CHEM 1210 Lecture Notes OpenStax Chemistry 2e

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COURSE ADMINISTRATIVE DETAILS

- o My office hours
- o Intro to my research
- o Introductory Quiz
- Grading details
 - · Exams 40, Final 20, Online Homework 15, Book Homework 15, Quizzes 10
 - \cdot Online homework
 - · Frequent quizzes
- o Importance of reading and learning on your own
- o Learning resources
 - · My Office Hours
 - · Tutoring services https://www.suu.edu/academicsuccess/tutoring/
- Show how to access Canvas
 - · Calendar, Grades, Modules, etc.
 - · Quizzes
 - · Textbook
- Introduction to chemistry
 - · Ruby fluorescence
 - · Levomethamphetamine
 - · Submerged salt crystals grow due dynamic equilibrium
 - · Rubber band elasticity
 - · Structure of the periodic table
 - · Salt on ice and purifying hydrogen peroxide

ESSENTIAL IDEAS

1.1 Chemistry in Context

- Modern chemistry is the end result of thousands of years of humans trying to explain and control the materials around them
- Early forays into chemistry (such as alchemy) had deep mystical roots and often relied on serendipity to make good progress
- Modern chemistry is a rigorous science, relying on falsifiability and the scientific methd (Figure 1.4)
- We sometimes refer to chemistry as "The Central Science" (Figure 1.3)
- To adequately describe and understand chemical phenomena, we often talk from different perspectives
 - **Macroscopic Domain** This is what we observe with bulk substances. Two chemicals react to produce a new chemical
 - **Microscopic Domain** We now understand that all microscopic effects are governed by the behavior of *microscopic* actors (molecules, atoms, electrons, etc.)
 - **Symbolic Domain** Effectively communicating chemical ideas requires new language. Chemical formulas, equations, and mechanisms are all symbolic representations

All three domains are on display in Figure 1.5

Quiz 1.1 - Scientific Method

Homework 1.1

- o 1: Thinking in terms of Chemistry
- o 3: The scientific method
- 5: Domains of inquiry

1.2 Phases and Classification of Matter

- Three primary phases of matter are shown in Figure 1.5 (and 1.6)
- o Plasmas are like a gas, but with electrically charged particles
- Mass vs Weight (for very fine measurements, the difference matters even on Earth due to buoyancy)

- Figure 1.8 illustrates the *law of conservation of matter*
- Classifying matter (Figure 1.11)
 - · Pure Substances
 - * Elements (Anything on the periodic table of the elements)
 - * Compounds (Combinations of elements can have very different properties from their constituent elements)
 - · Mixtures
 - * Heterogeneous mixtures (variable composition)
 - * Homogeneous mixtures (i.e. solutions, continuous composition)
- o Table 1.1 shows the abundance of many elements on Earth
- o Atoms are the smallest particle of an element that has the properties of that element
 - · Thought-experiment of dividing a sample in half ad-infinitum
 - · Ancient atomic theories and modern Dalton atomic theory (discussed in detail later)
 - · Atoms are *very* small; smaller than we could even detect until recently
- Molecules are collections of atoms held together with chemical bonds (more nuanced definition later)
 - · Many elements occur naturally as molecules, rather than atoms
 - · Figure 1.14 shows many molecular elements and compounds

1.3 Physical and Chemical Properties

- Physical Properties: Properties which can be observed without changing the chemical identity of the substance
- Chemical Properties: Properties which can only be observed through chemical reactions (e.g. flammability, acidity, electrochemical potential, etc.)
- Physical Changes: Any change which perserves the *chemical identity* of the substance (including phase changes)
- o Chemical Changes: Changes which alter the chemical identities of one of more substance
- Extensive Properties: Depend on the size of hte system (double the size, double the property measurement, such as mass or volume)
- Intensive Properties: Independent of system size (density, temperature, most chemical properties)
- The periodic table groups elements according to their properties (Figure 1.22)
 - · Metals conduct electricity and heat, are maleable and ductile
 - · Non-metals are very diverse, but generally poor conductors
 - · Metalloids exist at the boundary and share properties with both metals and non-metals
 - · There are many other ways to group the elements, which we will learn later

Quiz 1.2 - Matter, Properties, and Change

Homework 1.2

- 17: Classifying matter
- o 27: Classifying changes

1.4 Measurements

- All measurements are composed of three parts:
 - The magnitude of the measurement (the number itself)
 - The unit of measurement used (g, kg, lbs, etc.)
 - The degree of uncertainty in the measurement (this is usually implicit, and covered in the next section)
- Units are an essential part of any measuement. Develop a habit of *always* including units in your work
 - $u_{rms} = \sqrt{\frac{3RT}{M}}$ example of how units can guide problem solving and "unit purgatory"
 - SI units are a collection of fundamental units from which all other units can be derived (Table 1.2)
 - · Metric prefixes make it more convenient to discuss very large or very small numbers (Table 1.3)
 - · Scientific notation is an even more general and robust way of representing numbers
 - * The quantity is represented by a number with the decimal after the first digit
 - $\boldsymbol{\ast}\;$ The magnitude is represented by a power of 10
 - $\boldsymbol{\cdot}$ Practice converting between normal numbers, metric prefixes, and scientific notation
 - . For temperature, we use both K and ${^{\circ}C}$ (But not ${^{\circ}F})$ $T(K) = T({^{\circ}C}) + 273.15$
 - · Derived units will combine the fundamental units in some way

volume: m^3 , L, ml velocity: m/s

density: kg/m^3 , g/cm^3 (Table 1.4)

energy: $1J \equiv kgm^2/s^2$

1.5 Measurement Uncertainty, Accuracy, and Precision

- $\circ \ \ Countable \ quantities \ are \ considered \ to \ be \ \textit{exact} \ (no \ uncertainty)$
- o Measurements (and groups of measurements) always have some degree of undertainty

- · Accuracy is how close a measurement is to the *true value* (usually unknown, but approximated by calibration with a well-known standard)
- · Precision is how finely a measurment is made (What is the margin of error)
- · Figure 1.27 and Table 1.5 illustrate the differences between precision and accuracy
- · Accuracy is usually improved through calibration, and moving forward we will usually assume that measurements are as accurate as an instrument allows
- · Precision is represented in the way we write the number, and can be improved with a better instrument or with repeat measurements
- Significant figures are the way that we represent precision in a number
 - · The number of digits conveys the degree of precision
 - Example of me saying I'm $6ft\ 2in$ tall, vs me saying I'm $6ft\ 1.6241434in$ tall
 - · For graduated measurements, we record one digit beyond the lowest graduation (Figure 1.26)
 - · For digital measurements, we record the number as it is given by the instrument
 - For any given number, we should track both the *quantity* of significant figures, and the *position* of the least-significant digit
 - · In a written number, digits are considered significant according to the following rules:
 - * All non-zeros are significant
 - * All captive zeros (between two other significant digits) are significant
 - * Trailing zeros are always significant
 - * Leading zeros are never significant
 - * For scientific notation, only the digits of the quantity (not the magnitude) count
 - * Logarithmic quantities follow different rules which we will revisit in CHEM 1220 (chapter 14)
 - * Note that for some numbers scientific notation is *required* to convey the correct precision $(3.0 \times 10^3 m)$
- Errors propogate when multiple measurements are used in a mathematical operation
 - · For addition and subtraction, the least significant digit of the answer will be in the same position as the least significant digit of hte least precise input
 - · For multiplicationa and division, the quantity of significant digits in the answer will match the quantity of significant digits of the input with fewest significant digits
 - · When rounding an exact 5 (no further digits beyond the 5), round up or down to make the last digit even
 - · Compound problems involve multiple types of operations
 - * Solve the problem in steps, applying the correct rule to each step
 - * Track the significant figures (quantity and position) for each intermediate answer, but do *not* truncate or round any of these answers
 - * Only round after the last step \circ Practice $\frac{12.3g+34g}{12.0cm^3+7.7cm^3}=2.4g/cm^3$ (wrong answer with premature rounding)

Quiz 1.3 - Significant Figures

Homework 1.3

- o 45: Scientific Notation
- o 49: Counting Significant Figures
- o 53: Significant Figures and Calculations

1.6 Mathematical Treatment of Measurement Results

- Some quantities are calculated based on two or more measurements (such as velocity and density)
- \circ These formulas can be used to relate all three quantities together (i.e. $velocity = \frac{distance}{time}$)
- The derived quantity can be interpreted as a *comversion factor* between the other two quantitites
- o Conversion factors and unit conversions
 - · Elementary school perspective of ft to in conversions
 - · Conversion factors are a ratio between two identical quantities
 - · Converting units involves multiplying by 1 in the form of a conversion factor
 - · Units guide the problem solving
- o Dimensional Analysis is a problem-solving framework based on a series of unit conversions
 - · Don't dive straight into calculations and equations
 - · Identify the units you expect for the answer
 - · Identify the starting point
 - · Create a plan to convert units from the starting point to the answer
 - · Carry out the calculations
 - Practice converting 65.0 miles/hour into m/s
 - · The "railroad ties" or "picket fence" method can help organize your work
- Dimensional analysis is not the only way to solve problems, but it is versatile and robust; usually my preferred choice
- Practice a more abstract problem: Find the $^{miles}/_{gal}$ if a car consumes 8036~g of gasoline while driving for 40.0~min at $75~^{miles}/_{hour}$

Quiz 1.4 - Dimensional Analysis

Homework 1.4

- o 65: Simple unit conversion
- $\circ~87$: Density from volume and mass
- o 89: Mass from volume
- o 91: Volume from mass

ATOMS, MOLECULES, AND IONS

2.1 Early Ideas in Atomic Theory

- o 1807 Dalton's Atomic Theory: (1, 2 and 5 are not strictly true) (Figures 2.2-2.4)
 - 1. Matter is composed of atoms
 - 2. Atoms of a given element all have identical properties to each other
 - 3. Atoms of one element differ in properties from elements of a different element
 - 4. Chemical compounds consist of atoms of different elements combined in a specific ratio
 - 5. Chemical reactions *rearrange* the atoms which are already there, but cannot create or destroy atoms
- Development of Dalton's theory:
 - Dalton relied on prior work by Proust who demonstrated the law of definite proportions (Table 2.1)
 - · This was not at all expected my analogy with bread, or metal alloys
 - · Dalton further noted that ratios of these proportions followed the law of multiple proportions (Copper (I or II) Chloride example in the book)

2.2 Evolution of Atomic Theory

- About a century later, scientists discovered that atoms are made of even smaller components
- o J. J. Thomson discovered the electron, and its charge/mass ratio (Figure 2.6)
- Millikan's oil drop experiment found the fundamental charge (and thus mass) of an electron (Figure 2.7)
- Figure 2.8 shows some early ideas of how the positive and negative charges were distributed in an atom
- Ernest Rutherford discovered the atomic nucleus, consisting of very concetrated positive charge (Figures 2.9 and 2.10)
 - "It was quite the most incredible event that has ever happened to me in my life. It was almost as incredible as if you fired a 15-inch shell at a piece of tissue paper and it came back and hit you."
- Different *isotopes* of atoms were discovered with techniqes that produced isotopically pure samples
- Finally, the neutron itself was discovered in 1932, explaining what particle led to different isotopes

2.3 Atomic Structure and Symbolism

- o Atoms are made up of protons, neutrons, and electrons
- Figure 2.11 shows the small scale of the atom and nucleus
- Atomic units make discussions about atoms convenient
 - · The Atomic Mass Unit amu, Da, or $u=1.6605\times 10^{-24} g$
 - The fundamental charge $e = 1.602 \times 10^{-19} C$
 - The Angstrom Å= $10^{-10}m$
- Table 2.2 summarizes the properties of elementary particles in atoms
- We track the composition of an atom with three numbers:
 - The atomic number is the number of protons Z = p
 - The mass number is the number of protons and neutrons A = p + n
 - The number of neutrons is therefore n = A Z
 - The charge is the protons minus the electrons q = p e
 - · The number of electrons is e = p q
- o A positively charge atom is called a cation, and a negatively charged atom is called an anion
- Chemical symbols are a shorthand way of representing everything we need about an atom
 - There is a 1 or 2 letter symbol for each element (Table 2.3 shows some make sense, some don't)
 - · A is written as a left superscript
 - $\cdot \,\, Z$ is written as a left subscript, but can be left off
 - q is written as a right superscript with the magnitude first, then the sign. q is left off if q=0
 - Example: ${}_{6}^{13}C^{2+}$ (6 protons, 7 neutrons, 4 electrons)
- o Isotopes are different versions of elements with different mass numbers
 - · For the most part, different isotopes of an element behave exactly the same in chemisry
 - · Isotope abundance can be found by mass spectrometry, among other methods (Figure 2.15)
 - Table 2.4 shows the natural abundances of the isotopes of several light elements
 - · Atomic weight (atomic mass) is the weighted average of all the isotopes found on the Earth

$$M = \sum_{i} mass_{i} \times \frac{\%abundance_{i}}{100\%}$$

2.4 Chemical Formulas

- We can represent the actual structure and makeup of molecules at several levels of abstraction (Figures 2.16 and 2.17)
- o Molecular formulas
 - Each element is listed, with the number of atoms for each element written as a subscript (H₂O)
 - The order of elements follows certain patterns, with the least electronegative element often written first
- o Structural Formulas show how atoms are connected with covalent bonds represented as lines
- $\circ~$ Ball and Stick models show the three-dimensional geometry of a molecule
- o Space-filling models show the actual volume of space taken up by each atom in a molecule
- Figure 2.18 illustrates the difference between subscripts within a formula, and stoichiometric coefficients in front of formulas
- Empirical formulas show the mathematically simplified ratios of elements
 - · Some experimental techniques (especially early ones) could *only* give the empirical formula
 - · Ionic compounds are always reported with the empirical formula
 - · To find the empirical formula, divide all subscripts by their greatest common factor
 - · Some very different compounds share an empirical formula (carbohydrates CH₂O include formaldehyde, acetic acid, and sugar)
 - The molecular formula can be calculated from the empirical formula's weight and the molecular weight (Chapter 3)
- Arranging the same group of atoms in different ways produces different isomers
 - · Isomers share the same chemical formula, but can have very different properties
 - · Structural isomers differ in how the atoms are connected to each other (Figure 2.23)
 - · Optical isomers (or spatial isomers) are non-superimposable mirror images (Figure 2.24, glove analogy)

Quiz 2.1 - Atomic Theories

Homework 2.1

- 7: Properties of protons and neutrons
- 11: Atomic symbols from composition
- 19: Composition from atomic symbols
- 23: Atomic weight
- o 29: Molecular and empirical formulas

2.5 The Periodic Table

- As scientists discovered and studied more and more elements, they started to notice certain natural groupings according to physical and chemical properties
- Mendeleev arranged the atoms according to these groups and atomic weight, producing the first periodic table (Figure 2.25)
- Mendeleev even predicted the existence of and properties of yet-undiscovered elements based on gaps in his table
- Vocabulary around the periodic table:
 - · Rows are periods or series
 - · Columns are groups or families
- Figure 2.26 is a typical periodic table, showing the metals, non-metals and metalloids (contrast with my preferred table)
- o Figure 2.27 shows many of the names we use for important groups of elements
- The structure of the periodic table encodes rich information about the electrons in the elements, as we will learn in chapter 6

Quiz 2.2 - Periodic Table

Homework 2.2

- o 37: Classifying elements
- 41: Using group names

2.6 Ionic and Molecular Compounds

- An atom which gains or loses electrons (carries a charge) is called an *ion*
 - · Positively charged ions are called cations, and are smaller than their neutral atom counterparts (Figure 2.28)
 - · Negatively charged ions are called anions, and are larger than their neutral atom counterparts
 - We can predict which charge different elements will naturally tend to take based on their position in the periodic table (Figure 2.29)
 - · Many other elements can take two or more charges, especially the transition metals
- Some ions are composed of more than one atom and are called polyatomic ions
 - Table 2.5 gives some common polyatomic ions. Memorize the formula, name, and charge of these and their acids

- · Notice some trends in the names of oxyanions (per-ate, -ate, -ite, and hypo-ite)
- · We will learn about the acid names listed here (and more) in section 2.7
- Ionic compounds are held together by ionic bonds (coulombic attractions)
 - · Show dot diagram of how NaCl and CaCl form from elements
 - · Metal + non-metal is not an adequate definition of ionic compounds (NH,NO₂)
 - · Ionic compounds form an extended lattice of ions (covered more in CHEM 1220)
 - · Ions will combine to form neutral compounds
 - · Practice producing compound formulas from ions (include paranthesis for polyatomic ions where appropriate)
 - · Practice finding ions from formulas of compounds
- Molecular compounds are held together by covalent bonds (shared electrons)
 - Show a dot diagram of H_2 , H_2O , O_2 , and N_2
 - · Bonds between non-metals are covalent bonds
 - · Molecular compounds combine non-metals into discrete units called molecules
 - · Single, double, and triple bonds involve sharing two, four, and six electrons

2.7 Chemical Nomenclature

- Naming ionic compounds
 - Cation names are the name of the element, with the charge in roman numerals in parenthesis *if* the element could take multiple charges
 - · Anion names are the element name with an "-ide" ending (some element like P remove more than others)
 - · Polyatomic ion names are the same as you learned earlier
 - · For ionic compounds the name is simply cation name + anion name
 - · There is no indication of the quantity of atoms, that is inferred from charge neutrality
 - · Practice getting formulas from names and names from formulas
- Naming hydrates
 - · Some ionic compounds will incorporate water into their ionic lattice
 - · The formulas will have a \cdot then list the number of waters
 - · The names add the degree of hydration using prefixes from table 2.10 and adding "hydrate"
 - · The waters can be driven of with high temperature, producing the *anhydrous* form

Quiz 2.3 - Naming Ionic Compounds

Homework 2.3

- o 47: Predicting bond type in compounds
- o 49: Formulas from ions
- o 51: Names from formulas
- o 57: Names from formulas with transition metals
- o 59: Formulas from names

Resuming section 2.7 Chemical Nomenclature

- o Naming molecular compounds
 - · There are many ways to name molecular compounds, we will focus on just two here
 - · Naming binary molecular compounds
 - * # + name + # + name with "-ide" ending
 - * Least electronegative element (leftmost on the periodic table) goes first
 - * #s come from table 2.10
 - * Omit "mono-" for the first element
 - * Practice going from formula to name and vice-versa (Table 2.11)
 - · Naming molecular acids
 - * Molecular acid names are based on the name of the anion formed when all H⁺ are removed
 - * -ide ions form hydro-ic acids (Table 2.12)
 - * -ate ions form -ic acids
 - * -ite ions form -ous acids
 - * preserve the hypo- and per- prefixes
 - * Table 2.13 shows some oxyacid names

Quiz 2.4 - Naming Molecular Compounds

No textbook homework to accompany this quiz due to a lack of appropriate questions!

COMPOSITION OF SUBSTANCES AND SOLUTIONS

3.1 Formula Mass and the Mole Concept

- o For chemists, the amount of a substance we care about is not grams, but the quantity of atoms
- Analogy of hammers and feathers
- \circ The mole is defined as the relationship between AMUs and g

$$6.022\times 10^{23} AMU = 1g \longrightarrow 1 \\ mol AMU = 1g \longrightarrow 1 \\ AMU = 1 \\ \frac{g}{mol}$$

- Any time we would use the AMU we could replace it with g/mol, and this is called the *molar mass*
- Practice converting between grams and moles using atomic weight
- For compounds, we can find the molar mass, or formula mass by adding up the atomic weights of all the atoms in the formula
- We can count the literal number of atoms, but it is usually best to remain in units of moles
- Practice finding formula masses for compounds, and converting between grams and moles for compounds

3.2 Determining Empirical and Molecular Formulas

- o Empirical analysis is a method to determine empirical formulas
 - · In our text, the masses of C, H, and O are given directly
 - To get the formula, turn each mass into moles, and divide by the smallest mole number (Figure 3.11)
 - · Actually, this technique involves a careful combustion reaction
 - The mass of C comes from the amount of CO₂ produced
 - The mass of H comes from the amount of H₂O produced (remember the 2!)
 - The mass of O comes from subtracting those from the original amount
 - · Turn the masses to moles
 - Practice: 2.443g of unknown produce 5.368g of CO_2 and 2.930g of H_2O (M=60.096, propanol)
 - · Practice: 3.450g of unknown produce 5.057g of CO_2 and 2.070g of H_2O (M=180.16, hexose)
- % Composition gives the % by mass of each element in a compound

- · From masses, we can calculate the % mass composition easily
- · If % mass is given, we can just assume 100.0g and turn the %s into g to do emipirical analysis
- · From the empirical or molecular formula, we can get % mass composition: $\%_X = \frac{n M_X}{M_{Formula}}$
- We can determine the molecular formula from the empirical formula if we know the molar mass
 - The molecular formula will be an integer multiple of the empirical formula (could be 1)
 - That formula is the ratio $n = \frac{M_{molar}}{M_{empirical}}$
 - · Find the molecular formulas of the two examples above using their molar masses

Quiz 3.1 - Molar Mass

Homework 3.1

- o 3: Calculating molar mass
- o 17: Mass from moles
- o 18: Moles from mass (sorry, no solution!)

Quiz 3.2 - Empirical Analysis

Homework 3.2

- o 33: % composition from formulas
- o 37: Empirical formulas from % composition
- o 39: Molecular formula from % composition and molar mass

3.3 Molarity

- Molarity is the most common way of describing the *concentration* of a solute in a solution
 - \cdot The unit for molarity is M, which is unfortunate because that is also used for molar masses
 - Molarity is the moles of solute divided by the liters of solution $M=\frac{mol_{solute}}{L_{solution}}$
 - · Technically, the solution volume could be more or less than the solvent volume before dissolving. When we make a solution, we always describe the process as "add enough water to meet the mark"
 - · The molarity can now be used as a conversion between moles and volume
 - · Practice: Find molar concentration of solution made by dissolving $13.5g~{\rm NaCl}$ in 100.0ml of solution

- · Practice: How many moles of NaCl are contained in 35.62ml of the solution?
- · Practice: How many ml would be needed to provide $1.74 \times 10^{-3} mol$ of NaCl?
- o Solutions can be diluted or concentrated
 - The number of moles of solute is the same, just the volume of solution changes (by adding or removing solvent)
 - $\cdot C_1V_1 = C_2V_2$
 - Practice: Find the molar concentration when 36.43ml of 1.5M HCl are diluted to 100.0ml?
 - · Practice: How many ml of 0.125M NaNO $_3$ should be used to make 50.0ml of $2.4\times 10^{-3}M$?

3.4 Other Units for Solution Concentration

- There are other, less common units of concentration (including some not covered until chapter 11)
- o Mass %, or % by mass

$$\cdot \%m/m = \frac{m_A}{m_{total}} 100\%$$

o Volume %, or % by volume

$$\cdot \%v/v = \frac{V_A}{V_{solution}} 100\%$$

o m/V %

$$\cdot \%m/v = \frac{g_A}{V_{solution}} 100\%$$

o ppm and ppb

$$ppm = \frac{m_A}{m_{total}} 10^6 ppm$$

$$\cdot ppb = \frac{m_A}{m_{total}} 10^9 ppb$$

Quiz 3.3 - Concentration

Homework 3.3

- 49: Mass from molarity and volume
- 53: Calculating molarity
- o 61: Dilution
- o 73: ppm
- o 77: %m/m

STOICHIOMETRY OF CHEMICAL REACTIONS

4.1 Writing and Balancing Chemical Equations

- o A chemical equation relates the amounts of reactants and products in a chemical reaction
- \circ Consider the reaction: CH₄ + 2O₂ \longrightarrow CO₂ + 2H₂O (Figure 4.2)
- Reactants on the left, and products are on the right
- The large numbers are called *stoichiometric coefficients*
- o The subscript numbers are part of the chemical identity of the reactants and products
- The stoichiometric coefficients tell the right ratio to combine reactants, and the ratio of products produced (Figure 4.3)
- o Balancing chemical reactions:
 - To balance a chemical equation, we will add the proper coefficients to give the same numbers and types of atoms on both sides of the reaction arrow
 - · Start with any repeated atom moieties, such as polyatomic ions, and balance them as a unit rather than counting individual atoms
 - · Next balance elements that appear in only one compound on each side
 - Finally, balance any remaining elements (O is often best left to balance last)
 - · If necessary, multiply all coefficients by an integer

Practice: Balance the following chemical equation: $Fe_2O_2(s) + Al(s) \longrightarrow Al_2O_2(s) + Fe(s)$

$$\operatorname{Fe_2O_3}(s) + 2\operatorname{Al}(s) \longrightarrow \operatorname{Al_2O_3}(s) + 2\operatorname{Fe}(s)$$

Practice: Balance the following chemical equation: $C_8H_{18}(l) + O_2(g) \longrightarrow CO_2(g) + H_2O(g)$

$$2 C_8 H_{18}(1) + 25 O_2(g) \longrightarrow 16 CO_2(g) + 18 H_2O(g)$$

Practice: Balance the following chemical equation: $Ag_2SO_4(aq) + NaCl(aq) \longrightarrow AgCl(s) + Na_2SO_4(aq)$

$$Ag_2SO_4(aq) + 2 NaCl(aq) \longrightarrow 2 AgCl(s) + Na_2SO_4(aq)$$

- o Chemical reactions will sometimes include additional information
 - · Phases (s), (l), (g), (aq)
 - · Reaction conditions, such as temperature, or the presence of a catalyst (often written over the arrow)
 - Enthalpy of reaction or equilibrium constant (covered in later chapters)

- o Reactions with ionic compounds can often be written differently and simplified
 - The normal equation is called a *molecular equation* (despite involving ionic compounds) $CaCl_2(aq) + 2 AgNO_3(aq) \longrightarrow Ca(NO_3)_2(aq) + 2 AgCl(s)$
 - The *complete ionic equation* will break *soluble* ionic compounds into the separate cations and anions
 - This is a more accurate representation of soluble ionic compounds, which are not actually *to-gether* in any chemically meaningful way

$$Ca^{2+}(aq) + 2Cl^{-}(aq) + 2Ag^{+}(aq) + 2NO_{3}^{-}(aq) \longrightarrow Ca^{2+}(aq) + 2NO_{3}^{-}(aq) + 2AgCl(s)$$

- The *net ionic equation* removes *spectator ions*, which didn't actually participate in any chemical change
- · My silly analogy about party-goers and the social interactions in a party environment $Ag^+(aq) + Cl^-(aq) \longrightarrow AgCl(s)$

Quiz 4.1 - Balance Chemical Equations

Homework 4.1

- o 5: Balancing equations
- o 11: Net ionic equations

4.2 Classifying Chemical Reactions

- We can categorize chemical reactions in several ways, but for this class we will focus on the types of chemical changes caused by a reaction
- o Precipitation reactions form one or more solid ionic product from all aqueous ionic reactants
 - · First, predict the products by switching cations and anions
 - · Next, predict the phase by using solubility rules (Table 4.1)
 - · Identify and eliminate the spectator ions to get a net ionic equation
 - · Some combinations of reactants will have no reaction, and some may have two solid products
 - · More about precipitation reaction in CHEM 1220

Quiz 4.2 - Precipitation Reactions

Homework 4.2

o 28: Predicting products of reactions with ionic compounds

Resuming Section 4.2: Classifying Chemical Reactions

- Acid-Base reactions involve the transfer of a H⁺ between reactants
 - · Acids will react with water to produce hydronium ion
 - · Bases will react with water to produce hydroxide ions
 - · *Strong* acids/bases react with water to completion, while *weak* acids/bases only react partially (equilibrium)
 - Table 4.2 lists six common strong acids
 - · Alternative definition: Acids are proton donors and bases are proton acceptors
 - · A neutralization reaction produces water and a salt (ionic compound)
 - · More about acid/base reactions in CHEM 1220
- Redox reactions involve the transfer of one or more electrons
 - · Consdier the reaction: $2 \text{ Na(s)} + \text{Cl}_2(g) \longrightarrow 2 \text{ NaCl(s)}$
 - · OIL RIG (or LEO says GER) to remember which is oxidation, and which is reduction
 - · Oxidizing agents and reducint agents. "Agent" describes the effect on the reaction partner
 - · Sometimes the electron transfer is not as obvious: $CH_4 + 2O_2 \longrightarrow CO_2 + 2H_2O_3$
 - · We assign oxidation numbers to keep track of electrons
 - * Elements have ox. # of 0
 - * Monoatomic ions have ox. # equal to their charge
 - * Within compounds, O has ox. #=-2, and H has ox #=+1
 - * Oxidation #s add up to the total overall charge (this rule subsumes the first 2)
 - · Balancing redox reactions needs additional steps to account for the transferred electrons
 - 1. Split the reaction into half-reactions (need to find oxidation #s to do this)
 - 2. Balance all elements except H and O
 - 3. Add the electrons, based on the changes to oxidation #s
 - 4. Balance charge by adding H⁺ in acid, or OH⁻ in base
 - 5. Balance H and O together by adding water
 - 6. Multiply half-reactions to balance their electrons, then add them together
 - 7. Make any cancellations of water, H⁺, or OH⁻
 - · You can balance redox reactions as a whole instead of as half-reactions if you prefer
 - · Electrochemical series and predicting if redox reactions will be spontaneous or not (Not in the book!)
 - · More about Redox reaction in ... CHEM 1220!

Quiz 4.3 - Redox Reactions

Homework 4.3

- o 17: Assigning oxidation states
- o 19: Classify acid/base and redox reactions
- o 39: Balancing half-reactions

4.3 Reaction Stoichiometry

- Measured amounts should always be converted to moles in order to make comparisons to other chemical species in an equation
 - · We often want to calculate the corresponding amount of different chemicals in a chemical reaction
 - · Making direct comparisons will not work due to different molar masses, etc.
 - · Figure 4.11 shows how to convert to moles from different measurements
 - · My version of this figure ("Chemistry Land")
- Practice: $N_2(g) + 3H_2(g) \longrightarrow 2NH_2(g)$ Find other amounts for 0.75g of H_2

Quiz 4.4 - Stoichiometry

Homework 4.4

- o 49: Stoichiometry with solid reactants
- 57: Stoichiometry with aqueous reactions

4.4 Reaction Yields

- Often one or more reactants are provided in excess, with only one *limiting reactant*
 - · Pick a product, and calculate how much product would be produced for each reactant
 - · The lowest amount is the theoretical yield. The other amounts can be discarded
 - The reactant which lead to the lowest amount is the limiting reactant
 - · Calculate the rest of the corresponding amounts starting from the limiting reactant
 - · Double-check your work using the conservation of mass
- $\circ \ \ \text{Practice: 2 C}_4\text{H}_{\text{10}} + \text{13 O}_2 \ \longrightarrow \ 8 \text{CO}_2 + \text{10 H}_2\text{O with } 5.00g\text{C}_4\text{H}_{\text{10}} \text{ and } 9.00g\text{O}_2$
- Reactions rarely go perfectly, and the actual amount of product will be different from the theoretical yield

$$\circ \ \%yield = \frac{actual \ yield}{theoretical \ yield} \times 100\%$$

- \circ Yields below 100% can mean that the reaction didn't go to completion, or some product was lost during a purification stage
- $\circ~$ Yields above 100% usually indicate that some contaminants (including perhaps excess reactant) are mixed in with the product

Quiz 4.5 - Limiting Reactants

Homework 4.5

- o 61: Limiting Reactant Problem
- o 63: Percent Yield

4.5 Quantitative Chemical Analysis

- Quantitative analysis is the family of techniques which determine the amount of substance in a sample
- o Titrations find the concentration of an aqueous *analyte*
 - · React the unknown with a co-reactant of known concentration by adding titrant with a buret
 - · The chemical environment will change (pH, redox potential, etc.) once the reaction reaches the equivalence point
 - · Probes or color indicators can show when to stop the titration. This is called the end-point, and is ideally very close to the equivalence point

$$\cdot \ \frac{C_T V_T}{\nu_T} = \frac{C_A V_A}{\nu_A}$$

- Gravimetric analysis uses a chemical reaction to change the phase of the analyte so it can be separated and weighed
 - · Your % copper lab found the amount of copper by reducing it to a solid
 - \cdot Your hydrates lab found the amount of water by weighing it before and after the baking step
- o Combustion analysis We actually covered this earlier with % composition of compounds

Quiz 4.6 - Titrations

Homework 4.6

- o 79: Acid/base Titration
- 81: Precipitation Titration

THERMOCHEMISTRY

5.1 Energy Basics

- o Thermochemistry is the study of heat and energy changes in chemical reactions
- It also includes topics like entropy and spontaneity
- o Energy can come in two forms, kinetic and potential (Figure 5.3)
 - · Kinetic energy is the energy of motion: $KE = \frac{1}{2}mv^2$
 - For chemists, the kinetic energy that matters is the movement of individual atoms, molecules, etc., manifested as temperature (Figure 5.4)
 - · Heat will always flow from colder objects to warmer objects (Figure 5.6)
 - · Increases in temperature almost always cause objects to increase in volume (Figure 5.5)
 - · Potential energy is stored energy: Gravitaional, electrostatic, chemical, etc.
 - For chemists, the potential energy that matters is the energy of chemical bonds and intermolecular forces (Figure 5.2)
- Energy has several common units
 - The SI unit is the Joule: $1 J = 1 kg \frac{m^2}{s^2}$
 - The calorie: $1 \ cal = 4.184 \ J$
 - · The Calorie (kcalorie): 1 Cal = 1000 cal = 4184 J
- When we talk about changes and transfers of energy, we need to carefully define our system
 - The *system* is the part of the universe where the reaction occurs, such as a beaker or chamber with reactants
 - · The *surroundings* is the rest of the universe
 - · Open systems can exchange both heat and matter with the surroundings (an open beaker)
 - · Closed systems can exchange heat, but not matter with the surroundings (a closed chamber)
 - · Isolated systems cannot exchange either heat or matter with the surroundings (a closed, insulated flask)
- Heat from chemical and physical changes is usually associated with temperature changes or phase changes (Figure 5.7)
 - These examples all refer to heat from the *system's* perspective

- · Positive heat is observed by an "upward" phase change or a cold temperature
- · Holding ice in your hand will melt the ice, and make your hand cold
- · Negative heat is observed by a "downward" phase change or a hot temperature
- · Burning wood in a fire feels warm because of the negative system heat
- Heat and temperature change are related by several equations (Note that we are always measuring the temperature of the *surroundings*)
 - · When heat is added to a system, it will either undergo a phase change, or heat up
 - · The specific heat gives how much heat is required to warm a given substance
 - · For a complete system, $q = C\Delta T$ where C is the heat capacity
 - · For a pure substance, $q=mc_S\Delta T$ where c_S is the *specific heat* of the substance and m is the mass
 - Table 5.1 shows c_S for many common substances

 Practice: Find how much heat is required to raise the temperature of 15.0g of iron by $32^{\circ}C$ Practice: Find the temperature change when 42.5J of heat are added to 0.374g of aluminum

Quiz 5.1 - Energy

Homework 5.1

- 5: Heat capacity of samples
- \circ 9: $q = mc_S \Delta T$
- 11: $q = mc_S \Delta T$, but solve for c_S

5.2 Calorimetry

- Whenever heat is transferred, the total energy of the universe remains constant
- \circ For a heat transfer in an isolated system, $q_1 = -q_2$
- Consider a block of hot metal placed in a beaker of room temperature water. Heat will flow from the block into the water until the two temperature are equal to each other
- $\circ \ \ q_{1} = -q_{2} \ \text{becomes} \ m_{1}c_{1} \left(T_{f} T_{i,1} \right) = -m_{2}c_{2} \left(T_{f} T_{i,2} \right)$

Practice: A 10.0~g block of iron is heated to $93.5~^{\circ}C$ and placed in 25~ml of $23.0~^{\circ}C$ water. What is the final temperature? (25.9 $^{\circ}C$)

- We can also measure the heat transfer associated with a chemical reaction, called *calorimetry*
- Exothermic processes release heat into the surroundings (warm to the touch)
- Endothermic processes absorb heat from the surroundings (cool to the touch)

- o Figure 5.11 shows how a calorimeter would work for endothermic and exothermic processes
- Figures 5.12 and 5.13 show a constant pressure (coffee-cup) calorimeter
 - · The reaction is carried out in aqueous solution
 - · q_{rxn} is the heat released or absorbed by the reaction
 - · The heat of reaction is exchanged with the solution: $q_{rxn} = -q_{soln} = -mc\Delta T$
 - $\cdot m$ is the solution mass, which will include the water and any solutes
 - · c is the solution specific heat, but this is simplified by assuming $c_{soln} = c_{water} = 4.184 \frac{J}{q \circ C}$

$$\cdot \ \Delta H = \frac{q_{rxn}}{n_{rxn}} = \frac{-mc\Delta T}{n_{rxn}} \ \text{where} \ n_{rxn} \ \text{is the moles of reaction:} \ n_{rxn} = \left(\frac{n_A}{\nu_A}\right)$$

- Demo NaOH enthalpy of solvation $\left(10~g,100~ml,44.2~\frac{kJ}{mol}\right)$
- Figure 5.17 shows a constant volume (bomb) calorimeter
 - · The reaction is carried out in a chamber charged with high pressure O₂
 - · q_{rxn} is exchanged with the whole bomb-calorimeter apparatus
 - The calorimeter is calibrated to give a heat capacity (C_{cal}) with units $\frac{J}{\circ C}$

$$q_{rxn} = -q_{cal} = -C_{cal}\Delta T$$

 \cdot For constant volume, we measure U instead of H because w=0

$$\cdot \ \Delta U = \frac{-C_{cal}\Delta T}{n_{rxn}}$$

Quiz 5.2 - Calorimetry

Homework 5.2

- o 19: Reaching thermal equilibrium
- 25: Coffee cup calorimetry
- o 31: Bomb calorimetry

5.3 Enthalpy

- $\circ~$ The sum of all types of energy in a system is the $\it internal\, energy, U$
- \circ Any change in the internal energy must come from heat or work: $\Delta U = q + w$
- \circ Work is defined as $w = f \cdot d$ or $w = -P \cdot \Delta V$
 - · Positive work is when the system volume decreases
 - · Negative work is when the system volume increases, or a force moves part of the surroundings
 - \cdot This PV work is actually a bit problematic when trying to keep track of energy

- * T is easy to measure with a thermometer, but both P and (especially) V are more difficult to measure
- * Most of our work as chemists is done at constant pressure (open flask or in a balloon)
- * Under constant pressure, we can use Enthalpy(H) instead of internal energy (U)
- * Technically, H = U + PV
- * While U = q + w, H = q under constant pressure conditions (derivation in the book)
- * So, we only need to worry about heat when we deal with H
- Heat is usually associated with temperature changes or phase changes, as discussed previously
- First Law of Thermodynamics: The energy of the universe is constant
- Enthalpy in chemical reactions:
 - · A balanced chemical reaction may also include an enthalpy of reaction ΔH
 - · This tells how much heat is produced or consumed with one mole of reaction
 - · ΔH can be a conversion factor between heat and amounts of reactants or products Practice: Consider the reaction N₂(g) + 3 H₂(g) \longrightarrow 2 NH₃(g) $\Delta H = -92 \frac{kJ}{mol}$ If 2.25~g of H₂ are consumed in the above reaction, how much heat is released? (34.2 kJ)

If $54.6 \ kJ$ of heat are released, how many g of NH₃ will be produced? (20.2 g)

Quiz 5.3 - Enthalpy

Homework 5.3

- 41: Enthalpy of reaction from calorimetry
- 45: Heath from enthalpy of reaction
- 47: How much reactant from target heat output

Resume Section 5.3 – Enthalpy

- \circ The "Standard State" is notated by \circ and is at 1M concentration and 1atm (or 1bar), and technically doesn't include a temperature but values are often tabulated at $25^{\circ}C$
- \circ Standard Enthalpies of Combusion (ΔH_C°):
 - · Standard enthalpy of combustion is the energy released when something reacts with oxygen
 - · Table 5.2 Lists the enthalpies of combustion for many combustible substances
 - These values are not synonymous with energy density, due to molar mass and density complicating the values
- $\circ~$ Because enthalpy is a state function (Figure 5.20), we can calculate values of ΔH without measuring them

- Hess's Law: Any alternate path with the same starting and ending states will have the same overall ΔH
 - · Drawing an energy level diagram can help to illustrate Hess's law (Figure 5.24 as an example)
 - · Find ΔH for this reaction: C_{diamond} + O₂(g) \longrightarrow CO₂(g) $(\Delta H = -395.4 \, ^{kJ}/_{mol})$ $C_{diamond} \longrightarrow C_{graphite}$ $\Delta H = -1.9 \, ^{kJ/mol}$ $C_{graphite}$ + O2(g) \longrightarrow CO2(g) $\Delta H = -393.5 \, ^{kJ}/_{mol}$
 - The reverse of a reaction gives $-\Delta H$
 - Find ΔH for this reaction: C(s) + $\frac{1}{2}$ O2(g) \longrightarrow CO(g) $(\Delta H = -110 \, kJ/mol)$ $\mathrm{C(s)} + \mathrm{O_2(g)} \longrightarrow \mathrm{CO_2(g)} \qquad \Delta H = -393 \, ^{kJ/mol}$ $CO(g) + \frac{1}{2}O_2(g) \longrightarrow CO_2(g)$ $\Delta H = -283 \, kJ/mol$
 - · Double the reaction gives double the ΔH
 - · Consider trying to find ΔH for the reaction below:

• Find an alternate path using these reactions with known ΔH :

· The enthalpy of the first reaction can be found from the enthalpies of the other two

•
$$\Delta H_{\star} = \Delta H_A - 2\Delta H_B = -801 \frac{kJ}{mol}$$
 (Draw the energy level diagram)

Practice: Find the enthalpy of reaction (\star) using reactions (A), (B), and (C)

$$(\star) \operatorname{CS}_2(l) + 3 \operatorname{O}_2(g) \longrightarrow \operatorname{CO}_2(g) + 2 \operatorname{SO}_2(g) \qquad (\Delta H = -1075.0)$$

$$(A) \operatorname{C}(s) + \operatorname{O}_2(g) \longrightarrow \operatorname{CO}_2(g) \qquad \Delta H = -393.5 \, {^kJ/mol}$$

$$\Delta II = -393.3 \text{ mor}/\text{mor}$$

$$(B) S(s) + O2(g) \longrightarrow SO2(g) \qquad \Delta H = -296.8 \, {}^{kJ}/mol$$

$$(C)$$
C(s) + 2 S(s) \longrightarrow CS₂(l) $\Delta H = 87.9 \, ^{kJ}/_{mol}$

Quiz 5.4 - Hess's Law

Homework 5.4

- o 59: Hess's Law and enthalpy of reaction
- o 63: Hess's Law and enthalpy of reaction

Resume Section 5.3 – Enthalpy

- \circ Standard Enthalpies of Formation ΔH_f°
 - · To apply Hess's law to arbitrary reactions, you would need to devise an alternate path from an encyclopedia of known reactions – this would be *very* inconvenient

- Instead of using random reactions from one state to another, it is useful to devise a *standard state* for each element
- · The standard state is the most stable form of that element e.g. for O, it is $O_2(g)$, not $O_2(l)$ or $O_3(g)$
- Each compound will have a *standard formation reaction* which forms it from its elements in their standard state

For water, that's $H_2(g) + \frac{1}{2}O_2(g) \longrightarrow H_2O(l)$ (This is one time a $\frac{1}{2}$ coefficient is acceptable)

- · The enthalpy for this reaction is called the compound's Standard Enthalpy of Formation (ΔH_f°)
- Elements in their standard state have $\Delta H_f^\circ = 0$
- · Any reaction can be framed as a combination of standard formation reactions
 - * First, the reactants are broken down into their elements (the *reverse* of formation reactions)
 - * Then, the elements are reassembled into the products (formation reactions)
 - * The energy level diagram for any reaction is the same: reactants \rightarrow elements \rightarrow products
 - * This pathway doesn't need to be *practical*, it is enough to be theoretically *possible*

*
$$\Delta H_{rxn} = \sum_{products} \nu \cdot \Delta H_f^{\circ} - \sum_{reactants} \nu \cdot \Delta H_f^{\circ}$$

- * This formula is general, for any reaction at all
- * Instead of an encyclopedia of thousands of reactions, we only need a table of ΔH_f° values
- * Appendix G includes a large number of thermodynamic values like ΔH_f°

Practice: Find
$$\Delta H_{rxn}$$
 for $P_4O_{10}(s) + 6 H_2O(l) \longrightarrow 4 H_3PO_4(aq)$ $\left(\Delta H_{rxn} = 454.6 \frac{kJ}{mol}\right)$
Find ΔH_{rxn} for $C_3H_8(g) + 5 O_2(g) \longrightarrow 3 CO_2(g) + 4 H_2O(g)$ $\left(\Delta H_{rxn} = -2043.9 \frac{kJ}{mol}\right)$

Quiz 5.5 - Enthalpies of Formation

Homework 5.5

- $\circ~$ 67: Calculate ΔH_f° from reaction enthalpies
- o 69: Reaction enthalpies from enthalpies of formation

ELECTRONIC STRUCTURE AND PERIODIC PROPERTIES OF ELEMENTS

6.1 Electromagnetic Energy

- Light is an electromagnetic wave, which can be thought of like a sound wave or a wave on a lake (Figure 6.2)
- $\circ~$ Light has a wavelength, frequency, and speed according to the equation: $\nu\lambda=c$
- $\circ~$ The speed of light is a constant, $2.998\times 10^8\,\frac{m}{s}$
- \circ How far does light travel in 5.00 ms? (1.50 km)
- The electromagnetic spectrum is more than just visible light (Figure 6.3)
 - · Higher frequencies (shorter wavelengths) are UV light, X-rays, and gamma rays
 - · Lower frequencies (longer wavelengths) are infrared light, microwaves, and radio waves
 - · TV-remotes are flashy lights, and radio towers are flashlights-on-a-stick
 - There are important technical differences in how we can use these different kinds of light, but they are fundamentally the same thing (an alternating electromagnetic wave)
 - · Find the wavelength of your favorite radio station (MHz is a frequency of $10^6~s^{-1}$)
- Light-matter interactions were central to the discovery of modern physics
- The photoelectric effect was an important matter/light interaction (Figure 6.11)
 - · Sometimes light falling on a metal will eject an electron this is the photoelectric effect
 - · The kinetic energy of the ejected electron can be measured
 - The energy depended on the wavelength of light bluer light ejected electrons at faster velocities
 - · There was a threshold where electron ejection stopped, and redder light would have no effect
 - · This was surprising, because light intensity had *no* effect on the photoelectron energy
 - · Dim blue light would eject fast electrons, bright red light would have no effect
 - · This was eventually explained by the idea that light carries energy in small discrete packets
 - These packets of energy are called *photons* and the energy they carry depends on the wavelength
 - · The photoelectric effect could be described by the equation: $KE = h\nu \phi$
 - · Here, ν was a new constant, called Planck's constant, and ϕ was the metal's work function
 - The equation for the energy of light was: $E = h\nu$ where h is Planck's constant

- o Light can also be *absorbed* or *emitted* by matter
 - · Light is emitted by gases like Ne, Ar, or Na when high voltage passes through it
 - · Light can also be absorbed by gases and other materials
 - Each substance showed a unique fingerprint of wavelengths of light emitted or absorbed (Figure 6.13)
 - · He was first identified by its absorbance spectrum in sunlight
 - · The unique spectra arise from the particular energy levels of a substance
 - These spectra showed how matter can only store or release energy in certain, constrained amounts (or *quanta*)
 - · i.e. in addition to the energy of light, the energy of matter was *quantized* as well
- o Ultraviolet Catastrophe (Figures 6.10 and 6.9)
- o Wave/Particle duality (not in the textbook)
 - These experiments show that light behaves like both a *wave* (interference) and a *particle* (quantization)
 - It is wrong to say light is either of those things rather, it is a new thing with similarities to both (rhinosceros vs dragon + unicorn)
 - · On very small scales, matter behaves like both a wave and a particle as well!
 - · Electrons, in particular, are strongly wave-like, with a characteristic wavelength
 - · The IBM quantum corral image dramatically showed the real physicality of electron waves
 - This wave-like nature of electrons is important for understanding modern models of atomic structure
- All of these phenomena can be explained in terms of standing waves (Figures 6.7 and 6.8)

6.2 The Bohr Model

- The Rydberg Equation: (Figure 6.14)
 - · School teacher Rydberg recognized a pattern in the wavelengths of light in the H spectrum

$$\frac{1}{\lambda} = 1.097 \times 10^7 m^{-1} \left(\frac{1}{n_1^2} - \frac{1}{n_2^2} \right)$$

· His equation can be re-written in terms of energy

$$\cdot E = 2.179 \times 10^{-18} J \left(\frac{1}{n_1^2} - \frac{1}{n_2^2} \right)$$

- The Bohr model of the atom: (Figure 6.15)
 - There was no real explanation for *why* the absorption and emission spectra of different elements showed different discreet energies

- · Niels Bohr proposed that electrons orbit around the nucleus only at fixed distances
- · Absorption is when an electron shifts to a higher orbit, using a photon's energy
- · Emission is when an electron shifts to a lower orbit, releasing energy as a photon
- · The lowest energy state is the *ground state*, all others are *excited states*
- The discrete orbits represent states where the circumference of the orbit is equal to a number of wavelengths for the electron (Figure 6.17)

$$E_n = -2.179 \times 10^{-18} J\left(\frac{1}{n^2}\right)$$

Practice: What is the wavelength of the $2 \leftarrow 4$ transition in the H spectrum? (486.1 nm)

Quiz 6.1 - Light and the Bohr Model

Homework 6.1

- o 3: Frequency to wavelength
- 5: Wavelength to energy
- o 23: Energy of a Bohr state
- o 27: Energy of a Bohr transition

6.3 Development of Quantum Theory

- Quantum mechanics continued to develop
- Heissenberg Uncertainty Principle: We cannot simultaneously measure the position and velocity of an electron (or any other quantum mechanical particle)
- This means that we generally speak of where an electron is *most probable* to be found, rather than where it *actually is*
- o The Schrödinger wave equation describes matter starting from a wave-like perspective
 - DeBroglie began from the equation for the momentum of light: $p=\frac{h}{\lambda}$
 - · Light has no mass, but replacing p with mv give us: $mv=\frac{h}{\lambda}$ and rearranging gives: $\lambda=\frac{h}{mv}$
 - · This equation describes how anything with momentum has a characteristic wavelength
 - · Figure 6.18 illustrates one way electrons behave like waves
 - · Schrödinger went even further, adapting Maxwell's wave equations for matter with momentum (required some postulates)
 - The Schrödinger equation gives mathematical functions which describe the electron probability distribution
 - · Each solution is called an *orbital*, like the orbits of the Bohr model but 3-dimensional

- Orbitals are organized into shells and subshells
 - · Subshells are groups of orbitals with similar shapes and the same energy
 - · Subshells are named *s*, *p*, *d*, and *f*
 - · An s subshell has only one orbital (2 es), p has 3 (6 es), d has 5 (10 es), and f has 7 (14 es)
 - · Subshells are grouped into shells, which are indicated by numbers (1, 2, 3, etc.)
 - · These numbers are the numbers in the Rydberg equation, and are the principle energy levels
 - \cdot The first shell only has an s subshell, and each shell beyond that adds one type

Practice: How many electrons can be placed in a p subshell? (6)

How many electron can be placed in the 3rd shell? (18)

Which of the following subshells does *not* exist? 2s, 3f, 3p, 5d (3f)

- Subshells each have orbitals with different *shapes* (Figure 6.21)
 - · Because of the Heissenberg uncertainty principle, we describe region where an electron is likely to be found
 - · These regions have shapes based on the mathematical functions which form them
 - \cdot s orbitals are spherical, p orbitals are dumbells, d orbitals are clover-leafed
- Quantum Numbers
 - · Remember that orbitals are actually mathematical functions
 - · Certain parts of those functions depend on integer numbers (like n in the Rydberg equation)
 - · These integer numbers are called *quantum numbers*
 - · Quantum numbers can be thought as an "address" for each electron (Street, Buildling, Unit, Name)
 - * n Principal quantum number (1, 2, ...) gives orbital shell, energy, and size
 - * l Angular momentum quantum number (0, 1, ..., n-1) gives orbital subshell and shape
 - * l is why not all shells have all orbital types
 - * m_l Magnetic quantum number (-l, ..., l) gives orbital within a subshell (different spatial orientations)
 - * m_l is why subshells have different numbers of orbitals
 - * m_s Spin quantum number $\left(\pm \frac{1}{2}\right)$ gives "spin" of the electron (up- or down- arrow, Figure 6.23)
 - \cdot Each electron in an atom/ion must have a unique set of quantum numbers Pauli Exclusion Principle
 - · You should be able to point to the right electron given a set of quantum numbers, or give the 4 quantum numbers for an indicated electron in an energy level diagram

Practice: Give numbers or indicate electrons on an energy level diagram

Homework 6.2

- o 33: Properties associated with quantum numbers
- o 35: Identifying the subshell from quantum numbers
- 41: Orbital shapes and other properties

6.4 Electronic Structure of Atoms (Electron Configurations)

- o Electronic energy level diagrams
 - · The orbitals within each subshell are precisely equal in energy (degenerate)
 - · The subshells themselves differ in energy (their order will be explained shortly)
 - · Draw the subshells with one line for each orbital
 - · Each orbital can hold two electrons, drawn as up- and down-arrows
 - · Find the total number of electrons for the element or ion
 - · Aufbau Principle Fill up the orbitals with electrons from the bottom-up
 - Hund's Rule Fill a subshell with one electron in each orbital before pairing them up (like roommates in an apartment)
 - · This is the ground-state configuration of the element
- We can use the periodic table as a cheat-sheet to the order of the subshells and electron configurations (Figure 6.27)
 - · The P. T. is actually quite long Lanthanides and Actinides have been cut and pasted
 - · Each region of the P. T. represents a different subshell
 - · The rows represent different shells
 - · The number of elements in each block is the number of electrons each shell can hold
 - The *d*-block and *f*-block trail the row number by 1 and 2 (d(-1)) and f(-2)
 - · The order of the subshells is found by simply following the elements and noting in which block they reside
- o The arrangements of the electrons can be written as an electron configuration
- Write the subshells, with their number of electrons as a superscript Practice: Write the electronic configurations for O, Zr, and Bi
- o Especially for large elements like Pb, these configurations are very unwieldy
- We can shorten them by referencing the configuration of the *preceding* noble gas
- The electrons which make up this noble gas configuration are buried inside the atom, and called *core* electrons

- \circ For Bi, we get [Xe] $6s^24f^{14}5d^{10}6p^3$
- The outermost electrons (The ones we write) are called *valence* electrons, and are the ones involved in bonding and ion formation
- A few transition metals have anomalous configurations (memorize only Cr and Cu)
- The Lanthanum and Actinium boundary also shows some anomalies (don't memorize them)
- Figure 6.29 gives the valence configuration of every element's ground state
- \circ Technically, filled d and f subshells count as *core*, and only the outermost s and p electrons will always count as *valence*
- This means that only the main group elements have a reliable pattern in their number of valence electrons
- The number of valence electrons is the same as the "A" column names
- We can write the configurations of ions as well
 - · For most, simply add or remove electrons according to the normal pattern
 - Transition metals will lose the outermost s electrons before they lose any d electrons (This is why so many transition metals have a stable 2+ ion)
 - · Ions and atoms with identical configurations are called *isoelectronic* to each other

Practice: List several stable ions which are isoelectronic with Ar

Quiz 6.3 - Electronic Configurations

Homework 6.3

- o 49: Complete electron configurations
- o 53: Energy level diagram of valence electrons
- o 55: Identify an atom from its configuration
- 57: Identify an ion from its configuration
- o 63: Electron configuration of an ion
- 79: Isoelectronic configurations

6.5 Periodic Variations in Element Properties

- The sizes of atoms and ions is controlled by the attractive and repulsive forces between electrons and protons
 - · Electron and protons attract each other, shrinking the atomic size
 - · Electrons repel each other, increasing the atomic size
- \circ Effective nuclear charge (Z_{eff}) attempts to summarize these interactions
 - The actual nuclear charge (Z) is just the number of protons (quite high for larger elements)
 - · Core electrons will counteract much of the actual nuclear charge (called *shielding*)
 - $\cdot Z_{eff} = Z S$ S stands for "Shielding"
 - · S can be closely approximated by the number of core electrons Practice: Find Z_{eff} for Mg, S, and Br (2,6, and 7)
 - \cdot Slater's rules gives a more sophisticated and accurate value for S
 - · 1 for deep core electrons, 0.85 for V-1 electrons, and 0.35 for all but one V electron
 - · Note that Slater's rules are not in your textbook Practice: Find Z_{eff} for the same elements using Slater's rules (2.85, 5.45, and 7.6)
- o Atomic radius increases down a column because you are adding and entire new shell for each row
- \circ Atomic radius decreases across a row because of the increasing Z_{eff}
- This makes He the smallest element, and Fr the largest (Figure 6.30 and 6.31)
- Anions are much larger and cations are much smaller than their neutral counterparts (Figure 6.32)
- Ionization energy (IE) is the energy required to remove an electron
 - For example, it is the energy for this process: Li \longrightarrow Li⁺ + e⁻
 - · The two factors which control IE are radius, and Z_{eff}
 - · It is easier to remove electrons (smaller IE) from larger atoms
 - · It is easier to remove electrons (smaller IE) from atoms with lower Z_{eff}
 - · Opposite to radius, He has the highest IE, and Fr has the lowest IE
 - There are breaks in this trend at the beginning and middle of the p block (Figure 6.33 and 6.34)
- Second- and third- ionization energy is the energy to remove a second and third electron (Table 6.3)
 - · Each successive electron is harder to remove
 - · After the valence electrons are gone, removing a core electron is *much* harder to remove
- Electron affinity (EA) is the energy released when an electron is added
 - · Electron affinity is usually exothermic, so these values are mostly negative
 - · The *magnitude* of EA follows the same trend as IE
 - · There are lots of breaks in the trend (Figure 6.35 is a mess), so don't worry too much about EA

Quiz 6.4 - Periodic Trends

Homework 6.4

- o 67: Atomic radius trend
- \circ 71: Ionization energy trend
- $\circ~$ 75: Ranking atomic radii
- o 77: Ranking ionic radii
- \circ 85: Second ionization energy

CHEMICAL BONDING AND MOLECULAR GEOMETRY

- 7.1 Ionic Bonding
- **7.2** Covalent Bonding
- 7.3 Lewis Symbols and Structures
- 7.4 Formal Charges and Resonance
- 7.5 Strengths of Ionic and Covalent Bonds
- 7.6 Molecular Structure and Polarity

ADVANCED THEORIES OF COVALENT BONDING

- 8.1 Valence Bond Theory
- 8.2 Hybrid Atomic Orbitals
- 8.3 Multiple Bonds
- 8.4 Molecular Orbital Theory

GASES

- 9.1 Gas Pressure
- 9.2 Relating Pressure, Volume, Amount, and Temperature: The Ideal Gas Law
- 9.3 Stoichiometry of Gaseous Substances, Mixtures, and Reactions
- 9.4 Effusion and Diffusion of Gases
- 9.5 The Kinetic-Molecular Theory
- 9.6 Non-Ideal Gas Behavior

LIQUIDS AND SOLIDS

10.1 Intermolecular Forces

- Many physical properties of solids, liquids, and gases can be explained by the strength of attractive forces between particles (Figure 10.5)
- Phase changes happen due to the interplay between kinetic energy and intermolecular forces (Figure 10.2)
- o Pressure can also play a role in phase changes, as discussed later
- o These intermolecular forces come in different varieties
 - Dispersion Forces Non-polar molecules, impacted by polarizability, molecular weight, and surface area
 - * Dominant in non-polar molecules
 - * Created by induced dipoles (Figure 10.6)
 - * Impacted by polarizability (Table 10.1)
 - * Impacted by molecular weight (hydrocarbons from methane to wax)
 - * Impacted by molecule shape (Figure 10.7 compares the boiling points of pentane isomers)
 - · Dipole-Dipole Forces
 - * Dominant in polar molecules
 - * Results from attraction between permanent dipoles (Figure 10.9)
 - · Hydrogen Bonding
 - * Dominant only in molecules capable of hydrogen bonding
 - * Must contain a hydrogen-donor atom (H attached to N, O, or F)
 - * Must contain a hydrogen-acceptor atom (lone pair of electrons attached to N, O, or F)
 - * Hydrogen bonds are more than just particularly strong dipole-dipole forces. They have strong directionality according to VSEPR
 - * Figures 10.10, 10.14, and other figures on the Internet show water, DNA, and proteins all organized by hydrogen bonds
 - * Figures 10.11 and 10.12 illustrate how much hydrogen bonds exceed dipole-dipole forces in strength

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