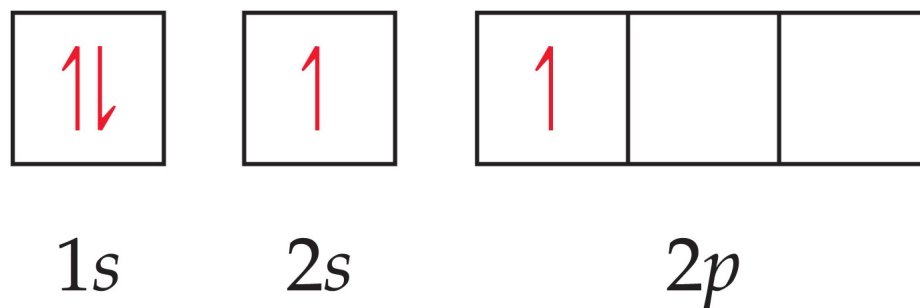
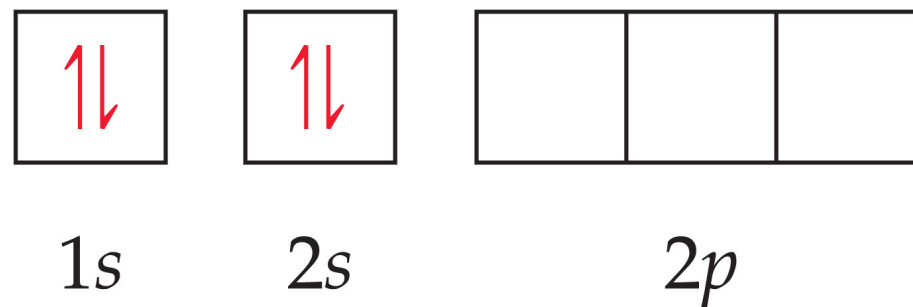
- Consider beryllium:
  - In its ground electronic state, beryllium would not be able to form bonds, because it has no singly occupied orbitals.



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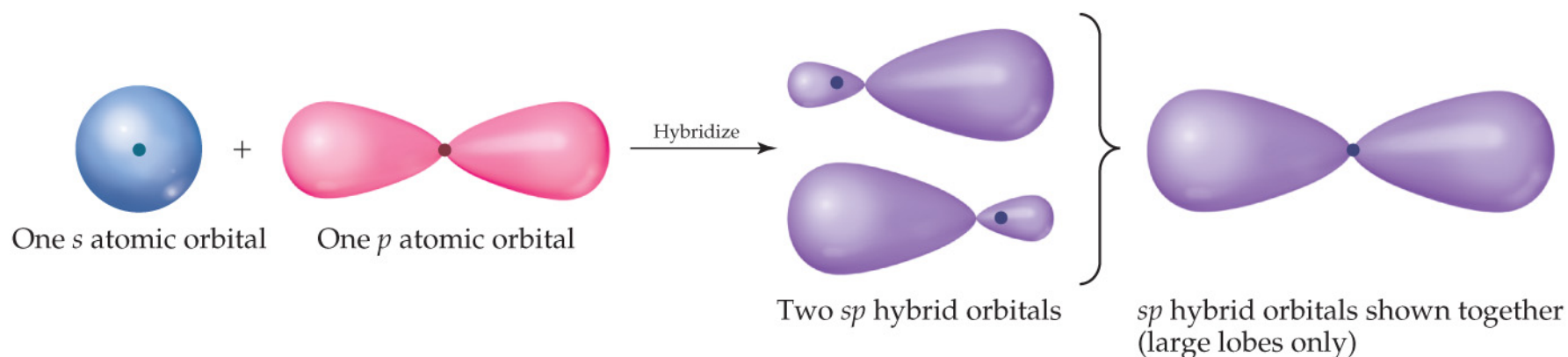
# Hybrid Orbitals

But if it absorbs the small amount of energy needed to promote an electron from the  $2s$  to the  $2p$  orbital, it can form two bonds.



# Hybrid Orbitals

- Mixing the  $s$  and  $p$  orbitals yields two degenerate orbitals that are hybrids of the two orbitals.
  - These  $sp$  hybrid orbitals have two lobes like a  $p$  orbital.
  - One of the lobes is larger and more rounded, as is the  $s$  orbital.



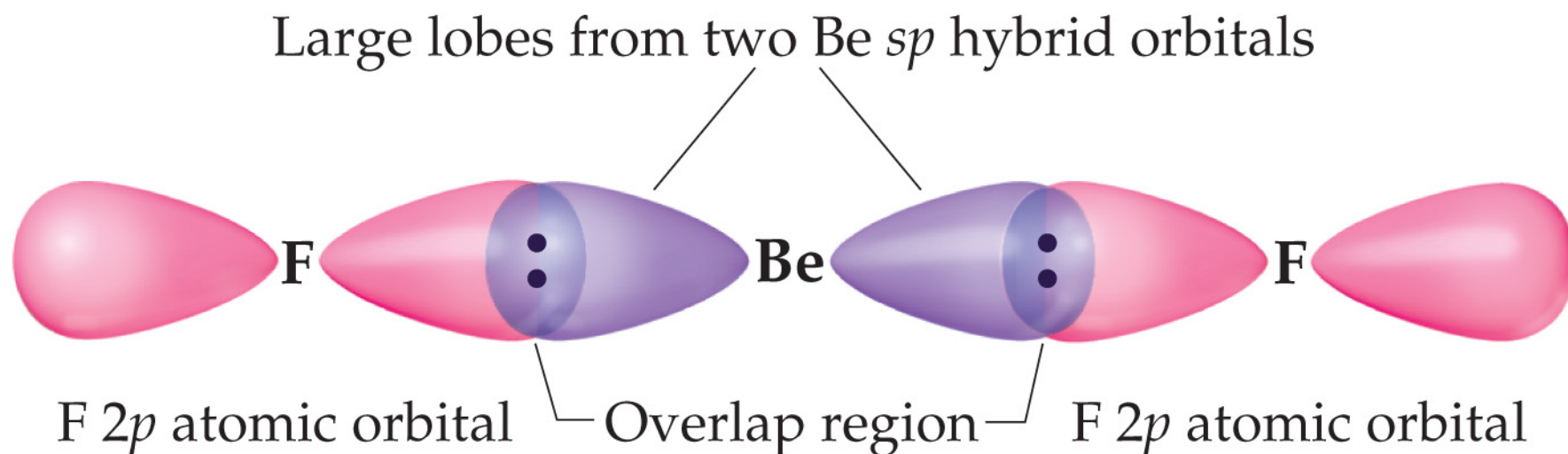
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# Hybrid Orbitals

- These two degenerate orbitals would align themselves  $180^\circ$  from each other.
- This is consistent with the observed geometry of beryllium compounds: linear.

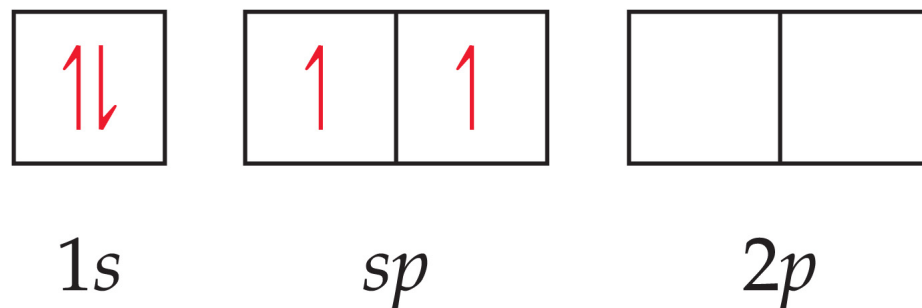


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# Hybrid Orbitals

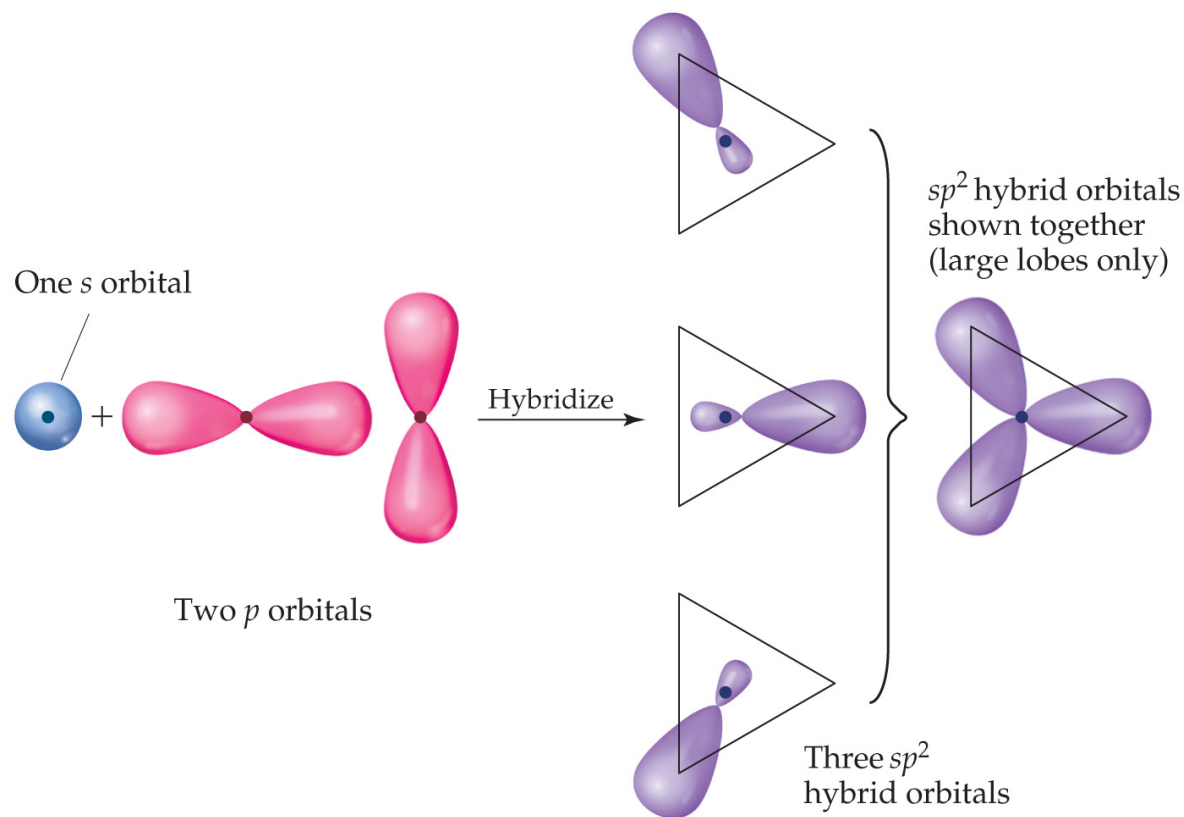


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- With hybrid orbitals, the orbital diagram for beryllium would look like this (Fig. 9.15).
- The  $sp$  orbitals are higher in energy than the  $1s$  orbital, but lower than the  $2p$ .

# Hybrid Orbitals

Using a similar model for boron leads to three degenerate  $sp^2$  orbitals.



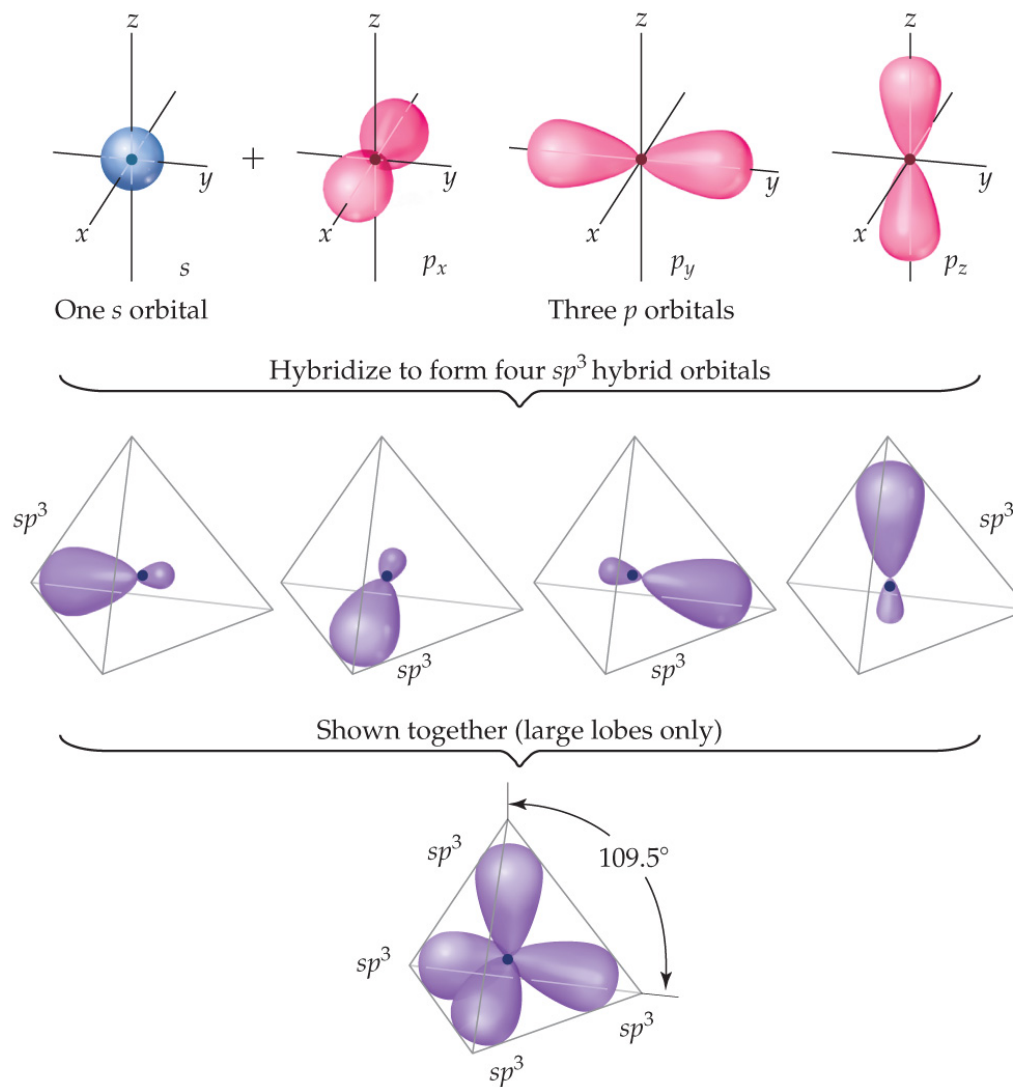
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# Hybrid Orbitals

With carbon, we get four degenerate  $sp^3$  orbitals.



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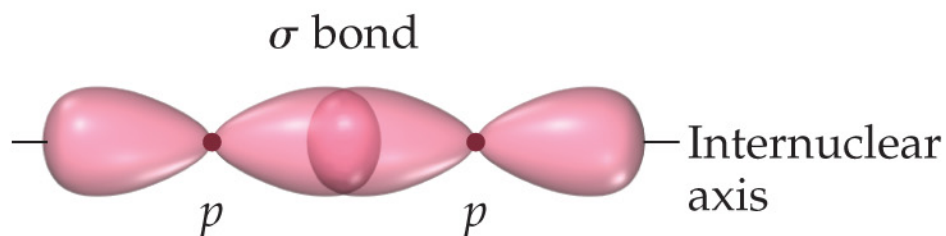
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# Valence Bond Theory

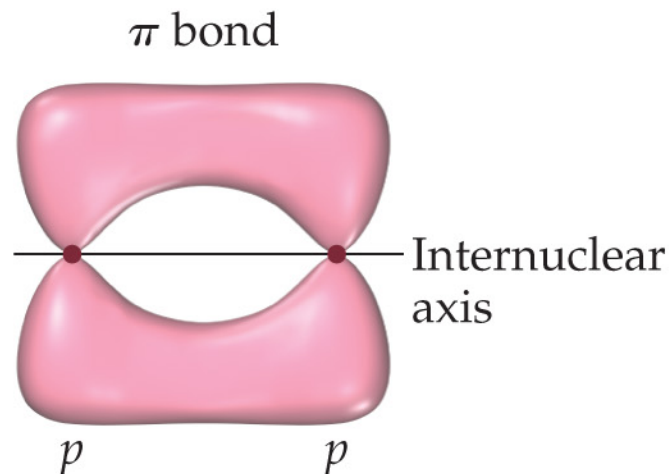
- Hybridization is a major player in this approach to bonding.
- There are two ways orbitals can overlap to form bonds between atoms.



# Sigma ( $\sigma$ ) Bonds

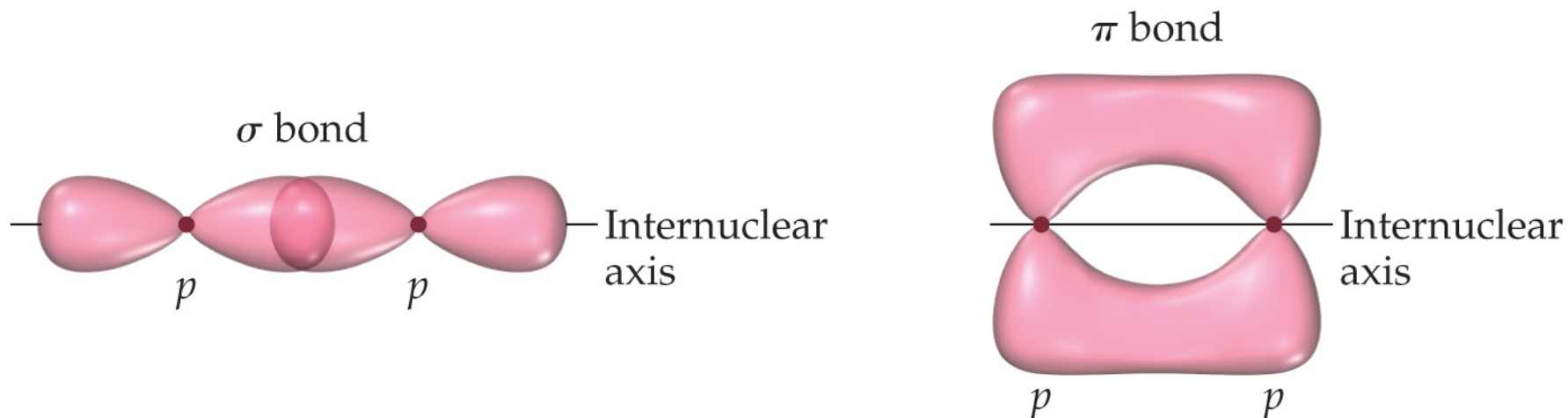


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- Sigma bonds are characterized by
  - Head-to-head overlap.
  - Cylindrical symmetry of electron density about the internuclear axis.

# Pi ( $\pi$ ) Bonds



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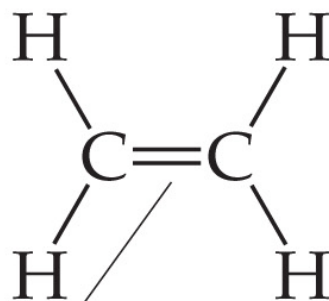
- Pi bonds are characterized by
  - Side-to-side overlap.
  - Electron density above and below the internuclear axis.

# Single Bonds

Single bonds are always  $\sigma$  bonds, because  $\sigma$  overlap is greater, resulting in a stronger bond and more energy lowering.



One  $\sigma$  bond



One  $\sigma$  bond plus  
one  $\pi$  bond



One  $\sigma$  bond plus  
two  $\pi$  bonds

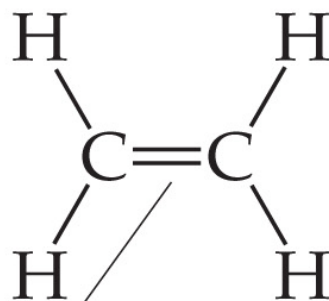
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# Multiple Bonds

In a multiple bond, one of the bonds is a  $\sigma$  bond and the rest are  $\pi$  bonds.



One  $\sigma$  bond



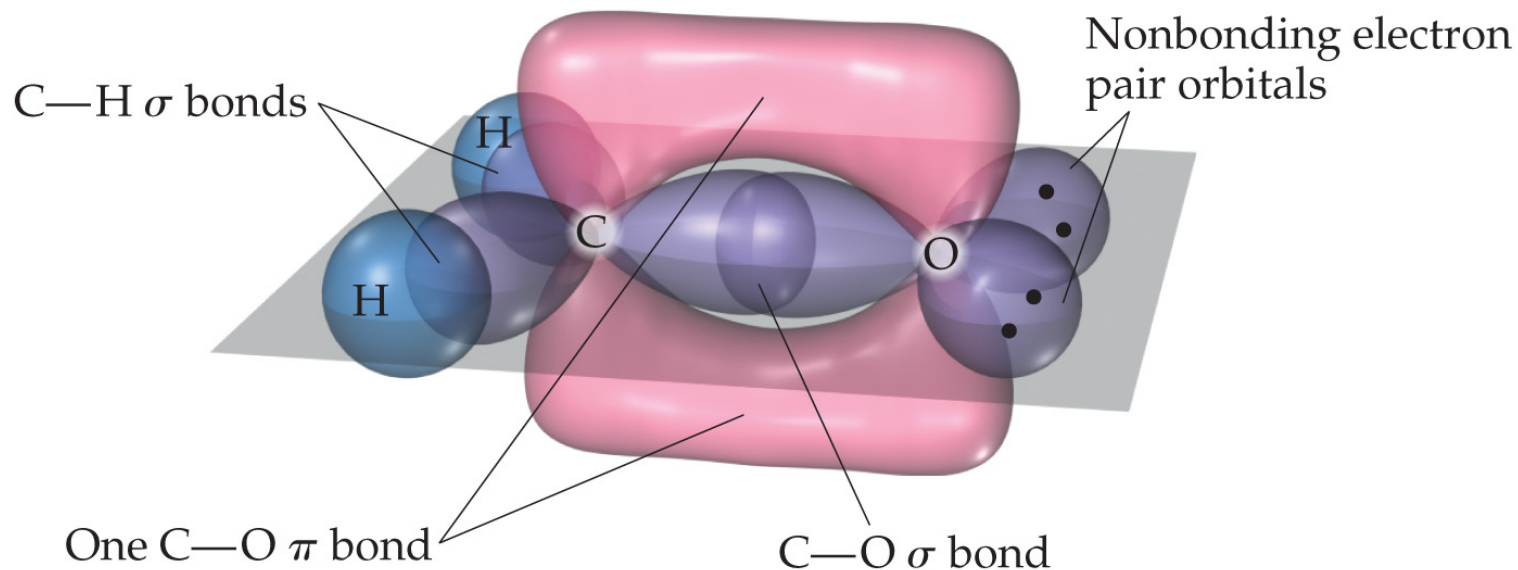
One  $\sigma$  bond plus  
one  $\pi$  bond



One  $\sigma$  bond plus  
two  $\pi$  bonds

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# Multiple Bonds

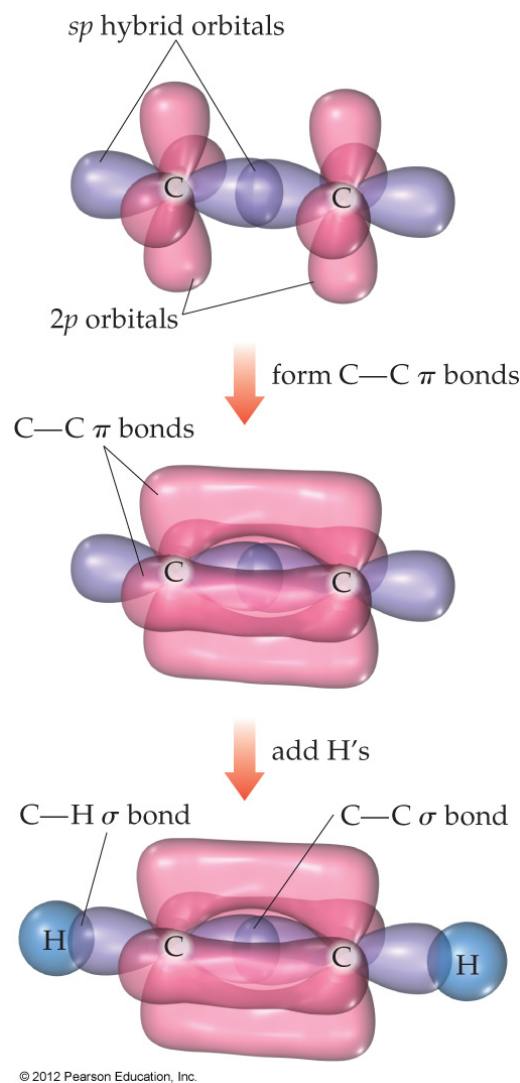


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- In a molecule like formaldehyde (shown at left), an  $sp^2$  orbital on carbon overlaps in  $\sigma$  fashion with the corresponding orbital on the oxygen.
- The unhybridized  $p$  orbitals overlap in  $\pi$  fashion.

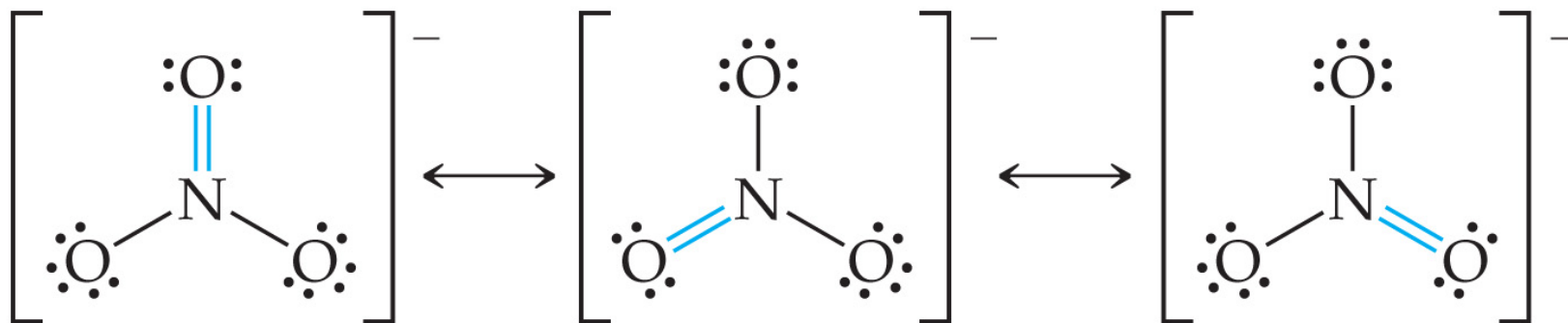
# Multiple Bonds

In triple bonds, as in acetylene, two  $sp$  orbitals form a  $\sigma$  bond between the carbons, and two pairs of  $p$  orbitals overlap in  $\pi$  fashion to form the two  $\pi$  bonds.



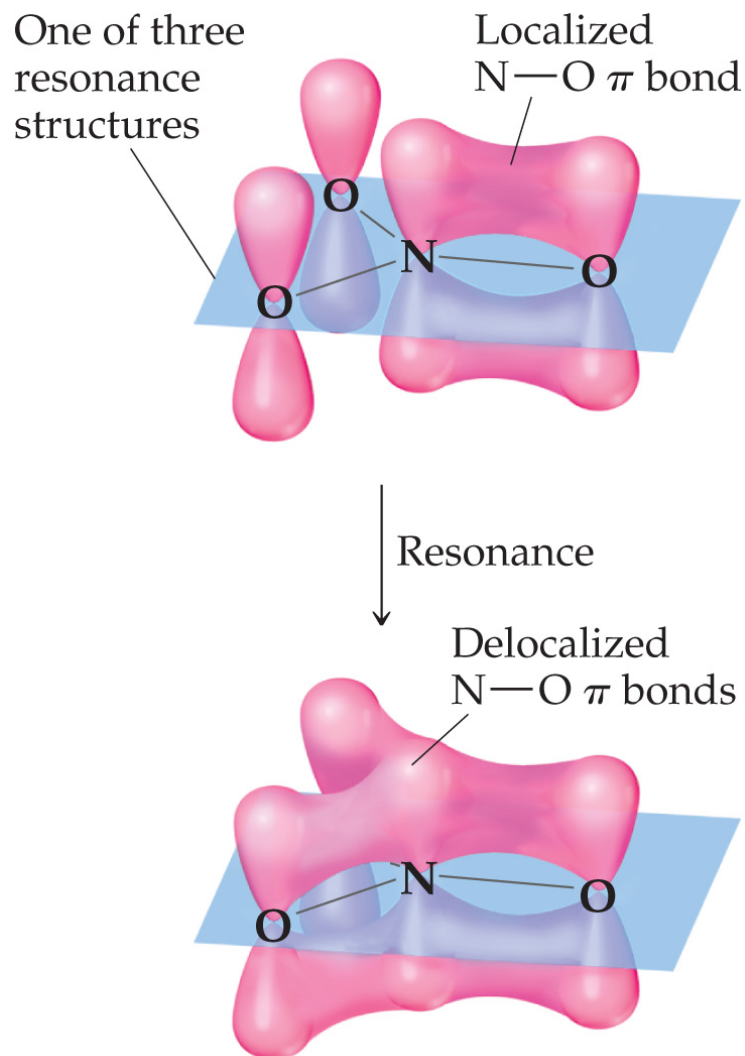
# Delocalized Electrons: Resonance

When writing Lewis structures for species like the nitrate ion, we draw resonance structures to more accurately reflect the structure of the molecule or ion.



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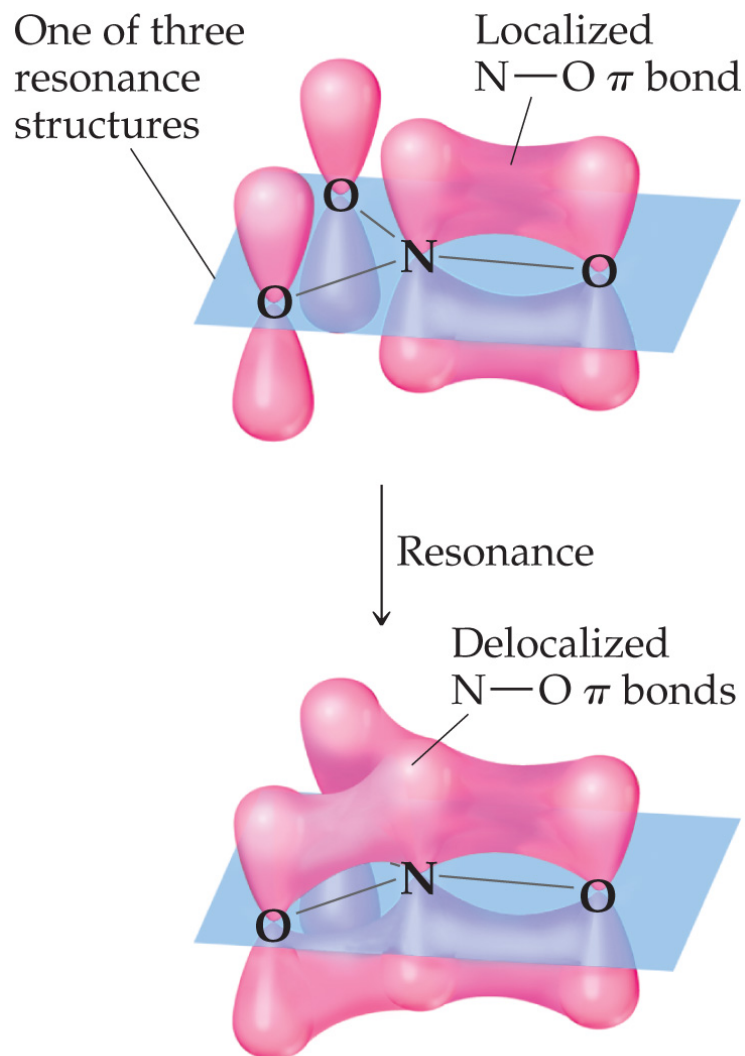
# Delocalized Electrons: Resonance



- In reality, each of the four atoms in the nitrate ion has a  $p$  orbital.
- The  $p$  orbitals on all three oxygens overlap with the  $p$  orbital on the central nitrogen.



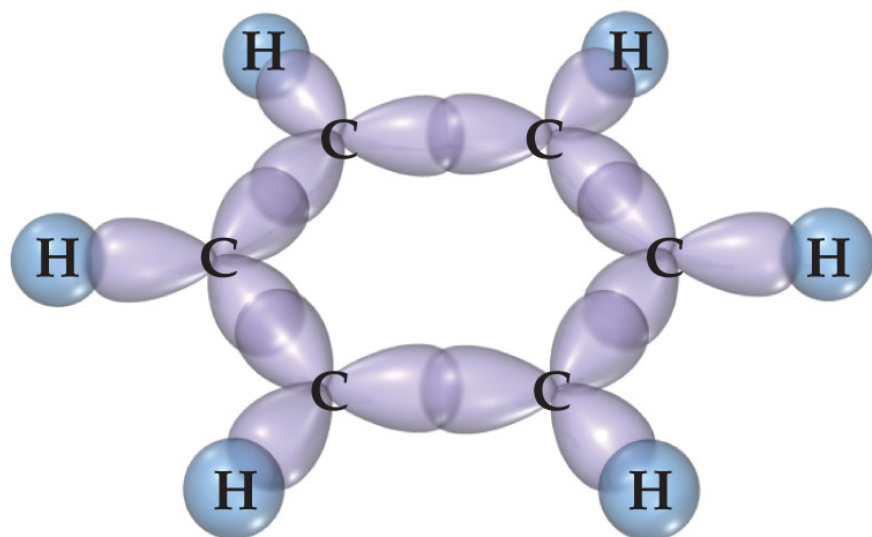
# Delocalized Electrons: Resonance



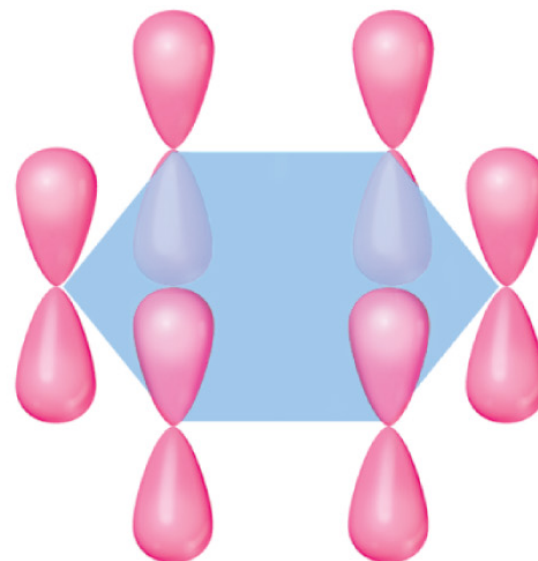
This means the  $\pi$  electrons are not localized between the nitrogen and one of the oxygens, but rather are delocalized throughout the ion.

# Resonance

The organic molecule benzene has six  $\sigma$  bonds and a  $p$  orbital on each carbon atom.



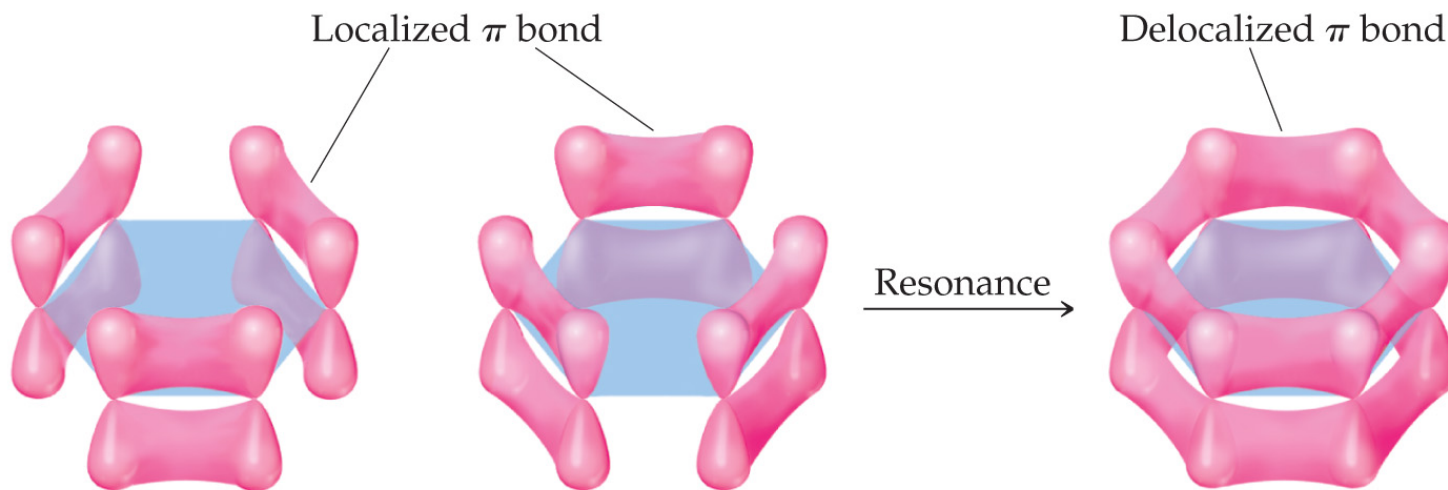
(a)  $\sigma$  bonds



(b)  $2p$  atomic orbitals

# Resonance

- In reality the  $\pi$  electrons in benzene are not localized, but delocalized.
- The even distribution of the  $\pi$  electrons in benzene makes the molecule unusually stable.



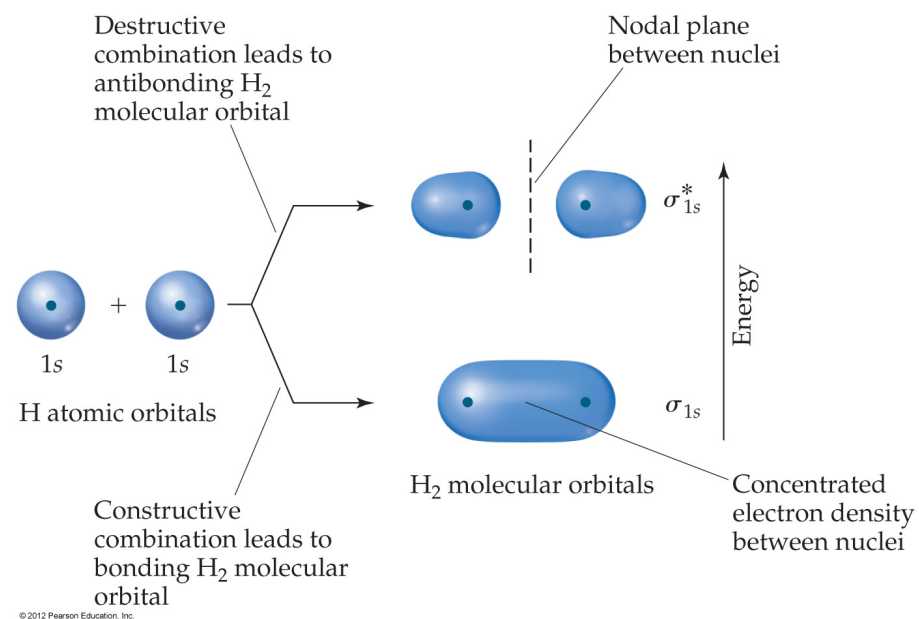
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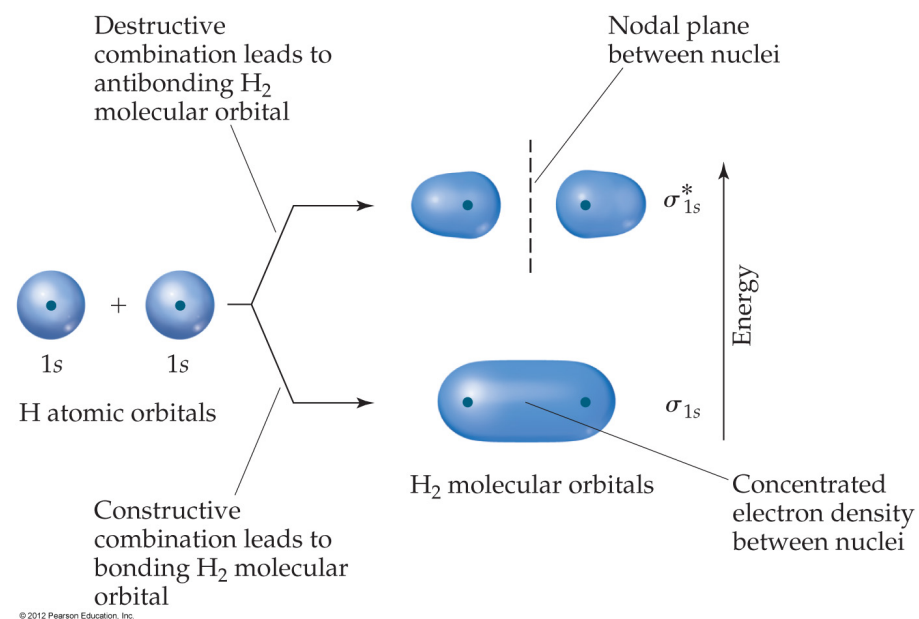
# Molecular-Orbital (MO) Theory

Though valence bond theory effectively conveys most observed properties of ions and molecules, there are some concepts better represented by molecular orbitals.



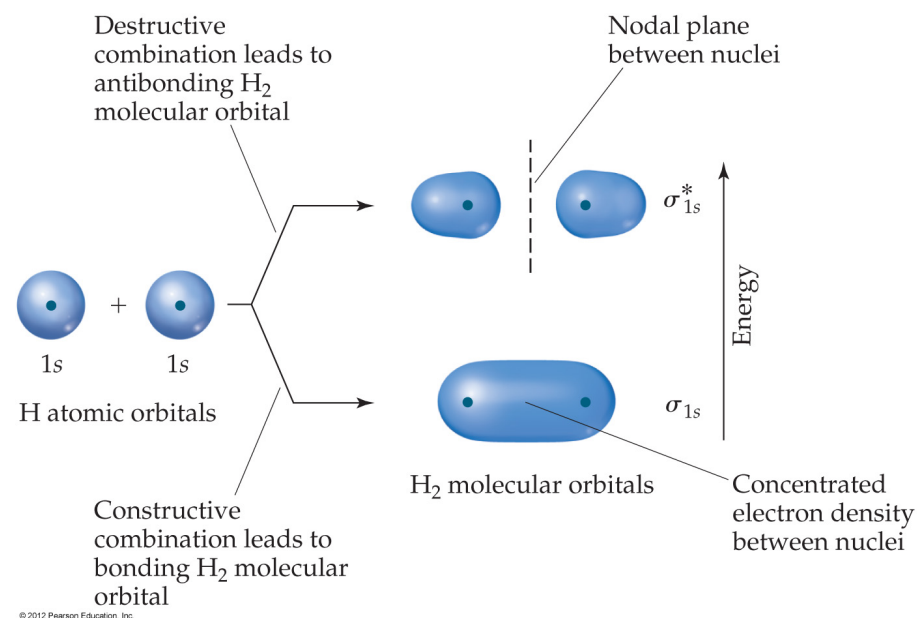
# Molecular-Orbital (MO) Theory

- In MO theory, we invoke the wave nature of electrons.
- If waves interact constructively, the resulting orbital is lower in energy: a bonding molecular orbital.

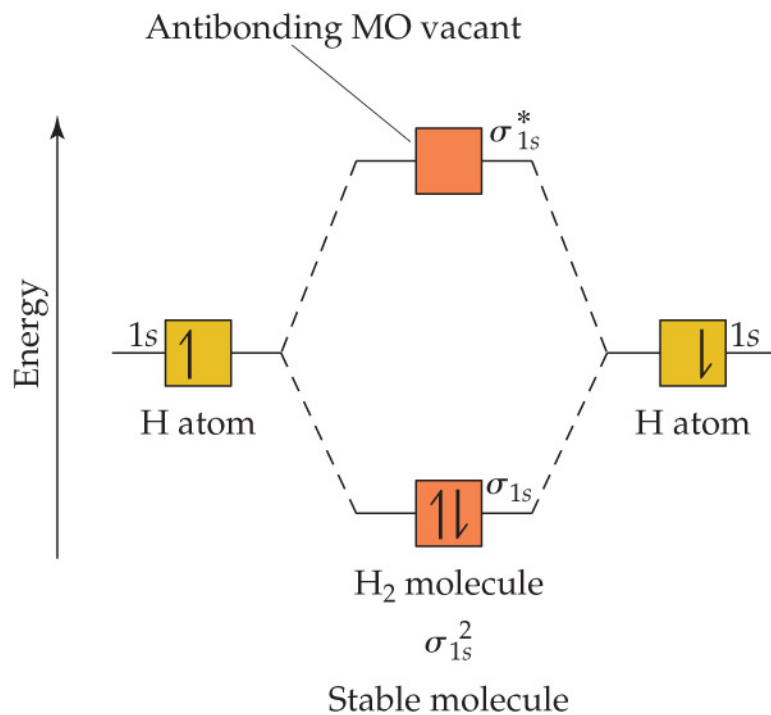


# Molecular-Orbital (MO) Theory

If waves interact destructively, the resulting orbital is higher in energy: an antibonding molecular orbital.



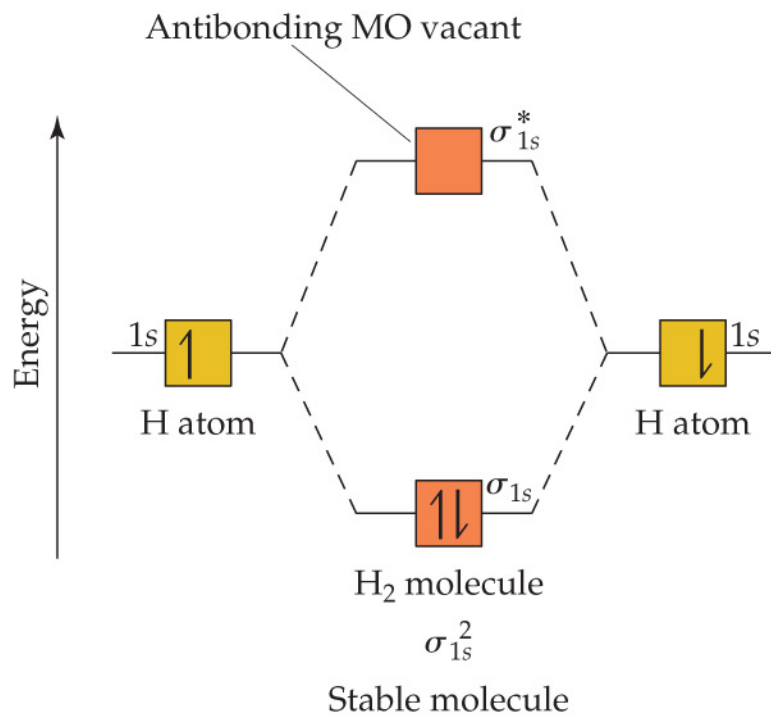
# MO Theory



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- In H<sub>2</sub> the two electrons go into the bonding molecular orbital.
- The bond order is one half the difference between the number of bonding and antibonding electrons.

# MO Theory



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For hydrogen, with two electrons in the bonding MO and none in the antibonding MO, the bond order is

$$\frac{1}{2} (2 - 0) = 1$$

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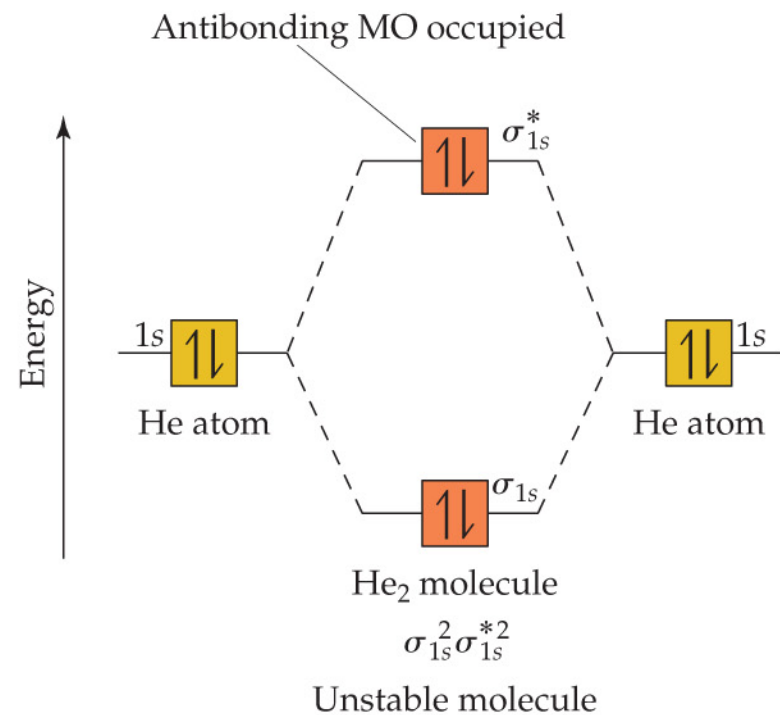


# MO Theory

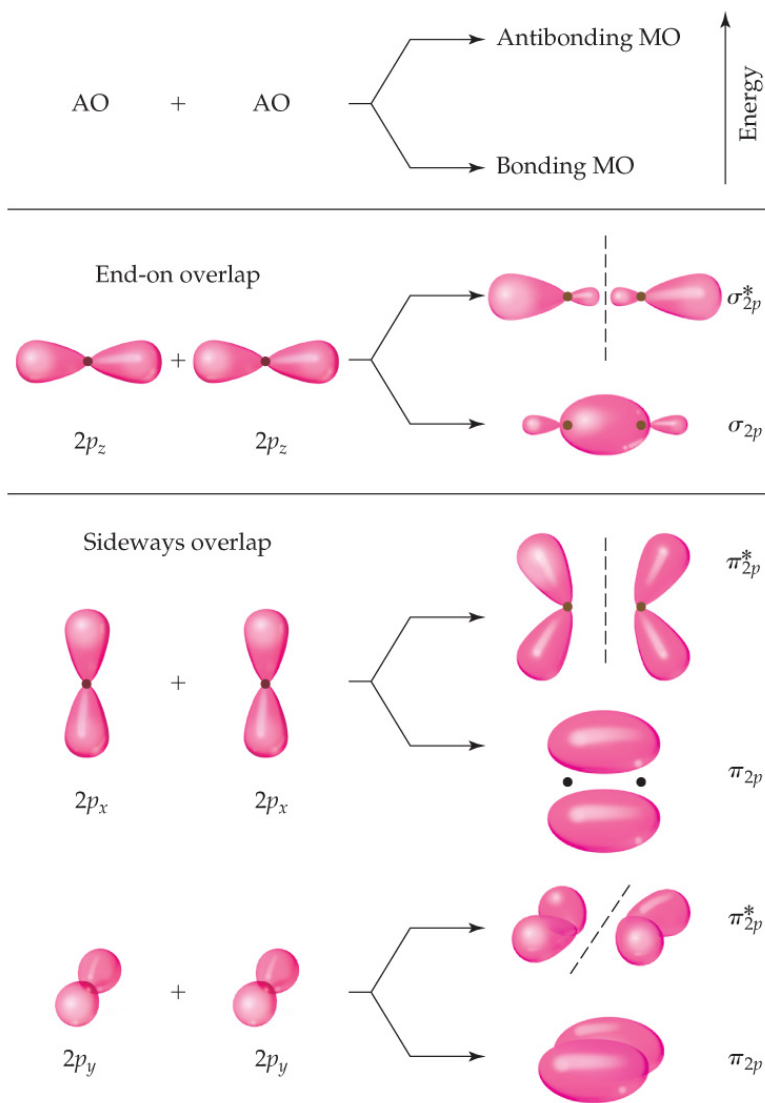
- In the case of  $\text{He}_2$ , the bond order would be

$$\frac{1}{2} (2 - 2) = 0$$

- Therefore,  $\text{He}_2$  does not exist.



# MO Theory

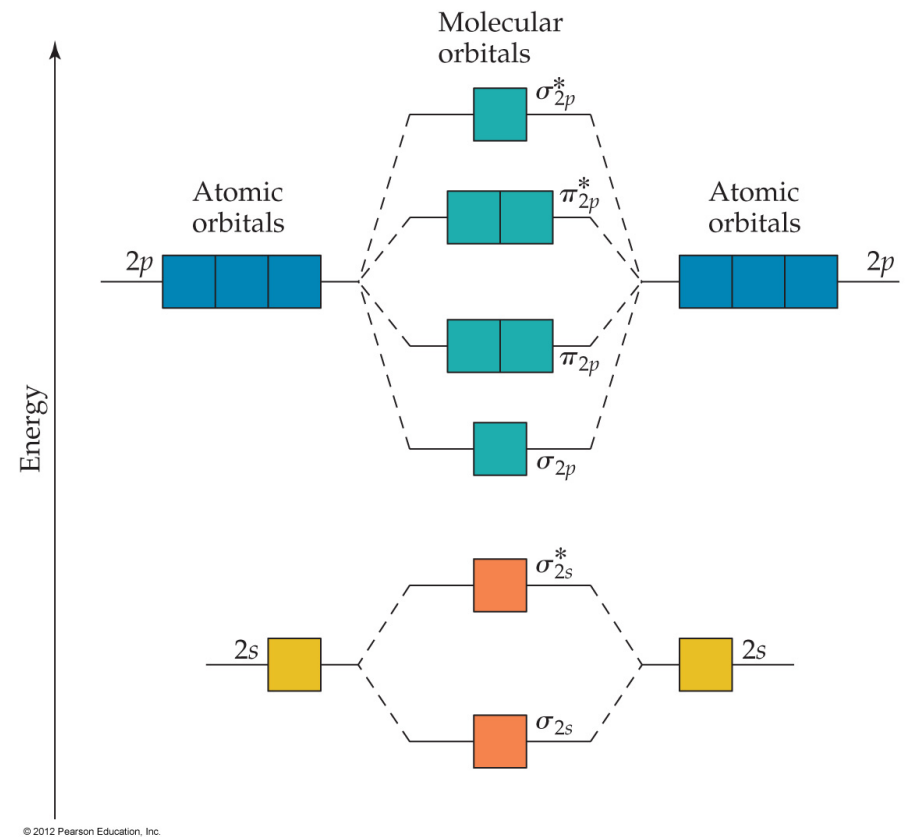


- For atoms with both  $s$  and  $p$  orbitals, there are two types of interactions:

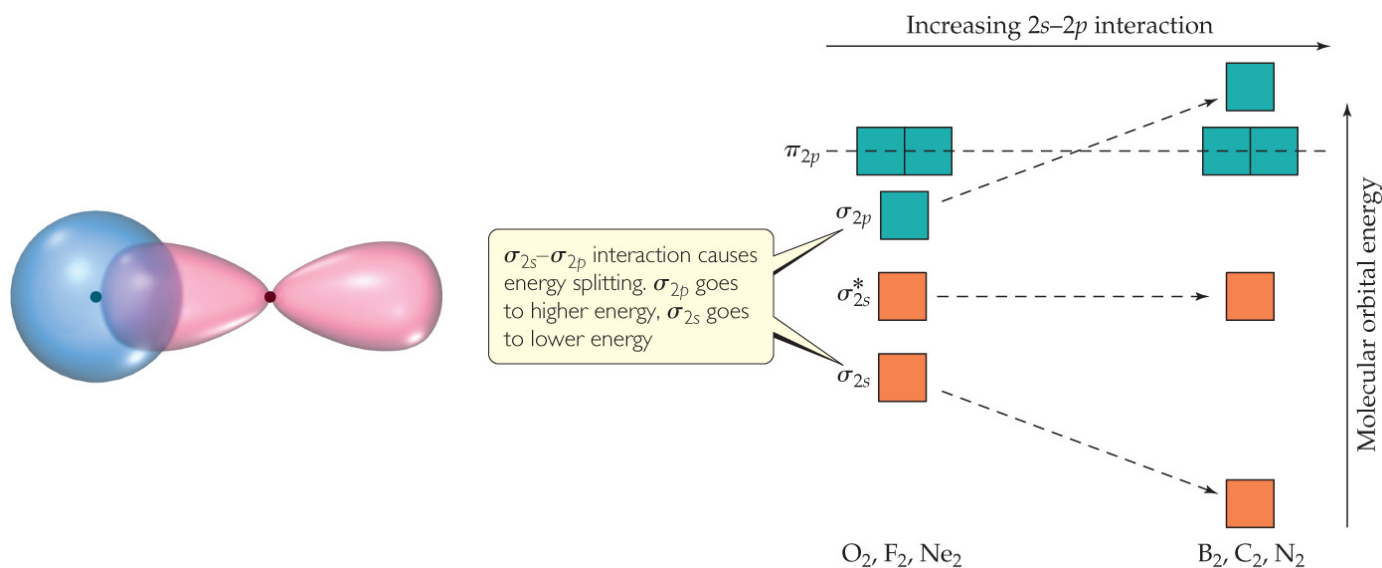
- The  $s$  and the  $p$  orbitals that face each other overlap in  $\sigma$  fashion.
- The other two sets of  $p$  orbitals overlap in  $\pi$  fashion.

# MO Theory

- The resulting MO diagram looks like this (Fig. 9.41).
- There are both  $\sigma$  and  $\pi$  bonding molecular orbitals and  $\sigma^*$  and  $\pi^*$  antibonding molecular orbitals.



# MO Theory



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- The smaller  $p$ -block elements in the second period have a sizable interaction between the  $s$  and  $p$  orbitals.
- This flips the order of the  $\sigma$  and  $\pi$  molecular orbitals in these elements.

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# Second-Row MO Diagrams

	Large 2s–2p interaction			Small 2s–2p interaction		
	B <sub>2</sub>	C <sub>2</sub>	N <sub>2</sub>	O <sub>2</sub>	F <sub>2</sub>	Ne <sub>2</sub>
$\sigma_{2p}^*$	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox" value="1↓"/>
$\pi_{2p}^*$	<input type="checkbox" value="1"/> <input type="checkbox"/>	<input type="checkbox" value="1"/> <input type="checkbox"/>	<input type="checkbox" value="1"/> <input type="checkbox"/>	<input type="checkbox" value="1"/> <input type="checkbox" value="1"/>	<input type="checkbox" value="1↓"/> <input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/> <input type="checkbox" value="1↓"/>
$\sigma_{2p}$	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/> <input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/> <input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/> <input type="checkbox" value="1↓"/>
$\pi_{2p}$	<input type="checkbox" value="1"/> <input type="checkbox" value="1"/>	<input type="checkbox" value="1↓"/> <input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/> <input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>
$\sigma_{2s}^*$	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>
$\sigma_{2s}$	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>	<input type="checkbox" value="1↓"/>
Bond order	1	2	3	2	1	0
Bond enthalpy (kJ/mol)	290	620	941	495	155	—
Bond length (Å)	1.59	1.31	1.10	1.21	1.43	—
Magnetic behavior	Paramagnetic	Diamagnetic	Diamagnetic	Paramagnetic	Diamagnetic	—

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