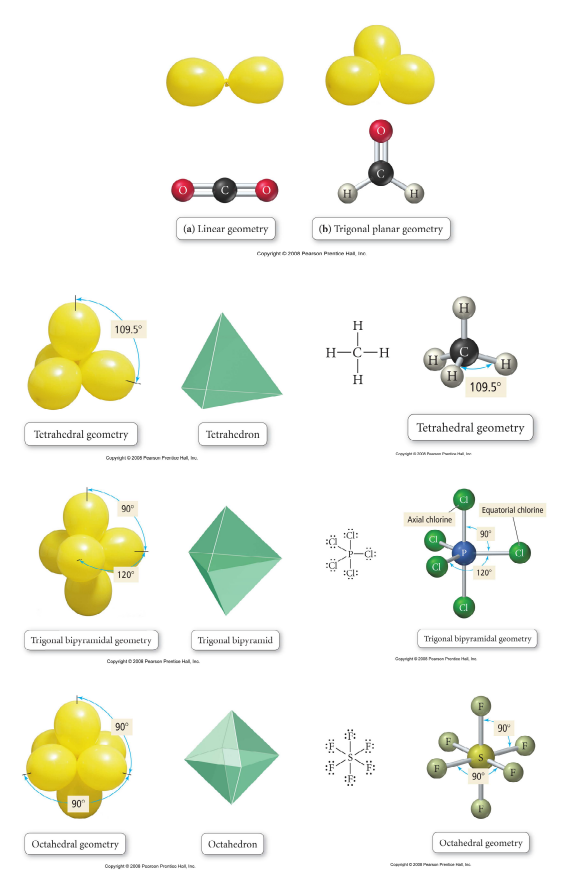
**<u>VSEPR Theory</u>**: Valence shell electron pair repulsion

- Electron groups repel one another through coulombic forces
- They will spread as far apart as possible on a molecule's central atom
- <u>1 electron group:</u>
  - 1 single bond
  - 1 double bond
  - 1 triple bond
  - 1 lone (unshared) pair of electrons

# The five major electron group geometries:

<u># e<sup>-</sup> groups</u>	<u>Geometry</u>	<u>Structure</u>	Ideal bond angle
2	Linear		
3	Trigonal planar		
4	Tetrahedral		
5	Trigonal bipyramidal		
6	Octahedral		

### Electron group geometries



Lone pairs and molecular geometry

**Electron group geometry** is the arrangement of the electron groups

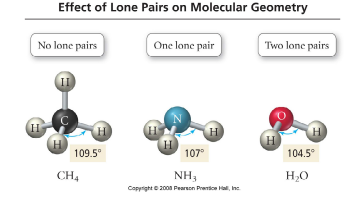
**Molecular geometry** is the arrangement of the atoms

These are the same if only **bonding** electron groups are attached to the central atom.

Nonbonding electrons (lone pairs) on the central atom will change the **molecular geometry**.

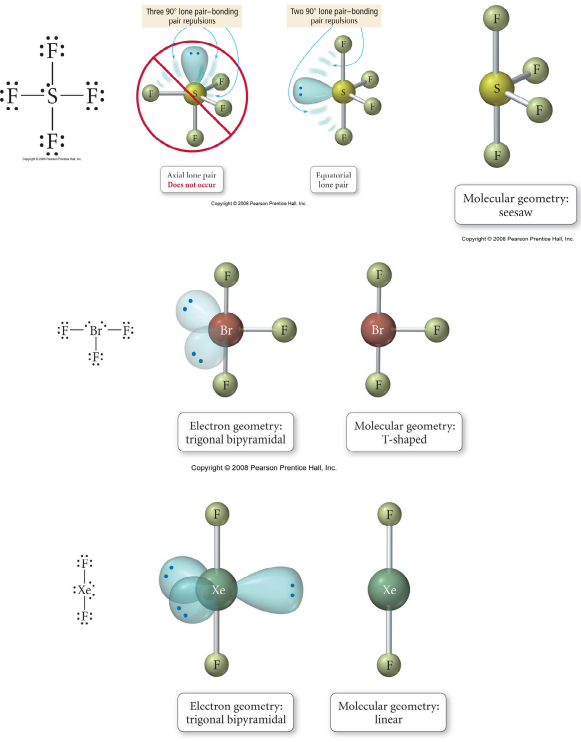
	_	e <sup>-</sup> group geom.	<u>structure</u>
CH4			
$NH_3$			
H <sub>2</sub> O			

Lone pairs will actually repel the bonds a little more than a bond would, decreasing bond angles slightly.



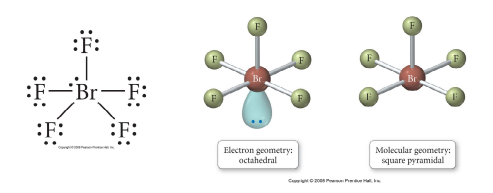
#### 5 electron groups with lone pairs

Lone pairs will **only** be placed in equatorial positions in the trigonal bipyramidal electron group geometry.

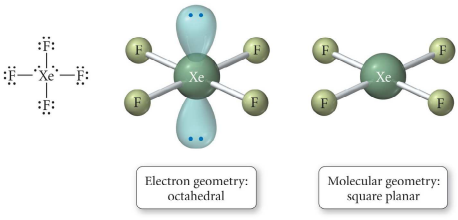


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6 electron groups with lone pairs



Two lone pairs in an octahedral electron group geometry will add across from each other to minimize lone pairlone par repulsions.



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Electron groups	Bonding groups	Nonbonding groups	Molecule shape	Example	Drawing
2	2	0	Linear	BeF <sub>2</sub>	F—Be—F
3	3	0	Trigonal planar	BF <sub>3</sub>	F   F <sup>_B</sup> _F
	2	1	Bent	$SO_2$	∷ ₀≈≈⊳₀
	4	0	Tetrahedral	CH4	H-COL
4	3	1	Trigonal pyramid	NH3	
	2	2	Bent	H <sub>2</sub> O	н_о
5	5	0	Trigonal bipyramid	PCl5	0-0-0 0-0-0
	4	1	Seesaw	$SF_4$	:s
	3	2	T-shaped	ClF3	н- <u>а</u> -н
	2	3	Linear	$XeF_2$	-Xe
	6	0	Octahedral	$SF_6$	F F F F
6	5	1	Square pyramid	IF <sub>5</sub>	
	4	2	Square planar	XeF4	F_Xe_F

Geometry practice

What is the electron group and molecular geometry of IBr<sub>5</sub>? Draw its flat Lewis structure and its 3-dimensional structure.

What is the electron group and molecular geometry of ICl<sub>2</sub>-? Draw its flat Lewis structure and its 3-dimensional structure.

Molecular shape and polarity

The dipoles of polar bonds will add together geometrically to form a <u>net dipole moment</u> for the molecule. Molecules with a net dipole moment are <u>polar</u>.

 $H_2O$ :

CO<sub>2</sub>:

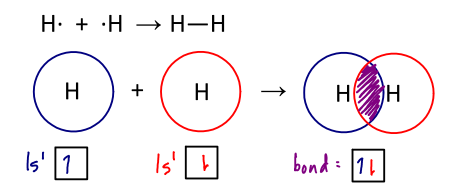
BF<sub>3</sub>:

 $CH_2F_2$ :

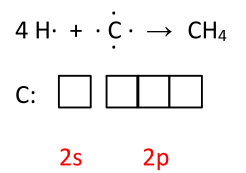
CO<sub>3</sub><sup>2-</sup>:

Valence bond theory

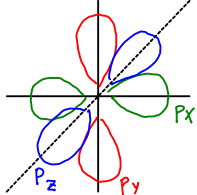
In valence bond theory, bonds are formed by the orbitals of two atoms overlapping.



But, many times the orbitals cannot combine as-is.



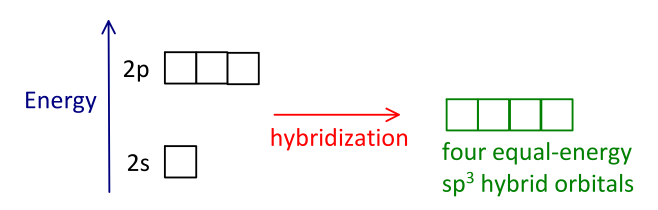
Remember, p orbitals are oriented on the x, y, and z axes:



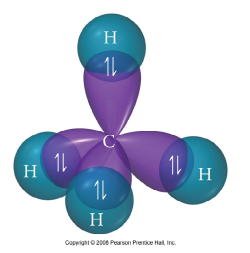
Using VSPER, what is the shape of the CH<sub>4</sub> molecule?

sp<sup>3</sup> hybrid orbitals

The one s and three p orbitals in carbon's valence shell combine together into 4 equivalent **hybrid orbitals** so carbon can make 4 bonds.



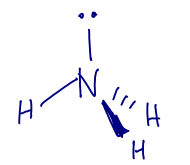
According to VSEPR, four equivalent hybrid orbitals (each containing one electron group) will best fit around a central atom with a \_\_\_\_\_\_ geometry.



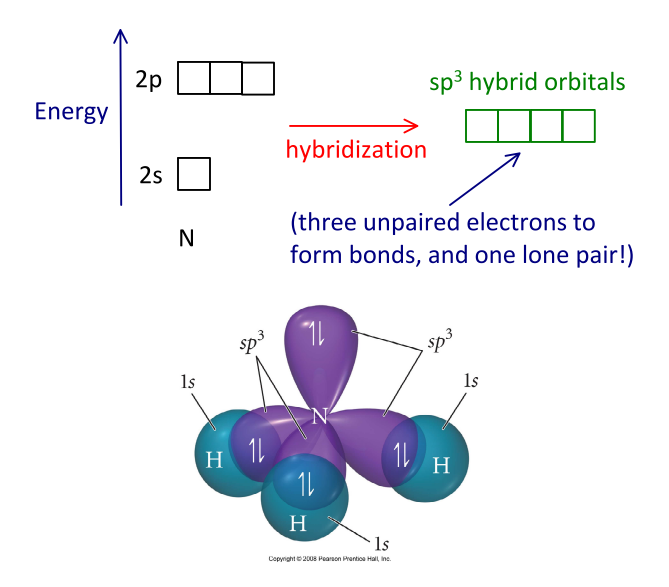
Any time there's a \_\_\_\_\_\_ electron group geometry, the hybridization of the central atom is \_\_\_\_\_.

sp<sup>3</sup> hybrid orbitals

NH<sub>3</sub>: electron group geometry:molecular geometry:# electron groups on central atom:

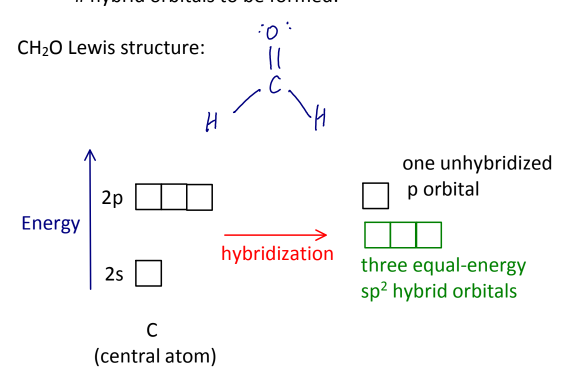


(The number of electron groups on the central atom is the number of hybrid orbitals that need to be formed!)

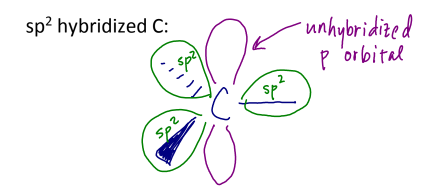


sp<sup>2</sup> hybrid orbitals

CH<sub>2</sub>O: # electron groups on central atom: # hybrid orbitals to be formed:

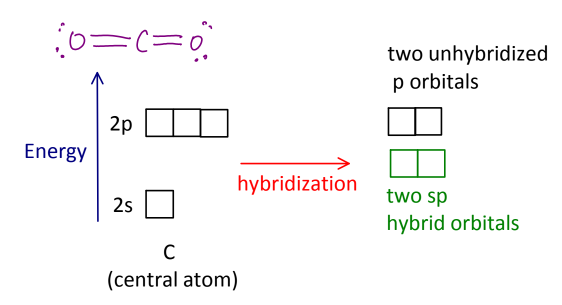


The three sp<sup>2</sup> hybrid orbitals will be \_\_\_\_\_\_ in shape. The unhybridized p orbital is perpendicular.



sp hybridization

CO<sub>2</sub>: 2 electron groups around C, so 2 hybrid orbitals



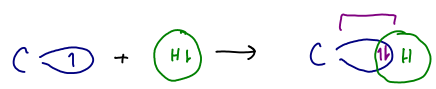
An sp hybridized central atom will be \_\_\_\_\_ in shape.

sp hybridized C:

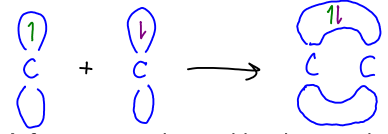
sp sp

 $\sigma$  and  $\pi$  bonds

The bonds in valence bond theory are classified by their positions relative to the two bonding atoms. If two half-filled orbitals combine straight between the two atoms, it's called a  $\sigma$  (sigma) bond.

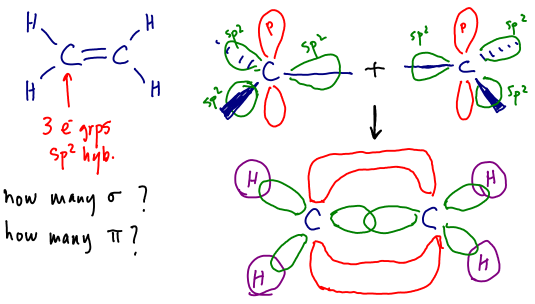


When two half-filled p orbitals combine side-by-side, it's called a  $\pi$  (pi) bond.

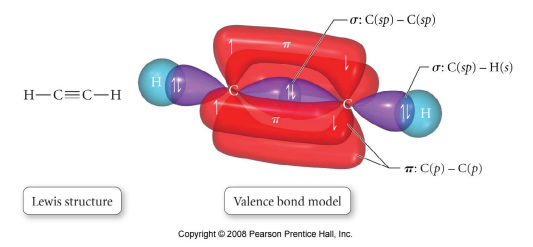


A <u>single bond</u> from Lewis theory like the C–H bonds in  $CH_4$  is made of a single  $\sigma$  (sigma) bond.

A <u>double bond</u> from Lewis theory like the C=C bond in  $C_2H_4$  is formed by one  $\sigma$  bond and one  $\pi$  bond.



If an atom is sp hybridized, it has 2 hybrid orbitals and 2 unhybridized p orbitals.



# Summary of $\sigma$ and $\pi$ bonding:

<b>Bond type:</b>	Lewis:	Valence bond theory:
Single		σ, π
Double	—	σ, π
Triple	$\equiv$	σ,π

## **Summary of hybrid orbitals**

<u>hybridization</u>	<u>unhybridized</u> <u>p orbitals</u>
	<u>hybridization</u>