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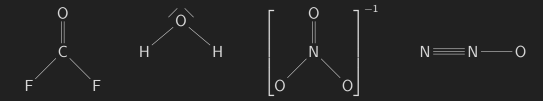
## 1 General Chemistry Review

## **Structural Theory of Matter**

- ▶ **Constitutional isomers**: same molecular formula, but different in the way the atoms are connect, i.e. their constitution is different.
- ▶ Each element forms a predictable number of bonds, from one to four.

#### **Electrons, Bonds, and Lewis Structures**

- ▶ **Covalent bond**: two atoms sharing a pair of electrons.
- ▶ The lowest energy (most stable) state of two atoms is determined both by bond length and bond strength.
- ▶ **Lewis structures**: drawings that show free electrons.
- ▶ Valence electrons are determined by the group, 1A-8A, of the periodic table.
- ▶ **Lone pair**: unshared, or nonbonding, electrons.
- $\triangleright$  Examples: COF<sub>2</sub>, H<sub>2</sub>O, NO<sub>3</sub>, N<sub>2</sub>O:



▶ Resonance structures: a set of two or more Lewis structures that collectively describe the electronic bonding of a single polyatomic species, including fractional bonds.

#### **Identifying Formal Charges**

- ▶ **Formal charge**: any atom that does not exhibit the appropriate number of valance electrons.
- ▶ Determing formal charge:

• Formula: 
$$FC = V - N - \frac{B}{2}$$

- ∘ V = valance electrons of element
- N = lone pair electrons
- B = bonded electrons
- ▶ Less than expected number of valence electrons result in positive charge.
- More than expected results in negative charge.

- The lower the magnitude of formal charge, the greater the stability of the whole molecule.
- Atoms that are more electronegative hold negative formal charges better, which results in greater stability vs when the negative charge is spread on less electronegative elements in a polyatomic species.
  - The dominant resonance structure will be that of the greatest stability.

#### **Induction and Polar Covalent Bonds**

- ▶ Bonds can classified into three categories: covalent, polar covalent, and ionic.
- ▶ The categories emerge from the electronegativity values of the atoms sharing a bond.
- ▶ **Electronegativity**: a measure of the ability of an atom to attract electrons.
- ▶ Electronegativity generally increases left to right, and from the bottom to top of the periodic table.
- ▶ **Covalent bond**: when the difference in electronegativity is less than 0.5.
- ▶ **Polar covalent bond**: when the difference in electronegativity is between 0.5 and 1.7, then the electrons are not equally shared and become polar.
- Induction: the withdrawl of electrons towards to more electronegative atom.  $\delta^+$  represents partial positive charged gained when electrons are pulled away, while  $\delta^-$  represents the partial negative charge pulled closer.
- ▶ **Ionic bond**: when the difference in electronegativity is greater than 1.7.
  - Electrons are not shared in this case, and attraction is insetsad just the result of oppositely charged ions.

#### **Atomic Orbitals**

- ▶ **Atomic orbital (AO)**: s(1), p(3), d(5), f(7).
- $\triangleright$  Locations where  $\psi$  (quantum wave function) is zero are called **nodes**.
- ▶ The more nodes that an orbital has, the greater it's energy.
- ▶ Degenerate orbitals: orbitals with the same energy level.
- > Order in which orbitals are filled is determined by three principles:
  - Aufbau principle: lowest energy orbital is filled first.
  - **Pauli exclusion principle**: each orbital can accommodate a maximum of two electrons that have opposite spin.

- **Hund's rule**: electrons are placed in each degenerate orbital before being paired up.
- ▷ Describing the nature of atomic orbital is done with two commoly used theories: Valence Bond Theory and Molecular Orbital Theory.
- ▶ The commonly used theories give a deeper understanding of covalent bonds, which is essentially just the overlap of atomic orbitals.
- ▶ Constructive/destructive interference: the result of two waves that approach each other, or overlap.
  - o Constructive interference produces a wave with larger amplitude.
  - Destructive interference cancel each other out and produes a node.

## **Valence Bond Theory**

- ▶ **Valence bond theory**: the sharing of electron density between two atoms is a result of the constructive interference of their atomic orbitals.
- ▶ Bond axis: the line that can be drawn between two hydrogen atoms.
- $\triangleright$  **Sigma bond** ( $\sigma$ ): a particular type of covalent bond that has circular symmetry with respect to the bond axis.
  - $\circ$  All single bonds are  $\sigma$  bonds.
  - The strongest type of covalent bond.
- $\triangleright$  **Pi bond (** $\pi$ **)**: covalent bonds where two lobes of an orbital overlap with two lobes of another atom.
  - Each atomic orbital has zero electron density at a shared nodal plane, passing through the two bonded nuclei.
  - $\circ$   $\pi$  bonds from double and triple bonds but generally do not form single bonds.

#### **Molecular Orbital Theory**

- ▶ Molecular orbital theory (MO): uses linear combinations of atomic orbitals to model and explore the consequences of orbital overlap.
  - The newly described orbitals are called molecular orbitals accroding to MO theory.
- Atomic orbitals refer to an individual atom, while molecular orbitals is associated with an entire molecular.

▶ In other words, MO theory states that atomic orbitals cease to exist when they overlap. Instead they are replaced with multiple molecular orbitals which span the entire molecule.

#### **Hybridized Atomic Orbitals**

- $hd problet sp^3$ -**hybridized orbitals**: produced by averaging one s orbital and three p orbitals.
  - Hybridized orbitals explains to geomtry of methane, which results form the now four degenerate orbitals pushing apart to achieve tetrahedral geometry.
  - Hybridized orbitals become unsymmetrical, producing a larger front lobe that is more efficient than standard p orbitals in the ability to form bonds.
  - $\circ$  All bonds in are  $\sigma$  bonds, and thus can be individually represented by the overlap of atomic orbitals.
- $\triangleright sp^2$ -hybridized orbitals: produced by averaging the s orbital with only two of p orbitals.
  - The remaining p orbital is unaffected, and free multiple p orbitals results in a  $\pi$  bond.
  - o This is done to expain geometry of compounds bearing a double bond.
  - $\circ$  A double bond if formed from one  $\sigma$  bond and one  $\pi$  bond.
  - Associated with trigonal planar geometry.
- ▷ sp-hybridized orbitals: produced by averaging of one s orbital and one p orbital.
  - Leaves two p orbitals and resulting in two  $\pi$  bonds.
  - A triple bond is formed with the addition of one  $\sigma$  bond due to the overlap of the sp orbitals.
  - o Geometry of a triple bond has linear geometry.
- ▶ Finding the hybridization of any atom can be done simply:
  - 1. Look at the central item.
  - 2. Determin groups (number of atoms and lone pairs attached) of atom.
  - 3. For groups 1-4:  $sp^x$ ; x = groups 1
  - 4. For groups 5-6:  $sp^3d^x$ ; x = groups 4
- ▶ Bond Strength and Bond Length:
  - Bond length decreases with more bonds.

o Bond strength increases with more bonds.

#### **Molecular Geometry**

- ▶ Valence shell electron pair repulsion (VSEPR) theory: enables the prediction of molecular geometry due to the pressumption that all electron pairs repel each other; resulting in a three-dimensional space that maximizes distance from each other.
- ▶ **Steric number**: the total number of electron pairs in a molecule. Can be bonds or lone pairs.
- $\triangleright$  **Tetrahedral geometry**: result of four  $\sigma$  bonds and zero lone pairs.
  - o produces a tetrahendron with bond angles of 109.5°.
- $\triangleright$  **Trigonal pyramidal geometry**: three  $\sigma$  bonds and one lone pair.
  - The lone pair occupy more space than bonded electron pairs, so the remaining angles are slightly less than a tetrahedral, at 107°.
  - The lone pair sits atop the base forming a pyramid like structure.
- $\triangleright$  **Bent geometry**: two  $\sigma$  bonds and two lone pairs.
  - VSEPR predicts the lone pairs to be in two corners of the tetrahedral, producing bond angles of 105°.
  - VSEPR predicts geometry H<sub>2</sub>O correctly, but for wrong reasons.
    - The lone pairs in  $H_2O$  have different energy levels, suggesting one pair occupies a p orbital with the other in a lower-energy hybridized orbital.
- ▶ VSEPR theory is best used for a first approximation and is mostly accurate for most small molecules.
- ▶ Trigonal planar geometry: three electron pairs forming three bond angles of 120° and lie on the same plan.
- ▶ **Linear geometry**: two electron pairs that oppose each other at 180°, forming a linear structure.
- ▷ General method of determining structure:
  - 1. 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.

## **Dipole Moments and Molecular Polarity**

 $\triangleright$  **Dipole moment** ( $\mu$ ): defined ast the amount of partial charge,  $\delta$ , on on either end of the dipole multiplied by the distance separtion, d:

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\circ \mu = \delta d
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- $\circ$   $\mu$  generally has an order of magnitude of  $10^{-18} \, \text{esu} \cdot \text{cm}$  due to general partial charge (esu) and distance (cm) values.
- $\circ$  1 debye (D) =  $10^{-18}$  esu·cm
- ▶ **Molecular dipole moment**: the vector sum of the individual dipole moments.
  - Lone pairs have significant effect on the molecular dipole moment.
  - Also called the net dipole moment.

## **Intermolecular Forces and Physical Properties**

- ▶ **Intermolecular forces**: the attractive forces between individual molecules that determed the physical properties of a compound.
- ▶ *Electrostatic*: forces that occur as a result of the attraction between opposite charges.
- ▷ Electrostatic interactions for neutral molecules (no formal charge) are often classified as into the following categories:
  - **Dipole-dipole interaction**: Compounds with net dipole moments.
    - In solid space these intereactions either repel or attract each other.
    - In liquid space these interactions tend to attract more often, raising melting/boiling point.

## Hydrogen bonding:

- Not actually a bond, just an interaction.
- When hydrogen bonds to a electronegative atom, then the hydrogen will have a  $\delta^+$ .
- F, O, N, Cl (Br, I). Most electronegative elements, from left to right, that hydrogen most often bonds too.
- Hydrogen bonding is strong due to size of hydrogen atom, resulting in very close partial charge interactions.
- The more hydrogen bonds, the higher the boiling point tends to be.

## Fleeting dipole-dipole interactions:

- Electrons are considered to be in constant motion, which restult in the center of negative charge to vary.
- On average, the dipole moment is zero, though can experience transient dipole moments, initiating fleeting attraction/repulsion.
- Heavier hydrocarbons generally experience a stronger force due to increased surface area, and thus greater chance for non-zero dipole moments, which results in higher boling points.
- Branched hydrocarbons generally have decreased surface area, decreasing boiling point relative to others of similar weight.
- ▶ When comparing boling points of compounds, look for following factors:
  - Any dipole-dipole interactions?
  - Formation of hydrogen bonds?
  - Number of carbon atoms. (surface area)
  - Degree of branching of compound. (surface area)

## **Solubility**

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