CHEM 22000 (Organic Chemistry I) Notes

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Week 1

The Basics: Bonding and Molecular Structure

1.1 Course Information

9/28:

- No labs this week.
- Virtual lab: Watch a video and record data in your notebook; answer embedded quiz questions.
- Collaborative Learning in Organic Chemistry (CLOC).
 - 2hr Sunday or Monday.
 - Contact Dr. Britni Ratliff (ratliff@uchicago.edu).
 - Pass/Fail grading (based on attendance).
 - You work on problems related to the lecture content under the supervision of someone who's taken the class before.
 - You can opt-in/out on a quarter-by-quarter basis.
- Review syllabus: Download alternate textbooks, put exam dates in the calendar, add office hours to calendar.
- Develop an understanding of how structure affects reactivity mechanistic principles.
- You don't have to memorize anything, but you have to remember everything.
 - Like learning a language.
 - Vocabulary, grammar (principles), apply to understand and predict.

1.2 Defining Organic Chemistry

- Organic chemistry: Traditionally, the chemistry of living organisms. Now, the chemistry of carbon compounds.
 - Carbon is of particular import because it can bond with itself, and it can form strong bonds with other elements (e.g., C, O, H, S, N, and P) as well.
 - Carbon is bound in simple molecules (such as CO₂ and CH₄), and highly complex ones (such as proteins, DNA, and RNA).
- Carbon compounds:

- Natural: Sugars, fats, gasoline, hydrocarbons, hormones, natural drugs, peptides, rubber, silk, starch, cotton, etc.
- Synthetic: Dyes, fragrances, soaps, drugs, medicines, plastics, materials, teflon, nylon, etc.
- OChem is a central science that feeds into fields such as biochemistry, molecular biology, molecular medicine, math/theory (e.g., buckyballs), engineering, and physics.

1.3 Gen Chem Review

- Today:
 - 1. Intro (done).
 - 2. Atomic structure and bonding (review from Gen Chem).
 - 3. Chemical bonds octet rule.
 - 4. Writing Lewis structures.
 - 5. Formal charges.
- Atomic structure and bonding.
 - Atoms \rightarrow elements \rightarrow compounds.
 - Nucleus (protons and neutrons) surrounded by electrons.
 - This year, we'll concern ourselves with the main group elements.
 - Electron configuration:
 - Aufbau principle: Electrons fill orbitals from lowest energy to highest energy.
 - Pauli exclusion principle: 2 elections/orbital with opposite spin quantum numbers (must pair $+\frac{1}{2}$ with $-\frac{1}{2}$).
 - Hund's rule: Orbitals with equivalent energy get partially filled first before more electrons are added.
 - Example: $1s^22s^22p^63s^1$ is Na.
 - Valence electrons are key in this class.
- Noble gas configurations and the octet rule.
 - Lewis noticed that there is a special stability associated with a filled outer shell.
 - Thus, we generally have 8 electrons in the filled outer shell.
 - For example, $Cl \xrightarrow{1 e^-} Cl^-$ and $Na \xrightarrow[-1 e^-]{} Na^+$.
 - Chemical bonds form because they allow the atoms to achieve a filled octet.
 - Two kinds of bonding: Ionic and covalent.
 - Ionic: Not covered much this year. Lose or gain an electron (forming cations and anions, respectively) to for a filled outer shell. Usually involves a metal and a nonmetal.
 - Covalent: Covered a lot this year. Sharing electrons to satisfy the need for an octet.
 - The atoms involved dictate whether bonding will be ionic or covalent.
 - Electronegativity: The ability of an atom to attract its valence shell electrons.
 - Defined by Pauling, who let Li = 1.0 and F = 4.0.
 - This is a very important concept for understanding bonding and reactivity.
 - EN increases across and up on the periodic table: More protons and a shorter distance away from the nucleus both mean a greater pull on the electrons.
 - Mnemonic (highest to lowest electronegativity): F O Cl N Br I S C H P.

- Non-polar covalent bonds form when $\Delta EN < 0.5$.
- Polar covalent bonds form when $\Delta EN \approx 0.5 1.9$.
- Exceptions to the octet rule: H wants 2 e[−]. Be wants 4 e[−]. B and Al want 6 e[−]. Molecule has an odd number of electrons (e.g., NO with 11 electrons is stable).
- Lewis structures.
 - General rules/procedure (there are exceptions).
 - 1. Determine the total number of valence electrons for the molecule. Add electrons for negative charges; remove for positive charges.
 - 2. Draw a skeleton and join atoms with single bonds. Put the atom that likes to make the most bonds in the center.
 - 3. Deduct 2 electrons from the count in step 1 for each single bond. Fill outside atoms with lone pair electrons.
 - 4. The remaining electrons go on the central atom.
 - 5. If you have too few electrons for every atom to have an octet, use lone pair electrons to convert single bonds to double bonds. We can also use triple bonds.
 - CH₄ and NH₃ presented as worked examples.
- 9/30: Today:
 - 4. Lewis Structures.
 - 5. Formal charges.
 - 6. Isomers.
 - 7. Structural formulas.
 - 8. Resonance.
 - 9. Orbitals and bonding.
 - Lewis structures:
 - H₂CO (formaldehyde) and CH₃COOH (acetic acid) presented as worked examples.
 - Formal charge determination:
 - If the number of valence electrons does not equal the total number of electrons on an atom, then
 you will have a formal charge.
 - Rule:

Formal Charge = normal valence
$$e^-$$
 - actual e^-
= valence e^- - $\left(\text{nonbonding } e^- + \frac{1}{2} \text{bonding } e^- \right)$
= valence e^- - $\left(\text{dots} + \text{lines} \right)$

- CH_3COO^- (acetate) has a formal charge of 6-7=-1 on its singly bonded oxygen.
- $CH_3NH_3^+$ (methyl ammonium) has a formal charge of 5-4=+1 on its nitrogen.
- Exceptions: Open shell Group III central atoms (e.g., B and Al).
 - BF₃ acts as a Lewis acid because it wants to grab $2e^-$ to form an octet.
 - It often acts in acid-base coupling reactions, grabbing a lone pair from an oxygen in an adjacent molecule and bonding through it.

1.4 OChem Basics

- Isomers:
 - Constitutional isomers: Same molecular formula but different bond connectivities.
 - Acetone vs. 3-propenol, yet both are C_3H_6O .
- Structural formulas:

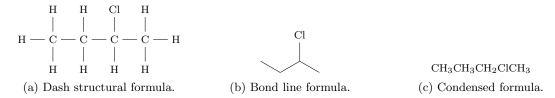


Figure 1.1: Structural formulas.

- Dash structural formula: A Lewis structure.
- Bond line formula: No C-H's, show a vertex for each carbon, show heteroatoms and heteroatom H's. Also known as line-angle structure, zig-zag structure.
- Condensed formula: All atoms written out with no bonds or lone pairs.
- 3D representation: A dash structural/bond line formula with wedges and dashes.
- **Resonance**: When a molecule or an ion can be represented by 2 or more Lewis structures, i.e., two or more structures with the same skeleton connected by different electrons.
 - Resonance structures or resonance contributors.
 - The actual molecule is somewhere between the contributors.
 - CO_3^{2-} presented as a worked example.
 - Guidelines:
 - 1. Only lone pairs or π electrons move (never move single bonds).
 - 2. No structure with greater than $8e^-$ on a 2nd row atom.
 - 3. The species with the maximum number of octets is the strongest contributor.
 - 4. Charge on suitable atoms (e.g., negative charge on the atom with the highest electronegativity).
 - Resonance stabilization comes from delocalization. When 2 or more resonance structures, the "real" structure is somewhere in between (the real is more stable than any contributor).
 - CH₃COO⁻ (acetate), CH₂CHCH₂⁺, and (CH₃)₂CO presented as worked examples.
 - You can also depict delocalization with a curving dashed bond and δ^{-} 's.

1.5 Bonding and Orbital Diagrams

- 10/5: Today:
 - 9. Orbital theory and bonding.
 - 10. Methane.
 - 11. Ethane.
 - 12. Ethylene.

- 13. Acetylene.
- 14. Comparison of sp^3 , sp^2 , sp orbitals.
- 15. VSEPR Model + Molecular Symmetry.
- Orbital theory and bonding:
 - Defines atomic orbitals.
 - Reviews s and p orbital shapes, positive and negative regions, and nodes.
 - Energy of orbitals diagram.
 - Phosphorous and sulfur can exceed the octet rule since they have d orbitals in which to stash extra electrons.
 - Filled with the Aufbau/Pauli Exclusion principles, and Hund's Rule.
 - Goes over bonding energy diagram.
 - Mathematically, we have a Linear Combination of Atomic Orbitals (or LCAO).
 - \blacksquare Electrons are represented as waves; thus, they have + and phases.
 - \blacksquare Opposite phases are destructive; this forms σ^* orbitals.
 - Same phases are constructive; this forms σ orbitals.
 - Goes over MO diagrams.
- Atomic orbital: A space where electrons are likely to be found 95% of the time.
- Degenerate (orbitals): Two orbitals with the same energy.
- Chemical bond: A favorable interaction between 2 atoms, i.e., one that helps to fill the outer orbitals to achieve a noble gas configuration.
- Bonding in methane:

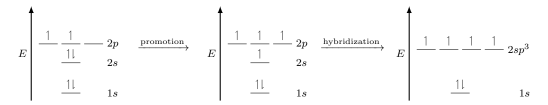


Figure 1.2: Bonding in methane.

- Draws an orbital diagram for carbon.
- Promotes an electron from $2s \rightarrow 2p_z$.
- Hybridizes $2s, 2p_x, 2p_y, 2p_z$ into 4 degenerate sp^3 orbitals of weighted average energy, each containing only 1 electron.
- Links each of these sp^3 electrons to the 1s electron in H_2 , forming σ orbitals.
- The new orbitals adopt a tetrahedral arrangement to be as far apart as possible.
- Bonding in ethane.
 - Two sp^3 electrons combine in a σ orbital; no electrons go into the σ^* MO.
- The structure of ethylene.
 - Side by side overlap of p orbitals forms a π bond.
 - The angle between the hydrogens in ethylene is slightly less than 120°.

- The bond is slightly shorter than in ethane (greater s character plus an additional type of bond).
- Features of the C=C double bond.
 - sp^2 -hybridized carbons making 3σ and 1π bond.
 - A π bond is weaker than a σ bond, but still strong.
 - \bullet $\sigma_{sp^2-sp^2}$ is stronger than $\sigma_{sp^3-sp^3}$.
 - Restricted rotation (hard to twist C_2H_2 by 90°).
 - cis-trans isomerism as a result of restricted rotation.
 - The π bond acts like a Lewis base with some systems since the π electrons are held relatively weakly. In other words, the π -electrons are exposed.
- Draws an MO diagram for the carbons.
- The structure of acetylene.
 - -2π bonds, 1σ bond.
 - Even greater strength, but not quite as much greater as the $\sigma_{sp^3-sp^3} \to \sigma_{sp^2-sp^2}$ difference.

1.6 VSEPR Theory

- 10/7: There's a special kind of electronegativity that relates to hybridization: An sp-hybridized carbon is more electronegative than an sp^3 -hybridized carbon, for instance.
 - VSEPR Model:
 - Electron pairs want to stay as far apart as possible in space.
 - Consider the bonding electrons (number of atoms bound) and nonbonding electrons.
 - Describe shape based on the position of nuclei.
 - Constructs VSEPR table for linear, trigonal planar, tetrahedral, trigonal pyramidal, bent

Week 2

Families of Carbon Compounds / Acids and Bases

2.1 Families of Carbon Compounds

- 10/7: Hydrocarbons:
 - Alkanes (C_nH_{2n+2}) and cycloalkanes C_nH_{2n} .
 - Alkenes (C_nH_{2n}) .
 - Alkynes (C_nH_{2n-2}) .
 - Aromatic:
 - Contains a benzene ring.
 - All bonds $\sim 140 \,\text{Å}$.
 - All carbons sp^2 .
 - Planar.
 - $-\pi$ electrons above and below the ring.
 - Special stabilization.
 - Covers drawing dipoles.
 - Polar and nonpolar molecules:
 - Dipole = distance \times change between charges.
 - $-\mu = r \times Q$
 - $-1D = 3.336 \times 10^{-30} \,\mathrm{Cm}.$
 - Analyzes molecules by drawing a Lewis structure, drawing a dipole along each bond, and drawing and labeling a net dipole, if applicable.
 - Goes through a number of examples.
 - Acetonitrile is a strong polar solvent.
 - Functional group: A common arrangement that determines shape, bonding physical and reactivity of organic compounds.
 - Families of carbon compounds:
 - Hydrocarbons: Aliphatic, aromatic.
 - Methyl, ethyl, propyl, R = alkyl groups.

- Phenyl: Ph- or ϕ -.
- Benzyl: $Ph-CH_2$ -, $C_6H_5CH_2$ -, Bn-
- Compounds with R-Z where Z is a heteroatom.
 - If Z is a halogen X, then the halogroup makes it an alkyl halide or haloalkane.
- Alkenyl halide: X-=.
- Aryl halide: Ph-X.
- Alcohols or phenols: R-OH.
- Ether: R-O-R'.
- Amines: NH₂R, NHRR', NRR'R".
- Thiols or mercathols: R-SH.
- Carbonyl group: R-CO-R'.
- Aldehyde: R-COH.
- Ketone: R-CO-R'.
- Carboxylic acid derivatives:
 - Acid: R-COOH.
 - \blacksquare Ester: R-COOR'.
 - Acid chloride: R-COCl.
 - Acid halide: R-COX.
 - Amide: $R-CONH_2$.
 - Acid anhydride: R-COOCO-R'.
- Nitrile: $R-C \equiv N$.
- Acrylonitrile: $=-C\equiv N$.

2.2 Discussion Section

- ACS in-text citations should be in superscripts as a list of number with no brackets or parentheses.
- Molecular formulas are C₂H₆O, not C₂H₅OH or CH₃CH₂OH.
- Make a table if you have a lot of data to put in (make it readable!).
- Distillation:
 - We need a boiling chip and stir bar inside the flask.
 - Vapor comes up from a round-bottomed flask, encounters a rubber stopper and gets diverted through a condenser instead.
 - Make use of countercurrent exchange and increase pressure by inflowing water in the gravitationally lower portion of the condenser.
 - Boiling chip is a coarse material with a lof of micropores inside.
 - The surface energy is reduced when the fluid is inside the micropores; within, it can more easily become a gas.
- As the mole fraction χ of a substance A increases...
- Raoult's law:

$$P_{\text{total}} = \frac{P_A \chi_A}{P_B \chi_B} = \frac{P_A \chi_A}{P_B (1 - \chi_A)}$$

• Dalton's law: The total pressure is equal to the sum of the partial pressures.

2.3 Intermolecular Forces and IR Spectroscopy

- 10/12: Intermolecular forces and physical properties.
 - **Boiling point**: The temperature at which the vapor pressure is equal to the pressure of the atmosphere above.
 - The stronger the intermolecular forces, the higher the boiling point.
 - The higher the molecular weight, the higher the boiling point.
 - Melting point: The temperature at which the crystalline solid and liquid are in equilibrium.
 - The stronger the intermolecular forces, the higher the melting point.
 - The more symmetrical, the higher the melting point.
 - Solubility.
 - Intermolecular forces.
 - All electrostatic attractions related to bond polarity.
 - 3 types: Dipole-dipole forces, hydrogen bonding, and dispersion forces.
 - Dipoloe-dipole forces: Attraction between opposite poles (1 kcal/mol to 3 kcal/mol).
 - Hydrogen bonding: Dipole-dipole interaction between H-atoms bonded to O, N, F (2 kcal/mol to 10 kcal/mol).
 - **Dispersion forces**: Weak (< 1 kcal/mol). Momentary distortion of the electron cloud (temporary dipole). Induces dipoles in surrounding molecules. *Also known as* **London forces**.
 - Depends on **relative polarizability**.
 - Dependes on the surface area of the molecule more surface area means more distance electrons can spread apart.
 - Relative polarizability: How far valence electrons are from the nucleus.
 - Solubility:
 - For something to be soluble, you need to have favorable forces between them.
 - Ionic compounds are soluble in water, less soluble in polar solvents, and insoluble in nonpolar solvents.
 - Organic compounds: < 3 carbons is soluble, 4-5 carbons is borderline, ≥ 6 is insoluble. More soluble in organic solvents.
 - Organic solvents:
 - CH₂Cl₂ methylene chloride.
 - HCCl₃ chloroform.
 - H₃CCOCH₃ acetone.
 - Diethyl ether.
 - THF.
 - Cyclohexane.
 - In TLC, the silica gel is very polar, so polar compounds will not move far up the plate. Nonpolar solvents will drag nonpolar compounds up pretty high.
 - HOMO and LUMO get closer as conjugation increases.
 - IR spectroscopy:

- The frequencies absorbed vary based on the type. Higher stretching frequencies for lighter atoms and stronger bonds.
- IR radiation causes transitions in vibrational modes of bonds.
- The stronger the bond and the lighter the atoms, the faster the vibration of the molecule and the higher the stretching frequency.
- The ΔE 's are inherent characteristics of the bonds and nuclei.
- Bonds absorb light of characteristic energy, frequency, and wavelength.
- We usually report IR spectra in terms of the wavenumber $\bar{\nu}$.
- The frequencies absorbed can indicate bond types and functional groups in the molecule.
- Anything above 1500 (of wavenumber less than 1500) is called the **fingerprint region** it may not tell you what a molecule is, but it will tell you if two molecules are the same.
- Sharp peaks at high wavenumbers are characteristic of N-H interactions.
- Make a line at $3000\,\mathrm{cm}^{-1}$. Things to the right of that indicate aliphatic C-H's. Things to the left indicate sp^2 C-H groups. Things more to the left indicate sp C-H groups.
- Not all bonds are visible stretching bands must change the dipole. Thus, for example, the C=H stretch in trans-but-2-ene is not IR active, but the C=H stretch in cis-but-2-ene is IR active.
- If you want to substitute D for H, the peak formerly associated with the R-H bond will move lower.

COMMON ABSORPTIONS											
Aromatic C-C	Two peaks usually in the range of $1500\mathrm{cm}^{-1}$ to $1600\mathrm{cm}^{-1}$										
C = C	$\sim 1650 {\rm cm}^{-1}$										
C = O	$\sim 1710\mathrm{cm^{-1}}$ (shifts to $\sim 1735\mathrm{cm^{-1}}$ for esters)										
C = C	$2100\mathrm{cm^{-1}}\ \mathrm{to}\ 2300\mathrm{cm^{-1}}$										
C=N	$2100\mathrm{cm^{-1}}\ \mathrm{to}\ 2300\mathrm{cm^{-1}}$										
C-H (aldehyde)	Two peaks at $2170\mathrm{cm}^{-1}$ and $2810\mathrm{cm}^{-1}$										
sp^3 C-H	Just to the right of $3000\mathrm{cm}^{-1}$										
sp^2 C-H	Just to the left of $3000\mathrm{cm}^{-1}$										
sp C $-$ H	$\sim 3300{\rm cm}^{-1}$										
N-H	$\sim 3300\mathrm{cm}^{-1}$ (one peak for $-\mathrm{NH}$, two peaks for $-\mathrm{NH}_2$)										
O-H (alcohol)	$\sim 3400\mathrm{cm}^{-1}$ (a broad, smooth peak)										
O-H (acid)	$\sim 2500\mathrm{cm^{-1}}$ to $3500\mathrm{cm^{-1}}$ (a very broad, ugly [not smooth] peak)										

Table 2.1: Common IR spectroscopy absorptions.