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## Week 1

### Friday, October 2

- ▶ Determing formal charge:
  - Formula:  $FC = V N \frac{B^1}{2}$
  - V = valance electrons of element
  - N = lone pair electrons; B = bonded electrons
- 1. What is the formal charge on P in the following structure? Each F and O has three lone pair of electrons.
  - $\circ$  P = 5 O 8(0.5); P = +1
- 2. What is the formal charge on O in the structure above?
  - $\circ$  O = 6 6 2(0.5); O = -1
- 3. What is the formal charge on P in the following structure? Each F still has three lone pairs of electrons, and O had the tow pairs indicated.
  - $\circ$  P = 5 0 10(0.5); P = **0**
- 4. Of the two structures shown for POF<sub>3</sub>, which is the most stable, and will, therefore, be the most abundant form?
  - Structure II
  - O has formal charge of **0** and is the most electronegative element with difference in charge between the resonance structures.
  - F has greater electronegativity, but remains the same between both structures, so it's not relevant.
  - Key difference: the double bond in structure II gives oxygen the lower magnitude formal charge between the two.
- 5. The fundamental concept upon which VSEPR, and hence molecular shapes, is based is that:
  - Electrons pairs repel each other;
    - negative charge repels other negative charges.
  - Electron repulsion is minimized by maximum angular separation;
    - in other words, angular separation maximizes distance between electrons.
  - Bonding pair electrons and lone pair electrons both occupy regions around the central atom:

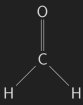
- if they didn't occupy the same space than they wouldn't interact and thus wouldn't affect shape.
- The electron dommain geometry and the molecular geometry is identical if there all of the electrons are bonding electrons;
  - the lone pairs are have a greater influence than bonded pairs, resulting in less space for bonded pairs.

#### All of the above

- ▷ General method of determining structure:
  - Count steric number—the total number of electron pairs in a molecule.
    Can be bonds or lone pairs.
  - 2. Determine predicted geometrical structure predicted (EDG) by VSEPR using steric number.
    - Octahedral:6, Bipyramid:5, Tetrahedral:4, Trigonal:3, Linear:2
  - 3. Determin impact (the MG) of lone pairs; more lone pairs results in less space between bonded pairs. Shape depends on EDG.
- 6. A resonance form of SOF<sub>2</sub>, completely consistent with the octet rule, is shown below. What is the electron domain geometry (EDG), and molecular geometry (MG) of this molecule?

#### Tetrahedral EDG and trigonal pyramidal MG

7. Draw a Lewis dot structure of formaldehyde ( $CH_2O$ ): what is the molecular shape of this molecule?



- Steric number = 3
  - Double bonds count as 1 for steric number.
- No lone pairs on central atom, C, so it's shape planar.
- Trigonal planar
- 8. The EDG for  $CH_3^-$  (a carbanion) is tetrahedral, and the MG is trigonal pyramidal. Why are the H-C-H bond angles less than 109.5° as in a perfect tetrahedron?
  - The lone pair electrons take up more space than bonding pair electrons.

## Week 2

#### Monday, October 5

1. The concept of orbital shapes comes directly from the wave model of the atom. What is the shape of an s orbital?

### > Spherical

- o S orbital is the most simple orbital, with only two electrons.
- Alternative shapes come from *nodes*; i.e. when *destructive interferences* cancels out the wave function.
- o Not circular, orbitals are three-dimensional.
- 2. What is the shape of a p orbital?

#### Dumbell shaped

- P orbital can hold 6 electrons (3 pairs).
- Each pair has one angular node, squeezing shape into dumbell in each direction (x, y, z).
- 3. When atomic orbitals overlap to form a covalent bond, the resultant bonding orbital is:

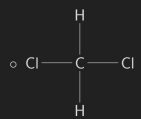
#### Lower in energy than the atomic orbitals from which it was formed

- Electrons in a covalent bond are in a more stable (lower energy) state due to multiple nuclei hodling them in place.
- Only when nodes are present do the electrons create a destabalized molecular orbit, incraseing the energy.
- 4. Why can't pure p orbitals be used in forming four equivalent bonds as in methane?
  - the three 2p orbitals can only hold 6 electrons.
    - True, we need to make four bonds for methane.
  - ▶ the bonds would have to be 90° apart.
    - If p had enough space then it would result in planar geometry with 90°, but the true arrangement is tetrahedral with angles of 109.5°
  - ▷ electron-electron repulsion would not be minimized.
    - Planar minimization would be 90°, but we have 3d space to work with, so it's not minimized.

#### > all of the above

- 5. When s and p orbitals combine to form hybrid orbitals, the resultant hybridized orbitals are:
  - ▷ lower in energy than the p orbitals
  - ▷ higher in energy than the s orbitals
  - > both of the above
    - It takes energy to move the electron up from the s orbital and hybridize the p orbitals.
    - The new hybridized sp orbital also has more energy than the sorbital.
- 6. What is the difference between a sigma bond and a pi bond?
  - $\triangleright$  in a  $\pi$  bond, electron density lies above and below the axis that conects the two nuclei; in a  $\sigma$  bond, the electronegative density lies along the axis that connets the two nuclei
    - $\circ$   $\sigma$  bond has circular symmetry with respect to the bond axis (axis that connets the two nuclei). i.e. it's along the axis.
- 7. What is the hybridization of the C in CH2Cl2?

 $\triangleright$  sp<sup>3</sup>



- Look at central atom C
- o Determin groups (number of bonds,  $\pi$  bonds count as 1, and lone pairs attached) 4
- o for groups 1-4;  $sp^x$ ; x = groups 1(3)
- This is also a trigonal pyramidal, due to lone pairs on Cl I guess?
  (lone pairs not drawn)
- 8. What is the hybridization of each C in benzene (shown below)?
  - $\triangleright sp^2$ 
    - Each carbon has 2H and  $\pi$  bond between, so groups = 3.
    - $\circ$  Groups 1 = 2, so sp<sup>2</sup>
    - $\circ$  Though, each  $\pi$  bond is delocalized, or free to spread across all the carbons. Still counts as 1 group.

## Wednesday, October 7

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## Friday, October 9

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