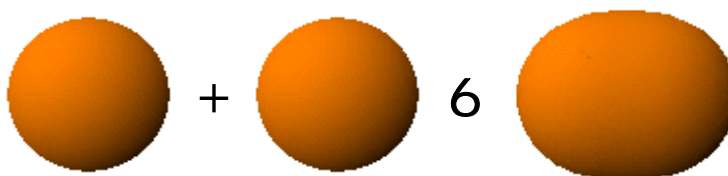


Quantum Mechanical Approaches to Molecular Bonding

- (In principle, it is possible to construct a Schrödinger equation, $\hat{H}\Psi = E\Psi$, to describe the electronic structure of a molecule.
- ; In practice, seeking exact solutions to the Schrödinger equation for molecules is an insurmountable mathematical problem.
- L Two principal approaches have been taken to construct approximate wave functions for molecules, starting with the atomic orbitals of the atoms comprising the molecules.
 1. Valence Bond (VB) theory - developed by Linus Pauling and co-workers, essentially puts the Lewis notion of electron pairs on a quantum mechanical footing, in which each shared or lone pair of electrons about an atom occupies a localized orbital.
 2. Molecular orbital (MO) theory, developed by Robert S. Mulliken and co-workers, constructs new orbitals that are *delocalized* (i.e., “spread out”) across the molecule.
- L VB and MO theories take different mathematical approaches to constructing wave functions for the molecule, but their results are often equivalent.

Valence Bond (VB) Theory

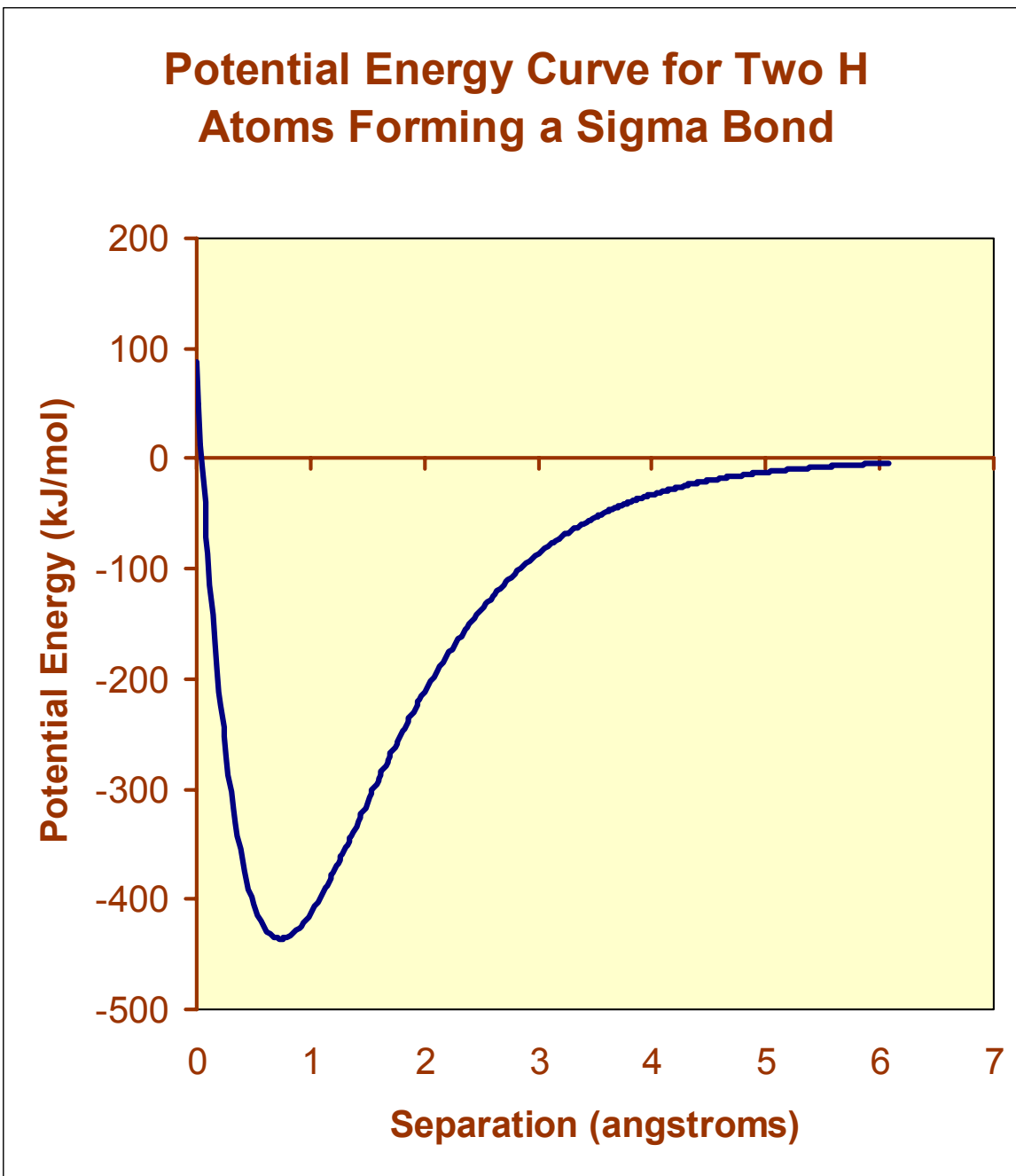
- L A chemical bond forms by sharing a pair of electrons through overlap of atomic orbitals on the bonded atoms.
- L When overlap creates an increase in electron density in the region between the two nuclei a **sigma bond** (σ bond) is formed.



Formation of sigma (σ) bond in H_2 from $1s$ orbitals

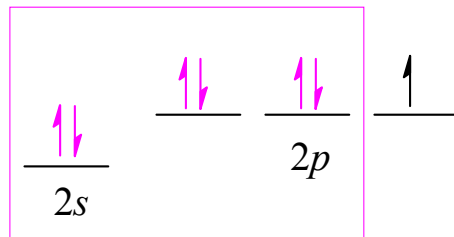
Potential Energy and Internuclear Separation

- Attraction of the electrons to two nuclei causes a lowering of the potential energy to a minimum at the normal bond distance.



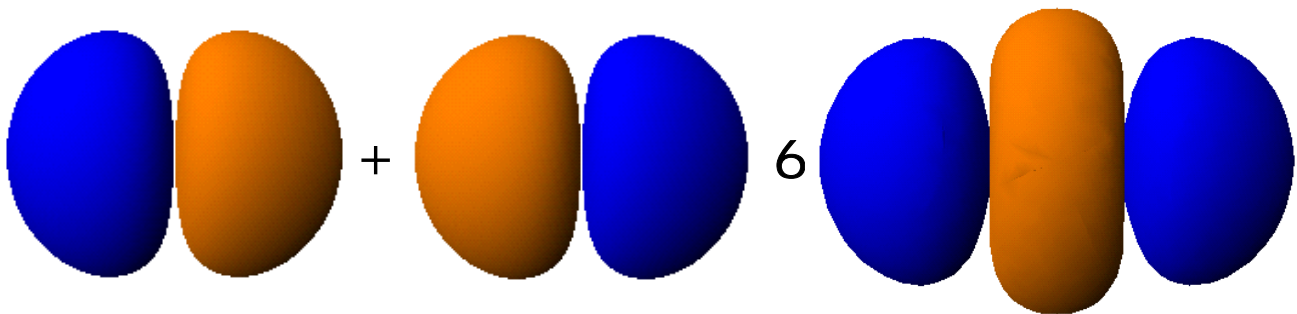
Sigma Bond Formation in F₂

Valence configuration:



Non-bonding electrons

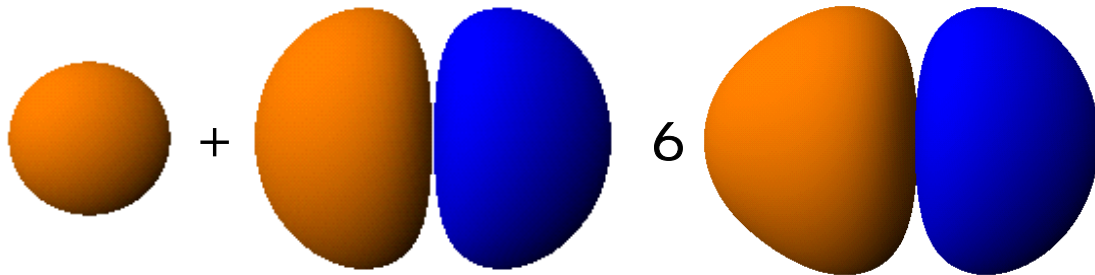
- The single bond in F₂ can be seen as a sigma bond formed by overlap of two 2p orbitals.



Formation of sigma (σ) bond in F₂ from 2p

Heteronuclear Diatomic Molecules

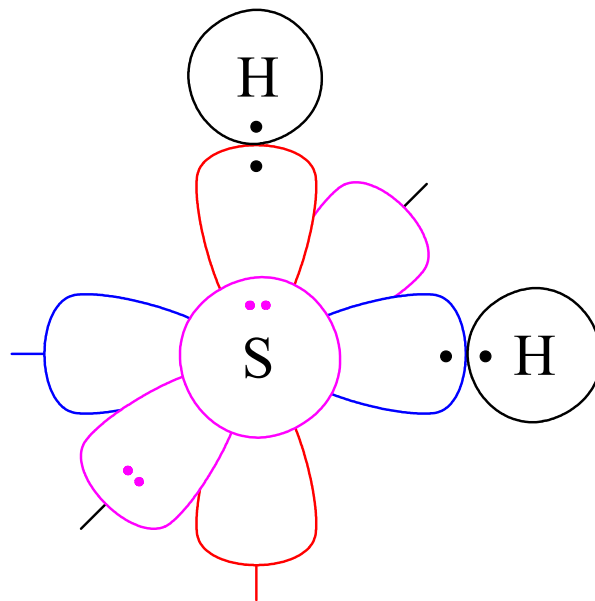
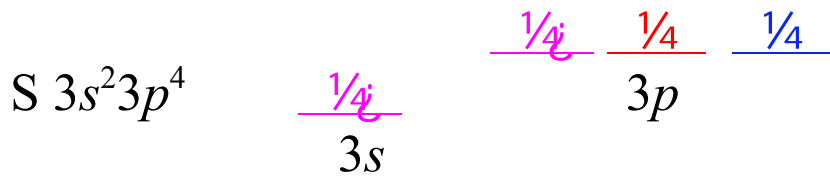
- In heteronuclear diatomic molecules, the overlap may involve two different types of orbitals.



Formation of sigma (σ) bond in HF from H $1s$ and F $2p$

Polyatomic Molecules

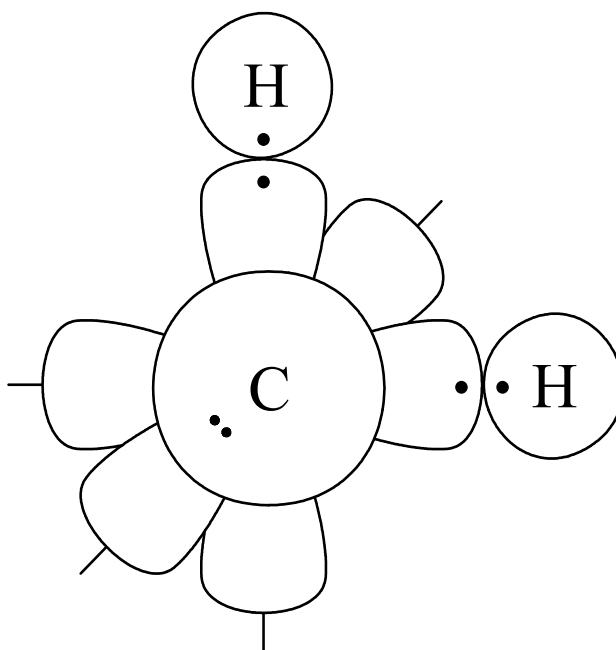
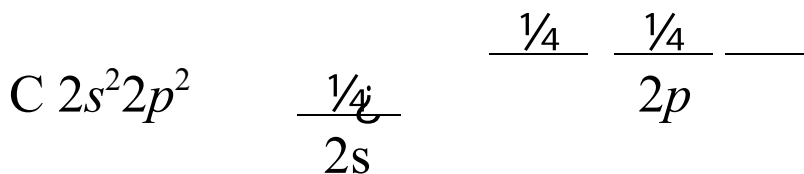
Example: H₂S



Polyatomic Molecules The Need for Hybridization

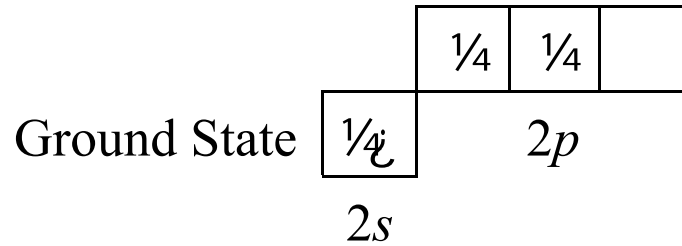
How can we form a VB model of four equal sigma bonds around carbon, starting with its valence configuration $2s^2 2p^2$?

- It seems that only two bonds could be made with an angle of 90° through overlap of carbon $2p$ orbitals with hydrogen $1s$ orbitals.

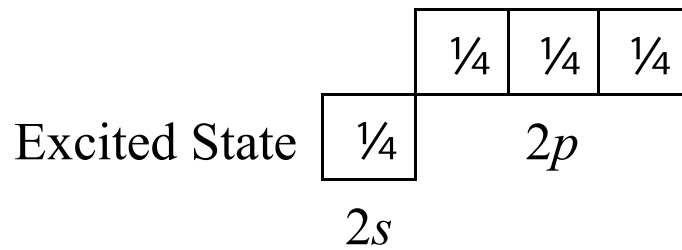


L But CH_2 does not exist as a stable molecule, and CH_4 is formed instead.

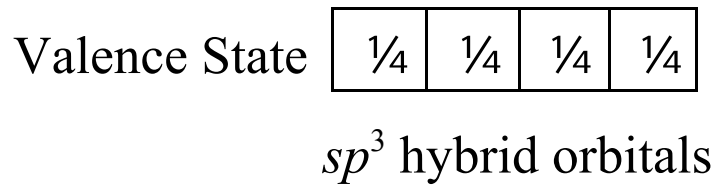
Hybrid Orbital Formation in CH₄ A Hypothetical Process



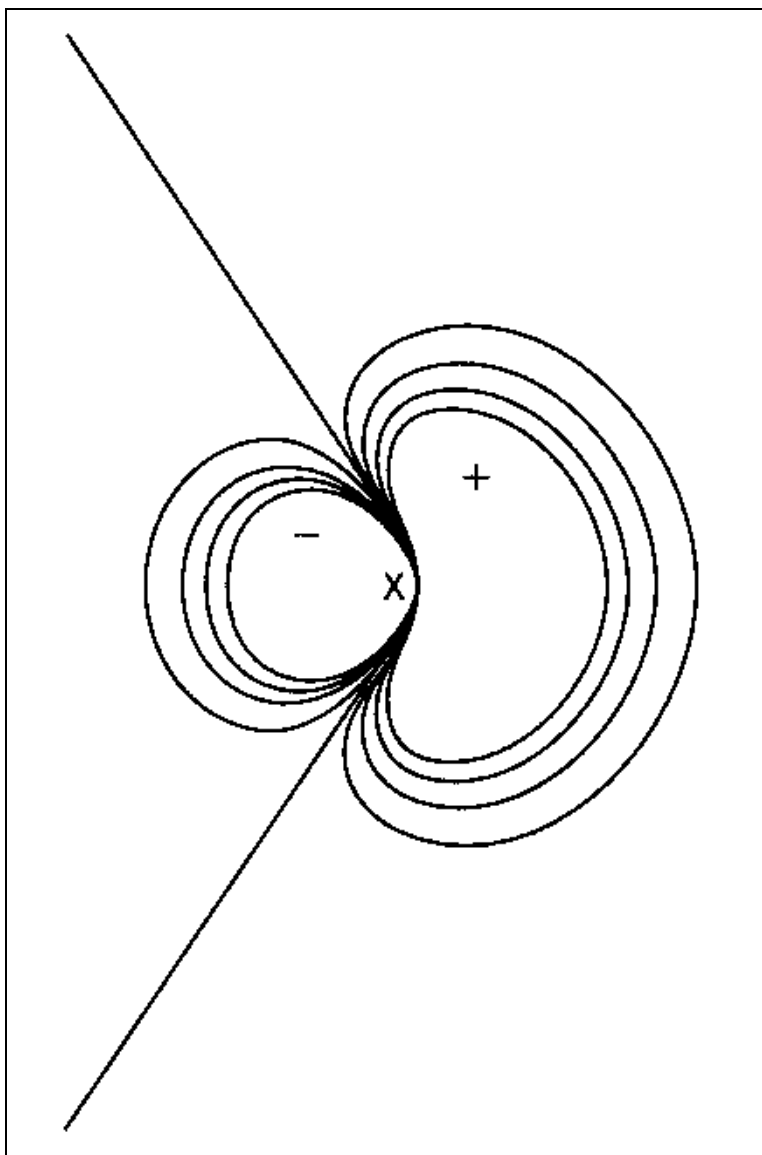
\



\



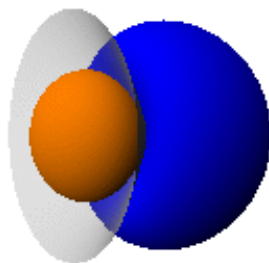
Contour Diagram of a Single sp^3 Hybrid Orbital



X marks the position of the nucleus

Boundary Surface Model of a Single sp^3 Hybrid

Three-dimensional model (rotated 30° about a vertical axis):

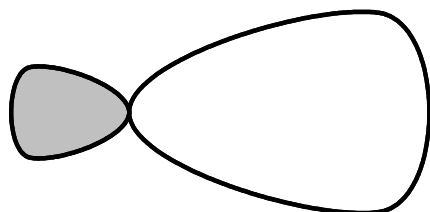


Cut-away rendering:

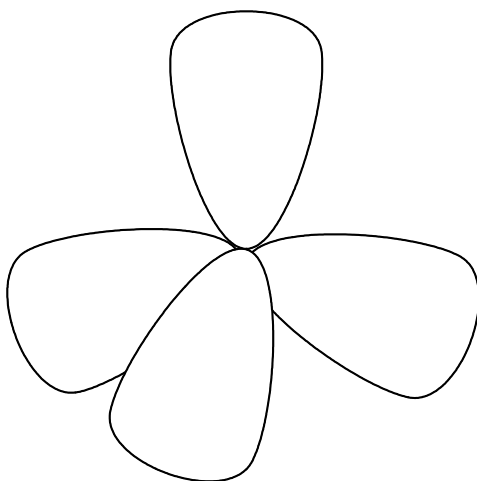


sp^3 Hybrid Orbitals - Simplified Sketches

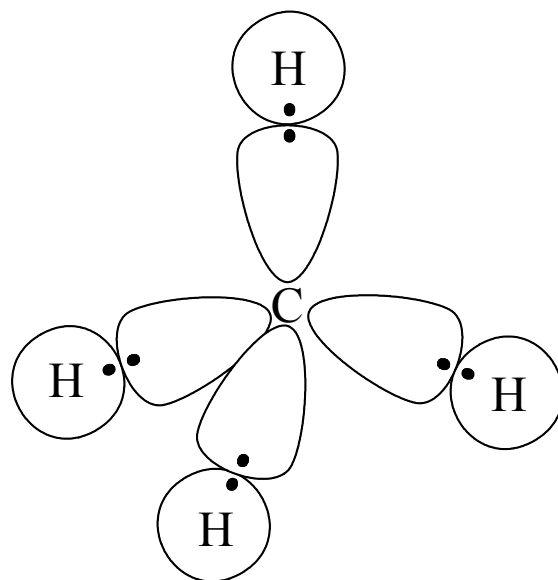
Shape of an individual sp^3 hybrid orbital:



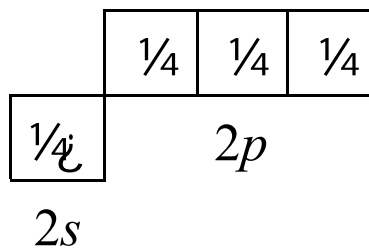
Set of four sp^3 hybrid orbitals in a tetrahedral arrangement:



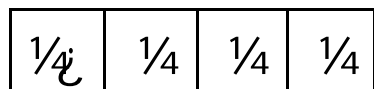
VB Model of CH₄ with sp^3 Hybrids on C



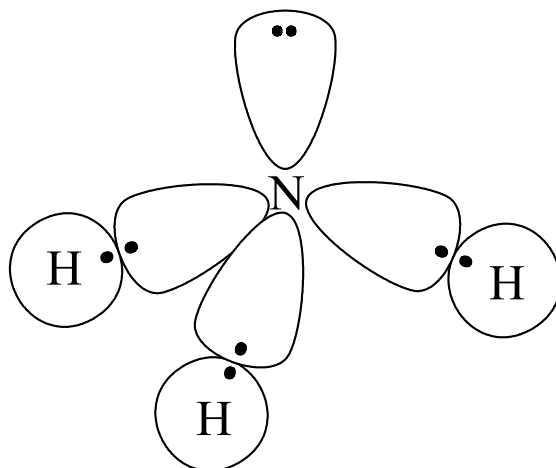
VB Model of NH_3 with sp^3 Hybrids on N



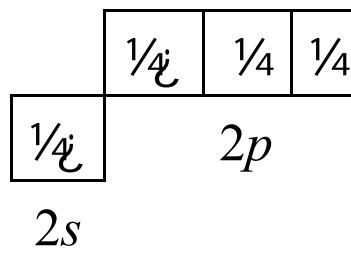
\



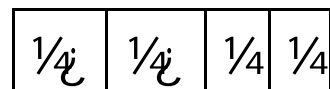
sp^3 hybrid orbitals



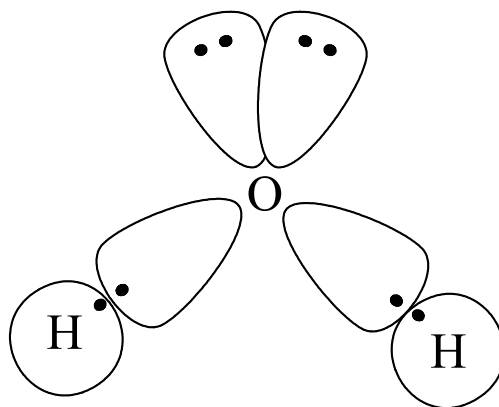
VB Model of H₂O with sp^3 Hybrids on O



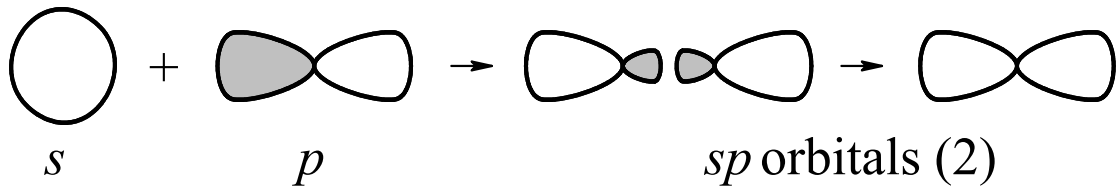
\



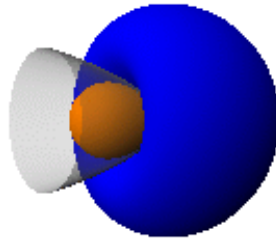
sp^3 hybrid orbitals



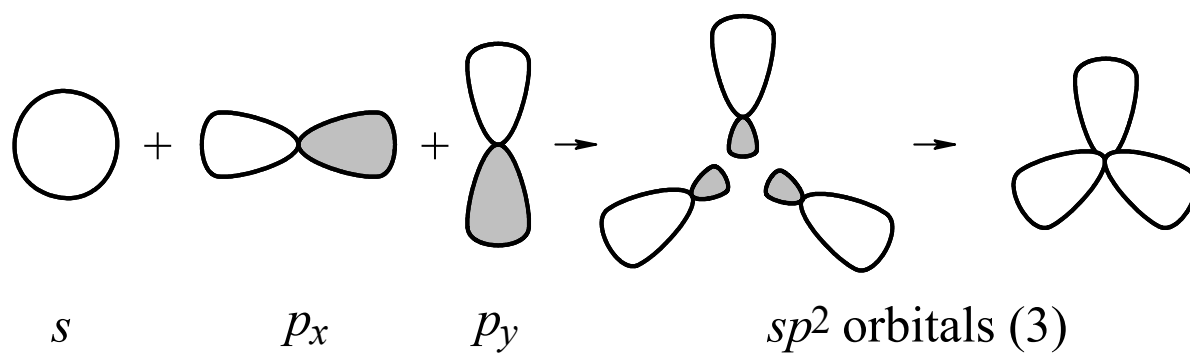
Hybrids for 2 Electron Domains



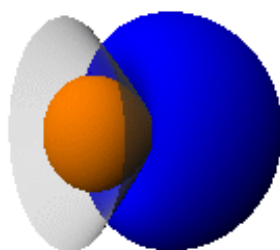
Boundary surface model of one *sp* hybrid:



Hybrids for 3 Electron Domains



Boundary surface model of one sp^2 hybrid:



Summary of Hybrid Orbital Types

Domains	Geometry	Orbitals Used	Hybrids
2	linear	s, p	sp
3	trigonal planar	s, p_x, p_y	sp^2
4	tetrahedral	s, p_x, p_y, p_z	sp^3
5	trigonal bipyramidal	$s, p_x, p_y, p_z, d_{z^2}$	sp^3d
6	octahedral	$s, p_x, p_y, p_z, d_{z^2}, d_{x^2-y^2}$	sp^3d^2