# CHEM 1110 Lecture Notes

Matthew Rowley

October 6, 2023

### COURSE ADMINISTRATIVE DETAILS

- o My office hours
- o Intro to my research
- o Introductory Quiz
- o Grading details
  - · Exams 40, Final 20, Quizzes 10, Achieve Homework 15, Textbook Homework 15
  - · Online Homework
  - · Textbook Homework
  - · Daily quizzes
- o Importance of reading and learning on your own
- Learning resources
  - · My Office Hours
  - · Tutoring services https://www.suu.edu/academicsuccess/tutoring/
- Show how to access Canvas
  - · Calendar, Grades, Modules, etc.
  - · Achieve Homework

#### MATTER AND MEASUREMENTS

#### 1.1 Chemistry: The Central Science

- My demos (Ruby, Periodic Table, NaAc hand warmers, Electroplating)
- Chemistry is the study of matter, its properties, and modes of changes
- o Chemistry is the "Central Science" because it ties in to other disciplines
- Figure 1.1 Web of disciplines
- Physical and chemical properties
- o Physical and chemical changes
  - · Pop balloon vs explode balloon

#### 1.2 States of Matter

Three phases of matter

Solid: Definite shape and volume – Particles are held rigidly in place

**Liquid:** Changing shape but definite volume – Particles flow past one another

Gas: Variable shape and volume – Particles fly apart and fill all available space

State changes: Fusion/Freezing, Vaporization/Condensation, Sublimation/Deposition

#### 1.3 Classification of Matter

- Figure 1.3 flow chart
- Pure substances vs mixtures
- Homogeneous vs heterogeneous mixtures
- Compounds vs Elements
- o Physical and chemical changes on the flow chart
- How do these changes look at the atomic level (un-numbered figure on p.6)
- Anatomy of a chemical reaction

### 1.4 Chemical Elements and Symbols

- Elements are given names and symbols (Some, like Fe, don't match in English)
- Table 1.2 Names and symbols of common elements
- Table 1.3 and 1.4 Elements in the Earth and in our bodies
- Chemical formulas Type and amount of each element in a compound (H<sub>2</sub>O, C<sub>12</sub>H<sub>22</sub>O<sub>11</sub>, etc.)

# 1.5 Chemical Reactions: An Example of Chemical Change

- Reactants, and Products
- o Reaction arrow and writing chemical equations
- o Demo: Zinc in copper nitrate solution

#### Homework 1.1:

- o 1.36 Physical vs. Chemical changes
- 1.38 Changes of state
- 1.41 Mixtures vs. Pure substances
- 1.44 Anatomy of chemical reaction
- 1.46 Element symbols
- 1.50 Counting atoms in a formula
- 1.52 Writing a formula from composition

# 1.6 Physical Quantities: Units and Scientific Notation

- Physical quantities are things that can be measured or calculated
- SI units (Table 1.5)
- Metric prefixes and  $10^n$  conversions (Table 1.6)
- Derived units
  - Squared and cubed units  $(cm^3)$
  - · Combined units (m/s)

- · Each prefix is a multiplication operation
- · Why would you use prefixes?
- $\cdot 100m$ , 0.1km, and 10,000cm
- Using units to check your formulas:  $B_e = \frac{h}{8\pi^2 \mu R_e^2 c} = cm^{-1}$
- o Scientific Notation
  - · Expressing numbers in scientific notation
    - \* Split the number into *quantity* and *magnitude*
    - \* Express the quantity as a number with MSD in the ones place
    - \* Express the magnitude as a power of 10
  - · Scientific notation and metric prefixes
    - \* For metric to scientific:  $(23.6kg \rightarrow 2.36 \times 10^4 g)$ First convert the metric prefix to a power of 10, then shift the decimal of the quantity to put its MSD in the ones place
    - \* For scientific to metric:  $(5.60 \times 10^{-7} m \rightarrow 560 nm)$ First shift the decimal to make the power of 10 match a prefix, then replace that power of 10 with the prefix

# 1.7 Measuring Mass, Length, and Volume

- Weight vs mass
- o kg,l (SI units) vs g,ml (More convenient in the lab)
- o SI vs other systems (Tables 1.7, 1.8, and 1.9)
- $\circ~$  Figure 1.6 shows the large impact of cubed units (i.e.  $1m^3=1000000cm^3)$

# 1.8 Measurement and Significant Figures

- Accuracy in measurements
- $\circ$  Statistics ( $\sigma$ ) in many measurements, simple rules for one measurement
- $\circ$  Consider my height: 6ft vs 74in vs 73.784672in
  - · More digits implies a more precise measurement
  - The first one is correct, if we allow that it is just approximate
  - · The last one is only appropriate if I was *actually* measured to the millionth of an inch
  - · If I were precisely 6ft (with high precision), how would I write that? (6.000ft)

- o Figures 1.6-7 show how we report measurements with balances and graduated glassware
- Significant figures
  - · Zeros in the middle of two numbers are significant
  - · Zeros at the beginning of a number are never significant
  - · Zeros at the end of a number *after* the decimal point are significant
  - · Zeros at the end of a number *before* the decimal point are not significant
- Scientific notation makes it easy to express precision (every 0 in scientific notation is significant)
- $\circ$  Consider 2, 300kg measured to  $\pm 1kg$  (2.300  $\times$  10<sup>3</sup>g)
- We should keep track of the total number of significant figures, and the position of the *least* significant figure in a number
- o Sig-fig practice: Give # of SFs and LSD position for several numbers

# 1.9 Rounding off Numbers

- o In science, we do not round off numbers haphazardly; Rounding reflects precision
- o Error analysis for addition and subtraction
  - · The LSD of the answer will match the LSD of the least precise input
  - # of SFs doesn't matter, and the answer might gain or lose SFs compared to the inputs
  - Practice: 96q + 43q = 139q and 23.4q 18.6q = 4.8q
  - · Consider adding  $1.00\mu l$  of water to a 8L Ace Hardware bucket The precision of the final amount is entirely determined by the low precision of the bucket
- o Error analysis for multiplication and division
  - The # of SFs in the answer will match the smallest # of SFs from the inputs
  - · Here, place value does not matter at all
  - Practice:  $45.7q \div 8.2cm^3 = 5.6g/cm^3$  and  $82.5miles/hour \times 53.24hours = 4.390miles$
- Compound problems
  - · Solve compound problems step-by-step, writing the intermediate answers
  - · For each intermediate answer, keep track of the # of SFs and LSD position
  - · Only round the final answer, don't round intermediates
  - · Practice:  $\frac{12.3g+34g}{12.0cm^3+7.7cm^3}=2.4^g/cm^3$  (wrong answer with premature rounding)

#### Homework 1.2:

- 1.58 Units and metric prefixes
- o 1.60 Scientific Notation
- o 1.62 Counting Significant Figures
- 1.66 +- error propagation
- 1.67 \*/ error propogation

### 1.10 Problem Solving: Unit Conversions and Estimating Answers

- Conversion factors
  - · Rearrange conversion equation to make unity ratios:  $1 = \frac{1in}{2.54cm} = \frac{2.54cm}{1in}$
  - Multiply a number by the right ratio to cancel out the starting units and leave the converted units
  - "Do I multiply or divide by the conversion factor?" is the wrong question. You are always multiplying by a ratio. The right question is "Where should the units be in my conversion factor?" The numbers go along with the units
  - · Most conversion factors have "1" with infinite significant figures, and the other number limits SFs in a calculation
  - · Some conversions are defined as *perfect* (such as 60s = 1min) and won't limit SFs at all
  - Take care to square or cube the first-power ratio for square or cubed dimensions:  $1in = 2.54cm \rightarrow 1^3in^3 = (2.54)^3 cm^3 \rightarrow 1in^3 = 16.4cm^3$
- Dimensional analysis is a method which frames any problem (even complex ones) in terms of multiple unit conversions
  - · The railroad ties or picket fence method makes it easy to organize a problem
  - · Set up all the unit conversions before doing any calculations
  - · Sort out significant figures at the end
  - · Practice:  $\frac{65.3miles}{h}\left|\frac{1h}{60min}\right|\frac{1min}{60s}\left|\frac{1.61km}{1mile}\right|\frac{1000m}{1km}=28.4m/_s$

# 1.11 Temperature, Heat, and Energy

- All chemical reactions involve a change in energy
- Energy is the capacity to supply heat or do work. It is in units of J
- Temperature is the measure of heat energy in an object

- · We don't use F (Booooooo, Fahrenheit!)
- · °C is split so that 0°C is freezing and 100°C is boiling
- $\cdot\,$  Absolute temperature, K, has spacing equal to °C, but starts at absolute zero
- Absolute zero is the temperature when all heat has been removed. The 0 in Kelvin is more fundamental than the 0 in  ${}^{\circ}\mathrm{C}$
- $T(^{\circ}C) = T(K) 273.15$
- The energy needed to raise the temperature of a substance is its specific heat (Table 1.10)
- 1cal = 4.184J, based on the heat capacity of water: 4.184J/gK
- $\circ q = mC\Delta T$ 
  - · How many J of heat are required to raise the temperature of 2.5 g of gold by  $8.0^{\circ}C$ ? (2.6 J)
  - 23.5 J of heat are removed from 44 g of iron. What is the temperature change?  $(-1.2^{\circ}C)$

# 1.12 Density and Specific Gravity

- $\circ \ density = \frac{mass}{volume}$  (Table 1.11), and can be used to convert mass and volume
  - · A classroom has a volume of  $8.00\times 10^3 ft^3$ . What is the mass of the air contained in that room? (268, 000g)
  - · A sample of gold weighs 8.5 g. What is the volume of this sample?  $(0.44 cm^3)$
- Density can be affected by temperature changes and phase changes. Water exhibits anomalous behavior in this respect
- Specific gravity relates the density of a substance to the density of water, and is measured by a hydrometer (Figure 1.10)

# Homework 1.3:

- o 1.78 Dimensional analysis word problem
- 1.82 Specific heat
- 1.88 Density measurement

#### ATOMS AND THE PERIODIC TABLE

#### 2.1 Atomic Theory

- o DEMO How many times can we subdivide a crystal of salt and have it still be salt?
- Postulates of atomic theory:
  - · All matter is composed of atoms
  - · The atoms of a given element differ from the atoms of all other elements
  - · Chemical compounds consist of atoms combined in specific ratios (H<sub>2</sub>O vs H<sub>2</sub>O<sub>2</sub>)
  - · Chemical reactions change the way that atoms are combined in compounds, but leave the atoms themselves unchanged
- Today we know that atoms are very small  $(m = 10^{-23} g \text{ and } r = 10^{-10} m)$
- o Atomic mass unit
  - · AMU is approximately the mass of a proton or a neutron
  - · Electrons are very small (less than a thousandth of an AMU)
  - Technically, <sup>12</sup>C has a mass of exactly 12amu
- Subatomic particles and their interactions (Table 2.1)
- Atomic structure (Figure 2.1)
  - · The protons and neutrons make up an extremely dense nucleus
  - · Electrons occupy the rest (more on this later)
  - · Most of the volume of atoms is empty (well, filled only with very low mass electrons)

#### 2.2 Elements and Atomic Number

- Atomic number (Z) is the number of protons and it distinguishes one element from another
- Neutral atoms will have the same number of electrons as protons
- Charged atoms (either more or fewer electrons than protons) are called ions
- Mass number (A) is the number of protons *and* neutrons

### 2.3 Isotopes and Atomic Weight

- Atoms with the same atomic number, but different mass numbers are called isotopes
- Hydrogen, deuterium, and tritium are the three named isotopes of hydrogen
- o Atomic symbols: 3H
  - · The mass number is in the top-left
  - · The atomic number (can be omitted) is in the bottom left
  - · The charge is in the top right (omitted if neutral)
- Atomic weight The non-integer number on the periodic table
  - · Atomic weight is based on the precise mass and natural abundances of the isotopes

· 
$$AW = \sum_{isotopes} \left( mass \times \frac{\% \ abundance}{100\%} \right)$$

 $\circ$  Li: 7.59% abundance and m=6.015122amu. Li: 92.41% abundance and m=7.016004amu

#### Homework 2.1:

- o 2.32 How elements differ from each other
- o 2.42 Composition from atomic symbol
- 2.46 Atomic symbols from composition
- 2.48 Calculate atomic mass from isotopes

#### 2.4 The Periodic Table

- o As more elements were discovered, scientists tried to find commonalities and patterns
- o Li, Na, and K were similar to each other, as were Cl, Br, and I
- Mendeleev arranged the elements in increasing mass, and saw a pattern
- Mendeleev even correctly predicted the properties of Gallium, Germanium, and Scandium yetundiscovered elements
- Rows are called "Periods" and columns are called "Groups"
- The periods get longer further down due to electronic structure We will talk about this soon
- Figure 2.2 shows two ways to divide the periodic table
  - · Main group vs. transition metals vs. inner transition metals
  - · Metals vs. Metalloids vs. Non-metals

### 2.5 Some Characteristics of Different Groups

- Periodic trend in atomic radius (Figure 2.3)
- Group 1A Alkali metals: Shiny, soft, and highly reactive
- o Group 2A Alkaline earth metals: Like Alkali, but less extreme
- o Group 7A Halogens: Colorful and corrosive in elemental form
- Group 8A Noble gases: Won't react with anything (\*almost\*)

#### 2.6 Electronic Structure of Atoms

- o Electronic structure largely determines physical and chemical properties
- o Electrons can only exist in quantized energy states because of their quantum wavelike nature
- These states have different *sizes* and characteristic *shapes* in 3-D space. The state defines where an electron can be found, and how much energy it has
- Each state of an electron can be characterized by *quantum numbers*, which can be thought of as the "address" for a given electron
  - · First is the shell number, *n* 
    - \* The shell determines the energy and size of a state
    - \* Shells can hold one or more subshells
    - \* Each successive shell can hold more electrons than the last (2, 8, 18, 32)
  - · Next is the subshell number, *l* 
    - \* Subshells come in 4 types: s, p, d, and f
    - \* The subshell type determines the shape of the electron state
    - \* Each subshell has a different number of individual states (orbitals), and each state can contain 2 electrons

Subshell	Shape	# of Orbitals	Max # of electrons
s	Sphere	1	2
p	Dumbell	3	6
d	Clover	5	10
f	Complex!	7	14

- Each successive shell adds a new subshell type (Table 2.2)
- · Spin differentiates the two electrons which share an orbital
- o Draw a hydrogenic energy level diagram, and practice identifying shells and subshells

# 2.7 Electron Configurations

- For multi-electron atoms, the subshells arrange themselves a bit differently
  - The first few subshells are: 1s, 2s, 2p, 3s, 3p, 4s, 3d, 4p (Draw the energy level diagram)
  - · We will show a way to remember this pattern in just a few minutes
- Electrons occupy orbitals according to the following rules:
  - · Electrons will occupy the lowest energy orbital available. This is the Aufbau principle.
  - Each orbital can hold only two electrons, which must have opposite spins. This is the Pauli exclusion principle
  - Degenerate groups of orbitals are all filled halfway with electrons of the same spin before any electrons are paired up. This is Hund's rule
- To find the configuration of an element, count the number of electrons and fill the orbitals according to the three rules above
- o To give the configuration, list the occupied subshells in order, and their occupancy as a superscript
- $\circ$  Practice: O:1 $s^22s^22p^4$  Sin
- Si: $1s^2 2s^2 sp^6 3s^2 3p^2$
- o Table 2.3 shows electron configurations for the first 20 elements
- Noble gas notation
  - · For longer configurations, we can reference the noble gas which comes previous to the element
  - The electrons represented by that noble gas are *core* electrons, and are not actually very important to bonding, ion formation etc.
  - The remaining electrons are in the outermost, *valence* shell, and are most important to chemistry
  - Practice: Cl:[Ne] $3s^23p^5$

# 2.8 Electron Configurations and the Periodic Table

- The periodic table is arranged according to properties, but properties are governed by electronic configuration. Ergo, the periodic table is arranged according to electronic configuration!
- o Figure 2.7 shows how the blocks of the periodic table represent subshells in electronic structure
- $\circ \:$  My figure includes d(n-1) and f(n-2) labels
- o Note that the width of each block matches the number of electrons each subshell type can hold
- The order of the subshells can be found by simply traversing up the elements in the periodic table
- $\circ~$  Practice: Pb:[Xe]  $6s^24f^{14}5d^{10}6p^2$

# 2.9 Electron Dot Symbols

- $\circ~$  It is useful to graphically represent the valence electrons with electron dot symbols
- Write the symbol for the element, then surround it with the correct number of dots to represent valence electrons
- Dots go on the four sides of the symbol, and pair up only when there are more than 4 electrons
- o Table 2.5 shows many of these diagrams

#### Homework 2.2:

- o 2.56 Classify elements (metal, non-metal, main group, etc.)
- o 2.66 How many subshells in each shell
- 2.70 Electron arrow diagrams
- o 2.73 Electronic configurations of elements

#### **IONIC COMPOUNDS**

### 3.1 Ions

- Metals and non-metals will combine to form ionic compounds
- Ionic compounds have high melting points, stable crystal structures, and a degree of water solubility
- When ionic compounds dissolve in water, they conduct electricity
- o Positive ions are called *cations* and negative ions are called *anions*
- Many reactions involve ions which are already charged, while other reactions generate ions through gaining or losing electrons

### 3.2 Ions and the Octet Rule

- o Elements will tend to gain or lose enough electrons to have a full *octet* of valence electrons
- This means that non-metals will gain electrons to reach the electron configuration of the next highest noble gas
- o Metals will lose electron to reach the next lowest noble gas
- Electron configurations for ions are written just the same as electron configurations for elements

### 3.3 Ions of Some Common Elements

- o Use the periodic table to predict the charge an element will take on as it forms an ion
- O Group 1A will be 1+
- o Group 2A will be 2+
- Group 6A will be 2-
- Group 7A will be 1-

### 3.4 Periodic Properties and Ion Formation

- Ionization energy is the energy required to remove an electron and create a cation
- o Electron affinity is the energy given off when an electron is gained, making an anion
- o Periodic trends in ionization energy and electron affinity
- Nonmetals such as C and N tend to not form ions

### 3.5 Naming Ions

- o For metals with just one ion (1A, 2A, and 3A groups), just add "ion" to the end of the name
- o For metals with multiple ions (transition metals, 4A, and 5A), state the charge and then add "ion"
- $\circ$  An old rule uses different endings (ferrous and chromous for 2+, and ferric and chromic for 3+)
- Anions change the end of the element name to "-ide"
- o See tables 3.1 and 3.2

# Homework 3.1:

- o 3.42 Identify ion from charge and electron number
- 3.44 Ions and the octet rule
- 3.46 Ion formation by gaining/losing electrons
- 3.50 Ionization energy
- 3.58 Naming Ions

# 3.6 Polyatomic Ions

- Polyatomic ions are composed of more than one atom
- These polyatomic ions are a covalently bound group, and should be considered as a single ion
- See table 3.3
- There are many series of polyatomic ions which differ in only the number of oxygen atoms
- $\circ~$  There are also series of polyatomic ions which differ in hydrogen atoms and charge (each H increases the charge by 1)

#### 3.7 Ionic Bonds

- Ionic compounds are held together by ionic bonds
- Positive cations are attracted to negative anions
- Ionic solids exhibit a regular crystal lattice on atomic scales

### 3.8 Formulas of Ionic Compounds

- To find the formula for an ionic compound, we need to balance the charges to give a neutral compound
- The ratio should be in a mathematically reduced form, making just one formula unit

### 3.9 Naming Ionic Compounds

- Simply combine the two ion names Cation first, then anion
- Do not state the ratio between them, like you do in molecular compounds
- Table 3.4 Some common ionic compounds and their applications

# 3.10 Some Properties of Ionic Compounds

- o Ionic compounds have different lattice structures, depending on the sizes and charges of the ions
- o Ionic solids are rigid but brittle, easily cleaved along crystal planes
- Ionic compounds have high melting and boiling points This is because ionic bonds are very strong
- Some ionic compounds dissolve in water by making many attractive interactions with water dipoles.
   Others are just bound too tightly, and are insoluble.

# 3.11 H<sup>+</sup> and OH<sup>-</sup> Ions: An Introduction to Acids and Bases

- The hydrogen cation (really just a bare proton) and hydroxide anion (OH⁻) are very important to aqueous chemistry
- $\circ$  When an acid dissolves in water, it donates a hydrogen cation to a water molecule forming  $\mathrm{H_2O^+}$
- When a base dissolves in water, it will accept a proton from water forming OH<sup>-</sup>
- Some acids can provide multiple H<sup>+</sup> in reactions (HCl, H<sub>2</sub>SO<sub>3</sub>, H<sub>3</sub>PO<sub>4</sub>), and some bases can accept multiple protons (NaOH, Ba(OH)<sub>2</sub>)

# Homework 3.2:

- o 3.62 Polyatomic Ions
- $\circ$  3.66 Ions combine to form compounds
- $\circ~$  3.68 Formulas from names for ionic compounds
- $\circ~$  3.70 Names from formulas for ionic compounds
- $\circ \ \ 3.76-Acid/base\ reactions\ with\ water$

#### MOLECULAR COMPOUNDS

### 4.1 Covalent Bonds

- A bond formed by the *sharing* of electrons is called a covalent bond. They occur between non-metals
- $\circ$  A group of atoms held together by covalent bonds is called a molecule (this can include compounds like  $CO_2$  or elements like  $O_2$ )
- Show a water molecule using dot diagrams
- o Repulsive and attractive forces in a covalent bond as a function of bond length
- Overlap of 1s orbitals for H<sub>2</sub> and 2p orbitals for F<sub>2</sub>

#### 4.2 Covalent Bonds and the Periodic Table

- o The periodic table can be used to predict how many bonds an element tends to make
- Consider HF, H<sub>2</sub>O, NH<sub>3</sub>, and CH<sub>4</sub>
- This pattern is the result of the octet rule. An atom makes as many bonds as it needs in order to reach a full octet
- Elements in the third row and below can form more bonds. See Figure 4.3

# 4.3 Multiple Covalent Bonds

- o Some molecules cannot satisfy the octet rule using only single bonds
- Consider F<sub>2</sub>, O<sub>2</sub>, and N<sub>2</sub>
- o Single, double, and triple bonds share 2, 4, and 6 electrons
- Multiple bonds in organic molecules

#### 4.4 Coordinate Covalent Bonds

- Coordinate covalent bonds are when both shared electrons came from one atom in the pair
- $\circ$  Identify them in molecular compounds by counting the electrons and comparing them to their valency: N<sub>o</sub>O
- o O3 and NH3BF3 are two more examples of coordinate covalent compounds
- o In acid/base chemistry, all acidic hydrogens form coordinate covalent bonds

#### Homework 4.1:

- 4.34 Identify covalent and ionic bonds
- 4.36 Predict # of bonds from periodic table
- 4.42 Identify coordinate covalent bonds

#### 4.5 Characteristics of Molecular Compounds

- Molecular compounds have a wide range of melting and boiling points, depending on their intermolecular forces (chapter 8)
- Few molecular compounds are soluble in water, but many are soluble in organic solvents
- Most molecular compounds do not conduct electricity, either as a pure substance or dissolved in water

#### 4.6 Molecular Formulas and Lewis Structures

- Molecular formulas tell the types and numbers of atoms found in a molecular compound
- Structural formulas use lines to represent covalent bonds and show the structure of a molecule
- Lewis structures add dots to represent lone pairs of electrons

### 4.7 Drawing Lewis Structures

- The first way to draw Lewis structures works for most simple organic molecules
  - · If the only atoms are C, H, O, X (Halogen), and H, then this approach might work
  - · Arrange the carbon atoms in a chain, or backbone
  - · Attach the heteroatoms with the appropriate number of bonds (based on the octet rule)

- · Make any multiple bonds in the carbon chain needed to complete their octets
- · If all of the elements have a full octet, and form the correct number of bonds, then it is a valid structure
- · Starting from a condensed structure makes this method particularly easy
- The next method is very useful for molecular compounds with more complex structure, or central atoms which aren't carbon
  - 1. Count up the total number of valence electrons (taking into account any net charge)
  - 2. Draw single bonds from the central atom to all peripheral atoms
  - 3. Fill the octets of the outer atoms with lone-pairs (H doesn't need any)
  - 4. Place any remaining electrons onto the central atom
  - 5. Satisfy the octet rule for the central atom, if needed, by converting outer lone-pairs into multiple-bonds with the central atom
- Practice: PCl<sub>3</sub>, HCN, SO<sub>4</sub><sup>2-</sup> (sulfate has single bonds for this class)

### 4.8 The Shapes of Molecules

- We know that molecules have a particular shape, defined by VSEPR theory
- We can predict the molecular geometry based on a molecule's Lewis structure
  - 1. Draw a valid Lewis structure
  - 2. Count the number of electron domains
  - 3. Count the number of bonds vs. lone pairs
  - 4. Consult Table 4.2 (or your memory)
- $\circ$  Bond angles can range from  $180^{\circ}$  to  $< 109.5^{\circ}$
- If lone pairs are present, they slightly reduce the bonding angles

#### Homework 4.2:

- 4.53 Condensed structural formulas
- 4.56 Drawing Neutral Lewis structures
- 4.62 Drawing Lewis Structures of polyatomic ions
- 4.66 Molecular geometry

### 4.9 Polar Covalent Bonds and Electronegativity

- Any covalent bond between different atoms will share electrons unequally
- o This unequal sharing leads to a polar bond
- The direction and intensity of the bond polarity is determined by comparing electronegativities of the two elements
- o F is the most electronegative, while Rb is the least
- o Electronegativity differences of o-o.4 are considered normal covalent bonds
- Electronegativity differences of 0.5-1.9 are considered polar covalent bonds
- o Electronegativity differences greater than 2 are considered ionic bonds

#### 4.10 Polar Molecules

- Polar bonds in a molecule can lead to a charge dipole over the entire molecule
- First find the polar bonds, then add up their dipole vectors to find the net dipole
- o Symmetry can often lead to a non-polar molecule despite having polar bonds

### 4.11 Naming Binary Molecular Compounds

- A compound made of only two types of atoms is called a binary compound
- When naming a binary compound, the less electronegative atom usually comes first
- The second element will have an "-ide" ending, just like in ionic compounds
- Use prefixes (Table 4.3) to indicate how many of each element are present

### Homework 4.3:

- 4.74 Polar bonds
- 4.76 Polar bonds in molecules
- 4.78 Polarity of molecules
- 4.82 Names from formulas
- 4.84 Formulas from names

#### CLASSIFICATION AND BALANCING OF CHEMICAL REACTIONS

### 5.1 Chemical Equations

- Chemical equations can be thought of as recipes which show what you start with and what you get out
- Coefficients indicate how many units of each reactant are needed, and how many units of each product are made
- These ratios are referred to by the name *stoichiometry*
- We find the coefficients by relying on the law of conservation of mass (balancing)

#### 5.2 Balancing Chemical Equations

- To balance chemical reactions:
  - · Write the unbalanced equation, with correct formulas for all reactants and products
  - · Identify an element which is unbalanced, and balance it with a coefficient
  - · Repeat the above step until all elements are balanced
  - · Reduce the coefficients if they share a common factor
- Combustion of C<sub>5</sub>H<sub>10</sub> gives an example where we have to double the coefficients

#### Homework 5.1:

- 5.24 Balance equations from chemical names
- 5.28 Balance general equations
- 5.30 Balance combustion reactions

#### **Classes of Chemical Reactions**

- This section was removed from the new edition, but it is worth providing an overview of the classes
  of reactions before we go over each in detail
- It is sometimes helpful to recognize different classes of chemical reactions

Precipitation reactions go from aqueous ionic reactants to solid ionic products:

$$Pb(NO_3)_2(aq) + 2 KI(aq) \longrightarrow 2 KNO_3(aq) + PbI_2(s)$$

Acid-base neutralization reactions result in water and a salt (H<sup>+</sup> is transferred)

$$HCl(aq) + NaOH(aq) \longrightarrow H_2O(l) + NaCl(aq)$$

Oxidation-reduction reactions change the oxidation state (or charge) of atoms:

$$Mg(s)+I_2(g) \longrightarrow MgI_2(s)$$

# 5.3 Precipitation Reactions and Solubility

- Solubility is a measure of how much a substance will dissolve in a solvent (usually water)
- In precipitation reactions, you must determine the solubility of *all* potential products
- Solubility rules (Table 5.1):
  - · A compound is probably soluble if its cation is ammonium  $(NH_4^+)$  or group 1A cations  $(Li^+, Na^+, Rb^+, or Cs^+)$
  - · A compound is probably soluble if it contains one of the following anions: Halide ions (except when paired with  $Ag^+$ ,  $Hg_2^{\ 2^+}$ , or  $Pb^{2^+}$  ions) Nitrate, perchlorate, acetate, or sulfate (except for sulfates paired with  $Ba^{2^+}$ ,  $Hg_2^{\ 2^+}$ , or  $Pb^{2^+}$  ions)
  - · Most other ionic compounds are not soluble
  - · Net ionic equation omit all species which are aqueous on both sides of the reaction

### **Net Ionic Equations**

- We are covering this topic early because of its relevance to precipitation reactions
- o Aqueous ionic compounds have cations and anions which are not actually bound together at all
- Separate aqueous ionic compounds into their ions this is the complete ionic equation
- Identify spectators species which are in exactly the same form on both the reactant and product sides
- Eliminate the spectators, leaving only the species which are involved in real chemistry This is the net ionic equation

#### Homework 5.2:

- 5.40 Ion solubility rules
- o 5.42 Predicition solubility of reaction products
- 5.46 Writing net ionic equations

#### 5.4 Acids, Bases, and Neutralization Reactions

- When a strong acid and base react, the product is *neutral* (neither an acid nor a base)
- The products can include water and a neutral ionic compound
- $\circ$  Some acid/base neutralization reactions produce gas:  $_2$  HCl(aq) +  $_2$ CO $_3$ (aq)  $\longrightarrow$  H $_2$ O(l) +  $_2$  KCl(aq) + CO $_2$ (g)
- $\circ$  Many bases yield OH<sup>-</sup> as a product, even if they dont have hydroxide as a component themselves: NH<sub>3</sub> + H<sub>2</sub>O  $\longrightarrow$  NH<sub>4</sub><sup>+</sup> + OH<sup>-</sup>

#### 5.5 Redox Reactions

- o Redox reactions involve changes in oxidation state, or the exchange of electrons
- OIL-RIG "Oxidation is losing, Reduction is gaining" electrons
- $\circ$  Oxidation leads to a higher oxidation state, while reduction leads to a lower (reduced) oxidation state
- Reduction and Oxidation always come together, since the electrons need a source and a destination
- We use language which can be confusing at first:
  - The oxidizing agent is the chemical which oxidizes its partner. It is reduced itself in the process. It is a destination for electrons
  - The reducing agent is the chemical which reduces its partner. It is oxidized itself in the process. It is a source for electrons
- Some redox processes we encounter in the world around us: Corrosion, Combustion, Respiration, Bleaching, Metallurgy

### 5.6 Recognizing Redox Reactions

- Determining oxidation states can get complicated, but for now we can simply relate oxidation state to charge
- o Elemental states have oxidation states of o
- o For a covalently bonded compound, the oxidation states sum up to the total charge
- Elements in a covalent bond will usually take the same oxidation state that they would in an ionic bond (C is an important exception, as well as non-metal oxides)

# 5.7 Net Ionic Equations

Covered out of order above

## Homework 5.3:

- $\circ$  5.34 Classify reactions by type
- $\circ \ \ 5.52-Assigning \ oxidation \ numbers$
- $\circ~5.54-Recognize$  oxidation and reduction
- 5.58 Oxidizing and reducing agents

#### CHEMICAL REACTIONS: MOLE AND MASS RELATIONSHIPS

#### 6.1 The Mole and Avogadro's Number

- The molecular weight or formula weight of a substance is the sum of atomic weights for its constituent atoms
- $\circ$  Different substances have different molecular weights, so 1.0g of  $C_2H_4$  has more molecules in it than 1.0g of HCl
- $\circ$  If we wanted to make  $C_2H_5Cl$  without any left-over reactants, then we would need more grams of HCl than of  $C_2H_4$
- $\circ$  Find the ratio by comparing the formula masses (28.0amu vs 36.5amu). If I have 1.0g of  $C_2H_4$ , how many grams of HCl do I need? (1.3g)
- The mole is an amount that relates amus (atomic scale) to grams (macroscopic scale)
- o The molar mass tells how many grams of a substance is one mole of molecules
- $\circ~N_A=6.022\times 10^{23}$  , Avogadro's number, is the number of amus per gram

#### 6.2 Gram-Mole Conversions

- We can convert from grams to moles and vice-versa using the molar mass
- How many moles is 2.5g of  $C_2H_5Cl$ ? How many molecules?

### **6.3** Mole Relationships and Chemical Equations

- o A balanced chemical equation relates the ratios of moles of reactants and products
- The coefficients tell how many moles are needed or produced
- We call these ratios *stoichiometry*
- Homework 6.1:
  - · 6.32 Calculating molar mass
  - · 6.36 Moles to mass and to molecules
  - · 6.39 Stoichiometric ratios in a balanced equation

### 6.4 Mass Relationships and Chemical Equations

- Mole to mole conversions rely on stoichiometric ratios (coefficients in a balanced chemical equation)
- o Mole to mass and mass to mole conversions rely on molar mass
- $\circ$  Mass to mass conversions must follow the path: mass<sub>1</sub>  $\longrightarrow$  moles<sub>2</sub>  $\longrightarrow$  moles<sub>2</sub>  $\longrightarrow$  mass<sub>2</sub>
- You should always check your calculated answer against a rough estimate to make sure it makes sense
- $\circ$  For  $H_2SO_4(aq) + 2 NaOH(aq) \longrightarrow Na_2SO_4(aq) + 2 H_2O(aq)$ :
  - · How many moles of NaOH needed to react with 0.490g H<sub>2</sub>SO<sub>4</sub>?
  - · How many grams of water and sodium sulfate would be produced?

### 6.5 Limiting Reagent and Percent Yield

- Unless the amounts of reactants are carefully chosen, one will run out first and the other will have left-over remaining after the reaction
- To find the limiting reagent, calculate how much product each one would yield. The reactant
  which yields less is the limiting reagent and the amount calculated is the theoretical yield
- o To find percent yield, you must have an actual mass of product:

$$\%Yield = \frac{ActualYield}{TheoreticalYield} \times 100\%$$

 $\circ \ CO_{_2} + H_{_2}O \ \longrightarrow \ C_6H_{_{12}}O_6 + O_{_2} \, (unbalanced)$ 

With 10.0g of  $CO_2$  and 5.00g of  $H_2O$ , how many grams of sugar are produced?

- $\circ$  if 4.8*g* of sugar are actually recovered, what is the % recovery?
- $\circ$  C<sub>6</sub>H<sub>8</sub>O<sub>7</sub> + NaHCO<sub>3</sub>  $\longrightarrow$  Na<sub>3</sub>C<sub>6</sub>H<sub>5</sub>O<sub>7</sub> + H<sub>2</sub>O + CO<sub>2</sub> (unbalanced) With 1.0g citric acid and 1.0g sodium bicarbonate how many grams of Na<sub>3</sub>C<sub>6</sub>H<sub>5</sub>O<sub>7</sub> are produced?
- $\circ$  You actually recover 0.56g of citrate. What is the % yield?
- Homework 6.2:
  - · 6.46 Finding reactant amounts from desired product yield
  - · 6.52 Limiting reactant problem

# CHEMICAL REACTIONS: ENERGY, RATES, AND EQUILIBRIUM

Some chemical reactions can be balanced, but meaningless because they will never occur naturally.

$$2 \operatorname{Au}(s) + 3 \operatorname{H}_2 O(l) \longrightarrow \operatorname{Au}_2 O_3(s) + 3 \operatorname{H}_2(g)$$

### 7.1 Energy and Chemical Bonds

- Potential energy is stored energy. The energy of chemical bonds is a form of negative potential energy
- o Kinetic energy is energy of motion. Motion on the atomic scale is observed as heat
- In a chemical reaction, energy is required to break chemical bonds, but energy is released when new bonds are formed
- If the energy released is greater than the energy required to break the bonds, then the reaction produces heat and the products are more stable than the reactants

# 7.2 Heat Changes During Chemical Reactions

- To predict if a reaction will give off heat, we must compare the energies of the bonds broken to the energies of the bonds formed
- Table 7.1 shows average bond enthalpies
- $\circ$  Consider the reaction:  $H_2(g) + Cl_2(g) \longrightarrow 2 HCl \Delta H_{rxn} \approx -43 \frac{kcal}{mol}$
- Since energy is conserved and the molecules lost energy overall, then that energy must have gone somewhere – It is released as heat!
- Reactions which release heat are called "exothermic," and they occur more readily in nature
- Reactions which take in heat (make the surroundings colder) are called "endothermic," and they are a bit harder to find in nature
- Demo Burning methanol and Ba(OH)<sub>2</sub>·<sub>8</sub>H<sub>2</sub>O(s) + 2 NH<sub>4</sub>SCN(s)  $\longrightarrow$  Ba(SCN)<sub>2</sub>(aq) + 2 NH<sub>3</sub>(g) + 10 H<sub>2</sub>O(l)
- $\circ~$  We have a fancy name for the heat of reaction carried under constant pressure conditions: "Enthalpy of Reaction"  $(\Delta H)$

### 7.3 Exothermic and Endothermic Reactions

- Practice an endothermic reaction:  $N_2(g) + O_2(g) \longrightarrow 2 NO(g)$   $\Delta H_{rxn} = 43 \frac{kcal}{mol}$
- Note that these quantities are -per mole. Table 7.2 compares combustion enthalpies on a -per gram basis

### 7.4 Why do Chemical Reactions Occur? Free Energy

- o Chemical reactions which occur in nature are called "spontaneous"
- o Non-spontaneous reactions can be made to occur if driven by some external input of energy
- $\circ$  Enthalpy is one governing factor in spontaneity, and entropy (S) is the other
- Entropy can be understood as the amount of disorder in a system
- A reaction will have  $\Delta S_{rxn} > 0$  if the products are more disordered than the reactants
- $\circ$  Entropy and Enthalpy are combined to form "free energy" ( $\Delta G_{rxn}$ ), which can be thought of as a measure of the spontaneity of a reaction
- $\circ \Delta G_{rxn} = \Delta H_{rxn} T\Delta S_{rxn}$
- $\circ~\Delta H_{rxn} < 0$  and  $\Delta S_{rxn} > 0$  both trend toward spontaneity Draw the quadrant diagram

#### 7.5 How do Chemical Reactions Occur? Reaction Rates

- $\circ \Delta G_{rxn}$  and free energy only tell us if a reaction can happen, but doesn't say how quickly a reaction will occur
- Some spontaneous reactions happen so slowly that they never really happen at all
- The rate of a reaction depends on different factors, like how the reaction actually occurs (from a molecular perspective)
- For reactions to occur, molecules must collide with enough energy and in the correct orientation (Figure 7.2)
- The energy required to start a reaction is called the activation energy, and it limits the rate of reaction
- Reaction coordinate diagrams can show factors which control both the energetics and the rate of a reaction(Figure 7.3)

#### Homework 7.1:

- · 7.30 Reaction enthalpies from bonds
- · 7.34 Predicting entropy changes
- · 7.46 Reaction coordinate diagrams

### 7.6 Effects of Temperature, Concentration, and Catalysts on Reaction Rates

- Temperature:
  - · Increasing the temperature increases the frequency and energy of molecular collisions
  - · Both of these factors will increase the rate of any reaction
- Concentration:
  - · Increasing the concentration increases the frequency of molecular collisions
  - · This will increase the rate of any reaction
- Catalysts:
  - · Catalysts change the way a reaction proceeds, requiring less activation energy (Figure 7.4)
  - · This will increase the rate of reaction, since more collisions will have the necessary energy

# 7.7 Reversible Reactions and Chemical Equilibrium

- Some chemical reactions can go in the forward or reverse direction dependig on the circumstances
- These reactions are called equilibrium reactions because they establish an equilibrium state where both reactants and products are present
- At equilibrium the forward and reverse reactions occur at the same rate, so the concentrations of products and reactants remains constant.

# 7.8 Equilibrium Equations and Equilibrium Constants

- o We can describe the equilibrium conditions for a reaction using math
- $\ \, \text{ The equilibrium constant:} \, K_{eq} = \frac{[M]^m [N]^n}{[A]^a [B]^b}$
- $\circ~$  The exponents in  $K_{eq}$  are from the coefficients in the balanced chemical equation
- We can therefore find the equilbrium constant if we measure the equilibrium amounts of reactants and products

- $\circ$  We could also find the equilbrium amount of one species if we know  $K_{eq}$  and the concentrations of all other species
- $\circ$  The value of  $K_{eq}$  can indicate whether thee are more products or reactants at equilibrium
- $\circ K_{eq} > 1$  favors products, while  $K_{eq} < 1$  favors reactants

# 7.9 Le Châtelier's Principle: The Effect of Changing Conditions on Equilibria

- Le Châtelier's principle describes how a system will respond after being perturbed away from equilibrium
- Effect of Changes in Concentration:
  - · When concentrations are changed, the system will respond to counteract the change
  - · Removing product will shift the reaction toward products
  - · Adding product will shift the reaction toward reactants
  - · etc.
- o Effect of Changes in Temperature
  - To understand the effect of a change in temperature we can consider heat as either a reactant or a product
  - · Heat acts as a product in an exothermic reaction, and as a reactant in an endothermic reaction
  - · Increasing the temperature is like adding heat, and decreasing the temperature is like removing heat
  - · When temperature changes, the value of  $K_{eq}$  actually changes
- Effect of Changes in Pressure (or volume):
  - First, count the number of moles on each side of the reaction (gas for pressure and aqueous for volume)
  - · Increasing pressure or decreasing volume will shift the reaction toward the side with fewer moles
  - · Decreasing the pressure increasing the volume will shift the reaction toward the side with more moles
- Homework 7.2:
  - · 7.48 Catalysts and activation energy
  - · 7.56 Equilibrium constant
  - · 7.68 LeChâtelier's principle

#### GASES, LIQUIDS, AND SOLIDS

#### 8.1 The States of Matter and Their Changes

- The three primary phases of matter: Solid, Liquid, and Gas
- o Phase changes can go either direction at the phase change temperature
- $\circ s \longrightarrow l$  and  $l \longrightarrow g$  are endothermic, while the reverse reactions are exothermic
- Phase change temperatures can be understood in the context of:  $\Delta G = \Delta H T\Delta S$
- Sublimation and dry ice

#### 8.2 Intermolecular Forces

- Intermolecular forces determine the physical phase of a substance
- London Dispersion Forces
  - · Figure 8.4
  - · Effect of molecular shape More surface area gives stronger forces
- o Dipole-Dipole Forces
  - · Figure 8.3
  - · Effect of polarity Stronger dipoles give stronger forces
- Hydrogen Bonds
  - · H-bonds special cases which go beyond normal dipole-dipole bonds
  - · A H atom is very strongly attracted to another molecule (almost like the H is shared between molecules)
  - · Page 221 un-numbered figure
  - · Electron donor must be an electronegative atom (N, O, or F)
  - · Hydrogen must be bound to an electronegative atom (N, O, or F)
  - · Acetone is a counter-example. It has O and H, but not bound together

# 8.3 Gases and the Kinetic-Molecular Theory

- Gas behavior can be mostly explained by assuming that the individual gas molecules follow these assumptions:
  - · A gas consists of many particles which move randomly and independently
  - · The individual gas particles exhibit no intermolecular forces
  - The volume occupied by the gas particles is vanishingly small compared to the volume filled by the gas collectively (i.e. the particles are very small)
  - · The average kinetic energy of the gas particles is proportional to the temperature (in K)
  - · Gas particles collide elastically
- Real gases deviate slightly from these assumptions, but we can ignore those deviations and treat them like an *ideal gas*

#### 8.4 Pressure

- o Pressure is the force a gas applies over an area
- Even ambient air is exerting a fairly high pressure atmospheric pressure
- We measure ambient pressure using a mercury barometer Figure 8.9
- $\circ~$  Pressure can be measured in many different units: 1atm=760mmHg=760torr=14.7psi=101325Pa

### 8.5 Boyle's Law

- $\circ$  Early physicists experimented with the fundamental state variables of gases (P, V, and T) and noticed simple relationships called gas laws
- o Boyle showed the relationship between pressure and volume
- $\circ~$  Demo a small balloon expanding in a vacuum:  $P \propto \frac{1}{V}$
- The pressure increases with decreased volume, which is consistent with the kinetic molecular theory of gases because there are more molecular collisions per second in the smaller volume
- $\circ$  Solve Boyle's law problems by relating the initial and final conditions:  $P_1V_1=P_2V_2$

### 8.6 Charles's Law

- $\circ~$  Charles's law relates the volume of a gas to temperature:  $V \propto T$
- o Demo a small balloon expanding as it moves from ice to boiling water
- This behavior is explained by the kinetic molecular theory of gases because the collisions are more forceful, pushing to greater volumes
- $\circ~$  Solve Charles's law problems with the following relation:  $\frac{V_1}{T_1} = \frac{V_2}{T_2}$
- Homework 8.1:
  - · 8.34 Intermolecular forces
  - · 8.42 Converting gas pressures
  - · 8.48 Boyle's Law
  - · 8.56 Charles's Law

### 8.7 Gay-Lussac's Law

- $\circ~$  Gay-Lussac's law provides the last relation:  $P \propto T$
- $\circ~$  This law is redundant, since the previous two laws could be combined to prove the Gay-Lussac law

#### 8.8 The Combined Gas Law

- Boyle's law, Charles's law, and the Gay-Lussac law all focus on the relationship between two state variables while keeping the third constant
- o These laws can be combined to give a law which allows all three variables to change

$$\frac{PV}{T} = k$$
 where  $k$  is a constant value

- $\circ~$  Solve these problems with the following relation:  $\frac{P_1 V_1}{T_1} = \frac{P_2 V_2}{T_2}$
- This law includes all the others, so it is really the only one that you need to know, except for the even better law we will teach you next class

### 8.9 Avogadro's Law

- o All of the above laws are for a constant (and unknown) number of moles of gas
- $\circ~$  More moles of gas will give greater volume :  $\frac{V_1}{n_1} = \frac{V_2}{n_2}$
- $\circ$  It is convenient to define a standard temperature (0°C) and pressure (1atm) or comparing different gas systems
- $\circ$  At STP, one mole of an ideal gas occupies a standard molar volume of 22.4L

#### 8.10 The Ideal Gas Law

- $\circ$  The ideal gas law combines everything we have covered so far: PV = nRT
- $\circ R$  is the gas constant, which can take on different values depending on the units:

$$R = 0.08206 \frac{L atm}{mol K} = 62.4 \frac{L Torr}{mol K}$$

- Find the pressure when 0.250mol of gas are placed in a 5.20L container at 15.0°C (1.14atm)
- $\circ$  How many moles are in a 3.15L helium balloon at 20.0°C and 0.820atm? (0.107mol)

#### 8.11 Partial Pressures and Dalton's Law

- Since gas particles don't interact at all, a mixture of gases doesn't introduce any new complexity
- Each gas exerts a partial pressure as if the other gases weren't there, and the total pressure is the sum of the partial pressures
- $\circ P_{total} = \sum P_{i,partial}$
- $\circ$  The air is mostly made of O<sub>2</sub> and N<sub>2</sub>. In Cedar City,  $P_{\rm O_2}=0.17atm$  and  $P_{\rm N_2}=0.64atm$ . What is the total barometric pressure? (0.81atm)
- Homework 8.2:
  - · 8.62 Ideal gas law
  - · 8.82 Ideal gas law
  - · 8.88 Dalton's law of partial pressures

#### 8.12 Liquids

- A liquid in a closed container will reach equilibrium with the vapor phase as vaporization and condensation rates come into balance (Figure 8.19)
- o The equilibrium pressure above a liquid is the vapor pressure
- Different liquids have different vapor pressures, and vapor pressure always increases with temperature (Figure 8.20)
- When the vapor pressure equals the external pressure ( $\approx 1 atm$ ), boiling will begin
- $\circ$  The lower external pressure at high altitudes is why water will boil at less than  $100^{\circ}C$  at those altitudes
- Viscosity is a measure of how freely a liquid flows. Some liquids, like honey, are very viscous
- Surface Tension is the resistance of a liquid to spreading out and increasing its surface area. This leads to curved droplets of liquids, since this shape minimizes surface area (Figure 8.21)

#### 8.13 Water: A Unique Liquid

- Water has many unique properties compared to other liquids
- Water has an unusually high heat capacity and enthalpy of vaporization due to its strong hydrogen-bonding intermolecular forces
- Water is nearly unique in having a solid phase which is less dense than the liquid phase. This is because ice has an unusually open crystal structure (Figure 8.23)

#### 8.14 Solids

- Many solids exhibit an ordered structure over long ranges. These are called crystalline solids
- Crystalline solids can be one of four types:
  - · Ionic solids, like NaCl, are made of a lattice of cations and anions
  - · Molecular solids, like sugar, are made of individual molecules arranged in a repeating lattice
  - · Covalent network solids, like diamond, are extended networks of covalent bonds. A diamond can *kind of* be described as one giant molecule!
  - Metallic solids are made of a lattice of metal atoms which share a sea of electrons across the whole solid. This is why metals conduct electricity
- Amorphous solids, like glass, show no long-range structure. They are formed when liquids cool too quickly for a proper lattice to form
- Amorphous solids slowly grow softer over a wide temperature range, rather than melting sharply at a single melting temperature

## 8.15 Changes of State

- When a substance undergoes a phase change, it will either give off or take in heat without changing the temperature
- o The amount of heat is called the heat of fusion or heat of vaporization
- Figure 8.24 shows a heating curve across two phase changes
- $\circ~$  Heat for changing temperature:  $q=mC_{p}\Delta T$
- $\circ~$  Heat for a phase change:  $q=m\Delta H_{forv}$
- $\circ~7.5g$  of ice begin at  $-6.00^{\circ}C$ . Find how many J are required to convert this ice into steam at  $125^{\circ}C$  (23,000J)

For water: 
$$C_s=2.11\frac{J}{g\circ C}$$
  $C_l=4.18\frac{J}{g\circ C}$   $C_g=2.00\frac{J}{g\circ C}$  
$$\Delta H_{fus}=334\frac{J}{g}$$
 
$$\Delta H_{vap}=2260\frac{J}{g}$$

- Homework 8.3
  - · 8.92 Pressure and boiling points
  - · 8.96 Amorphous and crystalline solids
  - · 8.98 Enthalpy of fusion problem

#### CHAPTER 9

#### **SOLUTIONS**

### 9.1 Mixtures and Solutions

- Recall the taxonomy of mixtures from chapter 1: Heterogeneous mixtures vs homogeneous mixtures, and solutions vs colloids
- $\circ$  The primary difference is particle size: Solutions have particles <2nm, colloids have particles 2-500nm, and heterogeneous mixtures have particles >500nm
- o In a solution, the solvent is the dominant component while any minority components are solutes
- o Solutions can be more than just solid-in-liquid Table 9.2

### 9.2 The Solvation Process

- In solvation, solute-solute interactions and solvent-solvent interactions are replaced by solutesolvent interactions
- o To determine solubility, consider the various interactions
  - · Stronger solute-solute interactions will make it hard to break apart the solute
  - · Stronger solvent-solvent interactions will make it hard to disrupt solvent to make room for solute
  - · Similar solute-solute, solvent-solvent, and solute-solvent interactions lead to high solubility
  - · This leads to the rule of thumb that "like dissolves like"
- The solvent will form a solvation sphere around the solute
- The net result can be exothermic or endothermic

### 9.3 Solid Hydrates

- $\circ~$  Some ionic compounds will naturally incorporate water molecules in their ionic lattice
- $\circ$  These compounds are called "hygroscopic," and are written with a dot like this: CaSO<sub>4</sub>· $\frac{1}{2}$  H<sub>2</sub>O

## 9.4 Solubility

- Solubility is the amount of one substance that you can dissolve in another substance
- o If two liquids can be mixed in any ratio, then they are said to be miscible
- o If a solution has dissolved as much of a solid as it can, then it is called "saturated"
- o A saturated solution can be identified if excess solute remains undissolved

### 9.5 Temperature Dependence of Solubility

- o Increasing the temperature will increase the solubility of most, but not all solids
- If solvation is endothermic, then solubility increases with increasing temperature. If solvation is exothermic, then solubility will decrease with increasing temperature
- A solution can be heated to dissolve more solute, then cooled to form a supersaturated solution
- Solute can be spontaneously precipitated from a supersaturated solution Demo with sodium acetate

### 9.6 Pressure Dependence of Solubility

- Pressure will affect the solubility of gases in liquids
- o The solubility is proportional to the gas pressure, according to Henry's Law

#### 9.7 Units of Concentration

- There are several different units used by chemists to describe concentration
- Percent concentrations:
  - · mass/mass percent mass of solute / mass of solution
  - · volume/volume percent volume of solute
  - · mass/volume percent mass of solute volume of solution
- Homework 9.1:
  - · 9.38 Water as a solvent
  - 9.42 Henry's law
  - · 9.44 Saturation and solubility
  - · 9.50 Preparing a solution of NaCl

### 9.8 Dilution

- o Dilution is when additional solvent is added to reduce the concentration of all solutes
- The number of moles of solvent doesn't change, so we can set the number of moles after dilution equal to the number of moles before the dilution
- $\circ \ C_1V_1=C_2V_2$  in general, or  $M_1V_1=M_2V_2$  for molar concentrations
- $\circ \frac{V_1}{V_2}$  is the dilution factor

## 9.9 Ions in Solution: Electrolytes

- o Substances which produce ions when dissolved in solutions are called electrolytes
- Electrolytes conduct electricity
- Substances fall into three categories:
  - · Strong electrolytes completely ionize (produce one or more ions for each formula unit)
  - · Weak electrolytes only partially ionize (often only a fraction of a percent)
  - · Non-electrolytes produce no ions

# 9.10 Electrolytes in Biology

- In our biochemistry, there are often many different ions in solution
- To talk about the concentration of complex solutions, we use two new units:
  - Equivalents (Eq): The number of ions that carry 1 mole of charge
  - · Gram Equivalents (g-Eq):  $\frac{\text{MolarMassofIon(g)}}{\text{ChargeonIon}}$
- More realistic concentrations are milliequivalents

# 9.11 Properties of Solutions

- Some properties of solutions depend on the concentration but not the identity of the solute
- These properties are called colligative properties
- o Vapor-Pressure Lowering Adding a solute will lower the solvent vapor pressure
- $\circ$  Boiling Point Elevation  $\Delta T_b = \kappa_b C_{molal}$
- $\circ~$  Freezing Point Depression  $\Delta T_f = \kappa_f C_{molal}$

## 9.12 Osmosis and Osmotic Pressure

- o Osmotic pressure is another important colligative property
- $\circ~$  Osmosis Pressure exerted across a semipermeable membrane  $\pi = \frac{nRT}{V}$
- o Isotonic solutions match the salinity of blood cells
- $\circ\;$  hypertonic solutions are more salty and will shrivel blood cells
- o hypotonic solutions are less salty and will swell (or lyse) blood cells
- o Homework 9.2:
  - · 9.68 Dilution
  - · 9.74 milliequivalents
  - · 9.82 Freezing point depression
  - · 9.86 Osmolarity

#### CHAPTER 10

#### ACIDS AND BASES

#### 10.1 Acids and Bases in Aqueous Solution

- Acid and base activity is mostly defined in the context of an aqueous solution
- o An acid will react with water to produce hydronium ions
- o A base will react with water to produce hydroxide ions

#### 10.2 Some Common Acids and Bases

o Acids and bases are common in products around us. Page 292 lists a few

### 10.3 The Brønsted-Lowry Definition of Acids and Bases

- There are several different definitions used for acids and bases, we will be using the Brønsted-Lowry definition:
  - · Remember that a H<sup>+</sup> ion is simply a proton
  - · An acid is a proton donor HCl reaction with water
  - · A base is a proton acceptor NH<sub>3</sub> reaction with water
- Under this definition, water acts as a base when reacting with an acid, and acts as an acid when reacting with a base More on this later
- Multiprotic acids can donate more than one hydrogen
- Every acid has a conjugate base, and every base has a conjugate acid They make up a conjugate pair

# 10.4 Acid and Base Strength

- Strong acids and bases will dissociate into H<sup>+</sup> or OH<sup>-</sup> completely (and are strong electrolytes)
- Weak acids and bases dissociate to different extents, less than 100%
- The relative strengths of a conjugate pair are inverse to each other (i.e. The stronger and acid, the weaker will be its conjugate base, and vice-versa)

- Table 10.1 shows conjugate pairs over a range of acid and base strengths
- Acid-base reactions are equilibrium reactions, and the relative strengths determine whether the reaction is reactant or product favored

### 10.5 Acid Dissociation Constants

- $\circ$  The strength of an acid is quantified by the acid dissociation constant  $K_a$
- o This is really just th equilibrium constant for the reaction of acid with water

$$\circ \ \ K_a = \frac{[\mathrm{H_3O^+}][\mathrm{A}^-]}{[\mathrm{HA}]}$$

- Multiprotic acids will have a seperate constant for each stage of dissociation
- $\circ$  Table 10.2 Gives the  $K_a$  values for several common acids

#### 10.6 Water as Both an Acid and a Base

- Water acts as a base when reacting with an acid, and acts as an acid when reacting with a base
- Substances like this are called *amphoteric*
- Water can also react with itself, acting as both an acid and a base to produce both hydronium and hydroxide
- $\circ$  The water dissociation constant  $K_w$  is very important to acid-base chemistry and has the value  $[{\rm H_3O^+}][{\rm OH^-}]=1.0\times10^{-14}$
- $\circ~$  In a neutral solution,  $[\mathrm{H_3O^{\scriptscriptstyle +}}] = [\mathrm{OH^{\scriptscriptstyle -}}] = 1.0 \times 10^{-7}$
- $\circ~$  As  $[\mathrm{H_3O^+}]$  goes up,  $[\mathrm{OH^-}]$  goes down and vice-versa to obey the  $K_w$  expression
- Homework 10.1:
  - 10.44 Identify strong acids
  - · 10.46 Identify acids and bases
  - · 10.54 Equilibrium expression  $K_a$
  - 10.56  $K_w$

### 10.7 Measuring Acidity: pH

- o pH is a way to quantify the acidity of a solution
- $\circ pH = -\log_{10}[H_3O^+]$
- Taking the log squeezes the very wide range of possible hydronium concentrations into a reasonably small range.
- o pH = 7 is neutral, pH < 7 is acidic, and pH > 7 is basic
- We can also define a pOH:  $pOH = -\log_{10}[OH^{-}]$
- $\circ~$  With these definitions we can convert  $K_w = [\mathrm{H_3O^{\scriptscriptstyle +}}][\mathrm{OH^{\scriptscriptstyle -}}]$  into 14 = pH + pOH

### 10.8 Working with pH

- We find pH by taking the negative log of the hydronium concentration
- We can find the hydronium concentration by taking ten to the power of -pH
- o Draw my box of conversions

## 10.9 Laboratory Determination of Acidity

- One simple way to measure acidity is with a color indicator (Figure 10.3)
- This can take the form of a drop to add to a solution, or a piece of litmus paper to drop the solution onto
- Indicators are chemicals that change color based on the acidity of their environment
- More accurate measurements can be made electronically using a pH meter

#### 10.10 Buffer Solutions

- Pure water will rapidly change pH if any acid or base is added
- o Buffer solutions can withstand changes in pH, even if strong acids or bases are added
- Figure 10.5 compares pH changes in pure water vs a buffer solution
- A buffer solution contains a conjugate acid/base pair, which reacts in place of the water to resist pH changes
- $\circ$  To make an ideal buffer, pick an acid/base pair whose  $pK_a$  equals the desired pH you want to buffer at

• The Henderson-Hasselbalch equation helps to solve for the pH of a buffer solution that doesn't have exactly equal amounts of the conjugate acid and base

$$\circ pH = pK_a + \log_{10}\left(\frac{[A^-]}{[HA]}\right)$$

 Our own bodies are regulated by buffer systems (carbonic acid and dihydrogen phosphate, among others)

### 10.11 Acid and Base Equivalents

 $\circ$  Just like with ion equivalents, it is often convenient to talk about acids and bases in terms of how many moles of  $H_3O^+$  of  $OH^-$  they will produce

$$\circ \ 1eq = \frac{1mol}{\# \ of \ \mathbf{H_3O^+} \ or \ \mathbf{OH^-} \ ions \ produced}$$

$$\circ \ 1g - eq = \frac{Molar Mass}{\# \ of \ \mathbf{H_3O^+} \ or \ \mathbf{OH^-} \ ions \ produced}$$

- Since acids and bases will partition between their conjugates, we can't properly talk about Molarity without doing further calculations
- $\circ~$  What we really want is the number of equivalents in a solution. This is called the Normal Concentration (N)

$$\circ \ N = \frac{eq}{L}$$

#### 10.12 Some Common Acid-Base Reactions

### 10.13 Titration

- o Titration is a technique used to determine the concentration of acids and bases
- To titrate an acid, a base of known concentration is added dropwise until the the acid is neutralized
- To titrate a base, an acid of known concentration is added dropwise until the the base is neutralized
- o At the equivalence point, the number of moles of acid and base should be equal
- The equivalence point can be identified by a color indicator, since the pH swings rapidly once the neutralization has occurred
- $\circ N_1V_1=N_2V_2$  This looks just like the dilution equation because it is based on the same assumption: That the number of moles is the same

## 10.14 Acidity and Basicity of Salt Solutions

- o When a salt dissolves, the cation and anion dissociate and are free to react with water
- The cation may be an acid, and the anion may be a base or an acid, so we need to consider them both
- Remember that a salt is the product of an acid-base reaction, so we can classify salts by the type of reactions which will produce them:
  - Strong acid + weak base = acidic solution
  - Weak acid + strong base = basic solution
  - Strong acid + strong base = neutral solution
  - · Weak acid + weak base = ? (it depends on the specific acid/base strengths)
- Homework 10.2:
  - · 10.68 Rough calculations of pH
  - 10.70 pH to  $\left[ H_3O^+ \right]$  and  $\left[ OH^- \right]$
  - · 10.78 Find pH of a buffer
  - 10.90 Titration

#### **CHAPTER 11**

#### **NUCLEAR CHEMISTRY**

#### 11.1 Nuclear Reactions

- Atomic symbols keep track of all subatomic particles:  ${}_{6}^{12}$ C,  ${}_{6}^{13}$ C,  ${}_{6}^{12}$ C<sup>+</sup>, etc.
- We can describe changes in nuclear reactions using these symbols:

$$^{14}_{6}C \longrightarrow ^{14}_{7}N + ^{0}_{-1}e^{-}$$

 $\circ$  In the reaction above,  $_{-1}^{\circ}e^{-}$  (simply an electron) does not really have an atomic number, but the  $_{-1}$  is useful for balancing reactions

## 11.2 The Discovery and Nature of Radioactivity

- Radioactivity was first observed in 1896 when photosensitive paper stored under a Uranium sample was exposed despite being kept in the dark
- $\circ~$  Radiation is categorized as  $\alpha$  particles,  $\beta$  particles, or  $\gamma$  radiation
- o Figure 11.1 shows how different types of radiation respond to an electric field
- o Table 11.1 shows the physical properties of different types of radiation

# 11.3 Stable and Unstable Isotopes

- o Radioactivity comes from unstable nuclei
- $\circ \ \ Some \ unstable \ nuclei \ are \ naturally \ occurring \ (U), \ while \ others \ (Pu) \ are \ generated \ artificially$
- o Figure 11.2 shows the "belt of stability"

# 11.4 Nuclear Decay

 $\circ \ \ Alpha \ Emission-Figure \ {\bf 11.3}$ 

$$^{238}_{92}U \longrightarrow {}^{4}_{2}He + {}^{234}_{90}Th$$

$$^{208}_{84}$$
Po  $\longrightarrow$  ?

○ Beta Emission 
$${}^{1}_{0}n$$
  $\longrightarrow$   ${}^{1}_{1}p^{+}$   $+$   ${}^{0}_{-1}e^{-}$ 
 ${}^{131}_{53}I$   $\longrightarrow$   ${}^{131}_{54}Xe$   $+$   ${}^{0}_{-1}e^{-}$ 
 ${}^{55}_{24}Cr$   $\longrightarrow$  ?

o Gamma Emission – Often present with other decay (Exception is Hafnium battery)

$$^{60}_{27}$$
Co  $\longrightarrow ^{60}_{28}$ Ni  $+ ^{0}_{-1}$ e<sup>-</sup>  $+ ^{0}_{0}\gamma$ 

$$\circ \ \ Positron \ Emission - \ _{_{1}}^{_{1}}p^{_{+}} \longrightarrow \ _{_{0}}^{^{1}}n \ + \ _{_{1}}^{^{o}}e^{_{+}}$$

$$_{19}^{40}K \longrightarrow _{18}^{40}Ar + _{1}^{0}e^{+}$$

$$\circ \ \ Electron \ Capture - \ _{_{1}}^{^{1}}p^{_{+}} + {\underset{_{-1}}{^{\circ}}}e^{_{-}} \longrightarrow \ _{_{0}}^{^{1}}n$$

$$^{^{197}}_{80}Hg + {^{\circ}}_{^{-1}}e^{^{-}} \longrightarrow {^{197}}_{79}Au$$

## 11.5 Radioactive Half-Life

- $\circ$  The half-life is the time required for half of a radioactive sample to decay
- o Figure 11.4 shows how the absolute decay rate decreases, but the half-life is constant
- $\circ$  Fraction Remaining =  $(0.5)^n$
- o Table 11.3 shows the half-lives and uses for common radioisotopes

## 11.6 Radioactive Decay Series

- o Radioisotopes often decay into other radioisotopes, creating a cascade of different reactions
- o Figure 11.5 shows the Uranium radioactive decay series

# 11.7 Ionizing Radiation

- Radioactive products are most damaging when they eject electrons off of other materials
- Table 11.4 shows the penetrating distance of different types of ionizing radiation

## 11.8 Detecting Radiation

# 11.9 Measuring Radiation

#### 11.10 Artificial Transmutation

#### 11.11 Nuclear Fission and Nuclear Fusion

- $\circ~$  Fission and fusion both release tremendous amounts of energy, which manifest as lost mass according to  $E=mc^2$
- o Fission Figure 11.7

$$_{0}^{^{1}}n + _{92}^{235}U \longrightarrow _{56}^{142}Ba + _{36}^{91}Kr + 3_{0}^{^{1}}n$$

- Note that the production of 3 neutrons can lead to a chain reaction
- Fusion Combining smaller nuclei to form larger ones

$$_{1}^{1}H + _{1}^{2}H \longrightarrow _{2}^{3}He$$

$${}_{2}^{3}\text{He} + {}_{2}^{3}\text{He} \longrightarrow {}_{2}^{4}\text{He} + 2 {}_{1}^{1}\text{H}$$

$${}_{2}^{3}\text{He} + {}_{1}^{1}\text{H} \longrightarrow {}_{2}^{4}\text{He} + {}_{1}^{0}\text{e}^{+}$$

- Homework 11.1:
  - 11.40  $\alpha$  and  $\beta$  decay
  - 11.44 Predicting products of  $\beta$  decay
  - · 11.48 Balancing fission reactions
  - · 11.56 Half-life calculations