AP QUICK REVIEW

STP = 0° C and 1 atm at STP 1mole = 22.4 L K = $^{\circ}$ C + 273

$$D = \frac{m}{V}$$

Density equation

Average Atomic Mass = $\sum \left(\frac{\% \text{ abundance}}{100}\right)$ (mass of isotope)

$$= \left(\frac{\text{\% abund. # 1}}{100}\right) \text{(mass of isotope 1)} + \left(\frac{\text{\% abund. # 2}}{100}\right) \text{(mass of isotope 2)} \dots$$

Solution Stoichiometry:

Determining Molarity:	Molarity = Moles Volume (in liters)
Molarity and Volume:	$M_1V_1 = M_2V_2$
	USE FOR DILUTION PROBLEMS

KEY SOLUBILITY RULES

- 1. Salts of ammonium (NH₄⁺) and Group I are always soluble.
- 2. All Cl⁻, Br⁻, I⁻ are soluble except with Ag⁺, Hg₂²⁺, and Pb²⁺ which are insoluble.
- 3. ClO₃, NO₃, and CH₃COO are soluble. 4. SO₄-² are soluble except with; Sr²⁺, Ba²⁺, Hg₂²⁺, Pb²⁺, which are insoluble≈(Ca²⁺)

Double displacement. Precipitation, neutralization, gas forming, H₂CO₃ in water = H₂O & CO₂

Single displacement or redox replacement: (metals displace metals and nonmetals displace nonmetals)

Combination or synthesis = two reactants result in a single product

- Metal oxide + water → metallic hydroxide (base)
- Nonmetal oxide + water → nonbinary acid
- Metal oxide + nonmetal oxide → nonbinary salt

Decomposition = one reactant becomes several products

- Metallic hydroxide → metal oxide + water
- Nonbinary acid → nonmetal oxide + water
- Nonbinary salt → metal oxide + nonmetal oxide
- Metallic chlorates → metallic chlorides + oxygen
- Electrolysis decompose compound into elements (water in dilute acids or solutions of dilute acids)
- Hydrogen peroxide → water + oxygen
- Metallic carbonates → metal oxides + carbon dioxide
- Ammonium carbonate \rightarrow ammonia, water and carbon dioxide.

Hydrolysis = compound reacting with water.

Watch for soluble salts that contain anions of weak acid the anion is a conjugate base and cations of weak bases that are conjugate acids.

Reactions of coordinate compounds and complex

- Complex formation by adding excess source of ligand to transitional metal of highly charged metal ion such as Al^{3+} Al =4 ligands others 2X ox #
- Breakup of complex by adding an acid \rightarrow metal ion and the species formed when hydrogen from the acid reacts with the ligand

Lewis acid base reactions = formation of coordinate covalent bond

Redox = change in oxidation state= a reaction between an oxidizer and a reducer.

Recognized:

- 1. Familiarization with important oxidizers and reducers
- 2. "added acid" or "acidified"
- 3. an oxidizer reacts with a reducer of the same element to produce the element at intermediate oxidation state

OXIDIZE	RS	RE	DUCERS
MnO ₄ in acid	Mn ²⁺	Halide ions	Halogens
MnO ₂ in acid	Mn ²⁺	Metal element	Metal ion
MnO ₄ in neutral or basic solution	MnO ₂	Sulfite	Sulfate
Cr ₂ O ₇ ²⁻ in acid	Cr ³⁺	Nitrite	Nitrite
HNO ₃ concentrated	NO ₂	Halogen element in dilute basic solution	Hypohalite ion an halide ion
HNO ₃ dilute	NO	Halogen element in concentrated basic solution	Halite ion
H ₂ SO ₄ , hot, concentrated	SO ₂	Metal ous ion	Metallic ion
Metal ic ions	Metal ous ions	H ₂ O ₂	O ₂
Halogens diatomic	Halide ions	C ₂ O ₄ ²⁻	CO ₂
Na ₂ O ₂	NaOH		
HCIO ₄	Cl		
H ₂ O ₂	H ₂ O		and II O (nearlies in formains the socials

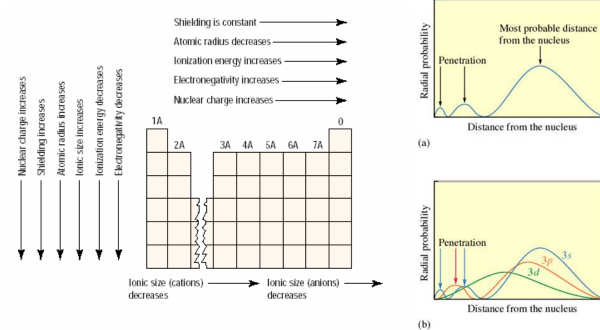
Combustion - complete combustion of hydrocarbons results in CO₂ and H₂O. (results in forming the oxide of the elements of the compound)

Gas Laws:

Kinetic molecular Theory: particles in constant random motion particles do not have volume	Ideal: particles 0 volume no attraction, deviate from ideal at low temperature and high pressure Van der Waal's Equation
particles do not attract T \tag{with \tau KE = \frac{1}{2}m \times v^2}	$[P_{obs} + a (\frac{n}{v})^2] \times (V - nb) = nRT$
	a corrects for attraction & b corrects for volume molecules that deviate from ideal gas behavior: large volume and large attraction between molecules ideal gases: molecules: have zero volume and do not attract or repel each other
The Ideal Gas Law: (P = pressure in atm, V = volume in liters, n = moles, R = 0.08201 L*atm/mol*K, T = temperature in Kelvin's)	PV = nRT
Combine Gas Law	DV DV
	$\frac{P_1 V_1}{T_1} = \frac{P_2 V_2}{T_2}$
Mole fraction; $\chi = \frac{\text{moles A}}{\text{total moles}}$	$P_A = P_{total} \chi_A$ $P_{total} = P_A + P_B + P_c + \dots$
Rate of Effusion:	$\frac{\text{Rate of Effusion for Gas 1}}{\text{Rate of Effusion for Gas 2}} = \frac{\sqrt{M_2}}{\sqrt{M_1}}$
Boyle's Law:	PV = K
Charles Law:	V/T = K

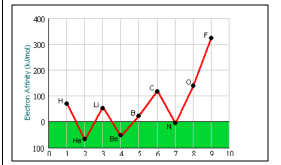
Root Mean Square: (R = 8.3145 J/ K * mol, M = mass of a mole of the gas in kg)	$u_{rms} = \sqrt{\frac{3RT}{M}}$
Atomic Theory:	
(n = integer, h = Planck's constant = 6.626 x 10 ⁻³⁴ J*s)	$\Delta E = nh\nu$
Energy per photon:	_ hc
$(c = 2.99979 \times 10^8 \text{ m/s})V =$	Energy _{photon} = $\frac{hc}{\lambda}$
de Broglie's equation:	$\lambda = \frac{h}{m v}$
Beer's Law: where A = absorbance, a = wavelength-dependent absorptivity coefficient, b = path length, and c = analyte concentration.	A = abc
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Parken rules (a, rules) $\Delta E = -2.178 \times 10^{-18} \mathrm{J} \left(\frac{1}{n_{\mathrm{final}}}^2 - \frac{1}{n_{\mathrm{initial}}}^2\right)$
Principle Quantum Number (n) has	Angular Momentum Quantum Number (t) has integral
integral values 1, 2, 3, The principle	values {0 to n-1}. This quantum number gives us the
quantum number is related to size of the	shape of the probability pattern and each shape is
electron cloud, n = 1, 2, 3,	referred to as a sublevel $\ell = \{0 \text{ to (n-1)}\}\$ the type of
determines the energy of the e	orbital (subshell) $0 = s$, $1 = p$, $2 = d$, $3 = f$
Magnetic Quantum Number (m _t) has	Magnetic Spin Quantum Number (m _s) differentiates the two electrons that can exist in an orbital and has the
integral values $\{-\ell \text{ to } +\ell\}$. The Magnetic	value of +1/2 and -1/2. s or $m_s = +\frac{1}{2}$ or -$\frac{1}{2}$ the
quantum number refers to the orientation of each sublevel. m or m _t =	"spin" of the electron
$\{-\ell \text{ to } + \ell\}$ which orientation of the	Diamagnetic = paired e
orbital (x, y, z for p orbitals)	Paramagnetic = unpaired e
	hode ray experiment / electron and proton,
•	- energy levels, Schrödinger- quantum
	ciple and deBroglie-dual nature) Milliken- charge of
	- atomic number, Pauli's- exclusion principle,
Mandalagy pariodic table	and the state of t

Mendeleev- periodic table



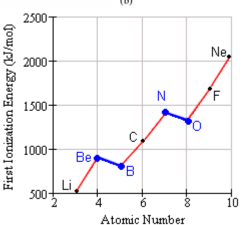
-Stating the trend is not an explanation:

- -ease of removing e: f > d > p > s because of penetration -s e spends a greater percent of time closer to nucleus than p and p more than
- -↑n or radius of atom ↓ nuclear attraction for valence e
- -↑Z same n ↑ nuclear attraction for valence e⁻ -↑shielding ↓ nuclear attraction for valence e (shielding is only a factor when comparing e from different n's)



Electron Affinity Exceptions

WHY? - $Xs^2 < 0$: <u>diamagnetic</u> atom with all paired electrons. Electron-electron repulsion. Xp³ < 0: atom with 3 unpaired p-orbital electrons each occupying its own subshell. The fourth electron will be the second electron in an orbital and will experience electron-electron repulsion. Xp⁶ < 0: next electron in energy level further from nucleus



Ionization energy Exceptions

1)
$$Xs^2 > Xp^1$$
 e.g. $^4Be > ^5B$

WHY? - . (s shields p ∴ p< further from nucleus than s) (s ¯e spends more time closer to nucleus than e in p)

Therefore, it requires less energy to remove the first electron in a p orbital than to remove one from a filled s orbital. Or :The energy of an electron in an Xp orbital is greater than the energy of an electron in its respective Xs orbital. Therefore, it requires less energy to remove the first electron in a p orbital than it is to remove one from a filled s orbital (further from nucleus)

2) $\rm Xp^3 > Xp^4~e.g.^7N > ^8O$ WHY? - After the separate degenerate orbitals have been filled with single electrons, the fourth electron must be paired. The electronelectron repulsion makes it easier to remove the outermost, paired

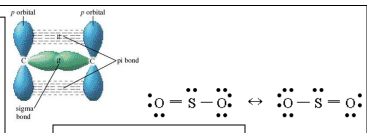
Subsequent ionization energies increase greatly once an ion has reached the state like that of a noble gas (what not why!!). Why is --. Fewer core electrons, closer to nucleus, greater Zeff → Takes more energy to remove.

	# of	# of Non-	Ma	lecular	Bond	
lybdridization	σ Bonds	Bonding Pairs		Shape	Angles	Example
sp	2	o	$\overline{}$	Linear	180°	BeH_2,CO_2
sp2	3	o	Υ	Trigonal planar	120°	SO ₃ , BF ₃
sp ²	2	1	\checkmark	Angular	<120°	SO ₂ , O ₃
sp ³	4	o	\downarrow	Tetrahedral	109.5°	CH ₄ , CF ₄ , SO ₄ ²⁻
sp ³	3	1	<i>~</i>	Trigonal pyramidal	<109.5°	NH ₃ , PF ₃ , AsCl ₃
sp3	2	2	<i>~</i> ~	Angular	<109.5°	H ₂ O, H ₂ S, SF ₂
sp ³ d	5	o	\Rightarrow	Trigonal bipyramidal	120°, 90°	PF ₅ , PCl ₅ , AsF ₅
sp ³ d	4	1	\Rightarrow	Sawhorse (irregular tetrahedron)	<120°, <90°	SF ₄
sp ³ d	3	2		T-shaped	<90°	CIF ₃
sp3d	2	3		Linear	180°	$XeF_2,I_{3^{\tilde{-}}},IF_2$
sp ³ d ²	6	o	*	Octahedron	90°	SF ₆ , PF ₆ -, SiF ₆ ² -
sp ³ d ²	5	1	$\stackrel{\cdot}{\times}$	Square pyramidal	<90°	IF ₅ , BrF ₅

Metallic bond: delocalized e

Ionic: transfer of e

Nonpolar Covalent: equal sharing of $\bar{}$ e Polar Covalent: unequal sharing of $\bar{}$ e σ overlap s/s, s/p, p/p - π overlap p/p single σ , double σ - π , triple σ - π - π single< double<triple: strength single>double> triple>: length



Resonance occurs when you have a *combination* of a multiple and single

Hybribs explains why bonds in molecules with different atomic orbitals behave as identical bonds ie. CH₄

Resonances explains why bonds in molecules with a combination of single, double or triple bonds behave identical ie. C₆H₆

0→0.4 NONpolar covalent 0.41→1.67 POLAR covalent above 1.67—IONIC

Calculating lattice energy (k is proportionality constant, Q is charge of ions, r = shortest distance between centers of the cations and anions)	Lattice energy = $k\left(\frac{Q_1Q_2}{r}\right)$ strength of ionic bond
	$\Delta H_{rxn} = \sum$ bond energy of reactants - \sum bond energy of products

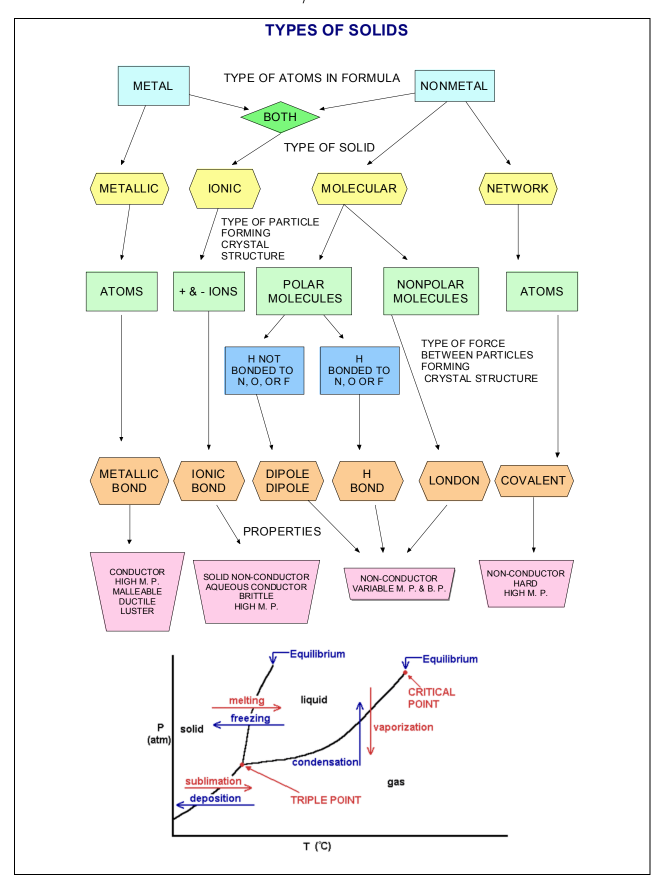
bond broken bonds formed

Bond Order: # of shared pairs linking X & Y ÷ # of X-Y links

Formal Charge = number of valence e^- – (# of lone e^-) – $\frac{1}{2}$ (# of bonding e-'s)

Intermolecular Forces:

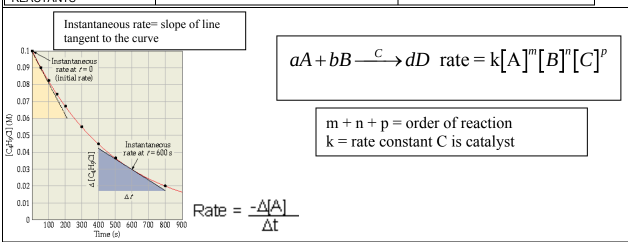
TYPE OF	CHARACTERISTICS OF	TYPE OF	MAGNITUDE	
INTERACTION	INTERACTION	SUBSTANCE	RANGE KJ/MOLE	
DIPOLE-DIPOLE	Positive end of a permanent dipole aligns itself with negative end of a permanent dipole on another molecule. {dipole moment}	TWO POLAR MOLECULAE	5-25	
HYDROGEN BOND	Special type of dipole-dipole. Is strong due to the lack of shielding around a hydrogen nucleus (when hydrogen is bonded to a highly electronegative atom.) {dipole moment}	TWO POLAR MOLECULES WITH HYDROGEN BONDED TO A N, F, OR O ATOM	5-25	
INDUCED DIPOLE- INDUCED DIPOLE (LONDON DISPERSION)	Attractive forces between molecules resulting from momentary mutual distortion (induced dipole) of electron clouds. (temporary dipoles) Increase in strength with increase molecular weight (electron cloud size) called polarizability.{induced dipole,-induced dipole}	TWO NONPOLAR MOLECULES	0.05 -40	



Solutions:	
Molality:	molality = \frac{moles of solute}{kg solvent}
Psoln: vapor pressure of the solution Xsolvent: mole fraction of solvent Psolvent: vapor pressure of pure solvent	Raoult's Law $P_{\text{soln}} = \chi_{\text{solvent}} P_{\text{solvent}}$
Mass Percent:	Mass percent = \frac{mass of solute}{mass of solution} \times 100
Osmotic pressure formula for electrolytes π: osmotic pressure M: molarity R: 0.0821 L*atm/K*mol	$\pi = iMRT$
Henry's Law P: partial pressure of gaseous solute C: concentration of dissolved gas k: constant	P = kC
Boiling point elevation m: molality Kb: constant	$\Delta \mathbf{T}_{\mathbf{b}} = i\mathbf{K}_{\mathbf{b}} \cdot \mathbf{m}$
Freezing Point Depression m: molality Kf: constant	$\Delta T_f = i K_f \cdot m$
van't Hoff factor	i = moles of particles in solution moles of solute dissolved

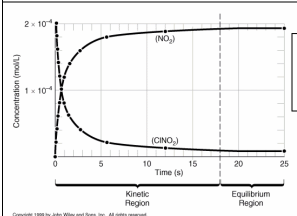
Chemical Kinetics:

FACTOR	AFFECT ON RATE	COLLISION THEORY
CONCENTRATION OF REACTANTS	increasing concentration increases rate (only reactants in the rate determining step affect the rate)	more molecules more collisions
TEMPERATURE	increasing temperature increases rate (approximately every 10°C increase in temperature doubles rate)	molecules have more kinetic energy ~ collide more often and harder ~ collisions more frequent and a greater % of the collisions are effective
CATALYST:	increase the rate of a reaction but not consumed during the reaction	lowers the activation energy therefore a greater % of collisions are effective
SURFACE AREA OF REACTANTS	increase surface area increases rate	more collisions



General Rate Law rate and [], Integrated rate law time and []					
Order in [A]	Rate Law	Integrated Form $y = mx + b$	Straight Line Plot	Half-Life t _{1/2}	
zeroth order (n=0)	$rate = k [A]^{0} = k$	$[A]_{t} = -k t + [A]_{0}$	$[A]_{t} \text{ vs } t$ $(slope = -k)$	$t_{1/2} = \frac{[A]_0}{2k}$	$\stackrel{[A]_0}{=}$ $\stackrel{I_{\text{fine}}}{=}$
first order (n=1)	$rate = k [A]^1$	$\ln[A]_t = -k t + \ln[A]_0$	$ln[A]_t vs t$ $(slope = - k)$	$t_{1/2} = \frac{\ln 2}{k} = \frac{0.693}{k}$	In[A]
second order (n=2)	$rate = k [A]^2$	$\frac{1}{[A]_t} = kt + \frac{1}{[A]_0}$	$\frac{1}{[A]_0} \text{ vs } t$ $(\text{slope} = k)$	$t_{1/2} = \frac{1}{k[A]_0}$	slope = k
Arrhenius Equation the whole e^(stuff) equation represents the fraction of collisions with sufficient energy to produce a rxn		k = Ae ^(-Ea/RT)	Ink slope = $\frac{1}{T}$	= <mark>-E_a</mark> R	
Derived Arrhenius Equation $\ln k = \frac{-E_a}{R} \left(\frac{1}{T}\right) + \ln A$				$\frac{C_a}{T} \left(\frac{1}{T}\right) + \ln(A) y = \frac{1}{T_1} \left(\frac{1}{T_1} - \frac{1}{T_2}\right) \text{ [plug in } 6$	mx + b [graphical method]
Reactants Reaction coordinate Reaction coordinate Energy Reaction coordinate Reaction coordinate Energy Endothermic					
				-	

Chemical Equilibrium:



To determine direction of reaction calculate Q: If

Q > K reaction \leftarrow If Q < K reaction \rightarrow

Calculating Equilibrium Constant:
(leave out solids and liquids)

$$zA + yB \xrightarrow{\bullet \bullet} xC + vD$$

$$K = \frac{[C]^{\bullet}[D]^{\bullet}}{[A]^{2}[B]^{y}}$$

 Δn = coefficients of gaseous products coefficients gaseous reactants

$$K_p = K_c(RT)^{\Delta n}$$

Changing the Coefficients Will Change K:

- Multiplying coefficients by a #, $K_{new} = K_{old}^{\#}$
- Reversing the equation, $K_{new} = 1/K_{old}$
- Adding several equations to get a net equation $K_{\text{new}} = (K_{\text{old}})(K_{\text{old}})...$

Acids and Bases:

pH = -log [H₃O⁺]
pOH = -log [OH⁻]

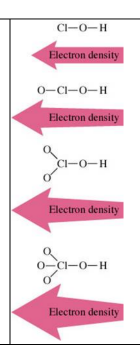
$$K_w = 1 \times 10^{-14} = [H_3O^+][OH^-]$$

 $14 = pH + pOH$
 $[H_3O^+] = 10^{-(pH)}$
 $[OH^-] = 10^{-(pOH)}$

Equilibrium Expressions:

HA + H₂O
$$\longleftrightarrow$$
 H₃O⁺ + A⁻ $K_a = \frac{\left[H_3O^+\right]\left[A^-\right]}{\left[HA\right]}$ < 1

$$B + H_2O \longleftrightarrow HB^+ + OH_- \quad K_b = \frac{\left[OH^-\right]\left[HB^+\right]}{\left[B\right]} \quad <1$$



pH of Salts:

- 1. Strong acid + strong base = neutral salt 3. Weak acid + strong base = basic salt
- Strong acid + weak base = acidic salt 4. Weak acid + weak base = ? (must look at K values to decide)

CB's (anions of weak acids) - react like nonhydroxide bases in water: NaF strong base weak acid salt anion is a CB: F⁻ + H₂O ←→ HF + OH⁻ CA's (cations of weak bases) - react like acids in water: NH₄Cl weak base strong acid salt cation is a CB: NH₄⁺ + H₂O ←→ H₃O⁺ + NH₃	$K_{w} = [OH^{-}][H^{+}] = 10^{-14} @25^{\circ}C$ $K_{w} = K_{a} \cdot K_{b}$ $pK_{w} = 14$
$HC1 + NH_3 \Longrightarrow NH_4^+ + C1^ CA$	amphoteric $ \begin{cases} $
% Dissociation	% dissociation = amount dissociated x 100

Summary of Buffer Reactions:

Weak Acid and Conjugate Base Buffer:

Buffer Reactions:

 $HA + H_2O \longleftrightarrow H_3O^+ + A^-$ (weak acid controls pH)

 $XA \rightarrow X^{+} + A^{-}$ (conjugate base A) **Neutralization Reactions:**

Strong acid added to buffer HCI + A

→ HA + Cl⁻

acid + base

Strong base added to buffer NaOH + HA→ Na+ + A+ H,O

base + acid

Weak Base and Conjugate Acid Buffer

Buffer Reactions:

 $M + H_2O \longleftrightarrow MH^+ + OH^-$ (weak base controls pH) $MHX \rightarrow MH^{+} + X^{-}$ (conjugate acid MH^{+})

Neutralization Reactions:

Strong acid added to buffer

 $HCI + M \rightarrow MH^{+} + CI^{-}$ acid + base

Strong base added to buffer

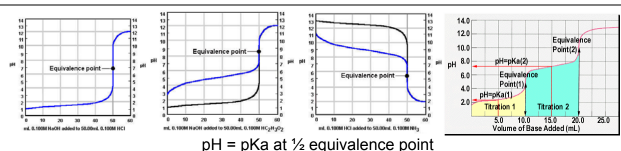
NaOH + $MH^{+} \rightarrow M + Na^{+} + H_{0}O$

base + acid

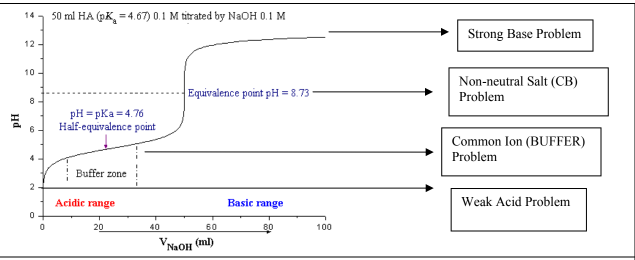
 igtimes RICE TABLE AFTER STRONG ACID OR BASE IS ADDED TO A BUFFER: CHECK igtimes

When a strong acid is added to buffer subtract M of strong acid from the base and add to the acid When strong base is added subtract the M of strong base from the acid and add to the base:

TITRATION



Titration: If an ACID & BASE are reacting write a neutralization reaction: (both strong) do Stoichiometry to find molarity of substance of interest; AT equivalence pt 7/ (one of them weak) SCAM TABLE then RICE TABLE, **AT** equivalence pt a **non neutral** salt will be CA or CB problem



Solubility Equilibrium:

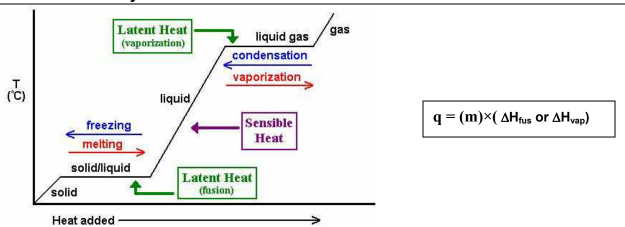
$$\operatorname{CaF}_{2(s)} \xrightarrow{\operatorname{H}_2 \circ} \operatorname{Ca}^{2+}(aq) + 2\operatorname{F}^{-}(aq) \qquad \qquad \operatorname{K}_{sp} = [\operatorname{Ca}^{2+}][\operatorname{F}^{-}]^2$$

 $\mathbf{Q}_{\mathrm{sp}} < \mathbf{K}_{\mathrm{sp}}$ - This means there are not enough ions in the solution. In order to return to equilibrium, more of the solid salt must dissociate into its ions.

 $Q_{sp} = K_{sp}$ - This means that the system is at equilibrium.

 $Q_{sp} > K_{sp}$ - This means that there are too many ions in the solution. In order to return to equilibrium, the excess ions will precipitate to form more solid.





standard conditions 1 atm and 25°C	
C = heat capacity	C _p also called specific heat m = mass of solution or total mass
$C_p = \frac{q}{\Delta t}$ or $C_p = \frac{q}{\Delta t \times m}$	m = mass of solution or total mass
ΔH = q coffee cup calorimetry	$q = mc\Delta T$
ΔH_f° for element = 0 \rightarrow Unit KJ/mole	$\Delta H^{o} = \sum n \Delta H^{o}_{f(products)} - \sum n \Delta H^{o}_{f(reactants)}$
ΔS° is <u>not</u> 0 \rightarrow Units in J//mole K	$\Delta S^{o} = \sum n \Delta S^{o}_{(products)} - \sum n \Delta S^{o}_{(reactants)}$
ΔG_f° for element = 0 \rightarrow Unit KJ/mole	$\Delta G^{\circ} = \sum n \Delta G^{\circ}_{f(products)} - \sum n \Delta G^{\circ}_{f(reactants)}$

$\Delta S_{surroundings} = \frac{-\Delta H}{\Delta T}$	$\Delta G^{\circ} = \Delta H^{\circ} - T \Delta S^{\circ}$	<u></u> _∆G°= rxn spontaneous
	$\Delta G = \Delta G^{\circ} + RT \ln Q$ (can use ΔG or ΔG°)	$\Delta G^{\circ} = -RT \ln K$
	(can use ΔG Of ΔG)	

TABLE	TABLE 20.1 Criteria for Spontaneous Change: $\Delta G = \Delta H - T \Delta S$				
Case	ΔH	ΔS	ΔG	Result	Example
1	_	+	_	spontaneous at all temp	$2 N_2O(g) \longrightarrow 2 N_2(g) + O_2(g)$
2	_	_	$\left\{egin{array}{c} - \\ + \end{array}\right.$	spontaneous at low temp nonspontaneous at high temp	$H_2O(1) \longrightarrow H_2O(s)$
3	+	+	{+ -	nonspontaneous at low temp spontaneous at high temp	$2 \text{ NH}_3(g) \longrightarrow N_2(g) + 3 \text{ H}_2(g)$
4	+	_	+	nonspontaneous at all temp	$3 O_2(g) \longrightarrow 2 O_3(g)$

	REACTION STATE	EQUILIBRIUM CONSTANT "K"	GIBBS FREE ENERGY "ΔG"	CELL POTENTIAL "ΔE"
	SPONTANEOUS	greater than one	negative	positive
	EQUILIBRIUM	zero	Zero	Zero
N	ON-SPONTANEOUS	Less than one	positive	negative

Electrochemistry:

LEO goes GER

lose electrons oxidize gain electrons reduction



the reactant reduced is the oxidizing agent and visa versa

$$1 \text{ V} = \frac{1 \text{ J}}{1 \text{ C}} = \frac{\text{(joule)}}{\text{(coulomb)}}$$

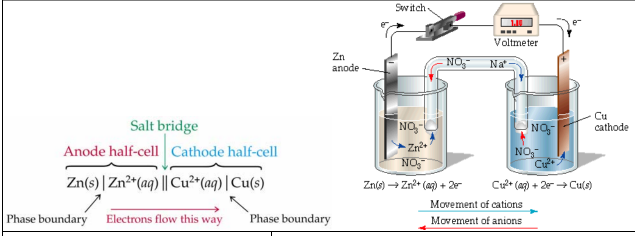
 $E_{cell\,is:}$ + spon rxn, – nonspon rxn, 0 rxn at eq

E°cell = E°ox + E°red

Largest E^o_{red} is reduced ∴ best oxidizing agent

Anode	Cathode
 Oxidation 	Reduction
 Anions to it from SB 	 Cations flow to it from SB
 - in voltaic cells 	+ in voltaic cells
 + in electrolytic cells 	- in electrolytic cells
 Electrons flow away from it 	Electrons flow towards it
 Cell notation Cl⁻ / Cl₂ 	 Cell notation Cu²⁺ → Cu_(s)





Standard conditions 1 atm 25°C		W,	max =	-qE	Δ	G =	-n	F E
							-	

Nernst Equation

F = faraday's constant n = number of electrons

At equilibrium
$$E_{cell} = 0$$

n = number of moles of electrons transferred

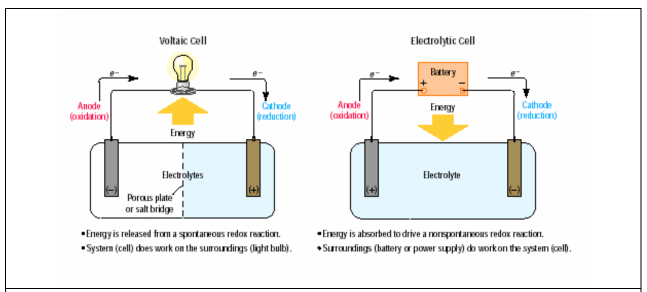
$$E_{\text{cell}} = E_{\text{cell}}^{0} - \frac{RT}{nF} \ln Q = E_{\text{cell}}^{0} - \frac{0.0592}{n} \log Q \text{ at } 25^{\circ}\text{C}$$

$$\log K = \frac{nE^{0}}{0.0592}$$

Electrolysis

Electrolysis involves forcing a current through a cell to produce a chemical change for which the cell potential is negative ~ electrical work causes an otherwise non-spontaneous chemical reaction to occur.

If more than one reactant is available for reduction, the one with the *highest* E°_{red} potential is most likely to be reduced. Metal ions or water can be reduced. When electrolysis occurs in aqueous solutions, if the metal has a reduction potential smaller than *0.8, then only water is reduced because water has the larger E°_{red}. Negative ions will be oxidized. If there are mixtures of metal ions, the metals will be reduced in order of largest E°_{red} to smallest E°_{red}.



<u>Faraday's Constant</u> - The charge on a mole of electrons: F = 96,500 C coulomb = (amp) (sec)

ORGANIC—carbon containing compounds

- -ane—CnH_{2n+2}; has all single bonds and is therefore saturated with hydrogens
- –ene--CnH_{2n}; has double bonds and is therefore unsaturated [dienes have 2 double bonds, trienes have 3, etc.; both are polyunsaturated]
- -yne--CnH_{2n-2}; have a triple bond and are also unsaturated
- ISOMERS—same structural formula yet different arrangement of atoms—can change physical properties;
- Alcohols have –OH—hydroxyl; not a base!!!!!!!!!
 - large IMF's due to H-bonding; increases BP, lowers VP, etc.
 - Increases solubility
 - -COOH—carboxylic acid group; weak organic acids

Alkane		Aldehyde	R,C,H
Alkene	C=C	Ketone	R CR'
Alkyne	—C≡C—	Carboxylic Acid	R, Ü, H
Alcohol	— C-ОН	Ester	0 R
Ether	-c-o-c-	Amine	-C-N

Flame Test Colors: Cu green, Li & Sr & red, Na yellow, Ba yellow green, K violet pink Colors of stuff: Cr₂O₇²⁻ orange, CrO₄²⁻ yellow, MnO₄ purple, Cr³⁺ Cu²⁺, Ni²⁺, green or blue, PbI₂ yellow, HF etches glass, NH₃ fertilizer, NaN₃ air bags, Zn added to Fe prevent rusting, Gases:O₂ is colorless and sustains combustion,H₂ is colorless and explodes!,Cl₂ is green, Br₂ is orange-brown; NO₂ is brown; lodine sublimes into a purple vapor

PROCESS	CHANGE I	CHANGE IN A		CHANGE IN Z		CHANGE IN UTRON/PROTON RATIO		
ALPHA EMISSIO	N -4			-2		INCREASE		
BETA EMISSION	V 0			1	DECREASE			
GAMMA	0			0		X		
POSITRON EMISSION	0		-1			INCREASE		
ELECTRON CAPTURE	0			-1		INCREASE		
ALPHA	BETA	GAMM	A	POSITRON		ELECTRON CAPTURE		
helium nucleus	fast moving electron	electro	energy omagnetic liation	antiparticle of an electron		inner orbital electron		
2 protons & 2 neutrons	electron's mass Very small	no	mass	electron's mass		electron's mass		electron's mass
+2 charge	-1 charge	no	charge	+1 charge		+1 charge		-1 charge
α , $\frac{4}{2}$ He	β, ⁰ ₋₁	(0 γ	0 e 1		e -1		
Do not travel far Not penetrating	More penetrating than alpha	Very o	langerous			Gamma rays are produced		
Common with	Nuclides with too		anies other	Nuclides wi		Nuclides with too		
heavy nuclides	many neutrons		f radiation	few neutro	ons	few neutrons		
A = -4	A = 0	N	lone	A = 0		A = 0		
Z = -2	Z = +1			Z = -1		Z = -1		

Test Taking Tips Multiple Choice:

- Do not spend too much time on difficult questions.
- . Go through entire test and answer all the easy questions first.
- When you come to a question that you can answer but it will take considerable time, mark the question and come back to it after you have gone through the entire test.
- You are penalized for wrong answers so do not randomly guess.
- You can mark in your test booklet so elimination works well. If you can eliminate several choices you may want to guess.

Free Response:

- Questions require you to apply and explain chemical concepts and solve multiple step problems.
- You do not have to answer in essay form and may save time using one of the following methods: bullet format, chart format or outline format.
- Write your answers in the space provided and number your answer clearly.
- There is a slight penalty for incorrect sig figs.
- Stating a Principle, Law, Theory or stating the name of the Principle law or theory is not an
 explanation or justification for an answer. Stating a trend is also not an explanation. State
 and apply the Principle, Law or Theory to the specific situation in the questions to explain
 your answer and use the reason for trends as explanations for trends not just stating the
 trend.

DESCRIPTION OF THE AP CHEMISTRY EXAM

Multiple Choice:

- 90 minutes
- 75 questions
- May not use a calculator
- 50% of total score
- Periodic Table and Reduction Potential Table are available
- No equation sheet
- This portion of the test covers more material than any individual student is expected to know.

Free Response:

- 95 minutes
- 50 % of total score
- Tables containing commonly used chemistry equations are provided with each examination for students to use when taking the free-response section.

Two parts: Part A: problems and Part B: concepts

• Part A:

55 Minutes for part A May use calculator

Part A; students will answer three problems—one problem involving chemical equilibrium and two other problems, one of which may involve quantitative analysis of data in a laboratory-based problem.

• Part B:

40 minutes for part B May not use calculators

Part B; three questions will be scored; Question four involves predicting equations.

Question 4 will now contain 3 required balanced net ionic equations and a question related to that equation. A Periodic Table & Standard Reduction Potentials are provided.

For example:

(i) A magnesium strip is added to a solution of silver nitrate.

Answer: $Mg + 2 Ag^{+} \rightarrow Mg^{2+} + 2Ag$ (ii) Which species is oxidized? Explain

Answer: Magnesium because it has lost 2 electrons

Two other free response questions will be answered. (Questions 5 and 6) requiring applying and explaining concepts, one of which may be based on laboratory in the case that no laboratory-based problem appears in Part A