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1 General Chemistry Review

Electrons, Bonds, and Lewis Structures

- ▶ **Covalent bond**: two atoms sharing a pair of electrons.
- ▶ **Octet rule**: main group elements that tend to bond in a way that each atom has eight electrons in it's valence shell.
 - Atoms that do not have eight will share electrons with other elements in order to maintain a stable state.
- ▶ **Main group elements**: sometimes called representative elements, are groups 1, 2 and 13–18 in periodic table.
 - Some elements in group 3 and 12 share properties between transition metals and the main group.
- ▶ The lowest energy (most stable) state of two atoms is determined both by bond length and bond strength.
- ⊳ Valence electrons are determined by the group, 1A–8A, of the periodic table.
- ▶ **Lone pair**: unshared, or nonbonding, electrons.
- ▶ **Lewis structures**: 2D model that represents covalent bonds as straight lines and lonpairs as dots.
- ⊳ Examples: COF₂, H₂O, NO₃⁻, N₂O:



▶ **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 results in a positive charge.
- More than expected number of valence electrons results in a 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.
 - **F, O, N, CI** (Br, I). Most electronegative elements, from left to right, that are often encountered.
- ▶ **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 1.9, then the electrons are not equally shared and become polar.
- ▶ **Induction**: the withdrawl of electrons towards to more electronegative atom. δ^+ represents partial positive charged gained when electrons are pulled away, while δ^- represents the partial negative charge pulled closer.
- ▶ **lonic bond**: when the difference in electronegativity is greater than 1.9.
 - Electrons are not shared in this case, and attraction is insetsad just the result of oppositely charged ions.

Dipole Moments and Molecular Polarity

- o **Dipole moment** (μ): defined as the amount of partial charge, δ , on on either end of the dipole multiplied by the distance separtion, d:
 - $-\mu = \delta d$
 - μ generally has an order of magnitude of 10^{-18} esu·cm due to general partial charge (esu) and distance (cm) values.
 - $1 \text{ debye (D)} = 10^{-18} \text{ esu} \cdot \text{cm}$
- **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.

Atomic Orbitals

- ▶ **Atomic orbital (AO)**: standing quantum wave (excitation in electron field) around an atom.
 - More energy leads to higher orbtails levels.
 - Gives principle quantum number, n, as is associated with distance from nucleus.
 - o Orbital levels: s(1 pair), p(3 pairs), d(5 pairs), f(7 pairs).
 - Angular momentum quantum number that describes three-dimensional region of space that the electron density occupies.
 - Magnetic quantum number descrices orientation in space of electron density.
 - $-m_l=0$; s orbital
 - $-m_{I}=-1, 0, 1; p_{x}, p_{y}, p_{z}$ orbitals.
 - \circ Locations where ψ (quantum wave function) is zero are called **nodes**.
 - The more nodes that an orbital has, the greater it's energy.
 - \circ Spin: allows an orbital to contain only two electrons, $\pm \frac{1}{2}$
- 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.
 - Constructive interference produces a wave with the vector sum of both waves.
 - 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.
- o Bond axis: the line that can be drawn between two hydrogen atoms.
- **Sigma bond (\sigma)**: a particular type of covalent bond that has circular symmetry with respect to the bond axis.
 - All single bonds are σ bonds.
 - The strongest type of covalent bond.
- **Pi bond** (π) : 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.
 - π bonds form double $(\sigma + \pi)$ and triple bonds $(\pi + \sigma + \pi)$.
 - Individual π bonds are weaker than σ 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.

- o 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.
- Molecular orbitals are more stable (lower energy) since electrons are attracted by both nuclei.
- \circ When there are nodes between the nuclei, then the resulting σ^* orbitals become antibonding, as they destabilize (increase the energy) of a molecular orbital.
- o Best used to produce a quantitative picture of bonding.
 - Describes strength, order, and polarity of bonds.
 - Allows for the presence of paired or unpaired electrons.
 - Has spectroscopic preperties.

Hybridized Atomic Orbitals

- **sp³-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.
 - All bonds in are σ bonds, and thus can be individually represented by the overlap of atomic orbitals.
- 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 π bond.
 - This is done to expain geometry of compounds bearing a double bond.
 - A double bond if formed from one σ bond and one π 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 π bonds.

- A triple bond is formed with the addition of one σ bond due to the overlap of the sp orbitals.
- Geometry of a triple bond has linear geometry.
- o Finding the hybridization of any atom can be done simply:
 - 1. Look at the central item.
 - 2. Determin groups (number of bonds, π bonds count as 1, and lone pairs attached) of atom.
 - · groups aka regions of electron density.
 - 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.
 - Bond strength increases with more bonds.
 - The more s character, the shorter and stronger the bond, and the larger the bond angle.
 - · s-character: contribution of the σ bond in a hybridization.
 - e.g. sp = 50%, $sp^2 = 33\%$, $sp^3 = 25\%$
 - · sp-sp bond is the strongest, sp³-sp³ is the weakest.

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 σ bonds and zero lone pairs.
 - o produces a tetrahendron with bond angles of 109.5°.
- \triangleright **Trigonal pyramidal geometry**: three σ 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 σ 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 H₂O 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.

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.
 - Ion-dipole: electrostatic interaction between an ion and a molecule with a dipole.
 - **Hydrogen bonding**: molecules with a hydrogen attached to an F, O, or N.
 - Not actually a bond, just an interaction.
 - When hydrogen bonds to a electronegative atom, then the hydrogen will have a δ^+ .

- 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.
- Stronger than dipole-dipole interactions.

• Fleeting dipole-dipole interactions:

- Electrons are considered to be in constant motion, which restult in the center of negative charge to vary.
- London Dispersion Forces (LDFs): On average, the dipole moment is zero, though it can experience transient dipole moments, initiating fleeting attraction/repulsion.
 - · All atoms and molecules have LDFs.
 - · Weakest, but the dominant force in non-polar molecules.
 - · Dispersion forces directly related to molar mass.
- 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? (increases boiling point)
 - Formation of hydrogen bonds? (increase boling point)
 - Number of electrons. (more electrons, higher boiling point)
 - Number of carbon atoms. (more surface area, higher boiling point)
 - Degree of branching of compound. (more branching, more surface area)

2 Molecular Representations

Types of Molecular Representations

▶ Partially condensed structures: the C−H bonds are not always drawn, saving space.

- ▶ Condensed structures: single bonds are not drawn and groups of atoms are clustered when possbile.
 - \circ CH₃CH₃CHOH \longrightarrow (CH₃)₂CHOH
- ▶ **Molecular formula**: simply shows number of each type of atom with no structural information.
 - o C₃H₈O
- ▶ Example of converting a condensed structure into a partially condensed structure:
 - (CH₃)₃CCH₂CH(CH₃)CH(CH₃)₂
 CH₃ H H H H
 CH₃ − C − C − C − C − C − CH₃
 CH₃ H CH₃ CH₃ CH₃
 - This shows just one isomer, more partially condensed structures are possible.

Bond-Line Structures

- **Bond-line structures**; aka skeletal structures; simplify drawing process of chemical structures and are easier to read.
- Each corner or endpoint represents a carbon atom.

- All examples have 6 carbon atoms
- Double bonds are shown with two lines, triple with three.
 - / | //
 - Triple bonds are drawn linearly due to sp-hybridization

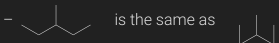
• Hydrogens are not shown; it is assumed that each carbon posses enough to satisfy octet rule.

Notes on Drawing Bond-Line Structrues

- Carbon atoms in a straight chain should be drawn in zigzag format in order to accurately show each carbon.
- Double bonds should be drawn as far apart as possible:

$$-$$
 = good \Rightarrow = bad

o Direction of a single bond is irrelevant:



- All heteroatoms (atoms other than carbon and hydrogen) must be drawn.
 - Hydrogens next to heteroatoms must be shown.
- Carbons cannot have more than four bonds.

Hydrogen Deficiency Index: Degrees of Unsaturation

Excerpt from Chapter 14: Infrared Spectroscopy and Mass Spectrometry

- ▶ **Saturated compounds**: the maximum number of hydrogen atoms possbile, relative to number of carbon present.
 - o Determining saturation using molecular formula: C_nH_{2n+2} n= carbon atoms
 - Halogens: takes the place of a hydrogen atom; add one H for each halogen.
 - Oxygen: no affect on saturation; ignore.
 - Nitrogen: needs an extra hydrogen; subtract one H for each nitrogen.
- \triangleright **Unsaturated compounds**: a compound that contains at least one π bond, resulting fewer than the maximum number of hydrogen atoms.
 - Compounds with rings also result in an unsaturated compound.
 - Degree of unsaturation: a number that represents half the "missing"
 number of hydrogen atoms when compared to a saturated compound.
- ▶ **Hydrogen deficiency index (HDI)**: the measure of degrees of unsaturation.
 - o e.g. two degrees of unsaturation results in a HDI of 2.

- Degrees of freedom help represent possbile structures, indicating possible double bounds, triple bounds, rings, or various combinations of each.
- o Only helpful when molecular formula is known for certainty.
- Formula: HDI = $\frac{1}{2}(2C + 2 + N H X)$
 - X: halogen atoms.

Identifying Functional Groups

- ▶ **Functional group (R)**: specific substituents or moieties within molecules that may be responsible for the characteristic chemical reactions.
 - Substituents: an atom or group of atoms which replaces one or more hydrogen atoms on the parent hydrocarbon chain.
 - **Moiety**: a part of a molecule which is typically found within other molecules and often given a specific name.

Characterizing Carbon Centers and Functional Groups

- Characterizing Carbon Centers
 - Primary 1°: a carbon with only one carbon-carbon bond.

- Secondary 2°: a carbon with two carbon-carbon bonds.

- Tertiary 3°: a carbon with 3 carbon-carbon bonds.

$$(CH)_3$$
-CH 3°

- Quaternary 4°: a carbon with four carbon-carbon bonds.

Characterizing Functional Groups

 Certain functional groups can be characters as 1°, 2°, or 3°, based on how many carbon bonds are attached to the carbon with the functional group.

Identifying Lone Pairs

- ▶ Formal charges must always be drawn on bond line structures, otherwise the resulting bond line structures would be inferred incorrectly.
- ▶ Lone pairs do not have to be drawn and usually are omitted.
- ▶ The formal charge allows you to determin lone pairs.

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

- V = valance electrons of element
- N = lone pair electrons
- B = bonded electrons
- Solve for lone pairs: $N = V FC \frac{B}{2}$
- ▶ Frequent usage will allow for intuition for lone pairs.

Common Patterns Between Formal Charge and Lone Pairs

- Associated Patterns for Oxygen
 - A negative (⊙) charge corresponds with 1 bond and 3 lone pairs.
 - The absence of charge corresponds with 2 bonds and 2 lone pairs.
 - A positive (⊕) charge corresponds with 3 bonds and 1 lone pair.

Associated Patterns for Nitrogen

- A negative charge corresponds with 2 bonds and 2 lone pairs.
- The absence of charge corresponds with 3 bonds and 1 lone pair.
- A positive charge corresponds with 4 bonds and 0 lone pairs.

Resonance

- ▶ **Resonance**: description of bonding in molecules or ions by the combination of multiple contributing streutres.
 - **Resonance structures**: each contributing structure of the resonance hybrid.
 - Formal charges are important to include when drawing resonance structures as it clarifies where locations of lone pairs and movement of electrons.
 - Total charge must remain the same between structures.
- ▶ Resonance does not describe any real process, rather it's a method to overcome inadequacy of bond-line drawings.
- Different from isomerism, which differs in arrangements of atomic nuclei in space, rather than how the electrons are assigned to the depictions.

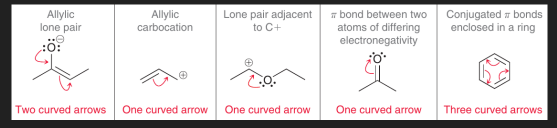
Resonance: Curved Arrows

- Curved arrows: a tool used to help draw resonance structres by representing electrons as if they were moving.
 - Somwhat different from curved arrow notation in reactions, which actually represent the flow of electron density.
 - Can help shows how to change the formal charge:
 - · Formal charges at the tail become more positive, since it's losing an electron.
 - · Formal charges at the head more negative, since it's gaining an electron.
- Avoid breaking a single bond.

- Structures must have atoms connected in same order, though there are minor exceptions that will be discussed later.
- This rule affects the placement of the tail of the arrow, as it represents distribution of previous electrons.
- Never exceed an octet for second-row elements.
 - Not a violation to have less than an octet.
 - This rule affects the placement of the head of the arrow, as it represents sharing of new electrons.
- Can only be used on adjcent atoms, though the electrons can be pushed multiple times.
- "Legal" moves:
 - $-\pi$ bond \rightarrow lone pair.
 - Lone pair $\rightarrow \pi$ bond.
 - $-\pi$ bond $\to \pi$ bond.
 - Every resonance structure can be built through a combination of above three moves.

Common Patterns of Resonance Structures

- Vinylic: the two carbon atoms bearing the double bond of a carbon-carbon double bond.
- Allylic: atoms connected directly to vinylic positions.



Resonance Hybrid

- Resonance hybrid: respresents the average of the contributing structures, with bond lengths and partial charges taking on intermediate values.
- No matter how many resonance structures are drawn, they collectively represent one entity.
- Drawn partial bonds and charges to illustrate the delocalization of electrons.

Delocalization

- Delocalization: the spreading of electrons between multiple atoms or covalent bonds.
 - Resonance stabilization: molecules and ions that are stabalized by the delocalization of electrons.
 - Plays a major role in the outcome of many reactions.
- When a lone pair participates in resonance, it will occupy a *p* orbtail rather than hybridized; important for 3d shapes of proteins.
- **Localized lone pair**: when a lone pair is not allylic to a π bond.
 - Whenever an atom posses both a π bond and a lone pair, they will not both participate in resonance.
 - Usually π bonds participate first.

Contributor Significance

- Some resonance structures may resemble the actual molecule more than another, in regards to energy and stability.
- Strcures with low potential energy are more stable compared to those of higher values and resemble the actual structure more.
- **Major contributors**: the most stable contributing structures.
- Minor contributors: less favorable contributing strcutres.
- Rules for contributing significance, descending:
 - The greatest number of filled octets.
 - The greatest number of covalent bonds.
 - Minimize formally charged atoms.
 - Separation of unlike and like charges, minimized and maximized respectively.
 - Negative charges placed on the most electronegativity atoms, positive charges placed on the less electronegative atoms.
 - Do not deviate substantially from idealized bond lengths and angles.
 - Maintain aromatic substructures locally while avoiding anti-aromatic ones.

3 Acids and Bases

Bønsted-Lowry Acids and Bases

- ▶ Acid: a proton donor; i.e., a H⁺ donor.
- ▶ **Base**: a proton acceptor; i.e., a OH^- (hydroxide ion), which wants a H^+ to form the more stable H_2O .
- - The strength of the acid/base is inversley proportional to the strength of the conjugate acid/base.
- ▶ Most acid-base reactions are reversible.
 - o Strong acids tend to be less reversible.
- ▷ Example using bond-line structures:



Quantitative Perspective

 Equilibrium: when there is no longer an observable change in concentrations of reactants and products.

$$- K_{eq} = \frac{[H_3O^{\dagger}][A^{-}]}{[HA][H_2O]}$$

– Water concentration is fairly constant and can be removed, giving K_a .

$$\cdot K_a = K_{eq} [H_2O] = \frac{[H_3O^{\dagger}][A^{-}]}{[HA]}$$

- K_a tends to be large, so it's converted to pK_a .

$$pK_a = -\log K_a$$

- · Generally ranges from -10 (strong acid) to 50 (strong base).
- $-pK_a(H^+)$ can be easily converted to $pK_b(OH^-)$:

$$pK_b = 14 - pK_a$$

- \circ Equilibrium favors formation of the weaker (higher pK_a) acid.
 - Reactions with vastly different pK_a values make the reverse process is negligible.
 - Can ignore the reverse reaction in such cases and treat it as a reaction in one direction.

Qualitiative Perspective

- Relative acid strength can be determind by comparing conjugate bases.
 - The more stable (weaker) the conjugate base, the stronger the acid.
 - Does not predict pK_a , just a means of comparing relative acid strenghts with out known pK_a
- **Stabilization factors**: (1) atom bearing the charge, (2) resonance, (3) induction, and (4) orbitals.
 - Generally follow decending order of significance; absence of difference in earlir factors allow for later factors to express more significance.
- **Atom bearing the charge**: Compare atoms bearing negative charge in each conjugate base after deprotonation.
 - First determin if atoms are in same row or column in the periodic table.
 - Row comparison: electronegativity is the dominant effect; stability is greater when the negative charge is on the more electronegative element.
 - Column comparsion: size is the dominant effect; stability is greater when the negative charge is on the larger element.
- **Resonance**: charge that is delocalized across multiple atoms will lead to more stable structures comapred to molecules with no resonance.
 - Helps determing relative stability when both molecules bare the same elements that have a difference in charge.
 - Again, more stability means it's the weaker conjugate base, meaning the proton removed from the atom creating the resonance hybrid will be more acidic.
- Induction: induction of other atoms can act to withdraw the negative charge away from the new electronegatively charged atom due to deprotonation.
 - Inductive effect diminishes the further the electronegative atom is away from the depronated atom.
- **Orbitals**: negative charges on atoms with lower hybridization result in greater stability due to proximity to positive nucleus, i.e., $sp > sp^2 > sp^3$
 - sp = triple bond, sp² = double bond, sp³ = three σ bonds.

Lewis Acids and Bases

- ▶ The lewis definition is more broad than the Brønsted-Lowry definition.
- ▶ Lewis describes acidity in terms of electrons, rather than protons.
- ▶ Lewis acid: electron-pair acceptor.
- ▶ Lewis base: electron-pair donor.
- ▷ All Bønsted-Lowry acids and bases are Lewis acid and bases, but the inverse is not always true.
- ▶ Most reactions are described in terms of lewis base and acids, since molecules without donatable protons are unable to be described by the Brønsted-Lowry definition.

Nucleophiles and Electrophiles

Excerpt from Chapter 6: Chemical Reactivity and Mechanisms \mapsto

Mechanisms and Arrow Pushing may also be useful.

- ▶ **Ionic reactions**, aka polar reactions: reactions that involve the participation of ions as reactants, intermediates, or products.
 - Most cases ions act as intermediates.
 - Radical reactions and pericyclic reactions are also major categories, but are typically not discussed in undergraduate courses.
 - lonic reactions occur when one reactant has a site of high electron density and the other reactant has a site of low electron density.
- ▶ **Nucleophiles**: an electron rich atom that is capable of donating a pair of electrons.
 - Lewis bases are nucleophiles, but not all nucleophiles are lewis bases.
 - Any atom that possesses a localized lone pair can be nucleophilic.
 - \circ π bonds can also function as nucleophiles due to their region of space having high electron density.
- ▶ **Electrophiles**: an electron-deficient atom that is capable of accepting a pair of electrons.
 - Lewis acids are electrophiles, but not all electrophiles are lewis acids.

Flow of Electron Density: Curved-Arrow Notation

- ▶ All reactions are accomplished via a flow of electron density.
- ▷ Electron density flow is illustrated with curved arrows.
 - Reaction mechanism: how the reaction occurs in terms of the motion
 - All ionic meachanisms, regardless of complexity, are combinations of four characteristic patterns of electron flow (discussed later).

Notes on Drawing Curved Arrows

- Tails must be placed on either a bond or a lone pair.
 - Shows the source, i.e., the electron donor (base).
 - Electrons can only be found in lone pairs or bonds, so never place the tail of a curved arrow on a positive charge.
- Heads must be placed so that it shows either the formation of a bond or the formation of a lone pair.
 - Shows the destination, i.e., the electron acceptor (acid).
 - Avoid drawing an arrow that violates the octet rule, so never draw an arrow that gives more than four orbitals to a second-row element.

4 Alkanes and Cycloalkanes

Nomenclature of Alkanes

- \triangleright **Alkane**: acyclic (linear structure) saturated hydrocarbons (no π bonds).
 - General chemical formula: C_nH_{2n+2}
- ▶ **Substituents**: branches connected to the parent chain.

Selecting the Parent Chain

• Parent chain: the longest carbon chain in an alkane.

Parent Names for Alkanes

Number of Carbons	Parent	Name
1	meth	methane
2	eth	ethane
3	pro	propane
4	but	butane
5	pent	pentane
6	hex	hexane
7	hept	heptane
8	oct	octane
9	non	nonane
10	dec	decane
11	undec	undecane
12	dodec	dodecane
13	tridec	tridecane
14	tetradec	tetradecane
15	pentadec	pentadecane
20	eicos	eicosane
30	triacont	triacontane
40	tetracont	tetracontane
50	pentacont	hectane
100	hect	hectane

- **Substituents**: branches connected to the parent chain, can be a single atom, groups of atoms, that replace one or more hydrogen atoms.
 - If there is competition between chains of equal length, then choose the chain with greatest number of substituents.
- Cycloalkanes (cyclo): presence of a ring in an alkane.

Naming Substituents

• **Alkyl groups**: Substituents that are named the same as the parents, but with the added letters ly.

ivames of Alkyl Groups					
Substituent Carbons	Terminology				
1	methyl				
2	ethyl				
3	propyl				
4	butyl				
5	pentyl				
6	hexyl				
7	heptyl				
8	octyl				
9	nonyl				
10	decyl				

Names of Alkyl Groups

- When a group is connected to the ring, then the ring is generally treated as the parent.
 - If the ring has fewer atoms the the rest of the structure, then it becomes a substituent.

Naming Complex substituents

- **Complex substituents**: branched alkyl substituents.
- Begin by numbering carbons going away from the parent chain, then name it as if its a parent chain itself.
 - Complex substituent are placed in parentheses, indicating it as a single substituent of the parent chain.
- Some complex substituents have common names that are so well established and allowed by IUPAC.
 - An alkyl group bearing three carbon atoms; only one way to branch it.
 - · Isopropyl group: (1-methylethyl):
 - Alkyl groups bearing four carbon atoms, which can be branched three different ways:
 - · **sec-butyl** (1-methylpropyl): ___

- isobutyl (2-methylpropyl): ______
 tert-butyl (1,1-dimethylethyl): ______
- Alkyl groups bearing five carbons, which can be branched many more ways. Two common ways:
 - · isopentyl (isoamyl) (3-methylbutyl): _____
 · neopentyl (2,2-dimethylpropyl): ____

Assembling the Systematic Name

- Locant: the location of a carbon numbered parent chain.
- Rules for assinging locant:
 - If one substituent is present, then assign the lowest number possbile.
 - When multiple substituents are present, then the first substituent receives the lowest number.
 - · If there is a tie, the second locant should be as low as possible.
 - · If tie cannot be broken, then lowest number is assigned alphabetically.
 - Prefixes are used when the same substituent appears more than once.
 - · di:2, tri:3, tetra:4, penta:5, 6:hexa
 - Hypens are used to separate numbers from letters, while commas are used to separate numbers from each other.
 - Substituents are alphabeticalized after all locants are correctly assigned.
 - · Prefixes are ignored during alphabeticalization.
- Summary of discrete steps:
 - 1. Identify parent chain
 - 2. Identify and name substituents
 - 3. Number the parent chain and assign a locant to each substituent
 - 4. Arrange the substituents alphabetically

Constitutional Isomers of Alkanes

- ⊳ For an alkane, the number of possible constitutional isomers increases with increaseing molecular size.
- Determing IUPAC name is the best way to tell if two alkanes are constitutional isomers, or just different representations of the same one.

		_		
\bigcap	1+100011	somers for	\	1 II / a /a a a
	uunnai i		various	DIKANAC

Molecular Formula	Constitutional Isomers			
C ₃ H ₈	1			
C_4H_{10}	2			
C_5H_{12}	3			
C_6H_{14}	5			
C ₇ H ₁₆	9			
C_8H_{18}	18			
C_9H_{20}	35			
$C_{10}H_{22}$	75			
$C_{15}H_{32}$	4,347			
$C_{20}H_{42}$	366,319			
C ₄₀ H ₆₂	4,111,846,763			

Newman Projections

- ▶ **Conformations**: the variety of possible three-dimensional shapes of a molecule that are interchangeable by low energy pathways.
 - Conformations vary in potential energy.
 - \circ Changes due to rotation about σ bonds.
- ▶ **Configurations**: refer to different orientations in space that require breaking of bonds (high energy pathway) to change.
 - Cis and trans isomers in alkenes (discussed later)
- ▶ **Newman projections**: a type of representation of compounds specially designed for showing the conformation of a molecule.
 - o Drawn from the angle of the observer, with the front carbon represented in front of the circle, and the back carbon behind the circle.
 - ***Chemmacros package is broken due to font usage, need to figure out how to fix that before inserting drawings***

Conformational Analysis of Ethane and Propane

- \circ **dihedral (torsional) angle**: the angle between substituents of front and back carbons as the σ bonds rotates.
- There are an infinite number of possbile conformations, but there are conformations of maximum and minium energy.
 - Staggered conformation: lowest energy conformation, when two substituents are at maximum dihedral angle from each other.
 - Eclipsed conformation: the highest energy conformation, when two substituents are at the minimum dihedral angle from each other.
- **Degenerate**: when all staggered conformations have the same amount of energy.
 - All staggered and eclipsed conformations of ethanes are degenerate.
- **Torsional strain**: the difference in energy between staggered and eclipsed conformations.
 - Recent quantum methods suggest conformation possesses a favorable interaction between occupied, bonding molecular orbitals and unoccupied, antibonding molecular orbitals.
 - An increase in potential energy occurs when the favorable overalp is broken.
 - A sample of ethane gas at room temperature will have \approx 99% of its molecules staggered.
- $\circ\,$ Ethane has total cost of 12 kJ/mol (4 kJ/mol/H), while propane has total cost of 14 kJ/mol.
 - Reasonable estimates of energy cost of an H eclipsing a CH₃ group must be 6 kJ/mol.

Conformational Analysis of Butane

- Butane has three eclipsed conformations that are not degenerate.
 - Dihedral angle of 0° has the highest eclipsed energy, while both conformations at $\pm 120^\circ$ are second highest in energy and degenerate.
 - Likewise, a dihedral angle of 180° has the lowest staggered energy, while both conformations at $\pm 60^\circ$ are second lowest in energy and degenerate.
- Anti conformation: the conformation with a dihedral angle of 180; the lowest staggered energy.

- Occurs when the methyl groups are farthest apart.
- **Steric interaction**: nonbonding intereactions that influences energy levels conformations.
- **Gauche interaction**: unfavorable intereaction between substituents, causing an increases in energy due to electron cloud repulsion.
 - Gauche intereaction is a type of steric intereactions present at $\pm 60^{\circ}$ of the next eclipsed conformation.
- o Costs of butane: 19 kJ/mol, 16 kJ/mol, 3.8 kJ/mol
 - Energy cost of eclipsing CH₃/CH₃: 11 kJ/mol
 - Energy cost for gauche interaction of CH₃/CH₃ 3.8 kJ/mol for butane.
 - Energy cost of eclipsing CH₃/H: 6 kJ/mol
 - Energy cost of eclipsing H/H: 4 kJ/mol

Cycloalkanes

- ▶ **Angle strain**: the increases in energy associated with a bond angle that has deviated from the preferred angle of 109.5°.
 - o Cyclic alkanes, excpet cyclopropane, are not planar.
 - Expected angels are different than origanally proposed by Adolph von Baeyer, which assumed rings were planar.
 - Angle strain is only one factor that contributes to the energy of various ring sizes.

▶ Cyclopropane:

- Under significant angle strain.
- Locked into an eclipsed conformation due to triangular structure;
 exhibiting significant torsional strain.
- Thus highly reactive and very susceptible to ring-opening reactions.

▶ Cyclopentane:

- Less angle strain than cyclopropane.
- More torsional strian than cyclopropane due to four sets eclipsing hydrogens.
- Adopts slightly "puckered" shape, which is the cause of reduced angle strain.

▷ Cyclopentane:

- Less total strain than both cyclopropane and cyclopentane.
- Can adopt a relatively low strained conformation.

Conformations of Cyclohexane

Chair conformation:

- Bond angles close to 109.5°; little angle strain.
- No torsional strain; all hydrogens are staggered.
- Least potential energy of cyclohexane conformations.
- Half-chair: highest potential energy, formed via interchange between alternate chair form; leads into twisted boat.

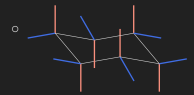
o Boat conformation:

- Bond angles also close to 109.5°; little angle strain.
- Two sources of torsional strain; many of hydrogens are eclipsed.
- One hydrogen on each side experiences a steric interaction called the flagpole intereaction.
- Second highest potential energy.
- Twisted boat: second lowest potential energy, a slightly less strained version of boat that avoids some of the flagpole interaction.
- Majority of cyclohexanes are found in chair form. All other forms are intermediates between alternate chair forms.

Drawing Chair Conformations



- Axial position: parallel to a vertical axis passing through the center of the ring.
 - less stable than equatorial due to steric strain.
- **Equatorial**: positioned approximately along the equator of the ring.



- The chair is more stable when the methyl (substituent) group is in the equatorial position.
 - The larger the substituent, the more equatorial-substituted conformer is favored.

5 Stereoisomerism

Overview of Stereoisomerism

- ▶ **Constitutional isomers**: aka structural isomers; same chemical formula, but different in the way the atoms are connect, i.e. their constitution is different.
- ▶ **Stereoisomers**: isomers that differ in spatial arrangement of atoms, rather than connectivity.
 - **Geometric isomerism**: aka cis-trans; locked into spatial positions due to double bonds or a ring structure.
 - Cis: functional groups that are on the same side of the carbon chain.
 - Trans: functional groups on opposite sides of the carbon chain.
 - Cis-trans terminology is used to describe disubstituted alkenes (carbon chain with π bond), even when the two substituents are different from each other.
 - Does not apply to disubstituted alkenes in which the substituents are connect in the same position.

Chirality

- **Superimposable (achiral)**: when an object's mirrored version is identical to the actual object.
- o Chiral: objects that are not superimposable.
- All three-dimensional objects can be classified as either chiral or achiral.
- **Enantiomer**: the nonsuperimposable mirror image of a chiral compound.
 - Can be used in speech the same way twin is used
 - Easiest way to draw enatiomers is to just change wedges and dashes, but there are multiples ways to mirror a molecule, so it can be more complex.
- The most common source of molecular chirality is the presence of a carbon bearing four different groups.
 - Chirality (chiral) center: a tetrahedral carbon that bears four different groups.
- **Diastereomers**: non-identical stereoisomers (nonsuperimposable) that are not mirror images of one another.
 - Enantiomers have the same physical properties, while diastereomers have different physical properties.

- Differences between enantiomers and diastereomers are especially relevant when comparing compounds with more than one chiral center.
- Maximum number of stereoisomers: 2ⁿ
 - · n: number of chiral centers
 - $\frac{2^n}{2}$: pairs of enantiomers.

Cahn-Ingold-Prelog System

- ▶ **Chan-Ingold-Prelog system**: a system of nomenclature for Identifying each enantiomer individually.
 - 1. Assign priorties to each of the four groups based on atomic number; the highest atomic number has the highest priority.
 - 2. Rotate the molecule so that the fourth priority group is on a dash (behind)
 - 3. Determin the configuration, i.e., sequence of 1-2-3 groups.
 - o clockwise (R) or counterclockwise (S).
- ▶ If there is a tie between the atoms connected, then continue outward until a difference is found.
 - Do not add the sum all atomic numbers attached to each atom, just the first in which the atoms differ.
 - Any multiple bonded atom, (2 or 3) is treated as if connected to multiple atoms equal to number of bonds.
- ⊳ Switching any two groups on a chiral center will invert the configuration, e.g.,



o Can be done twice with out changing configuration, e.g.,



Configuration in IUPAC nomenclature:

- The configuration of the chiral center is indicated at the begining of the name, italicized, and surrounded by parentheses.
- When multiple centers are present, then each must be preceded by a locant.

- ▶ Enantiomers exhibit identical physical properties, but different behavior to plane-polarized light.
 - **Polarization**: the orientation of electric field of the electromagnetic wave.
 - Plane-polarized light: a filter that only allows light of a particular polarization through.
- Description Optically active: property of compounds that rotate the plane-polarized light.
 - Can be measured using a polarimeter.
 - Chiral compounds are optically active, while achiral compounds are not optically active.
- \triangleright **Observed rotation (** α **)**: the rotation of light due to chiral compounds, which depends on the number of molecules light encounters.
 - Doubling the concentration or pathlength both double the observed rotation.
- \triangleright **Specific rotation** [α]: a standard concentration (1 g/mL) and a standard pathlength (1 dm) that allows for meaningful comparion between compounds.

$$\circ \ [\alpha] = \frac{\alpha}{cI}$$

- o c: concentration, l: pathlength
- Temperature (T) and wavelength (λ) both have a nonlinear relationship, so it is often noted as: $[\alpha]_{\lambda}^{T}$
- > Specific rotation for enantiomers are equal in magnitude but opposite in direction.
 - dextrorotaory: a compound exhibiting positive rotation.
 - levorotatory: a compound exhibiting negative rotation.
 - No direct relationship between R/S system of nomenclature, as that is independent of conditions, but dependent on observation angle.
 - The direction of polarized light, however, is dependent on conditions, and can change based on temperature or wavelength even with the same given configuration.

Enantiomeric Excess

- Optically (enantiomerically) pure: a solution containing a single enantiomer.
- **Racemic mixtrue**: a solution containing equal amounts of both enantiomers, resulting in an optically inactive appearance.
- **Enantiomeric excess** (*ee*): when a solution containing both enantiomers in unequal amounts, appearing optically active.

- %
$$ee = \frac{|\mathsf{observed}\;(\pmb{lpha})|}{|\mathsf{specific}\;[\pmb{lpha}]|} imes 100\%$$

Symmetry and Chirality

- ▶ Any compound with a single chiral center must be chiral, however, the same it not always true for two or more.
- ▶ **Reflectional symmetry**: when an object has a plane of symmetry that can be reflected across and still generate the same image.
 - Any compound that posses a plane of symmetry in any conformation will be achiral.
- ▶ **Rotational symmetry**: when an object has an axis of symmetry, i.e., it can be rotated around a single point and appear 2 or more times.
 - Order: how many times the object appears.
 - o Chirality is not dependent on rotational symmetry.
- ▶ **Point symmetry (inversion)**: when every part of an object has a matching part, i.e., equal distance from the central point, but in opposite direction.
 - o Absence of a plane of symmetry does not mean it is chiral.
 - o If a compound exhibits inversion, then it is achiral.
- ▷ Summary of relationship between symmetry and chirality:
 - The presence of absence of rotational symmetry is irrelevant to chirality.
 - A compound that has a plane of symmetry will be achiral.
 - A compound that lacks a plane of symmetry will most likely be chiral, with some exceptions.
- ▶ **Meso compounds**: achiral compounds (posses plane of symmetry or can be inverted) that have multiple chiral centers.
 - A family of stereoisomers cataining a meso compound will have less than 2^n stereoisomers.

Fischer Projections

- **Fischer projections**: two-dimensional representations of organic molecules by projection.
 - Limited to carbohydrates/sugars with multiple chiral centers.
- Horizontal lines are considered to be coming out of the page.
- Vertical lines are considered to be going behind the page.
- Helpful for quickly comparing relationship between stereoisomers with multiple chiral centers;
 - Enantiomers will have opposite configurations, while diastereomers will not.
 - Assuming the north is priority 2, and the south is priority 3:
 - · If the atom with priority 1 is on the right, then it will have an R configuration.
 - · If the atom with priority 1 is on the left, then it will have an S configuration.

6 Chemical Reactivity and Mechanisms

Enthalpy

- ▶ **Enthalpy**: the sum of the system's internal energy and the product of its pressure and volume.
 - A convenient state function preferred in measurements at constant pressure.
 - State function (function of state, point function): a system relating several state variables (variables of dynamic system) or quantities that depends on the current equilibrium of the system.
 - As a state function, enthalpy depeds only on the final configuration, not the path taken to achieve it.
- $hd \Delta H = q$ (at constant pressure)
 - \circ $\triangle H$ the exchange of kinetic energy, or heat (q).
 - Primary determined by the amount of energy necessary to break the bond homolytically.
 - Homolytic bond cleavage: genetrates two radicals (uncharged species), each which bears an unpaired electron.
 - Heterolytic bond cleavage: generation of charged species, i.e., ions.
- **Bond dissociation energy (\Delta H^{\circ})**: the energy required to break a covalent bond via Homolytic bond cleavage under standard conditions (pressure 1 atm and standard state—gas, liquid, or solid).
 - I.e., enthalpy.

Heat of Reaction

- **Heat of reaction**: the total change in enthalpy (ΔH°) .
 - The sign indicates the direction in which the energy is exchanged from the perspective of the system.
 - A positive ΔH° indicates that the system increased in energy due to transfer from the surroundings.
 - I.e., indicates an endothermic process.
 - A negative ΔH° indicates that the system decreased in energy due to transfer to the surroundings.
 - · i.e., indicates an **exothermic** process.
 - Physicists think of ΔH° as moving from system \longrightarrow surroundings

- Chemists think of ΔH° as moving from the surroundings \longrightarrow system.
- o Predicting sign and magnitude:
 - 1. Identify all bonds that are broken (gained energy from surroundings) or formed (lost energy to the surroundings).
 - 2. Then find the bond dissociation energies.
 - 3. Apply the sign to the bonds in respect to the system based on breakage (+) or formation (-).
 - 4. Sum to find ΔH° .

Entropy

- ▶ Entropy: informally defined as the measure of disorder associated with a system.
 - Essentially it represents the gradual decline towards disorder since random occurrences resulting order is less ultimately probable than disordered results.
- ▶ Spontaneous: a process that involves an increase in entropy, which will always occur given enough time.
 - \circ $\Delta S^{\circ} = \Delta S_{ ext{surr}} + \Delta S_{ ext{sys}}$
 - \circ ΔS° is the total change in entropy associated with the reaction.
 - Sometimes referred to as just S, ΔS , or ΔS_{total}
- ▶ Entropy of the system (reaction) can decrease, as long as the entropy of the surroundings increases by a greater amount.
- \triangleright Thus, $\triangle S^{\circ}$ must by positive for a reaction to be spontaneous.
- ▶ A larger number of possbile states corresponds with larger entropy.
 - Acyclic compounds can adopt more conformations, so they have more entropy than cyclic compounds.

Gibbs Free Energy

- \triangleright Under constant pressure and temperature, $\triangle S_{\text{surr}}$ can be mesesaured:
 - $\circ \Delta S_{\text{surr}} = -\frac{\Delta H^{\circ}}{T}$
 - o Both ΔH° and temperature (in Kelvin) can be easily measured, which means ΔS_{surr} can be mesesaured.
 - o Applying this definition:

$$\Delta S^\circ = -rac{\Delta H^\circ}{T} + \Delta S_{
m sys} \ -T\Delta S^\circ = \Delta H^\circ - T\Delta S_{
m sys} \ \Delta G^\circ = \Delta H^\circ - T\Delta S_{
m sys}$$

- ΔH is then associated with the change in entropy of the surroundings, as a transfer in energy to/from the surroundings from the system increases/decreases entropy of the surroundings.
- $-\Delta S^{\circ}$ is associated with the change of entropy of the system.
- ▶ **Gibbs free energy (** ΔG° **)**: the maximum amount of non-expansion work that can be extracted from a closed system.
 - Essentially a repackaged way of expressing entropy in a closed system.
 - \circ ΔG° is just ΔS° multiplied by the negative temperature in order to measure the entropy of the surroundings.
 - Thus, ΔG° must be negative for a reaction to be spontaneous. (second law of thermodynamics)
- \triangleright Often ΔH° has a much greater contribution than $T\Delta S^{\circ}$, so enthalpy is commonly an indicator of the spontaneity.
 - When more bond energy is broken $(+\Delta H^{\circ})$, then the ΔG° is decreased, which increases spontaneity.
 - When more bond energy is formed $(-\Delta H^{\circ})$, then the ΔG° is increased, which decreases spontaneity.
- ▶ However, temperature has a large influence on the entropy of the system:
 - o An **increase in temperature** (endothermic) leads to more bond energy being broken $(+\Delta S_{\text{sys}})$ since energy is transferred from the surroundings $(-\Delta S^{\circ})$, which is unfavorable, as it increases ΔG° .
 - **Endergonic**: nonspontaneous $(+\Delta G^{\circ})$ processes.
 - A **decrease in temperature** (exothermic) leads to more bond energy being formed $(-\Delta S_{\text{sys}})$ since energy is transferred to the surroundings $(+\Delta S^{\circ})$, which is favorable, as it decreases ΔG° .
 - **Exergonic**: spontaneous $(-\Delta G^{\circ})$ processes.

Equilibria

- ▶ **Reaction quotient** (Q_r, Q_c, Q) : a measurement of the relative quantities of products and reactants present in a reaction at a particular point in time.
 - Defined as the the ratio of the effective (molar) concentrations of the product over the reactants.
 - General chemical reaction with α moles of reactant A and β moles of reactant B to give ρ moles of product R and σ moles of product S:

$$-\alpha A + \beta B \iff \rho R + \sigma S$$

o Reaction quotient as a function of time:

$$-~Q_r(t)=rac{R_t^
ho S_t^\sigma}{A_t^lpha B_t^eta}$$

▶ **Equilibrium constant** (K_{eq}, K^{\odot}) : a chemical reaction's reaction quotient (Q_r) at chemical equilibrium given sufficient time.

$$\circ \ \textit{K}_{\text{eq}} = \frac{\textit{R}^{\textit{p}}\textit{S}^{\textit{\sigma}}}{\textit{A}^{\alpha}\textit{B}^{\textit{\beta}}} = \frac{\text{products}}{\text{reactants}}$$

- $\circ Q_r = K_{eq}$: the system is at equilibrium.
- $\circ Q_r < K_{eq}$: more reactants than products—reaction will shift towards the products. (right)
 - $Q_r = 0$: $Q_r \ll K_{eq}$ -reaction will shift towards products.
- $K_{eq} < Q_r$: more products than reactants—reaction will shift towards the reactants. (left)
 - $Q_r = \infty$: $Q_r \gg K_{eq}$ -reaction will shift towards reactants.
- \triangleright The ultimate direction of a reaction is governed by ΔG° .

$$\circ \ \Delta G^{\circ} = -RT \ln K_{eq}$$

- − R: gas constant 8.314 J/mol·K
- T: temperature in Kelvin (C° 273.15).
- \circ - ΔG° : the products are favored. (1 < K_{eq})
- \circ + ΔG° : the reactants are favored. ($K_{\rm eq} < 1$)
- o A small difference in ΔG° can have significant impact on the K_{eq} and ultimately determines the yield of products that can be expected from any reaction.
- \triangleright The relationship between $\triangle G^{\circ}$ and equilibrium falls under the realm of thermodynamics, i.e., the study of how energy is distributed under influence of entropy, i.e., the spontaneity/direction of reactions.

Kinetics

- ▶ **Kinetics (reaction kinetics)**: the investigation of experimental condictions that influence the speed (rate) of chemical reactions.
 - Spontaneity has nothing to do with the rate of a reaction.
 - Reveals information about reaction mechanisms and transition states, as well as the construction of models that describe reactions.
- ▶ Rate equation (general): rate = k[reactants]
 - *k*: a constant value that is specific to each reaction and dependent on a number of factors, such as, substrate type, physical state, surface area, concentration, temperature, catalysts, pressure, and light absorption.
 - Can be reduces to more general factors: concentration, energy of activation, temperature, and steric considerations.
 - Reaction speed is positively correlated with the rate of reaction.
 - The steric considerations, or the geometry and orientation, of colllisions have an impact as well, and will be discussed later.

Concentration

- Reactions are result of collision between reactants, thus increases concentration should the rate of reactions.
- Precise effect must be determined expirementally:
 - Rate = $k[A]^x[B]^y$
 - **Rate order**: the sum of exponents of the reactants.
 - E.g., kA = first, kAB = second, $kA^2B = \text{third}$.

Energy of Activition

- **Energy of activation** (E_a): the energy barrier between reactants and the products.
 - Represents the minimum amount of energy required for a reaction to occur between two reactants that collide.
 - Number of successful intereactions depends on tue number of molecules that have more than a certain threshold kinetic energy required to form the products.
 - Catalysts: a compound that can speed the reaction up with out being consumed by lowering the activation energy of a reaction.

Temperature

- Rasing the temperature causes the rate of the reaction to increases due to larger portion of molecules passing kinetic threshold for the reaction to occur.
- o Generally raising the temperature by 10 °C double the rate.

Energy Diagrams

- ▶ Kinetics refers to the rate of a reaction, while thermodynamics refers to the equilibrium concentrations of the reactants and products.
- ▶ Different sets of products can be favored from the reactants depending on activation energy, regardless of stability of the products.
- ▶ Products that require lower activation energy will be favored when environmental potential energy is low.
- ▶ More stable products will be favored when potential energy in the environment is high, or other means at overcoming the larger activation energy are present.
- ▶ Reactions often require multiple steps:
 - **Intermediates**: represent local minima of the reaction.
 - Have a certain, albeit short, lifetime.
 - Very common in reactions and widely observed.
 - **Transition states**: represent local maxima of the reaction.
 - Cannot be isolated.
 - Represents high-energy states where bonds are being simultanesously broken and formed.
- ▶ Hammond postulate: states that the transition state of a reaction resembles either the reactants or the products depending on proximity in terms of potential energy.
 - Endothermic reactions represent transition states that are closer to the products, since more energy remains in the system leaving less stable products.
 - Exothermic reactions represent transition statets that are closer the the reactants, since more energy leaves the system leaving more stable products.

Mechanisms and Arrow Pushing

- ▶ Previously excerpted section, Nucleophiles and Electrophiles, contains relevant information.
- ➤ There are four distinct characteristic patterns that make up all ionic mechansisms through various combinations; nucleophilic attacks, loss of a leaving group, proton transfers, and rearrangements.

Nucleophilic Attack

- When a nucleophile (electron dense atom/lewis base/donator) attacks (donates electrons) a electrophile (electron lacking/lewis acid/acceptor).
- Involves just one curved arrow, with tail on the nucleophile and the head on the electrophile.
- Often a nucleophilic attack causes a second resonance arrow to be drawn, showing the actual flow of electron density caused by the initial attack.
- \circ π Bonds often serve as nucleophiles, as they are relatively more electron dense compared to nearby σ bonds.

Loss of a Leaving Group

- When a substituent (group) is forced off (leaves) a molecule due to change in electron density.
- Often more than one arrow is drawn, again respresenting resonance arrows (or changes in electron density), that ultimately force the group to leave.

Proton Transfers

- o Previously discussed in terms of Brønsted acid and bases in chapter 3.
- Characterized by two curved arrows due to the combination of a new molecule that causes a compound to be protonated, or depronated, transfering the proton to a different atom.
- Sometimes the arrow from the proton is omitted.
- Somtimes there is more than two arrows due to resonance structures that can be created from the change, or again just the flow in electron density.

Rearrangements

- There are several kind of rearrangements, but carbocation rearrangements are focused here.
- **Hyperconjugation**: carbocations that can be stabilized by neighboring alkyl groups due to molecular orbitals that slightly overalp with empty *p* orbitals, placing some of its electron density there.
 - Primary, secondary, and tertiary: refer to the number of alkyl groups directly attached to the carbocation.
 - Tertiary are the most stable (more slight overlap) and primary are the least (less overlap)
- **Hydride shift**: involes the migration of a H⁻.
 - Involves the rearrangement of the carbocation to a more stable variant due to change migration of the H⁻.
- **Methyl shift**: similar to a hydride, except a whole methyl group is migrated instead.
 - The methyl group must be attached to the carbon atom that is adjacent to the carbocation for this to occur.
- Again, there are more, but hydride and methyl shifts are the most common.

Reversible and Irreversible Reaction Arrows

- Reversible arrows can be the result of nucleophilic attacks that leave the nucleophile being capable of a functioning leaving group (weak bases).
 - Non-reversible arrows show the opposite, i.e., poor leaving groups (strong bases).
- Likewise, reversible arrows can show when leaving groups, caused by any means, are capable of functioning as a good nucleophile.
 - Most leaving groups can also function as leaving groups, so most are reversible.
- All protons are technically reversible, however many have very minute reversible reactions.
 - A strong difference in pK_a represents this fact, while a small difference allows for protons to be transferred back and forth.
- Carbocation rearrangements tend to only occur when all possbile carbocations are present, which means they are hardly ever reversible.

7 Alkyl Halides: Nucleophilic Substitution and Elimination Reactions

Introduction to Substitution And Elimination Reactions

- ▶ **Alkyl halides**: compounds in which a halogen (Cl, Br, I) is connected to sn sp^3 hybridized carbon atom.
 - Alkyl halides are electrophiles since they contain such electron deficient halogens—making them easily accept new electrons.
- ▶ **Substitution reaction**: when a nucleophile replaces a halogen.
- ▶ **Elimination reaction**: when an alkyl halide is treated with a base, allowing fo formation of π bond, eliminating the halogen.
- ▶ Many reagents (e.g., OH⁻) can function either as a nucleophile or as a base, so substitution and elimination often compete with each other.
- ▷ Often the word substrate is used to refer to the alkyl halide since substitution and elimination can occur for a variety of substrates, not just alkyl halides.
- ▶ I an alkyl halide, the halogen serves two critical functions that render the alkyl halide reactive:
 - The halogen withdraws electron density via induction, rending the adjacent carbon atom electrophilic, and therefore subject to attack.
 - The halogen can serve as the leaving group for the compound, vital for substitution or an elimination to occur.
 - Good leaving groups are conjugate bases of strong acids, i.e., good groups are weak bases.
 - Generally an acid with a $pK_a < 0$ generates a stable enough base to be a good leaving group, which is why F (pK_a of HF is 3.2) is not one, despite being a halogen.
- ► The most common leaving groups are halides (I¯, Br¯, and Cl¯) and sulfonate ions (RSO₃¯)

Nomenclature and Uses of Alkyl Halides

- $\circ \ \alpha$ **position**: the position connected directly next to the halogen.
 - β **position**: positions connected to the α position.
 - As many as 3 β positions; following familiar naming structure of primary (1°), secondary (2°), or tertiary (3°) based on number of β connected to the α position.
- Naming Alkyl Halides:

- Halogens are simply treated as substituents, receiving fluoro-, chloro-, bromo-, and iodo-.
- Haloalkane: formal name for alkane containing a halogen substituent.
- Alkyl halide: the common name that treats the compound as with an alkyl substituent connected to a halide.
- Organohalide: any organic compound containing a halogen, including alkyl halides.
- Many organohalides are toxic and have been used as insecticides.
- Sucralose, which contains three chlorine atoms, is sold under the name of Splenda.
- Organohalides often serve as the synthesis of more complex molecules, especially aryl and vinyl halides (not discussed yet).
- ► Early investigation by Ingold and Hughes revealed concerted, i.e., simultanesous occurrences of reaction mechanisms, and stepwise, i.e., one mechanism that allows for a different mechanism(s) to take place.
- \triangleright Conditions that favor the concerted process (S_N2 , E2) will be explored first, while the stepwise (S_N1 , E1) will be explored later.

S_N 2 Reactions

- \triangleright **Kinetics of** S_N 2 **reactions**: a biomolecular (2) nucleophilic (N) substitution (S) reaction.
 - **Biomolecular**: a step that involves two chemical entities, such as when the alkyl halide and nucleophile collide during the reaction mechanism.
 - Rate: $v_0 = k$ [alkyl halide][nucleophile]
 - Second-order rate is means it shows a linear dependent relationship between rate and chemicals; any change in concentrations will have a proportional effect on the rate.

Stereospecificity

- **Inversion of configuration**: when an α position is a chiral center, then a change in configuration is generally observed after a reaction takes place.
 - back-side attack: a requirement for the inversion to take place; a nucleophile can only attack from the back side opposite to the leaving group.
 - Lone pairs of the leaving group act as regions of high electron density that block the front side of the substrate and thus the nucleophile attack.
 - The incoming electron pair must go through an empty orbital on the back, and cannot go through the occupied orbital in the front.
 - The result is like the chiral center with an umbrella top(filled orbital/halogen) flipping (leaving group) due to the wind (nucleophilic attack) hitting the back (empty orbital).
- **Stereospecific**: when the configuration of the product is dependent on the configuration of the starting material.

Effect of Substrate Structure on Rate

- \circ The rate of an S_N 2 reaction is extremely sensitive to the nature of the starting alkyl halide.
 - The presence of substituentsat the α or β positions can significantly reduce the rate.
- The rate of an S_N 2 reaction is more sensitive to the number of substituents at the α position.
 - Methyl halides and primary alkyl halides are the most effective.
 - t-butyl (a tertiary alkyl halide) is unreactive.
- \circ The effect of β substituents is not as great.
 - However it still has noticeable impact; even with just two substituents the rate is moderately reduced.
 - With three substituents the rate is too generally slow to be of any use.
- The more substituents, the more bonds that need to be broken/changed and the more steric intereactions there are during the transition state—leading to higher activation energy and thus a slower reaction.

Nucleophilic Strength and Solvent Effects

- Nucleophilicity: the rate at which a nucleophile will attack a suitable electrophile.
 - A high rate leads to a relatively fast reaction, while a slow rate leads to a slow one.
 - Generally means a strong nucleophile is required for practical $S_N 2$ reactions.
 - Nucleophilicity decreases as electronegativity increases.
- Polarizability: the ability of an atom to distribute its electron density unevenly in response to external influences.
 - Correlated with size of the atom, which increases the number electrons that are distant from the nucleus.
 - Has a significant effect on nucleophilicity, often more so than degree of negative charge.
 - Another contributing factor that makes flouride behave as a weak nucleophile.
- \circ S_N 2 reactions are generally ionic, as is the leaving group, so polar solvent is required in order to solvate the ionic species.
 - The transition state also often has ionic charater, which helps the stability of the state.
- Protic solvents: polar solvent that contains a hydrogen atom connected directly to an electronegative atom.
 - E.g., water, methanol (O-H), ethnol (O-H), ammonia (N-H), acetic acid (O-H).
 - Allows for stabilization of added nucleophiles in the solution, reducing potential energy and thus a higher activation energy for a reaction to take place.
 - $-S_N$ 2 reactions are generally slower in protic solvents.
- **Polar aprotic solvents**: protic solvents lacking a hydrogen atom connected to electronegative atoms.
 - E.g., Acetone, dimethylethylformamide, dimethylethylsulfoxide, acetonitrile; all of which π bonded oxygen or nitrogen with no hydrogen attached.
 - Lack of hydrogen fails to stabilize added nucleophiles, leaving compounds with higher potential energy and thus a lower activation energy for a reaction to take place.

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 $-S_N 2$ reactions are generally much faster in polar aprotic solvents.