# CHEM 1210 Lecture Notes

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#### **COURSE ADMINISTRATIVE DETAILS**

- $\circ \ \ \text{My office hours}$
- o Intro to my research
- o Introductory Quiz
- o Grading details
  - · Exams 40, Final 20, Homework 30, Quizzes 10
  - · 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/
- o Show how to access Canvas
  - · Calendar, Grades, Modules, etc.
  - · Achieve Homework
  - · Textbook
- $\circ$  Algebra Review Assignment  $\circ$
- Introduction to chemistry
  - · Ruby fluorescence
  - · Levomethamphetamine
  - · Rubber band elasticity
  - · Structure of the periodic table
  - · Salt on ice and purifying hydrogen peroxide

#### CHAPTER 1

#### SCIENCE AND MEASUREMENT

#### 1.1 Classification of Matter

- Matter is anything that has mass and occupies space
- o *Elements* are the basic building blocks of the stable matter around us
- Atoms are the smallest, indivisible pieces of matter (demo with a copper penny)
- Atoms bond together in chemical bonds (more on them later)
- When 2 or more elements combine in a definite ratio, they form *compounds* 
  - · Definite proportions and the analogy with bread recipes
  - · Chemical compounds are different from their elements (drowning in water 89% Oxygen)
- Pure substances vs mixtures (Figure 1.1)
- Heterogeneous vs homogeneous mixtures (Figure 1.2)
- o "Solution" is just another name for an homogeneous mixture

# 1.2 Properties of Matter

- o The first categories of properties are *physical* vs *chemical* 
  - Physical properties can be measured or observed without changing the chemical identity
  - · Chemical properties can only be measured or observed by changing the *chemical identity*
  - · Mass, density, color (absorption spectrum), specific heat, solubility, etc. are physical
  - · Flammability, oxidization potential, acidity/basicity, corrosiveness, etc. are chemical
- The next categories are *extensive* vs *intensive* properties
  - Extensive properties depend on the amount of a substance mass, volume, enthalpy of combustion, heat capacity, etc.
  - Intensive properties are independent of the amount of a substance density, temperature, reactivity, specific heat, etc.
- Finally, we categorize *physical changes* and *chemical changes* 
  - · Phsyical changes do not change the chemical identity of a substance, and can readily be undone dissolving, phase changes, deformations, temperature changes, etc.
  - · Chemical changes cannot be undone because the chemical identity of the substance has changed oxidation, combustion, corrosion, etc.

- · Chemical changes are accompanied by either the release or absorption of energy
- · Starting materials in chemical changes are called *reactants*, and the final materials are called *products*
- Demo with air and hydrogen balloons

### 1.3 Matter and Energy

- All chemical phenomena involve changes in *matter* and/or *energy*
- o Mass is the amount of substance. Contrasted with weight, mass doesn't change due to gravity
- o Energy is the capacity to do work
  - · Forms of energy: Heat, Chemical, Nuclear, Kinetic, Potential, Electrical, Sound, Light
  - · Energy can be transferred or change form, but total energy is always conserved

#### 1.4 The Scientific Method

- The scientific method is an orderly process for determining information about the natural world (Figure 1.8)
  - · Observation Something about the natural wold inspires a question
  - · Hypothesis The scientist suggests a plausible answer or explanation for the observations
  - · Experiment A carefully constructed experiment will test the validity of the hypothesis
    - \* Results inconsistent with the hypothesis force a revision, and retesting of the new hypothesis
    - \* Results consistent with the hypothesis suggest the hypothesis has merit
  - · Results should be corroborated by other scientists in independent experiments
  - · A well-supported hypothesis may become a scientific theory
- The scientific method is based on the principle of *falsifiability*. It proves bad hypothese false, but can never *prove* a hypothesis to be true
- As a hypothesis survives more and more experimentation, it becomes relieable enough to be considered a
  theory. Theories can later be proven wrong, but the new explanation is almost always a marginal extension
  of the older theory, which better explains certain edge cases
- Entirely revolutionary new theories, such as Einstein's general relativity, are exceptionally rare
- Figure 1.9 illustrates the differences between scientific laws, hypotheses, and theories
- o My story about magnets on string and the importance of following the scientific method
  - · Why can't we just make conclusions based on observations (as opposed to experiments)?
  - How was my bias displayed in this story?

# 1.5 The International System of Units

- o There are often many different units to describe the same quantity (quart, liters, cubic centimeters, etc.)
- o The International System of Units is a standardized set of units used by scientists globally
- o Base units are the fundamental units that can describe everything we measure (Table 1.3)
  - · Length m
  - · Mass kg (note that the SI unit is *not* the gram)
  - · Time s
  - · Electrical Current A
  - · Temperature K
  - · Luminous Intensity cd
  - · Countable Amount mol
- Large or small amounts can be described using metric prefixes (Table 1.4)
  - These prefixes are mostly based on powers of  $10^3$
  - · You must know the prefixes and their abbreviations in Table 1.4
  - · Introduction to simple conversions and scientific notation
- Derived units describe quantities that combine multiple base units together
  - · Volume can be  $l \times w \times h$ , or generalized to any shape
  - · Density is  $\frac{mass}{volume}$
  - · Some derived units are given a new name:  $J = \frac{kgm^2}{s^2}$
- When using the SI units, conversions tend to be easy (even for complex derived units)
- $\circ$  How much energy is required to accelerate a 1.75 kg object by 0.650  $m/s^2$  over a distance of 3.20 m? (3.64 J)

#### 1.6 Significant Digits

- Qualitative vs quantitative descriptions
- For quantitative descriptions, uncertainty comes in the form of accuracy and precision
  - · Accuracy refers to the average of a set of measurements corresponding to the true value (within a standard deviation)
  - · If inaccuracies are known, then they can be subtracted by calibration
  - · Precision refers to the standard deviation of a set of measurements on the same sample
  - · Figure 1.13 shows accuracy vs. precision on a dart board
  - · The precision of your tool should match the magnitude of the quantity to measure
  - · Estimate the position between the smallest graduation on marked rulers, burettes, etc.

- o Precision is communicated through significant figures
- $\circ$  Consider measuring something 1.5 cm vs 1.4973 cm
- $\circ$  Now consider 1.5 cm vs 1.5000 cm
- Identifying significant figures
  - · Any non-zero digits are significant
  - · Trailing zeros are significant if they come *after a decimal*
  - · Zeros between two significant figures are also significant
  - · Leading zeros are never significant
  - Practice:  $4010, 0.0034, 7.100, 639, 000, 6.390 \times 10^5$
- Scientific Notation not only makes it convenient to write very large or very small numbers, but it also communicates precision without any ambiguity
- o Propagating sig. figs through calculations
  - · Identify the # of sig figs and the position of the least significant digit for both numbers
  - $\cdot +-:$  The LSD of the answer will match the position of the LSD for the less precise input
  - · Practice: 120.7 g + 34 g = 155 g | 212 mm 210.95 mm = 1 mm
  - $\cdot \times$ : The answer will have as many sig. figs as the input with *fewer* sig figs.
  - Practice:  $56.3 \, miles/1.2 \, h = 47 \, miles/h$  |  $1.5 \, cm \times 3.62 \, cm = 5.4 \, cm^2$
- o Multi-step problems have two considerations:
  - · Do not round intermediate answers to avoid compounding rounding errors
  - · Keep track of both the # of sig. figs and the LSD for each intermediate answer
  - Practice:  $(0.0045 \times 20,000.0) + (2813 \times 12) = 34,000$  $863 \times [1255 - (3.45 \times 108)] = 762,000$

#### 1.7 Dimensional Analysis

- Every measurement will have its associated *unit(s)*, which is just as important as the quantity itself
- o For any calculations, the units can provide a guide and a check on your process
- Structuring your calculations around the units is a paradigm called *dimensional analysis* 
  - · Treat units like algebraic quantities, which can cancel in calculations
  - Conversions factors can be written as ratios. i.e. 1.00~in = 2.54~cm becomes  $1 = \frac{2.54~cm}{1.00~in}$
  - · Consider both the starting and ending units, and how to convert from one to the other
  - · Solve the problem with the "picket fence" method
  - Practice: How many m are in 1.00 ft? (0.305 m)

How many ¢ does someone make per second if their wage is 
$$\frac{\$12.00}{h}$$
?  $(0.3333c/s)$ 

 $\circ~$  Be careful with squared or cubed units! Convert  $0.05~m^3$  into  $cm^3~(50,000~cm^3)$ 

### 1.8 Density

- o Density relates how much mass is in a given volume, and is responsible for buoyant forces, etc.
- $\circ$  Table 1.7 gives the density of some common substances (add in  $\approx 0.001~g/ml$  for gasses)

$$\circ \ density = \frac{mass}{volume}$$

- $\circ$  Volumes can be found by  $l \times w \times h$ , or by measuring the volume of displaced water
- $\circ$  Practice: Find the density of a block with dimensions  $1.5~cm \times 6.4~cm \times 13.2~cm$ , and mass of 1.43~kg. Can guess the material? (11~g/ml Lead)

A gold nugget displaces  $4.52\ ml$  of water. What is its mass?  $(87.2\ g)$ 

# 1.9 Temperature Scales

- o There are three common temperature scales, Fahrenheit, Celsius, and Kelvin
- o Kelvin is essential to use in some formulas, but Celsius may be convenient at other times

$$T_{C} = \frac{5}{9} (T_{F} - 32)$$

$$T_{F} = \frac{9}{5} T_{C} + 32$$

$$T_{K} = T_{C} + 273.15$$

o Practice: Convert the current temperature from Fahrenheit to Celsius and Kelvin

#### CHAPTER 2

#### ATOMS AND THE PERIODIC TABLE

#### 2.1 Chemical Symbols

- o Each element has a Symbol, which starts with a capital letter
- o Chemical Formulas use the symbols and subscripts to show how many of each element are in a compound
- Parenthesis show atom groups: How many oxygen atoms are in Mg(ClO<sub>3</sub>)<sub>2</sub>

# 2.2 The Laws of Chemical Combination

- o Careful experiments show reactions follow several laws
- o The Law of Conservation of Mass
- $\circ$  The Law of Definite Proportions (For water, 1.00 g O combines with 0.126 g of H)
- $\circ~$  The Law of Multiple Proportions (For  $\rm H_2O_2,\,1.00~g$  of O combines with 0.063~g of H) Possible ratios of N:O 1:1.14 (NO), 1:0.57 (N<sub>2</sub>O), 1:2.29 NO<sub>2</sub>

# 2.3 The History of the Atom

- Based on these laws, Dalton formulated an atomic theory:
  - · Matter is made up of atoms
  - · Atoms of the same element are identical to each other. Atoms of different elements are different from each other
  - · Atoms combine to form molecules in whole-number ratios
  - · Different ratios of the elements give different chemical compounds
- $\circ\,\,$  J.J. Thompson discovered the electron in 1897
  - · His cathode ray experiment showed that electrons are negatively charged
  - $\cdot\,$  Thompson found the  $\frac{mass}{charge}$  ratio, and Millikan later found the electron mass itself
  - · Electrons were a component of atoms, but were first assumed to be distributed evenly through the atom
- $\circ~$  Ernest Rutherford discovered the nucleus in 1909
  - · Gold foil experimental details (firing a bullet into jello, vs. into a chainlink fence)
  - · The mass and positive charge of an atom are densely packed into the nucleus

- · Electrons exist diffusely outside of the nucleus (originally in orbits, now in clouds)
- · Atoms are mostly empty space (atom is  $\approx 1$  Å (100,000~fm) across, and nucleous is  $\approx 1~fm$  across )
- Later, neutrons were also discovered, and are a part of the nucleus

# 2.4 Subatomic Particles, Isotopes, and Ions

- The important subatomic particles are: electrons, protons, and neutrons (table 2.2)
- $\circ$  For neutral atoms, e = p, but n could vary
- $\circ$  The identity of an element depends entirely on p, so we call it the *atomic number* (Z)
- $\circ$  The sum p + n is the *mass number* (A)
- $\circ$  Atoms of the same element with different mass numbers are *isotopes* Practice: What is n for carbon with A=14? | What is n for lead with A=207?
- $\circ~$  The difference p-e is the  $\mathit{charge}~(Q).$  Atoms with  $\mathit{charge} \neq 0$  are called  $\mathit{ions}$
- $\circ$  Chemical symbols can show all these values like so:  ${}^A_Z X^Q$

#### 2.5 Atomic Masses

- o The mass of a single atom is too small to conveniently work with, so we use atomic mass units (AMUs)
- $\circ~$  The AMU is defined as  $\frac{1}{12}M_{^{12}\mathrm{C}}$  , and is  $1.00~g=6.022\times10^{23}~AMU$
- The mass of a single proton or a single neutron are about 1~AMU. An electron is about  $\frac{1}{1800}~AMU$
- o Individual isotopes have an actual, measured mass similar to their mass number
- o The masses we see on the periodic table are a weighted average, based on the isotope abundances
- $\circ \ \textit{Atomic Mass} = \textstyle \sum_{isotopes} (Fractional \ \textit{Abundance}) \times (Isotope \ \textit{Mass})$

_		<sup>6</sup> 3Cu	<sup>65</sup> Cu	
Practice: Give the atomic mass for Copper with:	Mass (amu)	62.929601	64.927794	
	Abundance (%)	69.17%	30.83%	

#### 2.6 The Periodic Table

- As the number of known elements grew, scientists began to organize them into categories
- $\circ$  F<sub>2</sub>, Cl<sub>2</sub>, Br<sub>2</sub>, and I<sub>2</sub> are highly reactive gases. Li, Na, K, and Rb are highly reactive metals
- Arranging the elements according to reactive trends and size essentially gives us the periodic table

- Dmitri Mendeleev is considered the father of the periodic table because he used his table to predict the existence and properties of undiscovered elements
- o Gallium and Germanium were later discovered to have properties very close to what Mendeleev predicted
- o Now the periodic table is arranged by atomic number, rather than by mass
- o We have discovered all the elements up to 118, which may be the only stable elements at all
- The periodic table is tremendously useful and information-rich in many ways which we will highlight in most chapters of the textbook
- Structure of the periodic table:
  - · The periodic table is really long, but we cut out a portion to fit it on a page
  - · Rows are called *periods* (but I will just call them rows)
  - · Columns are called *groups* or *families*
  - · Groups are labeled by two conventions: numbered 1-18, or 1A-8A
  - · Metals and non-metals are separated by the metalloids (Figure 2.19)
  - · Main group, transition metals, and inner-transition metals (Figure 2.18)
  - · Some families are named: Alkali Metals, Alkaline Earth Metals, Coinage Metals, Halogens, Noble Gases, and others you don't need to know (Figure 2.17)

#### CHAPTER 3

#### COMPOUNDS AND THE MOLE

# 3.1 Chemical Formulas

- We have already discussed how *Chemical Formulas* give the # and type of atoms in a compound
- o Another word for chemical formula is Formula Unit, which is especially used for ionic compounds
- o Molecular compounds are compounds composed of non-metals bonded together
  - · One formula unit is a *molecule*, which is a discrete particle with clear boundaries
  - · Atoms in a molecular compound are held together by covalent bonds more on them in chapter 10
- o Ionic compounds are compounds composed of two ions bound together
  - · The simplest ionic compounds are a metal bonded to a non-metal
  - · Ionic compounds form bonds in a large array called a lattice
  - · There is no clear boundary for a single formula unit, the lattice structure just goes on in all directions
  - · Different compounds will have different lattice structures
  - These atoms are held together by *ionic bonds*, which are simply the attraction between negative and positive charges
- o Metals bonded to metals for *metallic bonds*, which we will not cover in this class
- o Some elements for molecules in their natural state
  - The diatomic elements are: H<sub>2</sub>, N<sub>2</sub>, O<sub>2</sub>, F<sub>2</sub>, Cl<sub>2</sub>, Br<sub>2</sub>, and I<sub>2</sub>
  - · Some solid elements form other molecules:  $P_4$  and  $S_8$
  - · Some elements occur in more than one natural state, or *allotrope* Carbon can form coal, graphite, or diamond. Oxygen can form common  $O_2$  or ozone  $(O_3)$ . Phosphorous comes in two forms (red, and white)

#### 3.2 Naming Binary Covalent Compounds

- Binary molecular compounds are made up two non-metals
- Non-metals can combine in different ratios (NO, N<sub>2</sub>O, NO<sub>2</sub>, N<sub>2</sub>O<sub>3</sub>, and more!)
- Because of this, binary molecular compounds are named in a way that indicates how many atoms of each element are present

0	The num	ber of ato	ms is inc	licated l	by Greel	k prefixes (	(Table 3.	1)

С	Fi	irst Element Name	Second Elemen	it Name-	"ide"
		-	-		

- o The element futher left on the periodic table goes first in both the formula and the name
- o "Mono-" may be left off for the *first* element (i.e. Carbon Dioxide, not Monocarbon Dioxide)
- Binary compounds which contain H are usually called by their common names instead of using this convention (water, hydrochloric acid, and ammonia)

Practice: NO (nitrogen monoxide)  $N_2O$  (dinitrogen monoxide)  $NO_2$  (nitrogen dioxide)  $N_2O_3$  (dinitrogen trioxide)

### 3.3 Formulas for Ionic Compounds

- Positive ions are called *cations*, and negative ions are called *anions*
- Ions can be a single atom (monatomic), and the magnitude of the charge can often be inferred from the element's position in the periodic table (Figure 3.7)
- o Ions can also be made from a group of atoms which carry the charge collectively (polyatomic) Table 3.4
- o All ionic compounds are electrically neutral, so their positive and negative ions must cancel out
- Find the LCM between the positive and negative charges to know how many of each to include
   Practice: Give the formula unit for compounds made from the following pairs of ions:

$$\mathrm{Na^{+}}$$
 and  $\mathrm{Cl^{-}}(\mathrm{NaCl})$   $\mathrm{Ag^{+}}$  and  $\mathrm{S^{2-}}(\mathrm{Ag_{2}S})$   $\mathrm{Fe^{3^{+}}}$  and  $\mathrm{O^{2-}}(\mathrm{Fe_{2}O_{3}})$ 

 $\circ~$  For compounds with polyatomic ions, just be sure to include parenthesis

Practice: Give the formula unit for compounds made from the following pairs of ions:

$${\rm Na^{+} \, and \, CH_{3}COO^{-} \, (NaCH_{3}COO) } \qquad {\rm Ag^{+} \, and \, SO_{4}^{\, 2-} \, (Ag_{2}SO_{4}) } \qquad {\rm Ni^{2+} \, and \, PO_{4}^{\, 3-} \, (Ni_{3}(PO_{4})_{2}) } \\$$

Because of this charge balance, we can calculate the cation's positive charge based on the formula
 Practice: Give the positive charge of the cation in each ionic compound

$$\mathrm{VO}_{2}\left(\mathrm{V}^{4+}\right) \quad \mathrm{AuCl}_{3}\left(\mathrm{Au}^{3+}\right) \quad \mathrm{Al}_{2}\mathrm{S}_{3}\left(\mathrm{Al}^{3+}\right)$$

# 3.4 Naming Ionic Compounds

- o To name an ionic compound, simply give the cation name and the anion name
- Monoatomic cations are just the element name
- $\circ$  For elements with more than one common charge, the charge is indicated in (Roman Numerals) Practice: Give the name of the following cations: Ba<sup>2+</sup> (Barium) Cr<sup>2+</sup> (Chromium(II))
- o Ammonium and hydronium are the only common polyatomic cations
- o Monoatomic anions are just the element name with an "-ide" ending
- o Polyatomic anions follow some rules, but are best just memorized
  - · Oxyanions forms series' which follow the pattern shown in Table 3.3

- · Oxyanions with one or more hydrogens have "hydrogen-" "dihydrogen-" etc. added to their names
- · Table 3.4 lists common polyatomic ions (you can ignore the right column except chlorates)

Practice: Name the following ionic compounds: Na<sub>2</sub>SO<sub>3</sub> (sodium sulfite)

NH<sub>4</sub>NO<sub>3</sub> (ammonium nitrate) Fe(HCO<sub>3</sub>)<sub>3</sub> (iron(III) hydrogen carbonate)

- o Hydrates are ionic compounds which naturally incorporate water into their structure
  - · An integer number of water molecules are incorporated, called waters of hydration
  - · If a hydrated salt is heated sufficiently, the waters will evaporate and the salt become anhydrous
  - The formula indicates the number of water molecules per formula unit with a dot ( $CuSO_4 \cdot 5 H_2O$ )
  - To name the compound, give the anhydrous compound name followed by "\_\_\_\_\_ hydrate".

    Greek prefixes indicate the number of waters

Practice: Give the names of the following hydrated salts

MgSO<sub>4</sub>· H<sub>2</sub>O (magnesium sulfate monohydrate) MgCl<sub>2</sub>·6 H<sub>2</sub>O (magnesium chloride hexahydrate)

### 3.5 Naming Acids

- Acids are covalent compounds which will release H ions when dissolved in water
   (this is a white lie the actual mechanism for acids can get quite complex)
- $\circ~$  For example:  $\mbox{HNO}_{_3}$  will release its H in water to produce  $\mbox{H}^{\mbox{\tiny +}}(\mbox{aq})$  and  $\mbox{NO}_{_3}^{\mbox{\tiny -}}(\mbox{aq})$
- Not all H atoms will be released. For CH<sub>3</sub>COOH, only the last H is ionizable
- Often the ionizable hydrogen(s) will be written first. H<sub>2</sub>O<sub>2</sub> and acetic acid are common exceptions
- Acids with 1 ionizable hydrogen are *monoprotic*. Atoms with 2 or more are *polyprotic*
- Binary acids are named "hydro-Element-ic acid" (hydrochloric acid, hydrofluoric acid)
- Oxyacids are acids with H attached to an O-containing polyatomic anion
  - · The acid name depends on the anion name
  - · "-ate" ions become "-ic acid" (sulfate and sulfuric acid)
  - · "-ite" ions become "-ous acid" (hypochlorite and hypochlorous acid)

#### 3.6 Nomenclature Review

- Figure 3.15 gives a comprehensive flowchart for naming chemical compounds
- o Conventions for naming organic molecules will be covered in CHEM 1220

### 3.7 The Mole

- $\circ~$  We have already gone over how  $6.022\times10^{23}=1~mol$  , and 1~mol~AMU=1~g
- We can talk about moles of anything: formula units, atoms, etc.

Practice: How many moles of O atoms are in  $2.50 \, mol$  of  $\mathrm{Fe_2O_3?}$   $(7.50 \, mol)$ 

How many H atoms are in 1.40~mol of  $\mathrm{CH_4?}~(3.37\times10^{24}~atoms)$ 

# 3.8 Molar Mass

- The *Molar Mass* gives the mass for a mole of a given substance
- o The atomic weight (on the periodic table) is the mass for a mole of atoms
- o The molecular weight or formula weight is the mass of a mole of formula units for a compound
- $\circ$  I will use *Molar Mass* (M) as a general term for both
- o Many problems will involve converting between mass and moles, moles and mass
- o "Measurement Land" vs "Chemistry Land" vs "Atomic/Counting Land"

Practice: How many O atoms are in 1.35 g of  $CO_2$ ?  $(3.69 \times 10^{22})$ 

How many O atoms are in 1.35 g of CO?  $(2.90 \times 10^{22})$ 

What is the mass of  $0.675 \ mol \ of \ C_{_{12}}H_{_{22}}O_{_{11}}$ ?  $(23.1 \ g)$ 

### 3.9 Percent Composition

o Percent composition tells what percentage of a compound (by mass) is each element

$$\circ~\%_{Composition} = \frac{\nu M_{atomic}}{M_{formula}} 100\%$$

Practice: Find the % composition of O in  $H_2O$ . (88.81%)

Find the % composition of O in  $\operatorname{Fe_2(SO_3)_3}$ . (40.92%)

- Percent composition can be used to find the mass of a component in a compound
- $\circ$  The % composition is a conversion factor between g of the element and 100~g of the compound Practice: How many g of N are in 2.00~g of NH<sub>4</sub>NO<sub>3</sub>?

$$(M = 80.0434 \, g/mol, \%_N = 34.9978\%, \text{ and } 0.700 \, g)$$

### 3.10 Empirical Formulas

- Some analytical techniques can only give % composition, not the true molecular formula
- The *empirical formula* is the most mathematically reduced form of a chemical formula
- $\circ$  Many molecular formulas may share the same empirical formula (i.e.  $CH_2O$ ,  $C_2H_4O_2$  and  $C_6H_{12}O_6$ )
- o Finding empirical formulas requires converting masses to moles
  - · Sometimes you will be given masses (See combustion analysis below)
  - · If you only have % composition, assume a mass of  $100\ g$  and the %s become g
  - · Convert the masses to moles using atomic weights
  - · Reduce the moles of each atom to a ratio of whole numbers
  - · Table 3.6 shows the decimal ending for common fractional equivalents

Practice: Give the empirical formula for a  $20\ g$  sample with  $1.34\ g$  H,  $8.00\ g$  C and  $10.7\ g$  O.

 $(CH_2O)$ 

Practice: Give the empirical formula for a compound which is 69.94% Fe and 30.06% O by mass.

 $(Fe_2O_3)$ 

#### 3.11 Molecular Formulas

- The molecular formula gives the actual number of atoms of each type in a single molecule
- The molecular formula will be some integer multiple (possibly 1) of the empirical formula
- o The compound's molar mass will be the same multiple of the empirical formula's molar mass

# 3.12 Combustion Analysis

- o Combustion analysis is a laboratory technique for finding the empirical formula of combustible compounds
- It involves combusting a carefully weighed mass of compound in excess oxygen, and measuring the amount of water and carbon dioxide produced
- o The C in the carbon dioxide and the H in the water came exclusively from the unknown compound
- Any unaccounted-for mass is assumed to be oxygen
- Process for solving combustion analysis:
  - · Calculate moles of C atoms from CO<sub>2</sub> mass
  - · Calculate moles of H atoms from H<sub>2</sub>O mass
  - · Calculate mass of C and H atoms
  - · subtract C and H masses from sample mass This is the O mass
  - · Calculate moles of O from the O mass

- · Reduce all mole ratios to whole numbers (perhaps relying on Table 3.6)
- · This is the empirical formula

Practice: 4.24~g of an unknown are combusted to yield 6.21~g of  ${\rm CO_2}$  and 2.54~g of  ${\rm H_2O}$  (Empirical Formula is  ${\rm CH_2O}$ )

Practice: 2.50~g of an unknown are combusted to yield 5.79~g of  $\rm CO_2$  and 1.18~g of  $\rm H_2O$  (Empirical Formula is  $\rm C_8H_8O_3)$ 

Practice: The molar masses for the two compounds above are: 120.10~g/mol and 152.15~g/mol What are the molecular formulas?

(The formulas are:  $C_4H_8O_4$  and  $C_8H_8O_3$ )

#### **CHAPTER 4**

# CHEMICAL REACTIONS AND AQUEOUS SOLUTIONS

#### 4.1 Chemical Equations

- Chemical equations show how *reactants* (on the left) are converted into *products* (on the right)
- o When properly balanced, an equation also shows the proportions of reactants and products
- o Equations also can show the *phase* of each chemical and any reaction conditions
- o Aqueous solutions are when a substance is dissolved in water
- o Balancing chemical equations:
  - 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 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

Practice: Balance the following chemical equation:  $Fe_2O_3(s) + Al(s) \longrightarrow Al_2O_3(s) + Fe(s)$ 

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

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<sub>2</sub>SO<sub>4</sub>(aq) + NaCl(aq)  $\longrightarrow$  AgCl(s) + Na<sub>2</sub>SO<sub>4</sub>(aq)

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

# 4.2 Types of Chemical Reactions

- o Reactions can be grouped into broad categories
- o Synthesis or Combination reactions:
  - · Two or more reactants combine to form a single product
  - $\cdot 2 \text{Na(s)} + \text{Cl}_2 \text{g} \longrightarrow 2 \text{NaCl(s)}$
- o Decomposition reactions:
  - · One reactant decomposes into two or more products
  - $\cdot 2 H_2O(l) \longrightarrow 2 H_2(g) + O_2(g)$
- o Single and Double Replacement reactions:

- · One (or two) element or polyatomic ion is replaced by another in the product
- $\cdot \operatorname{Zn}(s) + 2\operatorname{HCl}(aq) \longrightarrow \operatorname{ZnCl}_2(aq) + \operatorname{H}_2(g)$
- $\cdot 2 \text{KI}(aq) + \text{Pb}(\text{NO}_3)2(aq) \longrightarrow \text{PbI}_2(s) + 2 \text{KNO}_3(aq)$
- Acid/Base reactions (A type of replacement reaction):
  - · One or more H<sup>+</sup> are exchanged
  - $\cdot$  HCl(aq) + NaOH(aq)  $\longrightarrow$  NaCl(aq) + H<sub>2</sub>O(l)
- Precipitation reactions (A type of replacement reaction):
  - · Aqueous reactants produce one or more solid products (called precipitates)
- Combustion reactions:
  - · A substance is reacted with O<sub>2</sub> to produce CO<sub>2</sub> and H<sub>2</sub>O
  - $\cdot \ \, C_{3}H_{8}(g) \, + \, 5\,O_{2}(g) \, \longrightarrow \, 3\,CO_{2}(g) \, + \, 4\,H_{2}O(g)$
- o Redox reactions (synthesis, decomposition, single replacement, and combustion reactions)
  - · We will see later how redox reactions involve the exchange of electrons
- o Not all reactions we can write down will actually proceed in nature
- Reactions actually happen because of driving forces which make products more thermodynamically stable than the reactants
  - · Precipitation reactions have stable solid products with strong ionic bonds
  - · Acid/Base reactions have stronger bonds to H after the neutralization
  - · Redox reactions transfer electrons to make more stable electronic configurations

# 4.3 Compounds in Aqueous Solution

- When ionic compounds dissolve in water, they dissociate into their cations and anions
- Figure 4.14 shows how ions are hydrated by the water molecules when they dissolve
- Solutions of ionic compounds and acids/bases are called electrolytes because they conduct electricity
  - · Strong electrolytes will dissociate completely and produce a lot of ions
  - · Weak electrolytes dissociate only partially, and produce few ions
  - · Soluble ionic compounds and strong acids/bases (Table 4.3) are strong electrolytes
  - · Weak acids/bases are weak electrolytes
- Solutions of other molecular compounds are non-electrolytes They don't dissociate in water

### 4.4 Precipitation Reactions

- Not every ionic compound will actually dissolve in water. For example, most rocks and crystals are ionic compounds which don't readily dissolve in water
- o Soluble compounds will dissolve in water, insoluble compounds will not
- o Solubility guidelines help predict if a compound is soluble or not
  - · Ammonium and Group I cations always form soluble compounds
  - · Nitrate, chlorate, perchlorate, and acetate *always* form soluble compounds
  - · Chlorides, bromides, and iodides are soluble *except* with Ag<sup>+</sup>, Pb<sup>2+</sup>, and Hg<sub>2</sub><sup>2+</sup>
  - · Sulfates are soluble *except* with Pb<sup>2+</sup>, Hg<sub>2</sub><sup>2+</sup>, and "heavy" group II ions
  - · Carbonate, sulfite, phosphate, and chromate generally form insoluble compounds
  - · S<sup>2-</sup> and OH<sup>-</sup> form *insoluble* compounds, except with group I, ammonium, and "heavy" group II ions
  - · Ag+, Pb2+, and Hg2+ generally form *insoluble* compounds
  - · Table 4.5 gives a comprehensive list, but you will be fine just memorizing the points above
- o Precipitation reactions:
  - · When two soluble ionic compounds react, there is the possibility of a precipitate forming
  - · First, identify the products when cations and anions switch partners
  - · Then determine if any of the products will be insoluble these are the precipitates
  - · If no precipitate forms, then no reaction really took place at all
- Demos: Mg(NO<sub>3</sub>)<sub>2</sub> with NaOH, and Ca(NO<sub>3</sub>)<sub>2</sub> with Na<sub>2</sub>CO<sub>3</sub>
- Net ionic equations:
  - · Ionic equations acknowledge that soluble ionic compounds do not exist with anions and cations paired together. Rather, the anions are solvated and the cations are solvated separately by themselves
  - To form an ionic equation, write the cations and anions separately for all aqueous (soluble) ionic compounds
  - · Insoluble compounds are still written with ions paired together because they actually are
  - · Some ions appear solvated on both sides of the equation they are called "spectators"
  - These spectators don't actually do anything at all they are as irrelevant to the reaction as the solvent, other trace solutes ( $N_2$  and  $O_2$ ), etc.
  - · To form a *net* ionic equation, simply eliminate the spectators

Practice: Write the net ionic equations for the two demonstrations above

$$Mg^{2+}(aq) + 2 OH^{-}(aq) \longrightarrow Mg(OH)_{2}(s)$$
  
 $Ca^{2+}(aq) + CO_{3}^{2-}(aq) \longrightarrow CaCO_{3}(s)$ 

### 4.5 Acid-Base Reactions

- Recognize acid/base reactions by the exchange of a H<sup>+</sup> ion
- When acids react with OH<sup>-</sup> (strong bases), water will be produced
- When acids react with weak bases, the H<sup>+</sup> switches over to the base
- o Balance acid/base reactions just like any other reaction
- o For ionic and net ionic equations, only strong acids should be written as dissociated ions

#### 4.6 Oxidation States and Redox Reactions

- o Oxidation states (or numbers) keep track of how many electrons reside on an atom (like a charge)
- o To determine the oxidation states in a compound, follow these rules:
  - · A neutral elements (not a part of a compound) has an oxidation number of 0
  - · Monoatomic ions have oxidation numbers equal to their charge
  - · The sum of oxidation numbers in any formula is equal to the total charge
  - · Oxygen tends to have an oxidation number of -2 in compounds
  - · Hydrogen tends to have an oxidation number of +1 in compounds
  - · All other elements can be determined from the rules above

Practice: Identify the oxidation states for all elements in the following compounds

- o Oxidation is the process of *losing* electrons. The oxidation number increases
- Reduction is the process of *gaining* electrons. The oxidation number decreases
- Two mnemonic devices to keep them straight:
  - · OIL RIG (oxidation is losing, reduction is gaining)
  - · Chemical reduction involves a mathematical reduction of the oxidation number
- To identify redox reactions involve the exchange of electrons
  - · To identify a redox reaction, first find the oxidation state of all reactants and products
  - · If the oxidation state of any elements changes, then this is a redox reaction
  - · The element whose oxidation number decreases is *reduced*
  - · The element whose oxidation number increases is oxidized
  - · Any electrons lost by the oxidized element have gone to the reduced element they are connected both chemically and mathematically by the electrons
  - The compound which contains the reduced element has oxidized its reaction partner. Therefore, it is the "oxidizing agent"

• The compound which contains the oxidized element has reduced its reaction partner. Therefore, it is the "reducing agent"

Practice: Identify the reducing agent and the oxidizing agent in each chemical reaction

$$2 \text{ Al(s)} + 3 \text{ Cl}_2(g) \longrightarrow 2 \text{ AlCl}_3(s)$$

$$(O.A. = \text{Cl}_2, \text{ R.A.} = \text{Al})$$

$$CH_4(g) + 3 O_2(g) \longrightarrow CO_2(g) + 2 H_2O(g)$$

$$(O.A. = O_2, \text{ R.A.} = \text{CH}_4)$$

o Balancing redox reactions can be quite complicated. You will learn how in CHEM-1220

# 4.7 Predicting the Products of Redox Reactions

- Synthesis and decomposition reactions are very often redox reactions
  - · Sythesis of ionic compounds from elements, and decomposition of ionic compounds to elements
  - · The oxidation states of metals in compounds will match the charges they can take as ions
- o Single-replacement reactions are also very often redox reactions
  - · Often, the replaced elements involve an element and an ion swapping places
  - · One element goes from neutral to positively charged it is oxidized it is a reducing agent
  - · The other element goes from positively charged to neutral it is reduced it is an oxidizing agent
  - · Some elements are easier to oxidize, and some ions are easier to reduce
  - · These processes are linked an element which is easy to oxidize has an ion which is difficult to reduce
  - · Table 4.7 shows an activity series for common elements in these reactions
  - · A reaction will proceed spontaneously if the reducing agent is higher in the table than the oxidizing agent
- o Demos: Copper wire in silver solution and Zn piece in copper solution

Practice: Predict whether each demonstrated reaction will proceed spontaneously

$$Cu(s) + 2 AgNO_3(aq) \longrightarrow Cu(NO_3)_2(aq) + 2 Ag(s)$$
  
(Spontaneous)  
 $Zn(s) + CuSO_4(aq) \longrightarrow ZnSO_4(aq) + Cu(s)$   
(Spontaneous)

#### CHAPTER 5

#### STOICHIOMETRY

# 5.1 Mole Calculations for Chemical Reactions

- o Stoichiometry refers to the ratios between substances in chemical reactions
- These ratios are expressed in the coefficients of a balanced chemical equation
- Use the coefficients to create a conversion factor between substances
- My problem-solving diagram ("Measurement Land" vs "Chemistry Land")

```
Practice: Consider the reaction 2 P + 3 Cl<sub>2</sub> \longrightarrow 2 PCl<sub>3</sub>

How many moles of Cl<sub>2</sub> are needed to react with 0.250 mol of P? (0.375 mol)
```

How many moles of  $PCl_3$  would be produced?  $(0.250 \ mol)$ 

# 5.2 Mass Calculations for Chemical Reactions

- We will usually be concerned about finding *masses* rather than moles
- o You cannot compare masses directly, but rather have to go through the molar stoichiometry
- $\circ$  Problems are structured as:  $mass_A \to mol_A \to mol_B \to mass_B$ Practice: Consider the reaction 3 O<sub>2</sub> + 2 KCl  $\longrightarrow$  2 KClO<sub>3</sub>

How many g of  $O_2$  are needed to react with 0.850~g of KCl? (0.547g)

How many g of KClO<sub>3</sub> would be produced? (1.40g)

o Note that the total mass on the reactant side should equal the total mass on the product side

# 5.3 Problems Involving Limiting Quantities

- o Reactants are often not mixed in the perfect, proper ratio
- One reactant will be consumed and run out first, it is called the *limiting reactant*
- The reaction stops once the limiting reactant runs out, so the limiting reactant controls how much product is produced
- There are many ways to solve limiting reactant problems. I suggest the following:
  - · Looking ahead, choose a product which you are interested in
  - · For each reactant, calculate the amount of product it could produce
  - · Compare the product amounts the smallest value is the only value you keep

- · Throw out all other values, they are just wishful thinking and will never really happen
- · The reactant which gave the smallest amount of product is the limiting reactant
- · Start with the limiting reactant for all future calculations
- o Sometimes you need to find the amount of excess reactant(s) which remains
  - · Starting from the limiting reactant, find the amount of excess reactant required
  - · Subtract that mass from the initial value this is the amount left over
- Here, the total mass at the end will match the total mass at the beginning

Practice: Consider the reaction 2 CH $_3$ OH(g) + 3 O $_2$ (g)  $\longrightarrow$  2 CO $_2$ (g) + 4 H $_2$ O(g)

If 16.0 g of  $O_2$  react with 48.1 g of  $CH_3OH$ , which is the limiting reactant?  $(O_2)$ 

How many g of each substance will be present after the reaction completes? (0 g O<sub>2</sub>, 37.4 g CH<sub>3</sub>OH, 14.7 g CO<sub>2</sub>, and 12.0 g H<sub>2</sub>O)

Practice: Consider the reaction  $_3$  Fe(s) +  $_4$  H<sub> $_2$ </sub>O(l)  $\longrightarrow$  Fe $_3$ O<sub> $_4$ </sub>(s) +  $_4$  H<sub> $_2$ </sub>(g)

If 5.00 g of Fe react with 5.00 g of H<sub>2</sub>O, which is the limiting reactant? (Fe)

How many g of each substance will be present after the reaction completes? (0 g Fe, 2.85 g H<sub>2</sub>O, 6.91 g Fe<sub>3</sub>O<sub>4</sub>, and 0.241 g H<sub>2</sub>)

# 5.4 Theoretical Yield and Percent Yield

- The amount of product you calculate above is the *theoretical yield*
- o In the lab, your *actual yield* may be more, or less due to errors and random factors
- We can calculate the % yield based on these two values

$$Percent \, Yield = \left(\frac{Actual \, Yield}{Theoretical \, Yield}\right) \cdot 100\%$$

Practice: If you run the first reaction above and recover 10.7~g of  ${\rm H_2O}$ , what is the % yield? (89.2%)

If you run the second reaction above and recover 7.05~g of  ${\rm Fe_3O_4}$ , what is the % yield? (102%)

# 5.5 Definition and Uses of Molarity

- The *solvent* is the substance doing the dissolving, the *solute* is the substance being dissolved, and the *solution* is the homogeneous mixture after dissolving
- o Concentrated solutions have a lot of solute, dilute concentrations have little solute
- $\circ~$  Molarity is sometimes notated with square brackets (i.e.  $\left[\mathrm{H_{2}SO_{4}}\right])$
- $\circ$  This can be rearranged to give:  $Moles_{Solute} = Molarity \cdot L_{Solution}$  or  $(n = M \cdot V)$

- $\circ$  For small volumes (ml), we can use millimoles (mmol) for convenience
  - Practice: What is the molar concentration when  $2.50\ g$  of NaCl are dissolved to make  $0.100\ L$  of solution?  $(0.428\ M)$
  - Practice: What mas of  $C_{12}H_{22}O_{11}$  should be dissolved to make 0.500~L of a 0.125~M solution? (21.4~g)
- o Dilution is when water is added to a solution to make it less concentrated
- o Because only solvent is added, the moles of solute is the same at the end as at the beginning
- This gives a simple equation for dilution:  $M_1V_1 = M_2V_2$ 
  - Practice: What is the final concentration when  $25.0\,ml$  of  $0.832\,M$  are diluted to  $150.0\,ml$ ?  $(0.139\,M)$
  - Practice: How much 0.650~M solution should be used to make 250.0~ml of a 0.100~M solution? (38.5 ml)
- o We can now add to the "Measurement Land" and "Chemistry Land" diagram
  - · Moles can be calculated from n = MV
  - · ml of the co-reactant can be found instead of g
  - · For finding concentrations of the product, consider the *additive* volumes of the reactants

Practice: Consider the following reaction: 2 HCl(aq) + Na<sub>2</sub>CO<sub>3</sub>(s)  $\longrightarrow$  2 NaCl(aq) + H<sub>2</sub>CO<sub>3</sub>(aq) How many ml of 0.250~M should be used to react with 0.125~g of Na<sub>2</sub>CO<sub>3</sub>? (9.43 ml)

#### 5.6 Molarities of Ions

- o When ionic compounds dissolve, the ions dissociate
- The concentration of the *compound* may be different from the concentration of a given ion
- Multiply the formula concentration by the number of ions in the formula

Practice: Consider dissolving 1.45~g of MgCl<sub>2</sub> to make 125~ml of solution

What is the concentration of  ${\rm MgCl_2}$ ? (1.22~M) What is the concentration of  ${\rm Mg^{2^+}}$  and  ${\rm Cl^-}$  ions? (1.22~M and 2.44~M)

#### 5.7 Calculations Involving Other Quantities

- o Volume is one way to express an amount of a substance
- $\circ$  We have so far dealt with volumes of solutions, where n=MV
- For a pure substance, the volume must be dealt with very differently

- · Volume must first be converted to mass through density:  $m = d \cdot v$
- · Then mass is converted to moles through the molar mass

Practice: Sodium metal has a density of  $0.97g/cm^3$ , and reacts with water according to the following reaction:  $2 \text{ Na(s)} + 2 \text{ H}_2\text{O(l)} \longrightarrow 2 \text{ NaOH(aq)} + \text{H}_2\text{(g)}$ If  $0.750 \text{ } cm^3$  of Na react with 200.0 ml of water, what is the final [NaOH]? (0.158 M)

 $\circ$  The subscripts in a chemical formula can also relate the number of atoms of a particular element Practice: In the reaction above, how many H *atoms* are released as gas?  $(1.91 \times 10^{22} \, \text{H} \, atoms)$ 

### 5.8 Calculations with Net Ionic Equations

- o Stoichiometric calculations with net ionic equations are just like with regular equations
- $\circ$  To calculate moles of ions from g of solid salt, you must know the full salt formula

# 5.9 Titration

- o *Titration* is a technique to determine the concentration of a solution
- The unknown is reacted with a solution with precisely known concentration (a standard solution)
- o A precisely measured volume of one reactant is placed in an erlenmeyer flask (for easy mixing)
- The other reactant (the titrant) is slowly added using a buret, to measure the volume added
- The *equivalence point* is the exact point when the unknown is completely consumed by the reaction
- The *end point* is when you actually stop the titration
  - · End point is usually identified by a color-changing indicator
  - · For acid/base reactions, the indicator changes color with pH
  - · Redox reactions also have appropriate color indicators
  - · The indicator is chosen so that the end point is as close as possible to the equivalence point
- $\ \, \text{ The stoichiometric coefficients } (\nu) \text{ are included in the titration equation: } \frac{M_A V_A}{\nu_A} = \frac{M_B V_B}{\nu_B}$

Practice: Find [NaOH] when  $25.00\,ml$  of NaOH are titrated with  $37.8\,ml$  of  $0.100\,M$  HCl  $(0.151\,M)$ 

Practice: Find <code>[HNO $_3]</code> when <math display="inline">50.00\ ml$  of <code>HNO $_3$ </code> are titrated with  $21.3\ ml$  of  $0.150\ M$  <code>Ca(OH) $_2$ </code>  $(0.128\ M)$ </code>

#### CHAPTER 6

#### THERMOCHEMISTRY

### 6.1 Energy and Energy Units

- 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
  - · Kinetic energy is the energy of motion:  $KE = \frac{1}{2}mv^2$
  - $\cdot\,$  Potential energy is stored energy: Gravitaional, electrostatic, chemical, etc.
- o 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 (k<br/>calorie): 1 $Cal = 1000 \ cal = 4184 \ J$

#### 6.2 Energy, Heat, and Work

- 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 (an closed, insulated flask)
- o Work and heat are the ways energy can enter or leave the system
- o We always talk from the perspective of the system, even though we are a part of the surroundings
- $\circ$  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
- Heat is usually associated with temperature changes or phase changes
  - · 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
- o First Law of Thermodynamics: The energy of the universe is constant
- o Internal energy is the sum of all kinetic and potential energy in a system
  - · Over a process, the sum of work and heat is the change in internal energy
  - $\cdot \Delta U = q + w$

Practice: A fire piston is a device for starting fires by rapidly plunging an airtight piston with a combustible material on its end.

If you push the plunger with a force of  $750\,N$  over a distance of  $10\,cm$ , what is w for the fire piston?  $(75\,J)$ 

If the fire piston is not perfectly insulated, and loses 5~J of heat to the surroundings, what is  $\Delta U$  for the system? (70 J)

# 6.3 Energy as a State Function

- o Some quantities are *Path Functions*, while others are *State Functions*
- To understand a state function, first consider a path function
  - · Path functions depend on the path taken
  - If I push a chair to one side of the room, then pull it back, it will be in the exact same state as it started, but the total w done on the system is not 0
  - · Work is a path function, and depends on the path (I might increase the work by taking a different path)
- o State functions do not depend on the path, but only on the state
  - · If I lift a chair up, then place it on the ground again, its final elevation will be the same, regardless of the path it took
  - · Temperature, elevation, and many thermodynamic quantities are state functions
  - · State functions are useful because you only need to know the current state to know the function's value
- $\circ$  While q and w are path functions, U is a state function
- Consider two very different ways to produce ethanol gas:
  - $\cdot C_2H_6O(s,-115 \,^{\circ}C) \longrightarrow C_2H_6O(g,79 \,^{\circ}C)$
  - $\cdot \ _{2}C(s) + _{3}H_{2}(g) + _{\frac{1}{2}}O_{2}(g) \longrightarrow C_{2}H_{6}O(g,79 \, ^{\circ}C)$
  - $\cdot$  Because the final states are identical, the final Us are identical
  - $\cdot$  Any other process (maybe with many steps) will have the same final U as long as it has the same final state

### 6.4 Energy and Enthalpy

- o All chemical reactions involve at least some exchange of heat, but many involve work as well
- o When gasses are produced or consumed, the pressure and volume will change, doing some work
- $\circ$  This work is measured by  $w = -P\Delta V$ , where  $1\ Latm = 101.325\ J$
- $\circ$  This PV work is actually a bit problematic when trying to keep track of energy
  - $\cdot T$  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)
  - · While U = q + w, H = q under constant pressure conditions
  - $\cdot$  So, we only need to worry about heat when we deal with H
- $\circ$  Processes with -H are called *exothermic*
- Examples of exothermic processes are downward phase changes, combustion reactions, etc.
- $\circ$  Processes with +H(+q) are called *endothermic*
- o Examples of endothermic processes are upward phase changes, ice-pack chemical reactions, etc.
- o Demo An endothermic reaction:  $8.5 g \text{ NH}_{2}\text{SCN}$  with  $16.0 g \text{ Ba}(\text{OH})_{2} \cdot 8 \text{ H}_{2}\text{O}$

#### 6.5 Specific Heat

- 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
- $\circ q = mc\Delta T$  where m is the mass, c is the specific heat, and  $\Delta T$  is the temperature change
- o Table 6.3 includes specific heats for a number of common substances

Practice: How much heat is required to heat 12.5 g of water by  $5.75 \,^{\circ}C$ ? (30.1 J)

If a 5.23 g block of Al at 22.0 °C is given 75.0 J of heat, what will its final T be? (38.1 °C)

#### 6.6 Calorimetry: Measuring Energy Changes

- o 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*
- o Constant-pressure (coffee-cup) calorimetry
  - · The reaction is carried out in aqueous solution
  - $\cdot q_{rxn}$  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  $(10 g, 100 ml, 44.2 \frac{kJ}{mol})$
- o Constant-volume (bomb) calorimetry
  - · The reaction is carried out in a chamber charged with high pressure O<sub>2</sub>
  - $\cdot q_{rxn}$  is exchanged with the whole bomb-calorimeter apparatus
  - · The calorimeter is calibrated to give a *heat capacity*  $(C_{cal})$  with units  $\frac{J}{\circ C}$

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

- For constant volume, we measure  ${\cal U}$  instead of  ${\cal H}$  because w=0

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

# 6.7 Enthalpy in Chemical Reactions

- $\circ$  A balanced chemical reaction may also include an enthalpy of reaction  $\Delta H$
- $\circ~$  This tells how much heat is produced or consumed with one mole of reaction
- $\begin{array}{l} \circ \ \Delta H \ {\rm can \ be \ a \ conversion \ factor \ between \ heat \ and \ amounts \ of \ reactants \ or \ products} \\ {\rm Practice: \ Consider \ the \ reaction \ N_2(g) + 3 \ H_2(g) } \longrightarrow \ {\rm 2 \ NH_3(g) \ } \Delta H = -92 \frac{kJ}{mol} \\ {\rm If \ } 2.25 \ g \ {\rm of \ H_2 \ are \ consumed \ in \ the \ above \ reaction, \ how \ much \ heat \ is \ released? \ (34.2 \ kJ) } \\ {\rm If \ } 54.6 \ kJ \ {\rm of \ heat \ are \ released, \ how \ many \ } g \ {\rm of \ NH_3 \ will \ be \ produced? \ (20.2 \ g)} \\ \end{array}$
- $\circ$  Because enthalpy is a state function, we can calculate values of  $\Delta H$  without measuring them
- $\circ~$  Hess's Law: Any alternate path with the same starting and ending states will have the same overall  $\Delta H$
- o Drawing an energy level diagram can help to illustrate Hess's law

$$\begin{array}{ll} \circ \ \, {\rm Find} \ \Delta H \ {\rm for \ this \ reaction:} \ {\rm C}_{\rm diamond} + {\rm O2(g)} & \longrightarrow {\rm CO2(g)} & (\Delta H = -395.4 \ ^{kJ}\!/mol) \\ \\ {\rm C}_{\rm diamond} & \longrightarrow {\rm C}_{\rm graphite} & \Delta H = -1.9 \ ^{kJ}\!/mol \\ \\ {\rm C}_{\rm graphite} + {\rm O2(g)} & \longrightarrow {\rm CO2(g)} & \Delta H = -393.5 \ ^{kJ}\!/mol \\ \end{array}$$

 $\circ$  The reverse of a reaction gives  $-\Delta H$ 

$$\begin{array}{lll} \circ & \mbox{Find } \Delta H \mbox{ for this reaction: } \mathrm{C(s)} + \frac{1}{2} \operatorname{O2(g)} & \longrightarrow & \mathrm{CO(g)} & (\Delta H = -110 \ ^{kJ}/_{mol}) \\ & \mathrm{C(s)} + \mathrm{O_2(g)} & \longrightarrow & \mathrm{CO_2(g)} & \Delta H = -393 \ ^{kJ}/_{mol} \\ & \mathrm{CO(g)} + \frac{1}{2} \operatorname{O_2(g)} & \longrightarrow & \mathrm{CO_2(g)} & \Delta H = -283 \ ^{kJ}/_{mol} \\ \end{array}$$

- $\circ$  Double the reaction gives double the  $\Delta H$
- $\circ$  Consider trying to find  $\Delta H$  for the reaction below:

$$(\star) \, \mathrm{C_2H_5OH(l)} + 2 \, \mathrm{O_2(g)} \, \longrightarrow \, 2 \, \mathrm{CO(g)} + 3 \, \mathrm{H_2O(l)} \qquad \Delta H = ?$$

• Find an alternate path using these reactions with known  $\Delta H$ :

- o The enthalpy of the first reaction can be found from the enthalpies of the other two
- $\circ \ \Delta H_{\star} = \Delta H_A 2\Delta H_B = -801 \frac{kJ}{mol} \quad \text{(Draw the energy level diagram)}$

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

$$\begin{picture}(B){\ S(s) + O_2(g)} &\longrightarrow {\ SO_2(g)} \\ \hline \end{picture} \Delta H = -296.8 \, {\it kJ/mol}$$

$$(C)$$
 C(s) + 2 S(s)  $\longrightarrow$  CS<sub>2</sub>(l)  $\Delta H = 87.9 \, kJ/mol$ 

### 6.8 Standard Enthalpies of Formation

- $\circ$  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
- $\circ$  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)

- $\circ~$  The enthalpy for this reaction is called the compound's  $\mathit{Standard}$   $\mathit{Enthalpy}$  of  $\mathit{Formation}$   $\left(\Delta H_f^\circ\right)$
- $\circ~$  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  $\Delta H_f^\circ$  values
- · Table 6.4 gives a few values, but appendix A2 is much more comprehensive

$$\begin{aligned} \text{Practice: Find } \Delta H_{rxn} \text{ for } & \text{P}_4 \text{O}_{10}(\text{s}) + 6 \, \text{H}_2 \text{O}(\text{l}) & \longrightarrow 4 \, \text{H}_3 \text{PO}_4(\text{aq}) & \left( \Delta H_{rxn} = -1042.6 \, \frac{kJ}{mol} \right) \\ & \text{Find } \Delta H_{rxn} \text{ for } & \text{C}_3 \text{H}_8(\text{g}) + 5 \, \text{O}_2(\text{g}) & \longrightarrow 3 \, \text{CO}_2(\text{g}) + 4 \, \text{H}_2 \text{O}(\text{g}) & \left( \Delta H_{rxn} = -2043.9 \, \frac{kJ}{mol} \right) \end{aligned}$$

**GASES** 

#### 7.1 Gas Pressure

- Figure 7.1 shows the different phases from a molecular perspective
- The force a gas exerts is called pressure
- Pressure can be measured in many ways, including with a barometer (Figure 7.2)
- o Atmospheric pressure depends on elevation, temperature, and humidity
- $\circ 1 atm$  is just a standard for pressure under certain conditions
- $\circ$  There are two other common units for pressure:  $1 \ atm = 760 \ torr = 760 \ mmHg$
- $\circ~$  And two less common units for pressure: 1~atm = 101.325~kPa = 1.01325~bar

# 7.2 Boyle's Law

- $\circ$  In the 17<sup>th</sup> and 18<sup>th</sup> century, scientists were studying how gases responded to temperature and pressure
- o Boyle studied how volume changed with pressure at constant temperature (Figure 7.4)
- Higher pressures lead to proportionately lower volumes
- $\circ \ V \propto \frac{1}{P} \qquad PV = k \qquad P_1 V_1 = P_2 V_2$

# 7.3 Charles's Law

- Charles studied how volume changed with temperature at constant pressure (Figure 7.10)
- Higher temperatures led to proportionately higher volumes
- $\circ \;$  Figure 7.9 shows how the V/T curve can be extrapolated back to absolute 0 Indeed, this method gave rise to the notion of absolute 0
- $\circ$  Consider doubling the temperature at 1 K and at 1  $\circ$  C. In the Kelvin scale, "double the temperature" has real, physical meaning (doubling the thermal energy)
- We *must* use the Kelvin temperature scale in these types of problems
- $\circ \ V \propto T \qquad \frac{V}{T} = k \qquad \frac{V_1}{T_1} = \frac{V_2}{T_2}$

### 7.4 The Combined Gas Law

 Boyle's law and Charles's law can be combined to compare states with different temperatures, volumes, and pressures

$$\circ \ \, \frac{P_1V_1}{T_1} = \frac{P_2V_2}{T_2}$$

Practice: A balloon with V=1.25~L at P=0.850~atm and T=284~k is released into the atmosphere where P=0.430~atm and T=243~K. What is the new volume? (2.11~L)

### 7.5 Avogadro's Law

- $\circ$  For the previous laws, the proportionality constants k were always different for different gases
- $\circ$  Once we established molar masses, scientists could investigate the relationship between n (# of moles) and P, V, and T
- $\circ$  Avogadro's law states that V is proportional to n at constant T and P

### 7.6 Ideal Gas Law

- $\circ\;$  Avogadro's law was the final piece to get a universal gas constant. k became R
- $\circ \ \frac{PV}{nT} = R \ \text{is usually rearranged as} \ PV = nRT$
- $R = 0.08206 \frac{L \, atm}{mol \, K}$ , but there are also other useful units (see Wikipedia page for R)
- This is called the *ideal gas law*. An *ideal gas* is one that follows this law perfectly
- $\circ$  Real gases will deviate from this law a little, but it is reliable at normal temperatures and pressures Practice: Standard temperature and pressure are defined as  $0.00\,^{\circ}C$  and  $1.00\,atm$  Find the volume of  $1.00\,mol$  of an ideal gas at STP (22.4 L)

# 7.7 Dalton's Law of Partial Pressures

- o In a mixture of gases, each component will exhibit a pressure the same as if the others weren't there
- o The total pressure is the sum of all these *partial pressures*
- This means that we can find the total pressure of a mixture of gasses by counting the total moles of all gases present
- We can find the pressure of one component from the total pressure and the relative composition
- $\circ P_A = \chi_A P_{total}$

Practice:  $\rm O_2$  accounts for about 21% of the molecules in our atmosphere. Find the partial pressure of  $\rm O_2$  based on today's barometric pressure

### 7.8 Molar Mass and Density in Gas Law Calculations

 $\circ$  We can modify the ideal gas law to relate the molar mass of a gas to its density

$$\circ PV = nRT \to \frac{n}{V} = \frac{P}{RT} \to \frac{nM}{V} = \frac{PM}{RT} \to d\left(\frac{g}{L}\right) = \frac{PM}{RT}$$

 $\circ~$  Note that this is a density in  $\frac{g}{L}.$  Gases are very diffuse and have very low densities

Practice: There are two ways we commonly create buoyancy in the atmosphere: He, and hot air. Calculate the density of regular air (assume pure N<sub>2</sub>) and He at STP, and the density of regular air at  $325\,^{\circ}C$  (0.625 g/L, 0.179 g/L, and 0.285 g/L)

Practice: An unknown gas is found to have density of  $0.575~\frac{g}{L}$  at  $22.4~^{\circ}C$  and 0.87~atm. Give a reasonable guess for the chemical identity of the unknown gas (CH<sub>4</sub>)

### 7.9 Gases in Chemical Reactions

o We can also use the ideal gas law to predict volumes or pressures of a product in a chemical reaction

Practice: 0.25 g of Na are reacted with acid according to the following equation:

$$2 \operatorname{Na}(s) + 2 \operatorname{H}^{+}(aq) \longrightarrow 2 \operatorname{Na}^{+}(aq) + \operatorname{H}_{2}(g)$$

If T = 295 K and P = 0.875 atm, what volume will the  $H_2(g)$  product occupy? (0.150 L)

Practice: Consider the reaction:  $2 \text{ Hg}_2 O(s) \longrightarrow 4 \text{ Hg}(l) + O_2(g)$ 

2.50~g of  ${\rm Hg_2O}$  are placed in a sealed 0.500~L chamber at 0.82~atm and decomposed through heat. Once thermal equilibrium is reached at T=297.5~K, what will be the final pressure inside the chamber? (0.97~atm)

#### 7.10 Kinetic Molecular Theory of Gases

- The kinetic molecular theory of gases is a model which can be used to derive the ideal gas law from first principles
- There are five postulates which define the theory:
  - 1. Gases are composed of small particles that are in constant, random motion (Video 7.20)
  - 2. The volume that is taken up by the molecules themselves is vanishingly small compared to the total volume of the gas
  - 3. Forces between the particles are negligible
  - 4. Molecular collisions are perfectly elastic
  - 5. The average kinetic energy of the gas molecules is directly proportional to the temperature
- These postulates are grounded in mathematical equations which can be analyzed through statistical dynamics to apply to large ensembles of particles and give observable state properties
  - · Pressure results from the billions of collisions between gas particles and the chamber walls
  - · Adding more gas particles will increase the frequency of collisions, raising the pressure
  - The average kinetic energy depends only on temperature, not chemical identity of the gas. This explains Dalton's law of partial pressures

# 7.11 Movement of Gas Particles

- $\circ$  The equation for the  $\flat$ 5 postulate is:  $KE_{avg}=\frac{3}{2}k_BT$
- o  $KE_{avg}$  is the average kinetic energy, and  $k_B$  is the *Boltzmann* constant
- $\circ$  We don't really need to use the Boltzmann constant because  $N_A k_B = R$ , and it is useful to talk about molar amounts anyway
- $\circ \;$  So, we get  $KE_{avg} = \frac{3}{2}RT$  You should actually use  $R = 8.314 \, \frac{J}{mol \; K}$
- $\circ \ \ \text{We also know that} \ KE = \frac{1}{2} M v^2 \ (\text{note that this is also a molar amount}), \\ \text{so we can write:} \ \frac{1}{2} M v^2 = \frac{3}{2} RT$
- $\circ~$  This can be solved to give the rms average velocity for a gas particle:  $v_{rms} = \sqrt{\frac{3RT}{M}}$
- $\circ~$  Note the "unit purgatory" issue here you must use  $J\equiv\frac{kg~m^2}{s^2}$  , and express M in kg
- o rms velocity is the square-root of the average of the squared velocities
- $\circ \ v_{rms}$  is used rather than mean velocity because the temperature is proportional to  $v^2$  , rather than v itself
- $\circ$  Figures 10.13 and 10.14 from BLMB show different velocity distributions for different gases and conditions Practice: Find  $v_{rms}$  for N<sub>2</sub>, SF<sub>6</sub>, and He at  $-20.00\,^{\circ}C$  and  $150.0\,^{\circ}C$

$$-20\,{}^{\circ}C(474.8\,{}^{m/s}\!,\,207.9\,{}^{m/s}\!,\,1256\,{}^{m/s}\!) \quad 150\,{}^{\circ}C(613.8\,{}^{m/s}\!,\,269.4\,{}^{m/s}\!,\,1624\,{}^{m/s}\!)$$

- $\circ~$  Diffusion is the spread of a gas from areas of high concentration to areas of low concentration
  - · Diffusion happens much more slowly than  $v_{rms}$  because the molecules collide so frequently
  - · Gas molecules take a random walk, deflected in a new direction with each collision
  - The *mean free path* is only about 70 nm for STP
  - · Higher pressures decrease the mean free path and slow down diffusion
  - · In a vacuum, the diffusion rate approaches  $v_{rms}$
- o Effusion is the slow leak of a gas through a hole
- $\circ$  These two processes both depend on  $v_{rms}$
- $\circ~$  Graham's law compares the effusion/diffusion rates of two gases:  $\frac{r_1}{r_2} = \sqrt{\frac{M_2}{M_1}}$

Practice: Effusion can be observed by reinforcing a balloon with tape, then poking a hole through the tape

Compare the effusion rate for a He and  $\rm N_{\rm 2}$  balloon  $\left(r_{\rm He}=2.65r_{\rm N_{\rm 2}}\right)$ 

### 7.12 Behavior of Real Gases

- o The ideal gas law is technically only valid for an ideal gas with no attractive/repulsive forces
- Real gas particles are attracted to each other, albeit weakly
- o These attractions are particularly important at high pressures and temperatures
- $\circ~$  For one mole of an ideal gas,  $\frac{PV}{RT}=1$  at all temperatures and pressures
- $\circ~$  Figure 7.26 shows how  $\frac{PV}{RT}$  varies with pressure and temperature
- Figures 10.19 and 10.20 from BLMB show the compression factor for different gases and conditions
- o The van der Waals equation is an improvement on the ideal gas law, which works better for real gases

$$\cdot \ \left[ P + a \left( \frac{n}{V} \right)^2 \right] (V - nb) = nRT \quad \text{ or- } \quad P = \frac{nRT}{V - nb} - a \left( \frac{n}{V} \right)^2$$

- $\cdot$  The a term accounts for attractions between gas particles and reduces the pressure of real gases
- $\cdot$  The b term accounts for the finite volume of gas particles and increases the pressure of real gases
- · Table 7.4 gives van der Waals a and b terms for many common gases

Practice: Use both the ideal gas law and the van der Waals equation to find the pressure for:

 $1.00\,mol$  of CCl  $_{\!\scriptscriptstyle A}$  at  $250\,K$  in  $10.0\,L$  and  $1.00\,L$ 

(2.0515 atm, 1.881 atm, 20.515 atm, and 37.79 atm)

### THE QUANTUM MODEL OF THE ATOM

## 8.1 A Brief Exploration of Light

- o Light is an electromagnetic wave, which can be thought of like a wave on a lake (Figure 8.2)
- $\circ$  Light has a wavelength, frequency, and speed according to the equation:  $\nu\lambda=c$
- $\circ~$  The speed of light is a constant,  $2.998 \times 10^8 \ \frac{m}{s}$
- How far does light travel in  $5.00 \, ms$ ?  $(1.50 \, km)$
- The electromagnetic spectrum is more than just visible light (Figure 8.4)
  - · 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  $10^6 \ s^{-1}$ )
- o Light-matter interactions were central to the discovery of modern physics
- The photoelectric effect was an important matter/light interaction
  - · 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,  $\nu$  was a new constant, called Planck's constant, and  $\phi$  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 8.5)
- · 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 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

### 8.2 The Bohr Model of the Atom

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

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

## 8.3 Electron Shells, Subshells, and Orbitals

- 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
  - 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
- o 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
  - · 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 3<sup>rd</sup> shell? (18)

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

- o Subshells each have orbitals with different shapes (Figure 8.14)
  - · 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

## 8.4 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)
- o Draw the subshells with one line for each orbital
- Each orbital can hold two electrons, drawn as up- and down-arrows
- o Find the total number of electrons for the element or ion
- o 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)
- o This is the ground-state configuration of the element

## 8.5 Electron Configurations

- We can use the periodic table as a cheat-sheet to the order of the subshells and electron configurations
  - · 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 (d(-1))
  - The order of the subshells is found by simply following the elements and noting in which block they reside
- 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)
- o The Lanthanum and Actinium boundary also shows some anomalies (don't memorize them)
- 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

## 8.6 Quantum Numbers

- o Remember that orbitals are actually mathematical functions
- $\circ$  Certain parts of those functions depend on integer numbers (like n in the Rydberg equation)
- o These integer numbers are called *quantum numbers*
- o 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
  - $\cdot \ l$  is why not all shells have all orbital types
  - ·  $m_l$  Magnetic quantum number (-l, ..., l) gives orbital within a subshell
  - $\cdot m_l$  is why subshells have different numbers of orbitals
  - $m_s$  Spin quantum number  $(\pm \frac{1}{2})$  gives "spin" of the electron (up- or down- arrow)
- o 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

### PERIODICITY AND IONIC BONDING

### 9.1 Valence Electrons

- $\circ$  Technically, filled d and f subshells count as *core*, and only the outermost s and p electrons will always count as *valence*
- o This means that only the main group elements have a reliable pattern in their number of valence electrons
- o The number of valence electrons is the same as the "A" column names

### 9.2 Atomic and Ionic Sizes

- 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
  - $\cdot$  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 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 V all but one V electron
  - · Note that this is different than what the textbook gives 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 9.3)
- o Anions are much larger and cations are much smaller than their neutral counterparts (Figure 9.4)

# 9.3 Ionization Energy and Electron Affinity

- o Ionization energy (IE) is the energy required to remove an electron
  - · For example, it is the energy for this process: Li  $\longrightarrow$  Li<sup>+</sup> + e<sup>-</sup>
  - · 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 9.6)
- Second- and third- ionization energy is the energy to remove a second and third electron
  - · 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 9.9 is a mess), so don't worry too much about EA

# 9.4 Ionic Bonding

- o Metal and nonmetal elemental atoms will react to form an ionic compound
- The number of electrons gained/lost will result in noble gas configurations for both elements
- o The cation and anion are now attracted to each other, and bind together in a lattice structure
- A *formula unit* is the smallest unit which builds the extended lattice (Figure 9.11)

### 9.5 Lattice Energy

- $\circ$  Consider the reaction: K(s) +  $\frac{1}{2}$  Cl<sub>2</sub>(g)  $\longrightarrow$  KCl(s)  $\Delta H_f^{\circ} = -437 \, \frac{kJ}{mol}$
- The energy of formation is more than just the IE of the metal and EA of the non-metal
- The Born-Haber Cycle is an alternate route for formation reactions of ionic compounds:
  - · The steps of the Born-Haber cycle are shown in Figure 9.13
  - · Sublimating the metal, breaking the diatomic bond, IE, EA, and lattice energy
  - · The lattice energy is directly related to the attractive force between the ions
- Lattice energy is controlled by two factors:
  - · Ions with larger charges have higher lattice energies (Table 9.2)
  - · Smaller ions have higher lattice energies (Figure 9.14)

#### COVALENT BONDING

#### 10.1 Formation of Covalent Bonds

- In both ionic and covalent compounds, bonds will form to complete the atoms' *octets* (*duets* in the case of H)
- o Instead of transferring electrons to form ions, covalent compounds will share electrons
- The electrons in a covalent bond will count toward the octets of both bonding partners
- o Drawing dot structures, we can see how many bonds an atom might need to form to fill its octet
- o We'll usually represent two shared electrons in a bond by a dash this is the beginning of Lewis structures
- Electron pairs not involved in a bond are called lone pairs
- o Single, Double and Triple bonds share 2, 4, and 6 electrons, respectively
- $\circ$  Double and triple bonds are shorter and stronger than single bonds Practice: Draw dot diagrams and Lewis diagrams for the diatomics  $F_2$ ,  $O_2$ , and  $N_2$  Identify the bonds by their type, as well as any lone pairs

### 10.2 Lewis Structures

- For more complex molecules, dot diagrams are not robust enough to predict the proper structure
- o Follow these 8 steps:
  - · Find the total number of valence electrons, considering any overall charge
  - · Write the peripheral atoms around the central atom
    - \* Central atom will be the least electronegative (except H)
    - \* This is often the first atom written in a formula (except H)
    - \* H will never be the central atom because it can only form 1 bond
  - · Connect each peripheral atom to the central atom with single bonds
  - · Place the remaining valence electrons around the peripheral atoms until their octets are complete
  - · If any electrons remain, place them on the central atom
  - · Convert peripheral lone pairs into double or triple bonds according to these two rules
    - \* Make multiple bonds until the central atom has a complete octet
    - \* Make multiple bonds in a way that minimizes *formal charges* (more on this later)
  - · Verify that the octet and duet rule are followed for all atoms in the structure
  - · If there is a charge, enclose the structure in square brackets and write the charge

Practice: CH<sub>4</sub>, H<sub>2</sub>O, NH<sub>3</sub>, HCN, CH<sub>2</sub>O, NO<sub>3</sub><sup>-</sup>, CO<sub>3</sub><sup>2-</sup>, and NH<sub>4</sub><sup>+</sup>

## 10.3 Resonance and Formal Charges

- o Often times we will have an arbitrary choice about which peripheral atom to form a double bond with
- $\circ$  Consider  $CO_3^{2-}$ , you could form the double bond with any of the three oxygens
- o In these instances, the molecules will exhibit a phenomenon called *resonance* 
  - · Resonance is when electrons in a double (or triple) bond are shared between two or more bonding locations
  - $\cdot \,\, \text{CO}_{\scriptscriptstyle 3}^{\,\, 2-}$  forms a bond with all three oxygen atoms
  - It is not nearly as strong as a normal double bond, because the electrons are spread between three bonding centers
  - This is called *delocalization*, and sometimes the trio of bonding locations are collectively called a delocalized bond
  - · The bond lengths are all the same, somewhere between a single and double bond length
- o We represent resonance two ways: First, resonance structures
  - · Draw a different structure for each bonding location
  - · Enclose each structure in square brackets (even for neutral structures)
  - · Draw double-headed arrows between the structures
- o The other method is hybrid structures
  - · Draw a single structure, with dotted lines for each bonding location
  - · These structures are closer to an accurate picture of reality
  - · Counting electrons in these structures is impossible (bonds and lone-pairs)

Practice: Draw the resonance structures and hybrid structure for  $NO_3^-$  and  $O_3^-$ 

- Electrons are not actually shared equally between bonding partners
  - · *Electronegativity* is the measure of how strongly an atom pulls on shared electrons
  - · Electronegativity increases across a row and up a column (Figure 10.7)
  - · Less electronegative atoms will be central atoms (except H)
- Formal charge compares how many electrons an atom has within an molecule to its number of valence electrons
  - · First draw a Lewis structure
  - · Count the electrons around an atom, dividing the bonding electrons between bonding partners
  - · Subtract this number from the number of valence atoms
- o Formal charge can be used to determine which structures are best when there are options
  - · Minimize the total formal charge
  - · Favor structures with negative formal charges on the more electronegative atoms

Practice: Find the best structures for OCN and N<sub>2</sub>O

## 10.4 Exceptions to the Octet Rule

- o A few elements will have *fewer* than 8 electrons
  - · Be has only 2 electrons, so it can form only 2 bonds
  - · B has only 3 electrons, so it can form only 3 bonds
- o Radicals are compounds with an unpaired electron
  - · Radicals tend to be very reactive
  - · Many radicals contain N (NO and NO<sub>2</sub>)
- Some elements can have *more* than 8 electrons
  - · Only elements in the 3rd row or below can exceed the octet rule
  - $\cdot$  This is because the extra electrons go into the d subshell
  - This can occur in order to minimize formal charges Practice: Find proper Lewis structures for  $SO_4^{\ 2-}$ ,  $PO_4^{\ 3-}$ ,  $ClO_3^{\ -}$ , and  $ClO_4^{\ -}$
  - It can also occur because there are simply too many outer atoms Practice: Find proper Lewis structures for SF<sub>6</sub>, ClF<sub>5</sub>, and PCl<sub>5</sub>

# 10.5 Polar Bonds and the Bonding Continuum

- o Figure 10.14 shows electrons being shared equally and unequally between bonding partners
- o The electrons will favor the element with greater electronegativity
- The uneven distribution of charge is called a *dipole* and the bond is called *polar*
- Polar covalent bonds exist along a continuum from purely covalent to purely ionic
- o Greater electronegativity differences give greater ionic character to the bond
- o Similar (or identical) electronegativities create non-polar bonds
- The dipole moment is  $\mu = qr$ , and can be measured
- $\circ$  Real dipole moments can be compared to the value for a complete transfer of electrons to give % ionic character
- $\circ \ \% \ ionic \ character = \frac{\mu_{measured}}{\mu_{ionic}} \times 100\%$

### 10.6 Bond Enthalpy

- o Bond enthalpy is the energy required to break a bond
- Really, not every C—H bond is the same. The surrounding atoms affect bond enthalpy
- We can take the average enthalpy of a given bond type over many molecules (Table 10.3)
- o This gives us a new way to calculate reaction enthalpies

- · Draw a Lewis structure for each reactant and product
- · identify the numbers and types of bonds
- · Consider first breaking all the bonds of the reactants to produce individual atoms
- · Then form new bonds between the atoms to make products

· 
$$\Delta H_{rxn} = \sum_{Bonds\ Broken} \Delta n \cdot H_{Bond} - \sum_{Bonds\ Formed} n \cdot \Delta H_{Bond}$$

- · This will give an *approximate* reaction enthalpy
- o Table 10.4 gives average values for bond lengths as well

Practice: Using Table 10.3, estimate the reaction enthalpy for the combustion of methane  $\left(-810 \frac{kJ}{mol}\right)$ 

### MOLECULAR SHAPE AND BONDING THEORIES

## 11.1 VSEPR and Molecular Geometry

- The three-dimensional shape of a molecule is important to its properties
- Isomers are molecules with the same chemical formula, but a different 3-dimensional arrangement of its atoms
- o Isomers might have quite different properties despite their identical composition
- Valence Shell Electron Pair Repulsion (VSEPR) model:
  - · VSEPR is based on the idea that electron pairs will arrange themselves to be as far apart from each other as possible.
  - · This model gives accurate geometries for covalent molecules
  - · First, draw a good Lewis structure
  - *Electron Domains* are the regions around the central atom where electrons group A single bond, a double bond, a triple bond, and a lone pair are all electron domains
  - The number of electron domains will give the *electron geometry* This electron geometry is the template on which molecular geometry is based
  - · Demo: Balloons naturally adopt the electron geometries
  - · Next, count how many domains are bonding vs lone pairs
  - · The number of bonds within the electron geometry determines the molecular geometry
  - · Table 11.1 sums it all up nicely Just memorize this table
  - · Trigonal Bipyramidal electron geometry
    - \* The 2 axial and 3 equatorial positions are different
    - \* Lone pairs will occupy the equatorial positions first
    - \* Linear molecules are symmetrical (this will matter later)
  - · Octahedral electron geometry
    - \* All the positions are equivalent
    - \* The second lone pair will be opposite the first one
    - \* Square planar molecules are symmetrical (this will matter later)
- For larger molecules, you can apply VSEPR to each bonding center (consider CH<sub>2</sub>CO<sub>2</sub>H)

## 11.2 Polar and Non-polar Molecules

- We already discussed electronegativity and its role in making polar *bonds*
- For molecules with many polar bonds, those dipoles might cancel each other out or work together to make a polar molecule
- The *molecular dipole* is the vector sum of all the bond dipoles
- o Factors that make non-polar molecules:
  - · No polar bonds like diatomic elements and O<sub>3</sub>
  - · Symmetry in the polar bonds (no lone pairs, or linear and square planar molecules)
- Factors that make polar-molecules:
  - Lone pairs which break symmetry
  - · Bonds with different atoms (CH<sub>2</sub>Cl<sub>2</sub>)

# 11.3 Valence Bond Theory: Hybrid Orbitals and Bonding

- Covalent bonding basics:
  - · Why should sharing electrons lead to a bond?
  - · Draw the attractive and repulsive forces in H<sub>2</sub>
  - · When the orbitals from both bonding partner overlap, these forces are optimized to form a bond
  - $\cdot$  Figure 11.17 shows how the energy changes with internuclear distance in  $H_2$
- Orbital overlap cannot explain the actual geometries we see in molecules. No atomic orbitals have a tetrahedral or trigonal planar geometry, for example
- Hybridization is the linear combination (mixing) of atomic orbitals
  - · Remember that orbitals are simply mathematical functions
  - · Consider the functions  $f_1(x) = x^2$ ,  $f_2(x) = x^3$ , and  $g(x) = f_1(x) + f_2(x)$
  - · Atomic orbitals can similarly be combined to form new orbitals, called hybrid orbitals
  - · Show my hybridization figures
  - · The number of hybrid orbitals is equal to the number of atomic orbitals mixed together
  - · We name the orbitals and the type of hybridization by the atomic orbitals used  $(sp, sp^2, and sp^3)$
  - · These hybridization types correspond to the electron geometries
- Hybrid orbitals and bonding
  - · Hybrid orbitals are used for 2 purposes: forming  $\sigma$  bonds, and housing lone pairs
  - $\cdot$   $\sigma$  bonds have electron density aligned along the bond axis
  - · Single bonds are  $\sigma$  bonds, and multiple bonds each contain a  $\sigma$  bond
  - · For sp and  $sp^2$  hybridized atoms, unhybridized p orbitals remain (Figures 11.24 and 11.26)

- These unhybridized p orbitals are used to form  $\pi$  bonds
- $\cdot \pi$  bonds have electron density along either side of the bond axis
- · Double bonds contain one  $\pi$  bond, and triple bonds contain 2  $\pi$  bonds
- · A double bond is like a hot dog in a bun a triple bond is like a hot-dog with 2 buns
- · Use BLMB Figures 9.23-9.28
- · Multiple p orbitals overlap in molecules which exhibit resonance (like benzene and nitrate)

## 11.4 Using Valence Bond Theory

- o Molecules can twist around their bonds without changing bond lengths or angles
- These rotations create different *conformations* (show with molecular models)
- $\circ$  The differences between  $\sigma$  and  $\pi$  bonds has significance to how molecules can move
- $\circ \ \sigma$  bonds have the same overlap no matter how you rotate them, so rotation does not break the bond
- $\circ$  This makes rotating around  $\sigma$  bonds energetically free and easy
- $\circ \pi$  bonds, however, would break if you twist them by  $90^{\circ}$  because they lose their overlap (Figure 11.3)
- $\circ \pi$  bonds are constrained, and cannot rotate without a substantial energy input

# 11.5 Molecular Orbital Theory

- In hybrid orbital theory, we combined atomic orbitals from the same atom to create new hybrid orbitals
- These hybrid orbitals had new shapes, which could explain bonding geometries
- We can apply the same trick to molecules combine atomic orbitals from *different* atoms across a molecule to create new orbitals
- These new orbitals span the whole molecule, and are called *molecular orbitals*
- Molecular orbitals are the closest approximations to the true Schrödinger equation solutions
- Molecular orbitals can combine in two ways (Figure 11.32):
  - · Adding orbitals together creates constructive interference between the orbitals
  - · Orbitals which combine constructively increase the electron density between the atoms
  - · This leads to stronger bonds, so these MOs are called *bonding* orbitals
  - · Subtracting orbitals creates destructive interference between the orbitals
  - · Orbitals which combine destructively decrease the electron density between the atoms
  - · This leads to weaker bonds, so these MOs are called antibonding orbitals (and get a \* label)
  - Both of the orbitals formed from s orbitals have electron density along the bond axis, so they are called the  $\sigma_{ns}$  and  $\sigma_{ns}^*$  orbitals
- Molecular orbital diagrams

- · When we draw molecular orbital diagrams, it is helpful to show the atomic orbitals which gave rise to the MOs
- · Draw the atomic energy level diagram for each atom on either side of the MO energy levels
- · Draw dotted lines showing how the atomic orbitals combine to produce new MOs
- · Fill the MO energy levels from the bottom-up as usual
- $\circ p$  orbitals can combine as well
  - Head-on combination creates two more  $\sigma$  orbitals (Figure 11.35)
  - · Sideways combination creates orbitals with density along either side of the bond axis, so they are  $\pi_{2p}$  and  $\pi_{2p}^*$  MOs
  - · Because there are two pairs of p orbitals that can combine this way, there are two degenerate  $\pi_{2p}$  and two degenerate  $\pi_{2p}^*$  orbitals
  - · The ordering of the  $\pi_{2p}$  and  $\sigma_{2p}$  orbitals is different for different elements (Figure 11.37)
- Figure 11.38 shows the configurations for all period 2 homonuclear diatomics (note the different orbital ordering)
- We can tell the *bond order* (single, double, or triple bond) of a diatomic from its MO diagram
  - · Electron pairs which occupy a bonding orbital increase the bond order by 1
  - · Electron pairs which occupy a non-bonding orbital decrease the bond order by 1
  - · Molecules with a bond order of o (Ne<sub>2</sub>) don't form a bond at all, and so don't occur naturally
- Magnetic properties are also indicated by the MO diagram
  - · *All* materials are either attracted to or repelled by a magnetic field
  - · *Diamagnetic* materials are repelled by a magnetic field, while *paramagnetic* materials are attracted by a magnetic field. Ferromagnetism (regular old magnetism) is like a cooperative paramagnetic effect in some metals
  - · Diamagnetism comes from having all electrons paired. C2, N2, and F2 are diamagnetic
  - · Paramagnetism comes from at least one unpaired electron. B<sub>2</sub> and O<sub>2</sub> are paramagnetic
- $\circ$  Molecules that exhibit resonance will have  $\pi$  bonding MOs which span all of the bonding locations
- o Some molecules without traditional resonance will still show delocalization (1,3-butadiene)
- Show some MOs in Avogadro on my laptop

### **ERRATA**

- Figure 1.13 identifies (c) as neither precise nor accurate, but it *is* accurate within the apparent precision
- $\circ \ \ \text{In Section 4.7, the decomposition of mercury} (II) \ \text{oxide should be: 2 HgO(s)} \ \ \overset{}{\longrightarrow} \ \ _{2} \ \text{Hg(l)} \ + \ O_{_{2}}(g)$
- o The formation of water given in Section 6.8 should produce only 1 mole of water as a product
- o Slater's Rules in section 9.2 should treat deep core and V-1 electrons differently
- $\circ$  Lattice energy is often referenced as positive values, instead of the negative values used here. This convention makes explaining the Born-Haber cycle a little easier because you only need to reverse one value  $\left(\Delta H_f^{\circ}\right)$