CHEM 123: Summary of Important Concepts Arranged by Hayden Scheiber - Resource Center TA

Thermodynamics

Equation/Concept	Info About Equation/Concept			
System	The system is defined as a group of material and/or radiative contents. It's properties can be fully described by thermodynamic state variables.			
Boundary	The boundary is the edge of a system. It may be a conceptual boundary or a real boundary.			
Surroundings	The entire universe outside of the system and its boundary.			
Isolated System	A system where neither energy nor mass can cross between the system and surroundings. The only truly isolated system is the entire universe, though we can produce approximately isolated systems (i.e. a well-insulated thermos).			
Closed System	A system where energy can exchange with the environment, but mass cannot. A good example of a closed system would be a car's cooling system: when the car is on, heat moves from the radiator into the environment, but no coolant leaves the system.			
Open System	A system where both energy and mass may freely exchange with the environment. Examples include open flask or cup of coffee.			
State Variable	State variables are variables that describe the thermodynamic state (i.e. the conditions) of a system. They are only well-defined for systems in equilibrium. State variables do not depend on the path taken to get to a given thermodynamic state. Examples: temperature, pressure, internal energy, enthapy, Gibbs free energy, entropy, volume, density, mass, particle number, and concentration.			
Path Variable	Path variables, unlike state variables, are non-zero only during a change in a system's thermodynamic state (i.e. when not at equilibrium). Path variables depend on the particular way in which a system evolves through time from one state to another. Heat and work are the only path variables used in this course.			
Intensive Variable	These are variables that are independent of the system size. Examples include: temperature, energy density, pressure, density, concentration, and specific heat capacity. The ratio of two intensive variables is still intensive.			
Extensive Variable	Extensive variables are variables that are proportional to the size of the system. If the system grows, then so do all extensive variables that describe the system. Examples include: internal energy, entropy, volume, mass, particle number, heat capacity. Dividing one extensive state variable by another will always produce an intensive variable because the system size dependence will cancel out.			
Equation of State	At equilibrium, state variables are not independent from one another. Ar equation of state defines the particular relationship between state variables Equations of state for real systems are often extremely complicated and may not have a closed form, but certain model systems can have fairly simple and useful relationships. For example, the ideal gas law is an equation of state that describes the relationship between state variables for a model gas system in which the particles of gas occupy zero volume (are point particles) and do not interact with each other. Although the ideal gas equation does not perfectly describe any gases in our universe, it becomes a very good approximation for dilute monatomic gases at high temperature.			
Internal Energy $(E \text{ or } U)$	Internal energy is the sum total of all forms of energy within a system. It includes all forms of potential energy, transnational energy, rotational energy, and vibrational energy. Generally, the total internal energy of a system is very difficult to measure, but changes in internal energy can be readily measured.			
Heat (q)	Heat is the quantity of energy that transfers between the system and surroundings due to a difference in temperature. When heat is absorbed by the system q is positive. When heat is removed from the system into the surroundings, q in negative. Heat only exists during a change in the system, it is a path variable			
Work (w)	Work is the energy transfer between system and surroundings due to a force applied over a distance. Work is associated with either an expansion (work done by the system, negative w) or a contraction (work done on the system, positive w) of the system. Work only exists during a change in the system, it is a path variable.			

Equation/Concept	Info About Equation/Concept		
Zeroth Law of Thermodynamics	Two systems that are separately in thermal equilibrium with a third system are also in thermal equilibrium with one another. This law was only formulated after the first three laws of thermodynamics were already written down, but is more fundamental in nature than the other three.		
$\Delta E = q + w$	This is the equation for the first law of thermodynamics. It states that the change in energy of the $system$ is equal to the heat flow q into the system (positive q is heat flow $into$ the system) plus the work done on the $system$ w . Work done on the $system$ corresponds to positive w . This is simply a statement of the conservation of energy and only applies $for\ a\ particular\ path$.		
$\Delta E = q_V$	Change in internal energy when the volume of the system remains constant. The subscript simply indicates that the heat is exchanged at constant volume.		
$dw = -P_{\text{ext}}dV$	The most general definition of the increment of work. This equation is for the $improper$ increment of work done on the system. It is improper because work is a path variable, so the increment of work done is not an exact differential. The definition of total work done on the system for a process. It is the integral from the initial volume to the final volume of the incremental amounts of work done along that path. This can be conceptualized as adding up the tiny increments of work dw from the initial state to the final state. This is the most general expression for work done.		
$w = -\int_{V_i}^{V_f} P_{\text{ext}} dV$			
$w = -nRT \ln\left(\frac{V_f}{V_i}\right)$	The is the work done on an <i>ideal gas</i> expanding/contracting reversibly and isothermally. In this situation, $P_{ext} = P_{sys} = \frac{nRT}{V}$ at every point along the path, so the above integral can be solved analytically to obtain the result shown.		
$w = -P_{\rm ext}\Delta V$	This is the work done on a system with constant external pressure. This follows directly from the definition of total work done for the case in which P_{ext} is a constant.		
Ideal Gas	An ideal gas is defined as a collection of point particles that do not interact with each other, but interact with the walls of their container elastically (i.e. energy and momentum is conserved for each collision). Ideal gas particles have masses, but the equation of state for ideal gases (ideal gas law) does not depend on particle mass. The internal energy of an ideal gas is $U = c_V nRT$, where c_V is the specific heat capacity at constant volume (approximately $\frac{3}{2}$ for a monatomic gas), n is the number of moles of gas, and T is the temperature. Hence the internal energy of an ideal gas only depends on T as long as the number of moles is held constant.		
PV = nRT	This is the ideal gas law. It is the equation of state for a gas of point particle that do not interact with each other. P is the pressure of the system, V is the volume of the system, n is the number of moles of gas, R is the gas constant and T is the absolute temperature. This equation works well to approximate the equation of state for real gases at low P and high T . The equation of state for a mixture of ideal gases. P_i is called the partial pressure of gas i . The sum of the partial pressures for all gases in a mixture equals the total pressure. This is strictly true only for ideal gases, but is a good approximation otherwise.		
$P = \sum_{i} P_{i} = \sum_{i} \frac{n_{i}RT}{V}$			
$x_i = \frac{P_i}{P} = \frac{n_i}{n}$	For ideal gases only, the partial pressure for a component i in a mixture of ideal gases is proportional to the number of moles n . The partial pressure P_i divided by the total pressure is just $\frac{n_i}{n}$, known as the mole fraction x_i .		
Isothermal Process	An isothermal process is a process that remains at a constant temperature along the entire path from its initial state to its final state.		
Constant T Process	A process whose initial and final states have the same temperature, but may change temperature along the path between states.		
Adiabatic Process	A process that occurs without exchange of heat at any point along the path. Adiabatic walls are system boundaries that do not allow exchange of heat. They are the walls of an isolated system. Inversely, Diathermal walls are boundaries that do allow heat exchange.		
Reversible Process	An idealistic process that moves from one state to another while remaining in equilibrium with the surroundings along the entire path. Reversible processes would necessarily occur over an infinite amount of time, and so are not actually possible. A reversible process represents the upper limit on the possible work that can be extracted by an expanding gas, as well as the lower limit on the amount of work required to compress a gas. Reversible processes are called "reversible" because they require the same but opposite amount of work and heat to run backwards as forwards.		

Equation/Concept	Info About Equation/Concept			
Heat Capacity C	Heat capacity is defined as the amount of energy required to raise the temperature of a system by 1 K. It is an extensive state variable, but may be made intensive by dividing by either the mass of the system or the number of moles.			
$c_{sp} = \frac{C}{m}$	c_{sp} is called the specific heat capacity. It is an intensive state variable because it is the ratio of two extensive state variables. Here m is the mass of the system.			
$c_m = \frac{C}{n}$	c_m is called the molar heat capacity. It is an intensive state variable because it is the ratio of two extensive state variables. Here n is the number of moles in the system.			
$q = C\Delta T = mc_{sp}\Delta T = nc_m\Delta T$	The heat flow into the system is the product of the heat capacity C and the change in temperature $\Delta T = T_f - T_i$. Note that C is an extensive variable. This is only truly applicable for small changes in temperature, as C a temperature dependent state variable.			
H = E + PV	Definition of enthalpy. Enthalpy is an extensive state variable which is most useful in constant pressure conditions and at equilibrium when $P = P_{ext}$. In this case, $\Delta H = q_p$ where q_p is the heat flow at constant pressure. Since pressure is often held constant in chemistry, enthalpy is a very useful state variable. The heat flow of a reaction is simply the change in enthalpy at constant pressure, so the enthalpy of formation for a reaction is quite easy to measure.			
$\Delta H = \Delta E + P\Delta V$	The change in enthalpy for a constant pressure process. When a process results in $\Delta H < 0$ it is called exothermic and when a process results in $\Delta H > 0$ it is called endothermic . Reaction enthalpy alone is not enough to determine whether or not the process will proceed spontaneously.			
Standard State	An arbitrary but agreed-up standard set of conditions for which many experimental state variables are measured: $P=1\mathrm{bar}\approx 1\mathrm{atm}$ and either a pure substance or a 1 M solution. The standard state does not specify a temperature, but a separate term, standard temperature, is defined as $T=273.15\mathrm{K}$ while standard ambient temperature is defined as $T=298.15\mathrm{K}$.			
ΔH_f°	ΔH_f° is the standard enthalpy of formation , sometimes called the standard heat of formation . It is defined as the change in enthalpy required to create one mole of a given substance at standard pressure (1 bar) starting from the compound's constituent elements in their standard states. The standard enthalpy of formation of any pure element in its standard state is then, by definition, $0 \mathrm{kJ} \mathrm{mol}^{-1}$. Because ΔH_f° depends on temperature, the temperature must be defined, although it is often assumed to be 298.15 K.			
$\Delta H_{\rm rxn}^{\circ} = \sum_{\rm P} \Delta H_{\rm f,P}^{\circ} - \sum_{\rm R} \Delta H_{\rm f,R}^{\circ}$	Hess's Law. This law applies equally well to any state variable, such as total energy, entropy, etc. It is a reflection of the fact that the change in a state variable does not depend on the path taken. Here, P refers to products and R refers to reactants. Hess's law tells us that we can sum up as many processes as we wish in order to "build up" the value of an unknown state variable (such as enthalpy) from a set of known ones.			
Entropy S	Entropy is an extensive state variable that can be thought of as a measure of the dispersal of energy within a system. Many people think of entropy as a measure of system disorder, but this is an imprecise and often wrong definition. Entropy is actually a measure of the number of possible microstates that would give rise to a given macrostate. For macrostates with a large number of potential microstates, entropy is high. From this we can see that very highly ordered macrostates will tend to (but not always) have relatively few microstates compatible with the macrostate, whereas disordered macrostates will have a high number of associated microstates.			
General Cases of $S > 0$	Entropy generally increases when: temperature increases; pure liquids or solutions are formed from solids; the number of molecules of gas increases from a reaction; there is a decrease in pressure; gases are formed from liquids.			
$\Delta S_{ m Universe} \geq 0$	The second law of thermodynamics. The equality is only satisfied for reversible processes or systems in equilibrium, for all non-reversible processes the inequality is true. Here $\Delta S_{Universe}$ is the change in entropy of the <i>entire universe</i> as a result of a process. Any process that proceeds must satisfy this inequality, but not every process that satisfies the inequality will proceed quickly (or at all). For example, the formation of graphite from diamond is favoured by entropy, but the reaction rate is essentially zero at ambient temperature.			

Equation/Concept	Info About Equation/Concept		
$dS = \frac{dq_{\text{rev}}}{T}$	This is one definition of entropy. It is calculable only along a reversible path, but since once calculated the resulting ΔS applies to any path between the initial and final state.		
$\Delta S = \frac{q_{\rm rev}}{T}$	True for constant T, reversible heat exchange only. This equation is often used to measure changes in entropy by following a pseudo-reversible path.		
G = H - TS	Definition of Gibbs free energy.		
$\Delta G^{\circ} = -RT \ln(K)$	The standard Gibbs free energy. ΔG° is the Gibbs free energy change per mole of reaction of pure reactants to pure products at standard conditions (298 K, 1 atm, solutions at 1 M concentration, pure liquids and solids). K is the equilibrium constant for that reaction. $\Delta G^{\circ} < 0$ indicates products are favoured; $\Delta G^{\circ} > 0$ indicates reactants are favoured.		
$S = k_B \ln(\Omega)$	The definition of entropy for an isolated system. Ω is the number of possible microstates and k_B is the Boltzmann constant, which is included to convert the units.		
$\Delta G = \Delta H - T \Delta S$	The change in Gibbs free energy for a process done at constant temperature and pressure.		
$\Delta G = \Delta G^{\circ} + RT \ln Q$	This equation can be used to predict the direction of a reaction. Q is called the "reaction quotient", and is the same as the equilibrium constant K , but calculated when a reaction has not yet reached equilibrium. $\Delta G < 0$ indicates that there are too many reactants and the reaction will favour products. $\Delta G > 0$ indicates that the reaction will proceed to the reactants side.		
$K = \frac{[\text{Products}]}{[\text{Reactants}]}$	The definition of the equilibrium constant. Square brackets indicate concentrations, and the concentrations should be raised to the power of their stoichiometric coefficients.		
$\Delta G = -T\Delta S_{\text{Universe}}$	This equation applies at constant temperature and pressure for a system. It is the reason that the change in Gibbs free energy is a measure of spontaneity under those conditions. $\Delta G < 0$ corresponds to an increase in the entropy of the universe when T and P are held fixed.		
$\ln K = \frac{-\Delta H^{\circ}}{R} \frac{1}{T} + \frac{\Delta S^{\circ}}{R}$	Van't Hoff plot: plot $\ln K$ vs $\frac{1}{T}$. The slope $m = \frac{-\Delta H^{\circ}}{R}$ and the y-intercept $b = \frac{\Delta S^{\circ}}{R}$. Using these plots assumes that the standard change in enthalpy and standard change in entropy are independent of T . This is only true for small changes in T .		
S°	The standard molar entropy. This is the absolute entropy of 1 mole of a substance in its standard state, usually measured at 298 K. By the third law, this is non-zero.		
$S_{0 \mathrm{K, perfect crystal}} = 0$	The third law of thermodynamics. Here $S_{0 \text{ K,perfect crystal}}$ refers to the entropy of a pure, perfectly crystalline solid at 0 K. The third law of thermodynamics gives us a reference point to define the zero of entropy. Basically, it states that there's only one possible microstate for a perfect crystal with no internal energy. The entropy for a system with only one microstate is $S = k_B \ln(1) = 0$.		

Thermodynamics of Acids and Bases

Equation/Concept	Info About Equation/Concept		
$K_a = \frac{[H_3O^+][A^-]}{[AH]}$	Definition of the acid dissociation constant. This is simply the equilibrium constant for acid reactions. AH is an arbitrary acid and A^- is its conjugate base. Concentrations should all be raised to the power of their stoichiometric coefficients.		
$K_b = \frac{[OH^-][BH^+]}{[B]}$	Definition of the base dissociation constant. This is simply the equilibrium constant for base reactions. B is an arbitrary base and BH^+ is its conjugate acid. Concentrations should all be raised to the power of their stoichiometric coefficients.		
$K_W = K_a K_b = [H_3 O^+][OH^-]$	Water self-dissociation constant. This is equal to 10^{-14} at 25 °C. It is the equilibrium constant for pure water undergoing dissociation.		
$pH = -\log_{10}[H_3O^+]$	Definition of pH , pOH is defined similarly, where $[H_3O^+]$ is replaced by $[OH^-]$		
$pK_W = pK_a + pK_b = pH + pOH$	Equal to 14 at 25 °C. The water self-dissociation constants in terms of negative logs.		
$[AH]_{\mathrm{Buffer}} \approx 0.1[A^-] \text{ to } 10[A^-]$	For effective buffers, the acid and conjugate base should be within 10 times concentration of each other.		
$pH_{\mathrm{Buffer}} \approx pK_a \pm 1$	A more succinct way of stating that buffers must be within 10 times concentration of each other.		
$pH = pK_a + \log_{10}\left(\frac{[A^-]}{[HA]}\right)$	The Henderson-Hasselbalch equation. Here AH and A^- are an acid base conjugate pair. This equation is useful in determining how the pH changes as a result of addition of an acid or base to a buffer solution. The equation implicitly assumes that the change in concentrations of the acid and conjugate base is negligible. This is only appropriate for solutions that begin as buffers and stay as buffers upon addition of some small amount of another acid or base.		

Kinetics

Equation/Concept	Info About Equation/Concept	
Factors That Increase Reaction Rates	1. Increase the temperature to increase the average kinetic energy per particle to overcome the transition state energy barrier. 2. Add a catalyst which lowers the transition state energy barrier of the reaction. 3. Increase the concentration of reactants (except for zero order reactions) allowing more collisions to occur per unit time.	
ΔG^{\ddagger}	The Gibbs free energy of activation. This is the free energy change required to reach the transition state, measured as the difference in Gibbs free energy between the transition state and the reactants.	
Rate Determining Step For a multi-step reaction, the rate determining step is the step with Gibbs free energy barrier, ΔG^{\ddagger} .		
Elementary Reaction	A reaction that proceeds through only one step, has only one energy barrier, and one transition state. For a general elementary reaction $aA + bB \rightarrow cC$ the rate law is always $Rate = k[A]^a[B]^b$ where k is the rate constant.	

Equation/Concept	Info About Equation/Concept			
Zero Order Reactions	These types of elementary reactions do not depend on the concentration of the reactants. Such reactions usually depend on some catalyst, which is limiting the rate of the reaction. Zero order kinetics are an artifact of the reaction conditions, and will collapse to higher order kinetics when the concentration of reactants becomes low enough to limit the rate of reaction (instead of the catalyst).			
$Rate = -\frac{b}{a} \frac{d[A]}{dt} = k$	The zero order differential rate law for a general reaction $a[A] \to b[B]$. This is used to calculate the rate constant k . $[A]$ is the concentration of reactants and a is its stoichiometric coefficient. b is the stoichiometric coefficient of the products. The "Rate" here is always the rate of product formation. Often times this equation will be written without the factor of $\frac{b}{a}$, with $k' = \frac{a}{b}k$.			
$[A](t) = -kt + [A]_0$	This is the integrated rate law for a zero order reaction, note that the rate constant here is really $\frac{a}{b}k$. This equation will provide the concentration of the reactants as a function of time. Plots of $[A]$ vs t will be linear for zero order reactions, with a slope of $-k$ and a y -intercept of $[A]_0$.			
$t_{\frac{1}{2}} = \frac{[A]_0}{2k}$	The half-life of a zero order reaction. This depends on the initial concentration of reactants.			
First Order Reactions	These types of reactions depend on the concentration of only one reactant, which a stoichiometric coefficient of 1. For example, the elementary reaction $A \to bB$.			
$Rate = -b\frac{[A]}{dt} = k[A]$	The differential rate law for a first order reaction $A \to bB$. Often the rate constant k will include the factor of $\frac{1}{b}$.			
$[A](t) = [A]_0 e^{-kt}$	The integrated rate law for first order reactions. This can also be written as $\ln[A] = -kt + \ln[A]_0$. This equation can be used to determine the concentration of reactants as a function of time. Plots of $\ln[A]$ vs t will be linear for first order reactions, with a slope of $-k$ and a y -intercept of $\ln[A]_0$.			
$t_{rac{1}{2}} = rac{\ln 2}{k}$	The half-life for a first order reaction. This does not depend on the initial concentration of the reactants.			
Second Order Reactions	These types of reactions depend on the concentration of one reactant with a stoichiometric coefficient of 2, or on the concentration of two reactants. For example, the elementary reaction $2A \rightarrow bB$.			
Rate $= -\frac{b}{2} \frac{[A]}{dt} = k[A]^2$ The differential rate law for a first order reaction $A \to bB$. Of constant k will include the factor of $\frac{2}{b}$.				
$\frac{1}{[A](t)} = kt + \frac{1}{[A]_0}$	The integrated rate law for second order reactions. This equation can be used to determine the concentration of reactants as a function of time. Plots of $\frac{1}{[A]}$ vs t will be linear for second order reactions, with a slope of k and a y -intercept of $\frac{1}{[A]_0}$.			
$t_{rac{1}{2}}=rac{1}{[A]_0k}$	The half-life for a second order reaction. This depends on the initial conce tration of the reactants.			
$k = Ae^{-\frac{E_a}{RT}}$	The Arrhenius equation. This applies to any reaction and describes the effect of temperature on the rate of reaction. A is the pre-exponential factor, and is usually assumed to be temperature independent over small changes in temperature; it describes the frequency of collisions that occur with the correct orientation (different for each reaction). E_a is the activation energy per mole, and is the minimum energy required to overcome the energy barrier for reaction. RT is the average thermal energy per mole. This equation is sometimes written $\ln k = -\frac{E_a}{R} \frac{1}{T} + \ln A$ so that plots of $\ln k$ vs $\frac{1}{T}$ are linear with a slope of $-\frac{E_a}{R}$ and a y -intercept of $\ln A$.			
$k = \frac{k_B T}{h} e^{-\frac{\Delta G^{\ddagger}}{RT}}$	The Eyring equation. This applies only to elementary reactions, and can be used to determine the Gibbs free energy of activation. k_B is the Boltzmann constant, which is just $\frac{R}{N_A}$ where N_A is Avagadro's number. This reaction comes about by assuming that formation of the transition state is kinetically controlled (i.e. is in equilibrium) but the formation of products is thermodynamically controlled (i.e. is irreversible).			
$\ln\left(\frac{k}{T}\right) = -\frac{\Delta H^{\ddagger}}{R} \frac{1}{T} + \frac{\Delta S^{\ddagger}}{R} + \ln\left(\frac{k_B}{h}\right)$	The linear form of the Eyring equation. Plots of $\ln \left(\frac{k}{T}\right)$ vs $\frac{1}{T}$ will be linear. The slope will be $-\frac{\Delta H^{\ddagger}}{R}$ and the <i>y</i> -intercept will be $\frac{\Delta S^{\ddagger}}{R} + \ln \left(\frac{k_B}{h}\right)$.			
Reaction Intermediates	These are species that are generated in one step of a reaction and consumed in a subsequent step.			
$\frac{d[\text{intermediates}]}{dt} = 0$	This is the steady-state approximation. Useful for reactions where the rate determining step is the production of an intermediate. This often greatly simplifies the rate laws for multi-step reactions.			

Equation/Concept	Info About Equation/Concept		
$V_0 = \frac{V_{\text{max}}[S]}{K_m + [S]}$	The Michaelis-Menton equation. Describes the kinetics of the enzyme equilibrium reaction $E + S \xrightarrow[k_{-1}]{k_{-1}} ES \xrightarrow[k_{-1}]{k_{-1}} E + P$. Here, $K_M = \frac{[E][S]}{[ES]} = \frac{k_{-1} + k_{cat}}{k_1}$ is the equilibrium constant for the enzyme-substrate binding; V_0 is the rate of product formation; $V_{\text{max}} = k_{\text{cat}}[E]_{\text{total}}$ is the reaction rate when all available enzymes are saturated. Derivation of this equation involves the steady state approximation $\frac{d[ES]}{dt} = 0$. Notice that in the limit where the concentration of substrate is large compared to that of the enzyme, $K_M \ll [S]$ such that $K_M + [S] \approx [S]$ and $V_0 \approx V_{\text{max}}$. In the limit of very low substrate concentra-		
	tion, $K_M \gg [S]$ such that $K_M + [S] \approx K_M$ and the kinetics become first order with respect to $[S]$.		

IUPAC Organic Nomenclature Notes

• The highest priority functional group is given the suffix, all other lower priority functional groups are named with prefixes.

Naming esters with IUPAC:

- If the ester is the highest priority functional group, assign the alpha carbon (the carboxyl carbon) to number 1.
- Name the other side of the ester as if it is a substituent group. Place the name of this substituent group in front. Name the primary chain as normal, but add "-oate" to the end (eg: ethyl methanoate).

Naming ethers with IUPAC:

- Determine which side of the ether is the higher priority side: the shorter carbon chain becomes the "substituent" chain.
- Name the shorter chain as normal, and then the "-oxy" suffix to the end. Place this name in front followed by a space.
- The higher priority chain is then named as normal, with no suffix.

Naming sulfides with IUPAC:

- Determine which side of the sulfide is the higher priority side: the shorter carbon chain becomes the "substituent" chain.
- Name the shorter chain as normal, and then the "-thio" suffix to the end. Place this name in front followed by a space.
- The higher priority chain is then named as normal, with no suffix.

Naming acyl halides (acid halides) with IUPAC:

- These functional groups have higher priority than amides, but lower than carboxylic esters.
- Name it as if its a carboxylic acid, but remove the "-oic acid" suffix.
- Add the suffix "-oyl [hal]-ide" where "[hal]-ide" is replaced by the halogen in question (fluoride, chloride, bromide, iodide).

Naming Carboxylic anhydrides with IUPAC:

- These functional groups have higher priority than esters, but lower priority than carboxylic acids.
- Name each component carboxylic acid separately.
- Alphabetically arrange them, separated by spaces followed by the word "anhydride" (eg ethanoic propanoic anhydride).

Organic Nomenclature Guide

Functional Groups (in order of decreasing priority)

Functional group	Formula	Shorthand	Principle name	Substituent name
Carboxylic acids	OH OH	R-COOH	-oic acid	carboxy
Esters	R OR'	R-COOR'	-oate	(R)oxycarbonyl
Amides	R NR ₂	R-CONR ₂	-amide	amido
Nitriles	R'——N	R'-CN	-nitrile	cyano
Aldehydes	R H	R-CHO	-al	охо
Ketones	R' R'	R'-CO-R'	-one	охо
Alcohols	R'-OH		-ol	hydroxy
Thiols	R'-SH		-thiol	sulfanyl
Amines	R'-NR ₂		-amine	amino
Hydrocarbons	R'-H		-e	(R)yl
Ethers	R'-O-R'		-ether	(R)oxy
Sulfides	R'-S-R'		-sulfide	(R)ylsulfanyl
Halides	R'-X		-(hal)ide	(hal)o
	(X=F, Cl, Br, I)		(e.g. bromide)	(e.g. fluoro)

Note: R=saturated C/alkyl or H; R'=saturated C/alkyl

Prefixes 1 (parent/substituent length)

Therixes I (parent) substituent tength,		
R (# of carbons)	Prefix	
1	meth-	
2	eth-	
3	prop-	
4	but-	
5	pent-	
6	hex-	
7	hept-	
8	oct-	
9	non-	
10	dec-	

Infixes

Infix
-an-
-en-
-yn-

Prefixes 2 (multiple substituents/infixes)

# of repeated elements	Infix
2	di-
3	tri-
4	tetra-
5	penta-

IUPAC Nomenclature Rules

- 1. Identify the highest priority functional group and corresponding suffix.
- 2. Identify the parent chain (longest carbon chain that includes (if C-containing) or is directly attached to (if non-C-containing) the highest priority functional group) and corresponding prefix.
- 3. Identify the infix based on the nature of the C-C bonds in the parent chain (single/double/triple)
- 4. Number the parent chain from end to end, giving the lowest # to (1) the highest priority functional group; (2) multiple bonds; (3) substituents closest to the end; (4) groups that come first alphabetically
- 5. Identify the position and name of each substituent.
- 6. If two or more substituents are present on the same carbon atom, use the same number twice.
- 7. If a substituent group or infix appears more than once, use the prefixes: di, tri, tetra, penta, hexa etc.
- 8. List the substituents alphabetically at the beginning of the name (use the root names from rule 5, not the prefixes from rule 7 to alphabetize).
- 9. Numbers are separated from letters by dashes (-). Multiple numbers are separated by commas (,).

Organic Chemistry: Bonding Theories

- Formal Charge: (Valence electrons) (Lone Pair electrons) $\frac{1}{2}$ (Bonding electrons)
- Electronegativity: The strength that an atom has to attract electrons. More electronegative atoms have lower energy orbitals. Electronegativity increases → and ↑ accross the periodic table.
- Lewis Dot Structures: The simplest bonding theory. Non-bonding valence electrons are represented by dots (eg H·) and bonding electrons are represented by lines (eg H—H). Good for connectivity, but does not show molecular geometry.
- VSEPR Theory: Valence Shell Electron Pair Repulsion theory. Is able to predict molecular geometry by assuming substituents take on the lowest possible energy arrangement around a central atom. Does not predict molecular orbitals, and so cannot be used for reactivity.
- Valence Bond Theory (VBT): Simplified version of molecular orbital theory, only involving the valence electrons. Predicts hybridization and good approximations for orbitals.
- Molecular Orbital (MO) Theory: Generates fully accurate molecular orbitals which accurately predict reactivity. Becomes extremely complicated for large molecules.
- Frontier MO theory: Remixes the hybrid orbitals generated in VBT using the same method as MO theory.

Organic Chemistry: Reactivity and Acid-Base Chemistry

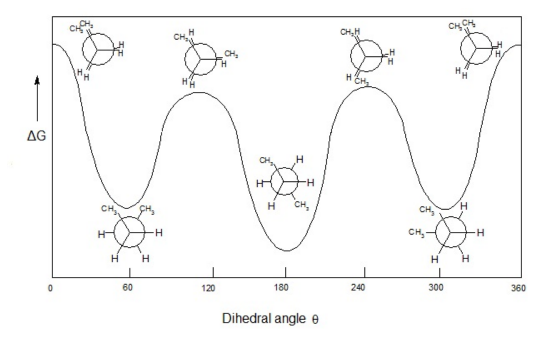
- Resonance: The concept in which two or more dot formulae for the same arrangement of atoms are necessary to fully describe the bonding in a molecule or atom. Basically an artifact of the inaccuracies inherent in our line-bond structures, which do not allow for partial bonding or electron delocalization to be shown from a single structure. Resonance allows movement of electrons through the π -system only, and always lowers the energy of the structure. Electrons flow form source (δ^- , -) to sink (δ^+ , +).
- Nucleophile: An electron-rich species that seeks positive charge to react with. Bases are often but not always good nucleophiles (counter example: extremely bulky bases are not good nucleophiles due to steric hindrance). Anions are usually good nucleophiles
- **Electrophile:** An electron-deficient species that seeks negative charge to react with. Positively or partially positively charged species make good electrophiles.
- $\mathbf{pK_a}$: Lower $\mathbf{pK_a}$ means stronger acid. Higher $\mathbf{pK_a}$ means stronger base. A better acid is better at donating protons, a better base is better at accepting protons.
- Strong Acids and Bases: These are unstable species, their conjugates will be much more stable and hence highly favoured at equilibrium.

• Factors That Effect Stability:

- 1. **Resonance:** Allows delocalization of charge, leading to stabilization. Resonance motifs: $\pi \sigma \pi$; $\pi \sigma$ -charge; $\pi \sigma$ -lone pair.
- 2. **Electronegativity:** Negative charge can be better stabilized by more electronegative species. Positive charge can be better stabilized by less electronegative (more electropositive) species. Comparison of electronegativity for stability is most important across a row on the periodic table.
- 3. **Polarizability:** More polarizable atoms can spread out charges better, and hence stabilize both negative and positive charges. Comparison of polarizability is most important down a column on the periodic table.
- 4. **Inductive Effects:** Nearby substituents can withdraw or donate electrons to help stabilize charge. This effect reduces with distance from the charge. More electronegative substituents are good electron withdrawing groups, helping to stabilize negative charge. Electron donating groups (alkyl groups, electron-rich groups) help stabilize positive charge.

• Electronegativity of Identical Atoms: Electronegativity increases with increasing s character, as s-orbitals are lower in energy. Electronegativity: $sp > sp^2 > sp^3 > sp^3d > etc$.

Organic Chemistry: Geometry and Stereochemistry

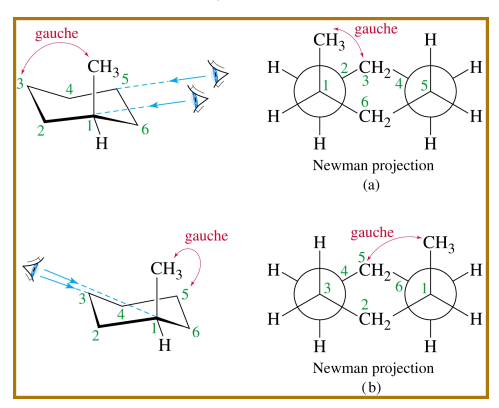


- Butane Conformations: For the dihedral angle between terminal methyl groups: Syn = 0°; Gauche = 60°; Eclipsed = 120°; Anti = 180°. Notice eclipsed is always higher energy than staggered. Energy is raised as the bulky substituents are forced closer together.
- Angle Strain: Always raises energy. If an sp³ hybridized carbon is force to take on a geometry with 90° bond angles as in cyclobutane, the energy of the system is raised.
- Cyclohexane Conformations: Two possible conformation types: chair or boat. Chair is always lower in energy. Chair flipping must pass through the boat conformation, and changes all equatorial substituents to axial and vice versa. The lowest energy chair conformation is the one that most reduces 1,3-diaxial interactions.
- 1,3-Diaxial Interactions: These are interactions of the three axial substituents in the chair conformation of cyclohexane on both faces. 1,3-Diaxial interactions are reduced when as few bulky substituents as possible occupy axial positions. Longer bonded groups like Br do not have strong 1,3-diaxial interactions.
- Cahn-Ingold-Prelog Priority Rules (CIP Rules): Assign highest priority to the highest atomic number atom at the first point of difference along substituents.
- **E/Z System:** Rank priority of substituents on both sides of the double bond separately using the CIP rules. If the highest priority substituents are cis (same side): Z configuration. If the highest priority substituents are trans (opposite side): E configuration.
- R/S System: Rank priority of substituents around an asymmetric carbon using the CIP rules. Point the lowest priority substituent backward and determine the direction from highest priority group to lowest $(1 \to 2 \to 3)$. If clockwise, then the group has R configuration. If anticlockwise, the group has S configuration.
- **E/Z** and **R/S** Nomenclature: Determine the underlying IUPAC root name as normal, then in brackets out in front of the name, list the positions of each stereocenter followed by its configuration in numerical order, separated by commas. (eg: (3S,5S)-5-bromo-3-chloro-5-fluoro-3-hexanamine).
- Stereoisomers: Isomers of identical constitution (connectivity) but differing in the arrangement of their atoms in space. Can refer to enantiomers, diastereomers, or conformers.

- Chirality: The property of any molecule or object of being non-superimposable with its own mirror image (e.g. hands are chiral). Chiral molecules cannot have a plane of symmetry or inversion center. Chiral molecules rotate plane-polarized light.
- Enantiomers: Two structures that that are non-superimposable mirror images of one another. Enantiomers must be chiral. Enantiomers have the (R/S) configuration at all asymmetric carbons flipped, but have the same (E/Z) configurations. Enantiomers have identical physical and chemical properties, except when interacting with other chiral compounds and in their interaction with plane-polarized light. Enantiomers rotate plane polarized light in opposite directions by the same amount.
- **Diastereomers:** Two structures that have the same connectivity (are stereoisomers) but have at least one difference in the (R/S) configurations at asymmetric carbons, but not at all asymmetric carbons.
- Meso Compounds: Compounds that would otherwise be chiral, but contain a plane of symmetry or inversion center which make them achiral. Meso compounds have asymmetric carbons arranged symmetrically, such that when the (R/S) configurations of all asymmetric carbons are flipped, an identical molecule is produced.

Newman Projections

Cyclohexane and Double Newman Projections



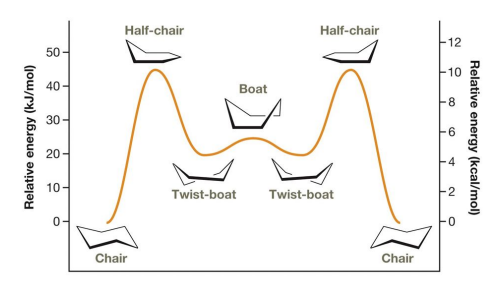
More Notes About Cyclohexane

- Conformations: There are a maximum of only two possible chair conformations for any substituted cyclohexane, and they can be switched through a chair flip. Chair flipping requires energy as it must pass through the boat conformation (see below). The boat conformation is always higher in energy than either of the chair conformations.
- 1,3-Diaxial Interactions: One of the two possible chair conformations of a substituted cyclohexane may be lower in energy, and therefore the preferred orientation. To determine which, look at the strength of the 1,3-diaxial interactions in each conformation.

• Factors That Affect 1,3-Diaxial Interactions:

- Bond length. Larger atoms like bromine or iodine have long bond lengths, and thus do not interact much through 1,3-diaxial interactions with hydrogen or methyl groups, but will interact with other groups that have long bonds!
- Degree of substitution. When comparing the strength of 1,3-diaxial interactions for alkyl groups, look at the number of non-hydrogen groups bound to carbons that are in the axial orientation. Alkyl groups that are highly substituted at the first carbon interact more strongly through 1,3-diaxial interactions and are thus less stable. Substitution at carbons beyond the first carbon in a chain are less important, but still may play a small role in determining relative energy.

Cyclohexane Chair Flip



Fischer Projections

		SN2	SN1
characteristics 人	mechanism	simultaneous bond making and breaking	bond breaking first then bond making
	key intermed or TS	crowded TS	carbocation intermed
	stereochemistry	inversion of stereochem	both inversion and retention of stereochem
	rate	= k [R-X] [Nuc ⁻]	= k [R-X]
ons	R-X	Methyl > 1° > 2°	3° > 2°
hat eff f reacti	Nuc	must be good Nuc	can be weak (is often the solvent)
factors that effect the rate of reactions	LG	good LG is better	must be good LG
	solvent	more polar is better (aprotic is better)	more polar is better (protic is better)

- \bullet More substituted electrophiles form more stable carbocations, and hence react more quickly through S_N1 reactions. More substituted electrophiles are less likely to react through S_N2 due to steric hindrance.
- In S_N1 reactions, the rate limiting step is the carbocation formation. Hence, the strength of the nucleophile is not very important.
- In S_N2 reactions, the rate limiting step is the formation of a concerted transition state. Stronger nucleophiles lower the energy of the transition state and hence improve reaction rates.
- Weak bases make good leaving groups for both types of substitution reaction. Weak bases are more stable as free species than strong bases, and hence do not raise the energy of the system too much upon dissociation.
- S_N1 can be catalyzed by protic solvents or acids in solution. A proton can be added to a bad leaving group, turning it into a better leaving group. Polar solvents also help to stabilize the carbocation.
- S_N2 reactions favor polar aprotic solvents, as these allow for stronger (charged) nucleophiles. Protic solvents will lose their proton to strong nucleophiles, making them weak. Polar protic solvents also tend to hydrogen bond with the nucleophile, creating a solvent shell around it and hindering its reactivity.
- Aprotic solvents do not solvate negatively charged nucleophiles that well. Therefore, negatively charged nucleophiles are more reactive in aprotic solvents.
- More polarizable nucleophiles react faster because they are not solvated as readily. Their charge is spread out more, so that Van der Waals bonds to solvent molecules are weaker. Small nucleophiles are highly solvated, and tend to react slowly in S_N2 reactions. Relative polarizability for nucleophiles is most important along the same columns of the periodic table.
- Adjacent π -systems help stabilize electrophiles in both S_N1 and S_N2 reactions through the possibility of resonance. In S_N1 reactions the carbocation is stabilized by resonance, whereas in S_N2 the concerted transition state is stabilized by resonance.
- S_N2 reactions only occur on sp^3 hybridized carbons.

- \bullet The rate determining step of S_N2 reactions is a bi-molecular step. Hence it has second order kinetics and the rate of reaction depends on the concentrations of both the nucleophile and electrophile.
- \bullet The rate determining step of S_N1 reactions is unimolecular. Hence it has first order kinetics and the rate of reaction only depends on the concentration of the electrophile.

Physical Constants and Periodic Trends (for Reference)

Constant	Value and Units		
Atomic mass unit (u)	$1 \mathrm{u} = 1.6605389 \times 10^{-24}\mathrm{g}$		
	$1\mathrm{g} = 6.022142 \times 10^{23}\mathrm{u}$		
Avogadro's number (N_A)	$N_A = 6.022142 \times 10^{23} \mathrm{mol}^{-1}$		
Boltzmann's constant (k_B)	$k_B = 1.38064852 \times 10^{-23}\mathrm{JK^{-1}}$		
	$k_B = R/N_A$		
Gas constant (R)	$R = 8.3144598 \mathrm{J}\mathrm{K}^{-1}\mathrm{mol}^{-1}$		
	$= 8.3144598 \times 10^{-3} \mathrm{kJ} \mathrm{K}^{-1} \mathrm{mol}^{-1}$		
	$= 8.3144598 \mathrm{kg} \mathrm{m}^2 \mathrm{s}^{-2} \mathrm{K}^{-1} \mathrm{mol}^{-1}$		
	$= 8.3144598 \mathrm{m}^3 \mathrm{Pa} \mathrm{K}^{-1} \mathrm{mol}^{-1}$		
	$= 0.082057338\mathrm{LatmK^{-1}mol^{-1}}$		
	$= 1.9872036 \times 10^{-3} \mathrm{kcal} \mathrm{K}^{-1} \mathrm{mol}^{-1}$		
	$= 8.2057338 \times 10^{-5} \mathrm{m}^{3} \mathrm{atm} \mathrm{K}^{-1} \mathrm{mol}^{-1}$		
	$=k_BN_A$		
Mass of electron (m_e)	$m_e = 5.485799 \times 10^{-4} \mathrm{u}$		
	$= 9.109383 \times 10^{-28} \mathrm{g}$		
Mass of proton (m_p)	$m_p = 1.0072765\mathrm{u}$		
	$= 1.6726217 \times 10^{-24} \mathrm{g}$		
Mass of neutron (m_n)	$m_n = 1.0086649\mathrm{u}$		
	$= 1.6749273\times10^{-24}\mathrm{g}$		
Planck constant (h)	$h = 6.626069 \times 10^{-34}\mathrm{Js}$		
Reduced Planck constant (ħ)	$\hbar = h/2\pi = 1.05457266 \times 10^{-34}\mathrm{Js}$		
Speed of light (c)	$c = 2.99792458 \times 10^8 \mathrm{ms^{-1}}$		

