AA Properties \& Changes of Matter
Physical properties: a quality of matter that can be observed or measured without changing it. Examples of physical
properties include: color, molecular weight, change of state and volume.
Physical Change: substances may change shapes or forms while undergoing a physical change, but no chemical reactions
occur and no new compounds are produced.
Examples of Physical Changes: crumpling a sheet of paper, melting an ice cube, , dissolving salt in water, casting silver in a

4B Extensive Properties: Depend on the amount of the substance NOT on the identity of the substance - Examples: Mass, Volume Length Intensive Properties: Depend on the identity but NOT dependent on the amount but - Examples: density, boiling point, solubility, specific heat



| 5A-History of Periodic Table |  |  |  |
| :---: | :---: | :---: | :---: |
| Contributor | Date | Contribution | Comment |
| Lothar Meyer | 1869 | Compiled a Periodic Table of 56 elements based on the periodicity of properties such as molar volume when arranged in order of atomic weight. | Meyer \& Mendeleev produced their Periodic Tables simultaneously. |
| Dmitri <br> Mendeleev | 1869 | He wrote out the properties of each element on a different card and spent a great deal of time arranging and rearranging them. He was looking for patterns or trends in the data on the cards. Mendeleev first arranged all the cards from lowest to highest atomic mass. He kept elements with similar chemical and physical properties grouped in families. His genius is shown in his ability to recognize the potential for missing data and to use existing data to predict the properties of these unknown elements. Mendeleev wrote the following Periodic Law: the properties of the elements are a periodic function of their atomic weights. | Mendeleev's Periodic Table was important because it enabled the properties of elements to be predicted by means of the 'periodic law': properties of the elements vary periodically with their atomic weights. |
| Henry Moseley | 1913 | In 1914, Henry Moseley determined the atomic number of each of the known elements. He realized that, if the elements were arranged in order of increasing atomic number rather than atomic weight, they gave a better fit within the 'periodic table'. He puts the elements tellerium and iodine, argon and potassium, cobalt and nickel in the right order. | He modified the 'Periodic Law' to read that the properties of the elements vary periodically with their atomic numbers. |




Atomic Radius:
DOWN a Group: Atomic radius INCREASES as you go DOWN a Group because each successive Period (row) has an additional occupied energy level. With each additional energy level the distance between nucleus and valence energy level increases. ACROSS a Period: Atomic radius DECREASES as you go ACROSS a Period because the net nuclear charge increases (Huh?). Remember, it's the protons (+) in the nucleus that pull on or attract the electrons in the orbitals. Across a Period you are adding more and more protons pulling on electrons occupying the same orbitals. The overall effect is more pulling power in the same basic space. This draws the electrons in closer, making the overall atomic radius smaller as you go left to right across a period. Ionic Radius:
Cations are positively charged ions formed when metals lose electron(s). Cations are smaller than the metal atom that forms the cation because it has fewer electrons and lose an outer energy level.
Anions are negatively changed atoms formed when a nonmetal gains an electron. The anion is larger than the nonmetal that forms the anion because it has more electrons.

## $1^{\text {st }}$ Ionization Energy: Amount of energy it takes to remove the most loosely held electron.

DOWN a Group: $1^{\text {st }}$ Ionization energy DECREASES as you go DOWN a Group because the farther the valence electrons are from the nucleus the weaker the attraction between the valence electrons and the nucleus. Less energy it costs for another atom to steal them.
ACROSS a Period: $1^{\text {st }}$ Ionization energy INCREASES as you go ACROSS a Period because the valence electrons are closer to the nucleus resulting in a greater attraction between the valence electrons and the nucleus. The atomic radius is getting smaller, so the protons do a great job of holding on tighter.
Electronegativity: the ability of the atoms of an element to attract the electron(s) of the atoms of other elements
DOWN a Group: Electronegativity DECREASES as you go DOWN a Group because the valence electrons are increasingly farther away from the attraction of the protons in the nucleus. Less pull, less "desire" to grab other electrons.
ACROSS a Period: Electronegativity INCREASES as you go ACROSS a Period because the number of protons (+ charges) in the nucleus increases. More protons in the nucleus means electrons are more strongly attracted to the nucleus.

## 6A Evidence of Atomic Theory

## Dalton's Atomic Theory:

1. All matter consists of tiny particles called atoms.
2. Atoms are indestructible and unchangeable. Atoms of an element cannot be created, destroyed, broken into smaller parts or transformed into atoms of another element. Dalton based this hypothesis on the law of conservation of mass and on centuries of experimental evidence.
3. All atoms of the same element are identical. Elements are characterized by the mass of their atoms. All atoms of the same element have identical weights,
4. Atoms combine in new ways during a chemical change. When elements react, their atoms combine in simple, whole-number ratios.

Two criteria are usually applied to any theory. First, does it agree with facts which are already known? Second, does it predict new relationships and stimulate additional observation and experimentation? The Theory has been modified to account for new observations.

1. Atoms are indestructible and unchangeable. Atoms of an element cannot be created, destroyed, broken into smaller parts or transformed into atoms of another element. We now know elements have subatomic particles and can be transformed by nuclear change but not by physical or chemical change.
2. All atoms of the same element are identical. Elements are characterized by the mass of their atoms. All atoms of the same element have identical weights. Atoms of the same element may have different numbers of neutrons but all atoms of the same elements have the same number of protons. Now an atom is defined as the smallest particle of an element that retains the properties of the element.


## Experiment J.J. Thompson

J. J. Thomson painted a substance known as "phosphor" so he could see exactly where the cathode rays hit because the cathode rays made the phosphor glow.
He placed a positively charged metal plate on one side of the cathode ray tube and a negatively charged metal plate on the other side of the cathode ray tube, as shown in the above figure. He put an object in the path of the cathode ray in order to observe the shadow that would result. Energy that travels as waves will bend around the object resulting in a shadow with fuzzy edges. If the ray is a stream of particles the ray will not bend around the object and the shadow will have sharp edges.


Observation J.J. Thompson
The Cathode ray traveled away from the negative cathode and towards the positive anode.

The flow of the cathode rays passing through the hole in the anode was bent upwards towards the positive metal plate and away from the negative metal plate.
The edges of the shadow were sharp (not fuzzy). No particles can be seen.

Thomson's experiment also revealed the electron has a very large charge-to-mass ratio.

Based on what was learned about electrons, two other inferences were made about atomic structure.

1. Because atoms are electrically neutral, they must contain a positive charge to balance the negative electrons.
2. Because electrons have so much less mass than atoms, atoms must contain other particles that account for most of their mass

Final Conclusion Plum Pudding model of the atom.
Ernest Rutherford


## Experiment Ernest Rutherford

Gold Leaf Scattering Experiment

1. fire massive alpha particles at the atoms in thin metal foil
2. alpha particles should pass like bullets straight through soft plum pudding atoms

## Observations Ernest Rutherford

 Most of the alpha particle went straight through the gold leaf. But to his great surprise a few alpha particles ricocheted!Conclusion Ernest Rutherford
Rutherford's Atom

- concentrated mass and ${ }^{\text {® }}$
 Most of the alpha particles went straight through the positive charge at the nucleus
- electrons roam empty space around the nucleus gold leaf because most of the atom was empty space. All the mass and positive charged occupies about $1 / 10,000$ the volume. (this is our nucleus).


## The electrons are in the empty space outside the

 center of very dense positive charge called the nucleusThe nucleus has to be extremely dense for the alpha particles to be deflected. For the nucleus to be dense enough to reflect the alpha particles it had to have very little volume and almost all the mass.
Niels Bohr Observations, Conclusions and Model of the Hydrogen Atom
Observation: that light emitted from samples of atom exposed to energy (energizing the atoms) when passed through a prism emit distinct wavelengths of light line spectrum instead of emitting all the colors of light. Each element emits the same line spectrum and are just as characteristic to that element as finger prints are to people.
Hypothesis: if energized atoms emit only discrete wavelengths, then electrons can have only discrete energies

## Conclusion:

 Bohr's Theory- Electrons occupy orbits at discrete distances from the nucleus called energy levels.
- Electrons in an non-energized atom occupy the lowest energy orbit (closest to nucleus this is called ground state.) - Electrons in energized atoms absorb just enough energy to move from a lower energy orbital to a higher energy orbit (further from nucleus) this is called excited state.

Electrons do not remain in excited state and return to ground state releasing energy that corresponds to all the possible difference in energy of the allowed orbits in the atom.




## 6B Electromagnectic Waves

Electromagnetic radiation can travel through empty space. Most other types of waves must travel through some sort of substance. For example, sound waves need either a gas, solid, or liquid to pass through in order to be heard.
The speed of light is always a constant. c= (Speed of light: $2.99792458 \times 10^{8} \mathrm{~m} \mathrm{~s}^{-1}$ or
$3.00 \times 10^{8} \mathrm{~m} / \mathrm{s}$ ) Wavelength $(\lambda)$ is the distance of one full cycle of the oscillation. Wavelengths are measured between the distances between two consecutive points on a wave which can be between consecutive troughs and crests. It is usually characterized by the Greek symbol $\Lambda$.
Frequency $(f)$ is defined as the number of cycles per second, and is expressed as sec ${ }^{-1}$ or Hertz ( Hz ).
Frequency is directly proportional to energy (High frequency =High energy) and can be express as: $\boldsymbol{E}=\boldsymbol{h f}$
Where $E$ is energy, h is Planck's constant and $f$ is frequency. Planck's constant: $\mathrm{h}=6.62607 \times 10^{-34} \mathrm{~J} \mathrm{~s}$

$$
C=\lambda f
$$

Where $\mathbf{C}$ is the speed of light, $\boldsymbol{\lambda}$ stands for the wavelength, and $\boldsymbol{V}$ (nu) stands for frequency. Shorter wavelength means greater frequency, and greater frequency means higher energy. Wavelengths are important in that they tell one what type of wave one is dealing with.

In general as the energy increases the wavelength decreases and the frequency increases. $\uparrow E \downarrow \lambda \uparrow f$
The opposite is also true as the energy decreases the wavelength increases and the frequency decreases. $E \uparrow \lambda \downarrow f$

6C Calculating Wavelength, Frequency and Energy

1) A laser used in eye surgery to fuse detached retinas produces radiation with a wave length of 640.0 nm . Calculate the frequency

$$
\begin{gathered}
c=\lambda f \\
\frac{c}{\lambda}=\frac{\lambda f}{\lambda} \text { or } f=\frac{c}{\lambda} \\
f=\frac{3 \times 10^{8} \frac{\mathrm{~m}}{\mathrm{~s}}}{640.0 \mathrm{~mm} \times \frac{1 \mathrm{~m}}{1 \times 10^{9} \mathrm{~nm}}}=4.69 \times 10^{14} \frac{1}{\mathrm{~s}} \text { or hertz }
\end{gathered}
$$

2) How much energy is in a photon of the laser light? (using the frequency calculated above)

$$
E=h f=6.626 \times 10^{-34} J s \times 4.69 \times 10^{14} 1 / s=3.11 \times 10^{-19} J \text { per photon }
$$

3) Visible light has frequency between about $4.0 \times 10^{14}$ hertz and $7.9 \times 10^{14}$ hertz or $1 / \mathrm{s}$. What is the wavelenght of the lowest frequency of light?

$$
\begin{gathered}
c=\lambda f \lambda=\frac{c}{f}= \\
\frac{3 \times 10^{8} \mathrm{~m} / \mathrm{s}}{4.0 \times 10^{14} 1 / \mathrm{s}} \\
=7.5 \times 10^{-7} \mathrm{~m}
\end{gathered}
$$

## 6D Calculating Average Atomic Mass

Relative abundance $=\frac{\text { \% abundance }}{100}$

## (atomic mass of isotope $\times$ relative abundance) $\times$ (atomic mass of isotope $\times$ relative abundance) $\cdots=$ average atomic mass Example:

The most abundant isotopes of silicon are silicon- 28 , silicon- 29 , and silicon- 30 . Given the atomic masses and abundances in the table below, what is the average atomic mass of silicon?

| Isotopic Composition of Silicon |  |  |
| :--- | :---: | :---: |
| Isotope | Atomic mass (amu) | Approximate abundance <br> (percent) |
| Silicon-28 | 27.97693 | 92.23 |
| Silicon-29 | 28.97649 | 4.68 |
| Silicon-30 | 29.97377 | 3.09 |

Remember we are changing percent abundance to relative abundance by dividing by 100 or moving the decimal two places to the left. The equation above will be used to calculate the atomic mass in amu.
$(27.97693 \mathrm{amu} \times .9223)+(28.97649 a m u \times .0468)+(29.97377 a m u \times .0309)=\mathbf{2 8 . 1} \mathbf{a m u}$

## 6E Electron Configurations \& Lewis Dot Structures

Atomic Orbitals and Electrons in Principal Energy Levels

| Principal <br> energy level | Type of <br> sublevel | Number of orbitals <br> in sublevels | Maximum number <br> of electrons |
| :---: | :--- | :--- | :---: |
| 1 | s | 1 | 2 |
| 2 | $\mathrm{~s}, p$ | $1+3=4$ | 8 |
| 3 | $s, p, d$ | $1+3+5=9$ | 18 |
| 4 | $s, p, d, f$ | $1+3+5+7=16$ | 32 |

Principal quantum numbers describe the size, shape, and energy of the orbital an orbital is probable location of an electron.

- Principal energy level (principal quantum number) $=n$ describes the size
- Sublevel $=\ell$ describes the shape of the probable location of an electron
- Orbital = each sublevel has a set number of orbitals of a given size and shape that can be occupied by a maximum of 2 electrons.
The location closest to the nucleus has the least amount of energy. This energy increases as you get further from the nucleus. Electrons occupy the lowest energy orbitals first.

lectron Configurations: Assigning each electron in an atom to the energy level and sublevel it occupies in the atom. Number of Electrons

Energy Level $\leftarrow$ $\qquad$ Sublevel
Each orbital can hold no more than two electrons. Two electrons in the same orbital must have
Example: Ni atomic number is 28 so it has 28 protons and 28 electrons because it does not have a charge. Electron 1 and 2 are in $1 s$, the $3^{\text {rd }}$ and $4^{\text {th }}$ electrons are in 2 s , the next six electrons which are the $5^{\text {th }}$ to the $10^{\text {th }}$ are in the $2 p$, the $11^{\text {th }}$ and $12^{\text {th }}$ electron are in the 3 s , the next six electrons which are $13^{\text {th }}$ to the $18^{\text {th }}$ are in the 3 p sublevels orbitals, the $19^{\text {th }}$ and $20^{\text {th }}$ electron occupy the 4 s orbital, the last 8 electrons will occupy the $3 d$

f block

| 4 f | 57 | 58 | 59 | 60 | 61 | 62 | 63 | 64 | 65 | 66 | 67 | 68 | 69 | 70 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $5 f$ | 89 | 90 | 91 | 92 | 93 | 94 | 95 | 96 | 97 | 98 | 99 | 100 | 101 | 102 |

## Lewis Dot Diagrams

- A Lewis dot structure is like a simplified electron energy level model.
- The Lewis structure contains the element symbol with dots representing electrons
- The only electrons shown are those on the outer energy level or valence electrons
- The electrons are placed around the element symbol, one at a time, clockwise or counterclockwise, and then grouped in pairs as more electrons are added.


## Hydrogen $1 s^{1} \mathrm{H}$.

Oxygen $1 s^{2} 2 s^{2} 2 p^{4} \cdot 0$.

$\begin{array}{lll}\text { Chloride } \\ \text { ion } \mathrm{Cl}^{-1} & 1 s^{2} 2 s^{2} 2 p^{6} 3 s^{2} 3 \mathrm{p}^{6} \quad: \quad \because 1:\end{array}$

$\bullet \mathrm{Na} \longrightarrow \mid \mathrm{Na}]^{+}+\mathrm{e}^{-}$

| 7A Naming Compounds |  |
| :--- | :--- |
| Must Memorize to Name Covalent <br> Compounds <br> Prefixes used for naming <br> covalent compounds  <br> Number Prefix |  |
| 1 | mono- * |
| 2 | di- |
| 3 | tri- |
| 4 | tetra- |
| 5 | penta- |
| 6 | hexa- |
| 7 | hepta- |
| 8 | octa- |
| 9 | nona- |
|   <br> 10  |  |
| *The prefix mono- is only used for the more <br> electronegative element. |  |


| Need to know the following to name acids |  |  |  |
| :--- | :--- | :--- | :--- |
| Anion ending | Formula | Acid name | Example |
| ide | HCl | hydro-(stem)-ic acid | hydrochloric acid |
| -ite | $\mathrm{H}_{2} \mathrm{SO}_{3}$ | (stem)-ous acid | sulfurous acid |
| -ate | $\mathrm{HNO}_{3}$ | (stem)-ic acid | nitric acid |

## Need to know when to add roman numerals to

 metal name| Ion Name | Ion | Ion Name | Ion |
| :--- | :--- | :--- | :--- |
| Iron(II) | $\mathrm{Fe}^{2+}$ | Silver(I) | $\mathrm{Ag}^{+}$ |
| Iron(III) | $\mathrm{Fe}^{3+}$ | Silver(II) | $\mathrm{Ag}^{2+}$ |
| Copper(I) | $\mathrm{Cu}^{+}$ | Mercury(I) | $\mathrm{Hg}^{+}$ |
| Copper(II) | $\mathrm{Cu}^{2+}$ | Mercury(II) | $\mathrm{Hg}^{2+}$ |

Three Types of Compounds to Name:

1. Ionic compounds: $1^{\text {st }}$ element a metal
2. Covalently bonded compounds: $1^{\text {st }}$ element a nonmetal
3. Acid: $1^{\text {st }}$ element is H

## Naming Ionic Compounds: First Element is a Metal

| POLYATOMIC IONS |  | O D D D |
| :---: | :---: | :---: |
| Acetate $\mathrm{C}_{2} \mathrm{H}_{3} \mathrm{O}_{2}^{-}$ | $\mathrm{CH}_{3} \mathrm{COO}^{-}$ | $\stackrel{1}{0}$ |
| Ammonium | $\mathrm{NH}_{4}^{+}$ | $\xrightarrow{\text { ¢ }}$ |
| Carbonate | $\mathrm{CO}_{3}^{2-}$ | ? |
| Chlorate | $\mathrm{ClO}_{3}^{-}$ | 0 |
| Chlorite | $\mathrm{ClO}_{2}^{-}$ | S |
| Chromate | $\mathrm{CrO}_{4}^{2-}$ | - |
| Cyanide | $\mathrm{CN}^{-}$ | $\bigcirc$ |
| Dichromate | $\mathrm{Cr}_{2} \mathrm{O}_{7}^{2-}$ | $\stackrel{-1}{0}$ |
| Hydrogen carbonate | $\mathrm{HCO}_{3}^{-}$ | - |
| Hydroxide | $\mathrm{OH}^{-}$ |  |
| Hypochlorite | $\mathrm{ClO}^{-}$ |  |
| Nitrate | $\mathrm{NO}_{3}^{-}$ |  |
| Nitrite | $\mathrm{NO}_{2}^{-}$ |  |
| Perchlorate | $\mathrm{ClO}_{4}^{-}$ |  |
| Permanganate | $\mathrm{MnO}_{4}^{-}$ |  |
| Phosphate | $\mathrm{PO}_{4}^{3-}$ |  |
| Sulfate | $\mathrm{SO}_{4}^{2-}$ |  |
| Sulfite | $\mathrm{SO}_{3}^{2-}$ |  |


| First word |
| :--- |
| lonic name of metal |
| (roman numerial is added only if the metal has more than one possible |
| charge) |

## Second word

name of anion( if it is an ion of one nonmetal or $\mathrm{OH}^{-}$or $\mathrm{CN}^{-}$the ion ends in ide) polyatomic ions other than $\mathrm{OH}^{-}$and CN - can end in ate or ite
$\mathrm{KNO}_{3}$ is named potassium nitrate.
Iron in FeO must have a +2 charge because oxygen has a-2 charge ( $\mathrm{O}^{-2}$ ). The compound's name is iron(II) oxide
Tin in $\mathrm{SnS}_{2}$ must have a +4 charge because $2 \mathrm{~S}^{-2}$ have a -4 charge. The name of the compound is tin(IV) sulfide
$\mathrm{BaCl}_{2}$ is named barium chloride
$\mathrm{Na}_{2} \mathrm{SO}_{3}$ is named sodium sulfite
$\mathrm{K}_{2} \mathrm{Cr}_{2} \mathrm{O}_{7}$ is named potassium dichromate
$\mathrm{Cu}_{3} \mathrm{PO}_{4}, \mathrm{PO}_{4}$ has a -3 charge so each of the 3 Cu must be +1 to $=+3$. The name of the compound is copper(I) phosphate

## Naming Covalently Bonded Compounds: First Element a Nonmetal

| First word <br> *Prefix + name of first element$\quad+\quad$Second word <br> *Prefix + usually first sylabol of second element + ide |
| :--- |

*The prefix mono- is only used for the more electronegative element. Prefixes are listed in a table on the page above.
$\mathrm{CO}_{2}$ Carbon dioxide $\quad \mathrm{As}_{2} \mathrm{Cl}_{5}$ diarsenic pentachloride $\quad \mathrm{CCl}_{4}$ carbon tetrachloride

## Naming Acids: First Element is H: Two Type of Acids Binary and Nonbinary

Binary Acid contains only 2 elements

## First word

hydro-(stem)-ic (the stem is usually the first syllable of second element)

```
Second word
acid
```

$\mathrm{HBr}(\mathrm{aq})=$ hydrobromic acid $\quad \mathrm{H}_{2} \mathrm{~S}(\mathrm{aq})=$ hydrosulfuric acid

## Nonbinary Acid: Hydrogen + Polyatomic lon

| First word |
| :--- |
| If the acid ends in polyatomic ion with sufix ite the ite becomes ous |$\quad+$

## Second word

If the acid ends in polyatomic ion with sufix ite the ite becomes ous
If the acid ends in polyatomic ion with sufix ate the ate becomes ic
$\mathrm{HClO}_{3}$ Chloric acid $\mathrm{HClO}_{2}$ chlorous acid $\mathrm{H}_{2} \mathrm{SO}_{4}$ sulfuric acid $\mathrm{H}_{2} \mathrm{SO}_{3}$ sulfrous acid

## 7B Writing Formulas

## All compounds other than covalent

Cross over the combining capacities from each of the ions so that they become the subscripts of the opposite ions. Reduce the subscripts to lowest terms if necessary. Subscripts of 1 are not written.
Remember:

1. Write positive ion first- write negative ion which maybe a monoatomic or polyatomic ion (more than one element)
2. Polyatomic do not change formula or subscripts- when adding a subscript ion is enclosed in parenthesizes.
3. Determine the charge on each ion

Write formula for hydrosulfuric acid
3. Crisis cross and reduce


Write the formula for tin (IV) sulfate


## Binary Covalent Compounds

The prefix on eaxh word of the name indicates the number of eachelement in the compound
Write the elements in the same order as they are in the name
Write the formula fo diphosphorous pentaoxide $\rightarrow \mathrm{P}_{2} \mathrm{O}_{5}$

## 7C Dot Diagrams

## Rules for Drawing Lewis Dot Diagrams

 of Covalent Compounds:1. Calculate the number of valence electrons for each atom and the total for the molecule or ion. For ions, add or subtract electrons as necessary to obtain the correct charge.
2. Write the skeletal structure; place correct number of electrons around each atom.
3. Replace pairs of electrons with a bond
4. Make certain each atom has a complete octet. Use multiple bonds if necessary to complete octet. Double check total number of electrons.

## Additional Hints:

H is always an end (terminal) atom. It has one electron and thus is only able to bond covalently with one other atom.
Halogens and Oxygen are often terminal atoms.
In binary compounds, the central atom has the lowest subscript and is usually listed first.
Carbon has four bonds in most compounds.

- Barium iodide- ionic:

- Carbon disulfide - covalent: $: \ddot{\mathrm{S}}: \mathrm{C}:: \ddot{\mathrm{S}}:$
- Arsenic triiodide - covalent:

$\mathrm{H}: \ddot{\mathrm{S}} \mathrm{e}: \mathrm{H}$
- Hydrogen selenide covalent:


## 7D Metallic Bonds Explain Properties of Metals

## Metallic Bonding



## Key Concepts

1. A metal is a lattice of positive metal 'ions'
in a 'sea' of delocalized electrons.
2. Metallic bonding refers to the interaction between the delocalized electrons and the metal nuclei.
3. The physical properties of metals are the result of the delocalization of the electrons involved in metallic bonding

## Examples:

1. Melting Points general, metals have high melting and boiling points because of the strength of the metallic bond. The strength of the metallic bond depends on the number of electrons in the delocalized 'sea' of electrons. (More delocalized electrons results in a stronger bond and a higher melting point.)
2. The delocalized electrons are free to move in the solid lattice.

These mobile electrons can act as charge carriers in the conduction of electricity or as energy conductors in the conduction of heat
3. The delocalized electrons are free to move in the solid lattice.

These mobile electrons can act as charge carriers in the conduction of electricity or as energy conductors in the conduction of heat 4. Metals typically have a shiny, metallic luster. Photons of light do not penetrate very far into the surface of a metal and are typically reflected, or bounced off, the metallic surface.
5. Metals are malleable and ductile.

The delocalized electrons in the 'sea' of electrons in the metallic bond, enable the metal atoms to roll over each other when a stress is applied.


## 8A Define \& Use Moles

The mole (mol) is a unit for counting objects. It is especially useful for counting tiny objects like atoms, molecules, ions, and formula-units.
1 mole of particles (atoms, ions or molecules) $=6.02 \times 10^{23}$ particles for any substance!
Other counting Units are:

A ream of paper contains 500 sheets of paper.


A dozen eggs contains 12 eggs.
$\checkmark 6.02 \times 10^{23}$ atoms ( particles) $=$ the atomic mass in grams
$\checkmark 6.02 \times 10^{23}$ atoms (particles) $=1$ mole
$\checkmark 1$ mole $=$ the atomic mass in grams

How many atoms make up 3.5 mol of Ag ?

$$
\frac{3.5 \mathrm{~mol} \mathrm{Ag}}{1} \times \frac{6.02 \times 10^{23} \text { atoms Ag }}{1 \mathrm{~mol} \mathrm{Ag}}=\mathbf{2 . 1 1} \times \mathbf{1 0}^{24} \text { atoms } \mathrm{Ag}
$$

How many molecules are there in 5.5 moles of carbon dioxide?

$$
\frac{5.5 \mathrm{~mol} \mathrm{CO}_{z}}{1} \times \frac{6.02 \times 10^{23} \text { molecules } \mathrm{CO}_{2}}{1 \mathrm{~mol} \mathrm{CO}_{z}}=\mathbf{3 . 3 1} \times \mathbf{1 0}^{\mathbf{2 4}} \mathbf{~ m o l e c u l e s} \mathrm{CO}_{2}
$$

How many moles are $7.0 \times 10^{15}$ atoms of sulfur?

$$
\frac{7.0 \times 10^{15} \text { atoms of } S}{1} \times \frac{1 \text { mole of } S}{6.02 \times 10^{23} \text { atoms of } S}=1.16 \times \mathbf{1 0}^{-8} \mathbf{~ m o l e ~} S
$$

## 8B Calculating Atoms, Ions, Molecules Using Moles



## How to Determine the Molar Mass of a Molecule:

To find the molar mass of a compound:

- Use the chemical formula to determine the number of each type of atom present in the compound.
- Multiply the atomic weight (from the periodic table) of each element by the number of atoms of that element present in the compound.
- Add it all together

Molar Mass of $\mathrm{Zn}\left(\mathrm{NO}_{3}\right)_{2}$ :
In this compound, we have 1 atom of zinc, 2 atoms of nitrogen ( $1 \times 2=2$ ) and 6 atoms of oxygen $(3 \times 2=6)$. The molar mass of zinc nitrate $=$
( 1 atom $\mathrm{Zn} \times 65 \mathrm{~g}$ of zinc) $=65 \mathrm{~g} \mathrm{Zn}$
( 2 atoms $\mathrm{N} \times 14 \mathrm{~g}$ of nitrogen) $=28 \mathrm{~g} \mathrm{~N}$
( 6 atoms $0 \times 16 \mathrm{~g}$ of oxygen) $=96 \mathrm{~g} \mathrm{O}$
$65+28+96=189 \mathrm{~g} \mathrm{Zn}\left(\mathrm{NO}_{3}\right)_{2}$

* $6.02 \times 10^{23}$ atoms ( particles) $=$ the atomic mass in grams
\& $6.02 \times 10^{23}$ atoms (particles) $=1$ mole
\& 1 mole $=$ the atomic mass in grams

Example 1: How many moles are present in 34 grams of $\mathrm{Cu}(\mathrm{OH})_{2}$ ?
Use conversion between moles and grams
1 mole = molar mass
determine molar mass $=(1$ mole $\mathrm{Cu} \times 63.55 \mathrm{~g} \mathrm{Cu})+(2 \mathrm{~mole} \mathrm{O} \times 16.00 \mathrm{~g} \mathrm{O})+(2 \mathrm{~mole} \mathrm{H} \times 1.00 \mathrm{~g} \mathrm{H})=97.55 \mathrm{~g} \mathrm{Cu}(\mathrm{OH})_{2}$

$$
\frac{34 \mathrm{~g} \mathrm{Cu}(\mathrm{OH})_{\mathrm{z}}}{1} \times \frac{1 \mathrm{~mole} \mathrm{Cu}(\mathrm{OH})_{2}}{97.55 \mathrm{~g} \mathrm{Cu(OH})_{z}}=\mathbf{0 . 3 5} \mathbf{~ m o l e ~} \mathrm{Cu}(\mathbf{O H})_{2}
$$

Example 2: How many atoms of aluminum are there in a 7.85 g sample of aluminum? (molar mass of Al is ? 26.982 g

$$
\frac{7.85 \mathrm{~g} \mathrm{Al}}{1} \times \frac{1 \mathrm{~mol} \mathrm{Al}}{16.982 \mathrm{~g} \mathrm{Al}} \times \frac{6.02 \times 10^{23} \mathrm{Al} \text { atoms }}{1 \mathrm{~mol} \mathrm{Al}}=\mathbf{1 . 7 5} \times \mathbf{1 0}^{\mathbf{2 3}} \mathrm{Al} \text { atoms }
$$

Example 3: 16.2 g of magnesium nitride contains how many magnesium ions?
write the formula for magnesium nitride $\rightarrow \mathrm{Mg}_{3} \mathrm{~N}_{2}$

- restate question $16.2 \mathrm{~g} \mathrm{Mg}_{3} \mathrm{~N}_{2}=$ $\qquad$ $\mathrm{Mg}^{+3}$ ions
- Identify conversions necessary to answer question. You can start by writing conversion you know and deciding if they help answer the question.
- 1 mole $\mathrm{Mg}_{3} \mathrm{~N}_{2}=$ molar mass of $\mathrm{g} \mathrm{Mg}_{3} \mathrm{~N}_{2}$
- Calculate molar mass (You should get $100.929 \mathrm{~g} \mathrm{Mg}_{3} \mathrm{~N}_{2}$ )
- How many $\mathrm{Mg}^{+2}$ ions are in 1 mole of $\mathrm{Mg}_{3} \mathrm{~N}_{2}$ ??? 1 mole $\mathrm{Mg}_{3} \mathrm{~N}_{2}=3$ mole of $\mathrm{Mg}^{+2}$
- 1 mole $\mathrm{Mg}^{+2}=6.02 \times 10^{23} \mathrm{~mol} \mathrm{Mg}^{+2}$

$$
\frac{16.2 \mathrm{~g} \mathrm{Mg}_{3} \mathrm{~N}_{z}}{1} \times \frac{1 \mathrm{~mol} \mathrm{Mg} \mathrm{~g}_{3} \mathrm{~N}_{z}}{100.929 \mathrm{~g} \mathrm{Mg}_{3} \mathrm{~N}_{z}} \times \frac{3 \mathrm{~mol} \mathrm{Mg}+2}{1 \mathrm{~mol} \mathrm{Mg}_{3} \mathrm{~N}_{z}} \times \frac{6.02 \times 10^{23} \mathrm{Mg}^{+2} \text { ions }}{1 \mathrm{~mol} \mathrm{Mg}^{+z}}=\mathbf{2 . 9 0} \times \mathbf{1 0}^{23} \mathbf{M g}^{+2} \text { ions }
$$

Example 4: 12.8 of dinitrogen tetraoxide contains howmany molecules of dinitrogen tetroxide?

$$
\frac{12.8 \mathrm{~g} \mathrm{~N}_{z} \mathrm{O}_{4}}{1} \times \frac{1 \mathrm{~mol} \mathrm{~N}_{z} \mathrm{O}_{4}}{92.010 \mathrm{~g} \mathrm{~N}_{z} \mathrm{O}_{4}} \times \frac{6.02 \times 10^{23} \mathrm{molecules}^{\mathrm{N}_{2} \mathrm{O}_{4}}}{1 \mathrm{molN}_{z} \mathrm{O}_{4}}=\mathbf{8 . 3 7} \times \mathbf{1 0}^{23} \text { molecules of } \mathrm{N}_{2} \mathbf{O}_{4}
$$

## 8C Calculating Percent Composition, Empirical Formulas and Molecular Formulas

Percent compostion is the \% be mass of each element in a compound

$$
\text { percent of an element in a compound }=\frac{\text { molar mass of the element }}{\text { molar mass of compound }} \times 100
$$

## Example 1: Calculate the percent of magnesium in $\mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2}$.

## Step 1

$1 \mathrm{Mg}=1 \times 24=24$
$2 \mathrm{~N}=2 \times 14=28$

$$
\begin{aligned}
& \text { Step } 2= \\
& \frac{24 \mathrm{~g} \mathrm{Mg}}{148 \mathrm{~g} \mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2}} \times 100=\mathbf{1 6 . 2} \% \mathbf{M g}
\end{aligned}
$$

$60=6 \times 16=96$
$24+28+96=148 \mathrm{~g} / \mathrm{mole}$

| Molecular Formula | Empirical Formula |
| :---: | :---: |
| $\mathrm{H}_{2} \mathrm{O}$ | $\mathrm{H}_{2} \mathrm{O}$ |
| $\mathrm{CH}_{3} \mathrm{COOH}$ | $\mathrm{CH}_{2} \mathrm{O}$ |
| $\mathrm{CH}_{2} \mathrm{O}$ | $\mathrm{CH}_{2} \mathrm{O}$ |
| $\mathrm{C}_{6} \mathrm{H}_{12} \mathrm{O}_{6}$ | $\mathrm{CH}_{2} \mathrm{O}$ |

Empirical Formula - Simplest formula for a compound; it is the smallest whole-number ratio of the atoms present

Molecular Formula - Expression of the formula for a compound; it shows the actual number of atoms of each element present in one molecule of the compound.

Empirical Formula: Simplest formula for a compound; it is the smallest whole-number ratio of the atoms present.

## Here's a rhyme:

## Percent or data to mass <br> Mass to mole <br> Divide by smallest <br> Multiply 'til whole

Example 2: Calculate the empirical formula for a compound composed of $26.6 \%$ potassium, $35.4 \%$ chromium, and $38.1 \%$ oxygen.
Percent or data to mass: First, assume that you are given a 100-g sample, which would mean that the mass of each element in grams is equal to the percent composition value. If given data in grams you will need to calculate grams of each element using the data.
From grams we find moles of each element in the formula.

## Mass to mole

Find moles potassium.
$\frac{26.6 \mathrm{~g} \mathrm{~K}}{1} \times \frac{1 \mathrm{~mol} \mathrm{~K}}{39.10 \mathrm{~g} \mathrm{~K}}=0.6803 \mathrm{~mol} \mathrm{~K}$
Next find moles of chromium.
$\frac{35.4 \mathrm{~g} \mathrm{Cr}}{1} \times \frac{1 \mathrm{~mol} \mathrm{Cr}}{52.00 \mathrm{~g} \mathrm{Cr}}=0.6808 \mathrm{~mol} \mathrm{Cr}$
Next find moles of oxygen.
$\frac{381 \mathrm{~g} \theta}{1} \times \frac{1 \mathrm{~mol} \mathrm{O}}{16.00 \mathrm{~g} \theta}=2.381 \mathrm{~mol} \mathrm{O}$

Divide by smallest Next, set up a mole ratio that relates the moles of each element to the moles of the element that is least present.

$$
K=\frac{0.6803}{0.6803}=1 \quad \mathrm{Cr}=\frac{0.6808}{0.6803}=1.0007=1 \quad 0=\frac{2.381}{0.6803}=3.4999=3.5
$$

Multiply 'til whole Because a chemical formula must have only whole numbers we multiply by 2 to yield:
$\mathrm{K}=1 \times 2=2 \quad \mathrm{Cr}=1 \times 2=2 \quad \mathrm{O}=3.5 \times 2=7 \quad$ These numbers are the subscripts $\mathrm{K}_{2} \mathrm{Cr}_{2} \mathrm{O}_{7}$ as the empirical formula.

To determine the Molecular Formula you need to be given:

1. Empirical formula
2. Molar mass of compound

Molecular Formula:

1. Calculate empirical formula if not given
2. Divide the molar mass by the empirical formula mass.
3. Multiply all the subscripts in the empirical formula by the number you get when you divide the molar mass by the mass of the empirical formula


Example 3: A 1.000 g sample of red phosphorus powder was burned in air and reacted with oxygen gas to give 2.291 g of a phosphorus oxide. Calculate the empirical formula and molecular formula of the phosphorus oxide given the molar mass is approximately $284 \mathrm{~g} / \mathrm{mol}$.

1. Calculate empirical formula:

Percent of data to mass \% not given use data to determine the mass of P and O :
given that $P=1.00 \mathrm{~g}$ and $P_{x} \mathrm{O}_{x}=2.291 \mathrm{~g}$. You know the mass of $\mathrm{O}=$ mass of $\mathrm{P}_{\mathrm{x}} \mathrm{O}_{\mathrm{x}}-$ mass of P

$$
2.291 \mathrm{~g} \mathrm{P}_{\mathrm{x}} \mathrm{O}_{\mathrm{x}}-1.000 \mathrm{~g} \mathrm{P}=1.291 \mathrm{~g} \mathrm{O}
$$

## Mass to mole:

$\frac{1.00 \mathrm{~g} P}{1} \times \frac{1 \mathrm{~mol} \mathrm{P}}{30.97 \mathrm{~g} \mathrm{P}}=0.032289 \mathrm{~mol} \mathrm{P}$
$\frac{1.291 \mathrm{~g} \theta}{1} \times \frac{1 \mathrm{~mol} \mathrm{O}}{16.00 \mathrm{~g} \theta}=0.0806875 \mathrm{~mol} \mathrm{O}$

## Divide by smallest:

0.032289 mol P
$\frac{0.032289 \mathrm{~mol} \mathrm{P}}{}=1$
$\frac{0.0806875 \mathrm{~mol} \mathrm{O}}{0.0806875 \mathrm{~mol} \mathrm{O}}=2.48989 \approx 2.50 \mathrm{~mol}-$ not close enough to round

## Multiply 'til whole

$2 \times 2.500$ is 5 which is whole so multiply each by 2
$1 \mathrm{P} \times 2=2 \mathrm{P} \quad 2.500 \times 2=50$

## Empirical formula is $\mathrm{P}_{2} \mathrm{O}_{5}$

2. Divide the molar mass by the empirical formula mass:

Empirical mass $(5 \times 30.97 \mathrm{~g} P)+(2 \times 16.00 \mathrm{~g} \mathrm{O})=141.943 \mathrm{~g}$ empirical formula mass
The "empirical formula weight" $=141.943 \mathrm{~g}$

$$
\frac{284}{142}=2
$$

3. The molecular formula is a multiple of 2 times the empirical formula:
$\mathrm{P}=2 \times 2=4$
$\mathrm{O}=2 \times 5=10$
molecular formula is $\mathrm{P}_{4} \mathrm{O}_{10}$


## $2 \mathrm{H}_{2}+\mathrm{O}_{2} \rightarrow 2 \mathrm{H}_{2} \mathrm{O}$

Equations must be balanced because: Law of Conservation of Matter: Atoms can be neither created nor destroyed in an ordinary chemical reaction, so there must be the same number of atoms on both sides of the equation. The mass of all the reactants (the substances going into a reaction) must equal the mass of the products (the substances produced by the reaction).

H H


Two types of numbers are found in a chemical equation:
Subscripts: $\mathrm{H}_{2} \mathrm{O}$
The small numbers to the lower right of chemical symbols. Subscripts represent the number of atoms of each element in the molecule.

## Coefficients: $2 \mathrm{C}_{2} \mathrm{H}_{6}(\mathrm{~g})+7 \mathrm{O}_{2}(\mathrm{~g}) \rightarrow 4 \mathrm{CO}_{2}(\mathrm{~g})+6 \mathrm{H}_{2} \mathrm{O}(\mathrm{g})$

The large numbers in front of chemical formulas. Coefficients represent the number of molecules of the substance in the reaction.

## Steps in Balancing a Chemical Equation An Equation is balanced when there are the same numbers of atoms of each element on both sides of the equation.

Balancing a chemical equation involves a certain amount of trial and error. In general, however, you should follow these steps:

1. Count each type of atom in reactants and products. Does the same number of each atom appear on both sides of the arrow? If not, the equation is not balanced, and you need to go to step 2.
2. Place coefficients, as needed, in front of the symbols or formulas to increase the number of atoms or molecules of the substances. Use the smallest coefficients possible. Warning! Never change the subscripts in chemical formulas. Changing subscripts changes the substances involved in the reaction. Change only the coefficients.
3. Repeat steps 1 and 2 until the equation is balanced.

Example: Balance this chemical equation for the reaction in which nitrogen ( $\mathrm{N}_{2}$ ) and hydrogen $\left(\mathrm{H}_{2}\right)$ combine to form ammonia $\left(\mathrm{NH}_{3}\right): \quad \mathrm{N}_{2}+\mathrm{H}_{2} \rightarrow \mathrm{NH}_{3}$
1: First count the nitrogen atoms on both sides of the arrow. There are two nitrogen atoms in the reactants so there must be two in the products as well.
Place the coefficient 2 in front of $\mathrm{NH}_{3}$ to balance nitrogen: $\mathrm{N}_{2}+\mathrm{H}_{2} \rightarrow \mathbf{2} \mathrm{NH}_{3}$
Now count the hydrogen atoms on both sides of the arrow. There are six hydrogen atoms in the products so there must also be six in the reactants. Place the coefficient 3 in front of $\mathrm{H}_{2}$ to balance hydrogen: $\mathrm{N}_{2}+3 \mathrm{H}_{2} \rightarrow 2 \mathrm{NH}_{3}$.


| $4 \mathrm{NH}_{3}+5 \mathrm{O}_{2} \rightarrow$ | $4 \mathrm{NO}+6 \mathrm{H}_{2} \mathrm{O}$ |
| :---: | :---: |
| 4 N (nitrogen atoms) | 4 N (nitrogen atoms) |
| 12 H (hydrogen atoms) | 12 H (hydrogen atoms) |
| 10 O (oxygen atoms) | 10 O (oxygen atoms) |

Mole Y
Stoichiometry

## BASED UPON MOLE－MOLE RATIOS．

Basic Pattern：
1．If quantity given is grams Convert to moles
2．Perform mole $\leftarrow \rightarrow$ mole conversion
3．Convert to desired units
Stoichiometry Example：If iron pyrite， $\mathrm{FeS}_{2}$ ，is not removed from coal，oxygen from the air will combine with both the iron and the sulfur as coal burns．If a furnace burns an amount of coal containing $100 \mathrm{~g}^{\mathrm{g}} \mathrm{FeS}_{2}$ ，how much $\mathrm{SO}_{2}$（an air pollutant）is produced？The following example will discuss each conversion．The math is done when all the conversions are done
1．Write a balanced equation showing the formation of iron（III）oxide and sulfur dioxide．

$$
4 \mathrm{FeS}_{2}+11 \mathrm{O}_{2} \rightarrow 2 \mathrm{Fe}_{2} \mathrm{O}_{3}+8 \mathrm{SO}_{2}
$$

The equation provides conversion factor to convert from moles of $\mathrm{FeS}_{2}$ to moles of $\mathrm{SO}_{2}$

$$
4 \mathrm{moles}^{\text {of } \mathrm{FeS}_{2}=8 \mathrm{moles} \text { of } \mathrm{SO}_{2} \text { or } 1 \mathrm{~mole} \mathrm{FeS}_{2}=2 \mathrm{~mole} \mathrm{SO}_{2} .42}
$$

2．Write the mass information given in the problem．

$$
\begin{gathered}
\frac{100 \mathrm{~g} \mathrm{FeS}_{2}}{1} \\
\frac{100 \mathrm{~g} \mathrm{FeS}_{2}}{1} \times \frac{1 \mathrm{~mol} \mathrm{FeS}_{2}}{120 \mathrm{~g} \mathrm{FeS}_{2}}
\end{gathered}
$$

3．Convert grams of $\mathrm{FeS}_{2}$ to moles of $\mathrm{FeS}_{2}$ ．

4．Changes moles of $\mathrm{FeS}_{2}$（reactant）to moles of $\mathrm{SO}_{2}$（product）．
This ratio comes from the coefficients in the balanced equation．
Notice that the ratio could be reduced from $8: 4$ to $2: 1$ when placed in the factor－label form．Reducing is not necessary．

$$
\frac{100 \mathrm{~g} \mathrm{FeS}_{2}}{1} \times \frac{1 \mathrm{~mol} \mathrm{FeS}_{2}}{120 \mathrm{~g} \mathrm{FeS}_{2}} \times \frac{8 \mathrm{~mol} \mathrm{SO}_{2}}{4 \mathrm{~mol} \mathrm{FeS}_{2}}
$$

5．Convert moles of $\mathrm{SO}_{2}$ to grams of $\mathrm{SO}_{2}$

$$
\frac{100 \mathrm{~g} \mathrm{FeS}_{z}}{1} \times \frac{1 \mathrm{~mol} \mathrm{FeS}_{z}}{120 \mathrm{~g} \mathrm{FeS}_{z}} \times \frac{8 \mathrm{~mol} \mathrm{SO}_{z}}{4 \mathrm{~mol} \mathrm{FeS}_{z}} \times \frac{64 \mathrm{~g} \mathrm{SO}_{2}}{1 \mathrm{~mol} \mathrm{SO}_{z}}=107 \mathrm{~g} \mathrm{SO}_{2}
$$

LIMITING REAGENT：is the reagent that is totally consumed during the reaction and therefore determines or limits the amount of product formed．
EXCESS REAGENT：is the reactant that is not totally consumed during the reaction and therefore some remains when the reaction has stopped．
THEORETICAL YIELD：is the calculated amount of product that could be produced if no error occurred．Error always occurs so the theoretical yield never equals the measured amount of product produced．

Limiting Problem: A 50.6 g sample of magnesium hydroxide is reacted with 45.0 g of hydrochloric acid according to the reaction:
$\mathrm{Mg}(\mathrm{OH})_{2}+2 \mathrm{HCl} \rightarrow \mathrm{MgCl}_{2}+2 \mathrm{H}_{2} \mathrm{O}$ what is the theoretical yield of $\mathrm{MgCl}_{2}$ ?
Is this a limiting reagent problem? One way to find out is to write down what is known about any component of the reaction below that component:
Notice how quantities of both reactants are known. Which one will be used up first? You can't tell, nor should you jump to any conclusions. Just because it looks like there is less $\mathrm{Mg}(\mathrm{OH})_{2}$ present does not automatically mean it will be used up before all of the HCl is consumed. This is a limiting reagent problem.
To determine which reactant is the limiting reagent:

1. Calculate the amount of product using the amount of each reactant.
2. The reactant that yield the least amount of product is the limiting reagent.
3. The smallest amount of product calculated is the theoretical yield of this product.

$$
\begin{aligned}
& \frac{45 \mathrm{~g} \mathrm{HCl}}{1} \times \frac{1 \mathrm{~mol} \mathrm{HCl}}{36.5 \mathrm{~g} \mathrm{HCl}} \times \frac{1 \mathrm{~mol} \mathrm{MgCl}}{z}{ }_{2} \times \frac{95.3 \mathrm{~g} \mathrm{MgCl}_{2}}{1 \mathrm{~mol} \mathrm{HCl}}=58.7 \mathrm{~g} \mathrm{MgCl}_{2}
\end{aligned}
$$

HCl is limiting reagent because it produced the least amount of product because it was consumed first. The HCl will be run out before the magnesium hydroxide and thereby limit the amount of product formed. For this reason, use the moles of HCl to calculate the theoretical yield of magnesium chloride:

$$
\frac{1.23 \mathrm{~mol} \mathrm{HCl}}{1} \times \frac{1 \mathrm{~mol} \mathrm{Mg}(\mathrm{OH})_{z}}{2 \mathrm{~mol} \mathrm{HCl}} \times \frac{95.3 \mathrm{~g} \mathrm{Mg}(\mathrm{OH})_{2}}{1 \mathrm{~mol} \mathrm{Mg}(\mathrm{OH})_{z}}=\mathbf{5 8 . 6} \mathbf{g ~ M g}(\mathbf{O H})_{2}
$$

## PERCENT YIELD PROBLEMS:

- Theoretical yield - this is what you calculate from stoichiometry - only in a perfect world
- Actual yield - what you get in a lab in the real world (no yield will be 100\%-useful in lab reports)

$$
\text { Percent yield }=\frac{\text { Actual Yield }(\text { measured amount of product produced })}{\text { Theoretical Yield }(\text { calculated possible amount of product produced })} \times 100
$$

## Percent Yield Example:

Consider the combustion of 215.0 g of butane, C 4 H 1 o with plenty of oxygen. If you are able to collect 295.3 g of water in the laboratory, what was your percent yield?

$$
\begin{gathered}
\frac{215.0 \mathrm{~g} \mathrm{C}_{4} \mathrm{H}_{10}}{1} \times \frac{1 \mathrm{~mol} \mathrm{C}_{4} \mathrm{H}_{10}}{59.084 \mathrm{~g} \mathrm{C}_{4} \mathrm{H}_{10}} \times \frac{10 \mathrm{~mol} \mathrm{H}_{z} \mathrm{O}}{2 \mathrm{~mol} \mathrm{C}_{4} \mathrm{H}_{10}} \times \frac{10.015 \mathrm{~g} \mathrm{H}_{2} \mathrm{O}}{1 \mathrm{~mol} \mathrm{H}_{2} \mathrm{O}}=333.4 \mathrm{~g} \mathrm{H}_{2} \mathrm{O} \\
\frac{295.3 \mathrm{~g} \mathrm{H}_{2} \mathrm{O}}{333.4 \mathrm{~g} \mathrm{H}_{2} \mathrm{O}} \times 100=\mathbf{8 8 . 6} \%
\end{gathered}
$$

## 9A Describe Gas Laws and use to Calculate

## Boyle's Law

This law says: The pressure and the volume of a gas are inversely proportional to each other, when temperature and moles of gas are kept constant. In other words, if you keep the temperature and moles of gas constant, and you increase the pressure on the gas, the volume will decrease, and if you lessen the pressure on a gas, the volume will increase.
$\mathrm{P}_{1} \mathrm{~V}_{1}=\mathrm{P}_{2} \mathrm{~V}_{2}$


## Charles' Law

This law says: The temperature and the volume of a gas are directly proportional, assuming pressure and moles of gas are kept constant Another way to say this is: if the temperature goes up the volume will go up when pressure and number of moles are constant. The opposite also happens; if temperature is lowered, volume will be less. TEMPERATURE IN KELVIN ONLY $\frac{\mathrm{V}_{1}}{\mathrm{~T}_{1}}=\frac{\mathrm{V}_{2}}{\mathrm{~T}_{2}}$


## Avogadro's Law

Equal moles of gases at the same temperature and pressure will occupy equal volumes. Volume and moles are directly proportional if temperature and pressure are kept constant. Simply, more moles results in a larger volume and fewer moles results in a smaller volume if T and P are not changing.
STP $=0^{\circ} \mathrm{C}$, and at 1 atm . At STP, one mole of gas, any gas, will occupy exactly 22.4 liters.
$\underline{V_{1}}=\underline{V_{2}}$
$\begin{array}{ll}\mathrm{n}_{1} & \mathrm{n}_{2}\end{array}$
Dalton's Law of Partial Pressures total pressure = sum of all partial pressures

- partial pressure: pressure exerted by one gaseous component in a mixture
-Therefore, the pressure of a mixture of gases is equal to the sum of the partial pressures of the individual gases.
C

V
P
T
M
N
g

|  | 22.4 L | 22.4 L | 22.4 L |
| :--- | :--- | :--- | :--- |
| Volume | 1 atm | 1 atm |  |
| Pressure | 1 atm | 10 | $0^{\circ} \mathrm{C}$ |
| Temperature | $0^{\circ} \mathrm{C}$ | $0^{\circ} \mathrm{C}$ | 16.0 g |
| Mass of gas | 4.00 g | 28.0 g | $6.02 \times 10^{23}$ |
| Number of <br> gas molecules | $6.02 \times 10^{23}$ | $6.02 \times 10^{23}$ |  |



Combine Gas Law ; $\frac{\mathrm{P}_{2} \mathrm{~V}_{1}}{\mathrm{~T}_{1}}=\frac{\mathrm{P}_{2} \mathrm{~V}_{2}}{\mathrm{~T}_{2}}$ or $\frac{\mathrm{P}_{2} \mathrm{~V}_{1}}{\mathrm{~T}_{1} n}=\frac{\mathrm{P}_{2} \mathrm{~V}_{2}}{\mathrm{~T}_{2} n}$ USE WHEN CONDITIONS CHANGE

## Ideal Gas Law

PV =nRT USE WHEN CONDITIONS DO NOT CHANGE. All Units must be the same units that are in R
P - pressure (in atm).
V - volume (in liters). Type equation here.
n - quantity of gas (in moles).
R - the gas constant . $0821 \mathrm{~atm} \mathrm{~L} / \mathrm{mol}$ K.
T - temperature, IN KELVINS ONLY ALWAYS

## Example Problems:

1. A small 2.00 L fire extinguisher has an internal pressure of 506.6 kPa at $25^{\circ} \mathrm{C}$. What volume of methyl bromide, the fire extinguisher's main ingredient, is needed to fill an empty fire extinguisher at standard pressure if the temperature remains constant?
This problem is solved by inserting values into (change in conditions)

$$
\begin{gathered}
\frac{P_{1} V_{1}}{T_{ \pm} n_{I}}=\frac{P_{2} V_{2}}{T_{z} n_{z}} \text { OR P } P_{1} V_{1}=P_{1} V_{1} \operatorname{PLUG} \operatorname{IN}(506.6 \mathrm{Kpa})(2.00 \mathrm{~L})=(101.3 \mathrm{Kpa})\left(\mathrm{V}_{2}\right) \\
V_{2}=\frac{506.6 \mathrm{Kpa})(2.00 \mathrm{~L})}{101.3 \mathrm{Kpa}}=\mathbf{1 0 . 0 L}
\end{gathered}
$$

2. If a gas has a volume of 3.00 L , a pressure of 740.0 Torr and a temperature of 310.0 K , what will the temperature of the gas be in ${ }^{\circ} \mathrm{C}$ if the pressure is increased to 1.50 atm while keeping the volume constant?

$$
\left.\begin{array}{cc}
\mathrm{V}_{1}=3.00 \mathrm{~L} & \mathrm{~V}_{2}=3.00 \mathrm{~L} \text { constant so } \mathrm{V} \text { can be deleted from equation } \\
\mathrm{P}_{1}=740 \mathrm{Torr} \quad \mathrm{P}_{2}=1.50 \mathrm{~atm}=\frac{1.50 \mathrm{~atm}}{1} \times \frac{760 \text { Torr }}{1 \text { atm }}=1.14 \times 10^{3} \mathrm{Torr} \\
T_{1}=310 \mathrm{~K} \quad \text { Change in conditions }
\end{array}\right] \begin{gathered}
\mathrm{T}_{2}=? \mathrm{C}^{\mathrm{o}}
\end{gathered}
$$

3. A sample of 6.9 moles of gas is placed in a container with a volume of 30.4 L at $62^{\circ} \mathrm{C}$. What is the pressure of exerted by the gas?
$\mathrm{PV}=\mathrm{nRT}$ (no change in conditions) All units must match the units in the R you use. 1 am using $.0821 \mathrm{~atm} \mathrm{~L} / \mathrm{mol} \quad \mathrm{K}=\mathrm{C}^{\circ}+273=62^{\circ} \mathrm{C}+273=335 \mathrm{~K}$

$$
\mathrm{PV}=\mathrm{nRT} \quad \mathrm{P}=\frac{\mathrm{nRT}}{\mathrm{~V}} \text { plug in } \mathrm{P}=\frac{(6.9 \mathrm{~mol})\left(.0821 \frac{\mathrm{~atm} \mathrm{t}}{\mathrm{~mol} \mathrm{~K}}\right)(335 \mathrm{~K})}{30.4 \mathrm{t}}=6.2 \mathbf{~ a t m}
$$

4. Three gases $A, B$, and $C$ are contained in a tank. What is the pressure in atmospheres in the tank if the $P_{A}$ is 3.5 atm., the $P_{B}$ is . 724 atm, and the $P_{C}$ is .450 atm? $\mathrm{P}_{\mathrm{T}}=\mathrm{P}_{1}+\mathrm{P}_{2}+\mathrm{P}_{3}+\ldots=\mathrm{P}_{\mathrm{T}}=3.5 \mathrm{~atm}+.724 \mathrm{~atm}+.450 \mathrm{~atm}=4.67 \mathrm{~atm}$


Nitrogen monoxide and oxygen gas combine to form the brown gas nitrogen dioxide, which contributes to photochemical smog. What volume (in L) of nitrogen dioxide gas is produced when 34 L of oxygen gas react with an excess of nitrogen monoxide? Assume conditions are at STP.

$$
\begin{aligned}
& 2 \mathrm{NO}+\mathrm{O}_{2} \rightarrow 2 \mathrm{NO}_{2} \\
& 34 \mathrm{~L} \mathrm{O}_{2}= \\
& \mathrm{L} \mathrm{NO}_{2} \\
& \text { What do we know? } \\
& 1 \mathrm{~mol}=22.4 \mathrm{~L} \\
& 1 \mathrm{~mol} \mathrm{O}_{2}=2 \mathrm{~mol} \mathrm{NO}_{2} \\
& \frac{34 L O_{z}}{1} \times \frac{1 \mathrm{~mol} \theta_{z}}{22.4 L \theta_{z}} \times \frac{2 \mathrm{~mol} \mathrm{~N}_{z}}{1 \mathrm{~mol} \mathrm{O}_{z}} \times \frac{22.4 \mathrm{~L} \mathrm{NO}_{2}}{1 \mathrm{~mol} \mathrm{~N}}=68 \mathrm{~L} \mathrm{NO}
\end{aligned}
$$

## 9C Kinetic Molecular Theory

THE KINETIC MOLECULAR THEORY is used to explain the behavior of gases and is based upon the following postulates:

1. Gases are composed of a many particles that behave like hard spherical objects in a state of constant, random motion.
2. These particles move in a straight line until they collide with another particle or the walls of the container.
3. These particles are much smaller than the distance between particles, therefore the volume of a gas is mostly empty space and the volume of the gas molecule themselves is negligible.
4. There is no force of attraction between gas particles or between the particles and the walls of the container.
5. Collisions between gas particles or collisions with the walls of the container are elastic. That is, none of the energy of the gas particle is lost in a collision.
6. The average kinetic energy of a collection of gas particles is dependent only upon the temperature of the gas.


## In short, the key parts are:

1. constant random motion
2. straight line motion
3. negligible volume
4. no forces of attraction
5. elastic collisions
6. KE $\alpha \mathrm{T}$



## 10C Molarity

The definition of molarity ( M ) is $M=\frac{\text { mole of solute }}{\text { Lof solution }}$

## Example:

Calculate the molarity of a solution containing 8.61 g potassium hydroxide, dissolved in enough water to form 750 mL of solution. You to use M formula 5.61 g of KOH will need to be converted to mol and 750 mL will need to be converted to L .

$$
\begin{gathered}
\frac{86.1 \mathrm{~g} \mathrm{KOH}}{1} \times \frac{1 \mathrm{~mol} \mathrm{KOH}}{56.105 \mathrm{~g} \mathrm{KOH}}=1.53 \mathrm{~mol} \mathrm{KOH} \\
\frac{750 \mathrm{ml} \text { of solution }}{1} \times \frac{1 \mathrm{~L}}{1000 \mathrm{~mL}}=.750 \mathrm{~L} \text { of solution } \\
M=\frac{\text { mole of solute }}{L \text { of solution }}=\frac{1.53 \mathrm{~mol} \mathrm{KOH}}{.750 \mathrm{~L} \mathrm{solution}} \\
=\mathbf{0 . 0 1 4 ~ M \text { or } \frac { \text { mol } } { \boldsymbol { L } }}
\end{gathered}
$$

10D Dilutions

## $M_{1} V_{1}=M_{2} V_{2}$

Concentrated $\mathrm{H}_{2} \mathrm{SO}_{4}$ is 18.0 M . What volume is needed to make 2.00 L of 1.00 M solution?

$$
M_{1} V_{1}=M_{2} V_{2}
$$

$$
18.0 \mathrm{M} \times \mathrm{V}_{1}=1.00 \mathrm{M} \times 2.00 \mathrm{~L}
$$

$$
\mathrm{V}_{1}=\frac{1.00 \mathrm{M} \times 2.00 \mathrm{~L}}{18.0 \mathrm{M}}=. \mathbf{1 1 1} \mathrm{L} \text { or } 111 \mathrm{~mL}
$$

10E Types of Soltions
Solubility is defined as the maximum amount of solute dissolved by a given amount of solvent at a specified temperature. The solubility of the given substances in a given solute is temperature-dependent.
Unsaturated: If the solution contains less quantity of solute than what can be dissolved, at a specified temperature it is unsaturated. When more solute is added into solution the solute dissolves.
An unsaturated solution is one in which more of the solute could dissolve at the same temperature.
Saturated: A saturated solution is one in which no more of the solute will dissolve at a specific temperature. Solution containing the maximum amount of solute at a specified temperature is saturated. When more solute is added into the solution the solute will no longer dissolve.
Supersaturated: A supersaturated solution is when a solution which contains more solute than would normally dissolve at a certain temperature. The solution becomes supersaturated by elevating the temperature and the solution contains the maximum amount of solute at an elevated temperature is carefully cooled to a lower temperature and is now supersaturated. When more solute is added into the solution, crystals will form.
10F Solubility is the maximum amount of a substance that can be dissolved in a given amount of solvent. Solubility depends on temperature and pressure, as well as on the type of solvent being used. Solubility is not affected by an increase in the surface area of the solute or by agitation of the solution although the rate of solution formation (dissolution) is affected.


## Factors Affecting Solubility:

Temperature affects solubility of solids differently than the solubility of gases.

Solid solubility: $\uparrow$ temperature often (not always) $\uparrow$ solubility
Solid pressure has no effect on solubility

Gas solubility: $\uparrow$ temperature $\downarrow$ solubility
Gas solubility: $\uparrow$ pressure $\uparrow$ solubility


## Factors Affecting Rate of Solution Formation (dissolution) of Solids:

$\uparrow$ temperature $\uparrow$ rate of dissolution
$\uparrow$ agitation (stirring or shaking) $\uparrow$ rate of dissolution
$\uparrow$ surface area (grinding or crushing a solid) $\uparrow$ rate of dissolution

## 10G Acid Base Theories

| ARRHENIUS DEFINITION | BRØNSTED-LOWRY DEFINITION |
| :---: | :---: |
| An acid is a substance that contains a hydrogen and ionizes to produce hydrogen ions in aqueous solutions. <br> Acid-produces a proton $\left(\mathrm{H}^{+}\right)$in water $\mathrm{HCl} \xrightarrow{\mathrm{H}_{2} \mathrm{O}} \mathrm{H}^{+}+\mathrm{F}^{-}$ | $\mathrm{Br} \varnothing$ nsted-Lowry acid is defined as a hydrogen-ion donor Acid-donates ( gives up)a proton $\left(\mathrm{H}^{+}\right) \mathrm{HF}+\mathrm{H}_{2} \mathrm{O} \leftrightarrow \mathrm{H}_{3} \mathrm{O}++\mathrm{F}^{-}$ |
| A base is a substance that contains a hydroxide and dissociates to produce hydroxide ions in water ( $\mathrm{OH}^{-}$) <br> Base-produces a hydroxide ion in water $\left(\mathrm{OH}^{-}\right)$ $\mathrm{NaOH} \xrightarrow{\mathrm{H}_{2} \mathrm{O}} \mathrm{Na}^{+}+\mathrm{Cl}^{-}$ | $\mathrm{Br} \varnothing$ nsted-Lowry base is defined as a hydrogen-ion acceptor. Base-accepts (bonds to $\mathrm{H}^{+}$) a proton ( $\mathrm{H}^{+}$) $\mathrm{NH}_{3}+\mathrm{H}_{2} \mathrm{O} \leftrightarrow \mathrm{NH}_{4}^{+}+\mathrm{OH}^{-}$ |
| require acid-base reactions to occur in aqueous solutions |  |
| limited to substances with those "parts"; ammonia is a MAJOR exception! | Brønsted-Lowry definition includes all Arrhenius bases that produce hydroxide ions that <br> can accept a proton. But the Brønsted-Lowry definition also covers other bases, such as ammonia, $\mathrm{NH}_{3}$, that do not contain hydroxide ions. |

## 10H Recognizing Neutralization, Precipitation and Oxidation and Reduction Reactions

Acid-base reactions are reactions that occur between an acid and a base in which a proton is transferred. The product includes a salt and, often, water.
3.Precipitation reactions are reactions that occur when two aqueous solutions react and produce a solid precipitate
An oxidation-reduction (redox) reaction, also called a redox reaction, is any reaction that involves the exchange of an electron or oxygen. In general, when the oxidation number of a certain atom is different in one of the reactants than it is in one of the products, it means that at least one electron(s) were exchanged.

Reactant an acid (formula starts with an H ) and a base (formula end in OH or is $\mathrm{NH}_{3}$ ). If the base ends in OH the products are a salt (first element a metal) and water. Salt has the + ion of the metal and - ion of the base. If the base is $\mathrm{NH}_{3}$ product will only be a salt that has a formula that starts with $\mathrm{NH}_{4}$ and the last ion in the formula will be the anion in the acid.
Use solubility chart: If one product is insoluble the reaction is a precipitation reaction.

If the reaction if not an acid base or precipitation reaction it will most likely be a redox reaction. Assign oxidation numbers to all the elements in the equation if an elements oxidation number changes the reaction is a redox reaction.

## 10H

Oxidation Number Rules

1. The oxidation number for an atom in its elemental form is always zero. The oxidation number of a monoatomic ion $=$ charge of the monatomic ion.
2. The oxidation number of all Group 1 A metals $=+1$ (unless elemental) and The oxidation number of all Group 2A metals $=+2$ (unless elemental).
3. In ionic binary compounds the oxidation number is the same as the oxidation number used to write the formula of the compound
4. Hydrogen $(\mathrm{H})$ has two possible oxidation numbers:
+1 when bonded to a nonmeta
1 when bonded to a metal
5. Oxygen ( $O$ ) oxidation numbers is -2 :
6. The oxidation number of fluorine ( F ) is always -1
7. The sum of the oxidation numbers of all atoms (or ions) in a neutral compound $=0$.
. The sum of the oxidation numbers of all atoms in a polyatomic ion = charge on the polyatomic ion.

| $\mathrm{K}_{\mathrm{w}}=\left[\mathrm{H}_{3} \mathrm{O}^{+}\right]\left[\mathrm{OH}^{-}\right]$ | $\mathrm{K}_{\mathrm{w}}=1 \times 10^{-14}$ at $25^{\circ} \mathrm{C}$ |
| :--- | :--- |
| $\mathrm{pH}=-\log \left[\mathrm{H}_{3} \mathrm{O}^{+}\right]$ |  |
| $\mathrm{pOH}=-\log \left[\mathrm{OH}^{-}\right]$ |  |
| $1 \times 10^{-14}=\left[\mathrm{H}_{3} \mathrm{O}^{+}\right]\left[\mathrm{OH}^{-}\right]$ |  |
| $14=\mathrm{pH}+\mathrm{pOH}$ |  |
| $\left[\mathrm{H}_{3} \mathrm{O}^{+}\right]=10^{-(\mathrm{pH})}$ |  |
| $\left[\mathrm{OH}^{-}\right]=10^{-(\mathrm{pOH})}$ |  |

Example 1: calculate the pH of a solution with a hydrogen ion concentration of $6.3 \times 10^{-4} \mathrm{M}$

$$
\mathrm{pH}=-\log \left(6.3 \times 10^{-4}\right)
$$

$$
=-(-3.2)=3.2
$$

Example 2 if a solution has a hydroxide ion concentration of $1.3 \times 10^{-10}$, then you can calculate the pOH and find the pH .

$$
\begin{aligned}
& \mathrm{pOH}=-\log \left(1.3 \times 10^{-10}\right)=9.9 \\
& \text { Therefore, } \mathrm{pH}=14-9.9=4.1
\end{aligned}
$$

10J The strength of acids and bases depends on the degree to which they dissociate. The greater the degree of dissociation, the greater their strength.
In general, strong acids and strong bases dissociate, or ionize, completely in aqueous solution. In other words, almost 100 percent of a strong acid or strong base interacts with water and ionizes. A strong dissociation is described using one arrow pointing to the right, as shown.


## Strong Acid $\rightarrow$ All of HCL becomes $\mathrm{H}_{3} \mathrm{O}^{+}$and $\mathrm{Cl}^{-}$

Weak Acid $\rightarrow$ very small amount of $\mathrm{HC}_{2} \mathrm{H}_{3} \mathrm{O}_{2}$ becomes $\mathrm{H}_{3} \mathrm{O}^{+}$and $\mathrm{C}_{2} \mathrm{H}_{3} \mathrm{O}_{2}^{-}$

## 11A Concept of Energy

1. The Usual Definition of Energy: the ability to do work and to transfer heat.
work is moving an object against an opposing force
work $=$ distance $\times$ opposing force $w=d \times f$
SI unit of work or energy: the joule (J)
2. Two Basic Forms of Energy: potential energy: energy of position examples: boulder on a ledge, chemical bonds kinetic energy: energy of motion
examples: pool balls, molecules
Forms of Energy:
Energy is found in different forms including light, heat, chemical, and motion. There are many forms of energy, but they can all be put into two categories: potential and kinetic.

## 11B Conservation of Energy

Whether a reaction absorbs energy or releases energy, there is no overall change in the amount of energy. Energy cannot be created or destroyed. This is the law of conservation of energy. Energy can change form - for example, from electricity to light - but the same amount of energy always remains.
If energy cannot be destroyed, what happens to the energy that is absorbed in an endothermic reaction? The energy is stored in the chemical bonds of the products. This form of energy is called chemical energy. In an endothermic reaction, the products have more stored chemical energy than the reactants. In an exothermic reaction, the opposite is true. The products have less stored chemical energy than the reactants. The excess energy of the reactants is released to the surroundings when the reaction occurs. The following graphs compare the energy changes in endothermic and exothermic reactions.


11C Calculate Enthalpy of Reaction $\Delta \mathrm{H}_{\mathrm{rxn}}$ from Bond Energies If not given you will need to write the Lewis structures to use bond energy data to determine $\Delta \mathrm{H}_{\mathrm{rxn}}$ The bond enthalpy is the energy needed to break one mole of bonds in gaseous molecules under standard conditions ( 298 K and 1 atm ).
The bond enthalpy is the energy needed to break one mole of bonds in gaseous molecules under standard conditions ( 298 K and 1 atm )

| Heat Flow | Sign of $\boldsymbol{q}$ | Type of Reaction | Products | Surroundings |
| :--- | :---: | :---: | :---: | :---: |
| from surroundings to system | + (positive) | endothermic | gain heat | lose heat (get colder) |
| from system to surroundings | - (negative) | exothermic | lose heat | gain heat (get hotter) |

## Example

The complete combustion of propane can be represented by the following equation:
$\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{3}(\mathrm{~g})+5 \mathrm{O}_{2}(\mathrm{~g}) \rightarrow 3 \mathrm{CO}_{2}(\mathrm{~g})+4 \mathrm{H}_{2} \mathrm{O}(\mathrm{g})$
or we should redraw the equation for the reaction to represent the bonds present

$$
\mathrm{H}-\underset{\mathrm{C}}{\mathrm{C}-\stackrel{\mathrm{C}}{\mathrm{C}} \mathrm{C}-\stackrel{\mathrm{C}}{\mathrm{C}} \mathrm{C}-\mathrm{H}+5 \mathrm{O}=\mathrm{O} \longrightarrow 3 \mathrm{O}=\mathrm{C}=\mathrm{O}+4 \mathrm{H}-\mathrm{O}-\mathrm{H}}
$$

Total endothermic change for bond breaking:
$8 \times \mathrm{C}-\mathrm{H} \quad 2 \times \mathrm{C}-\mathrm{C} \quad 5 \times \mathrm{O}=\mathrm{O}$
$(8 x+413)+2(+347)+5(+498)$
Total exothermic change for bond forming:
$6 \times \mathrm{C}=\mathrm{O} \quad 8 \times \mathrm{O}-\mathrm{H}$
(6x-805) + (8x-464)
$\Delta H=[(8 \times 413)+(2 \times 347)+(5 \times 498)]-[(6 x-805)+(8 x-464)]=-2054 \mathrm{~kJ} / \mathrm{mol}$
$-\Delta \mathrm{H}$ so the reaction is exothermic releasing energy to the surrounding's

## Calculate Enthalpy of Reaction $\Delta \mathrm{H}_{\mathrm{rxn}}$ from Standard Enthalpies of Formation

- Molar Heat of Formation $\left(\Delta \mathrm{H}_{\mathrm{f}}\right)$ : is the amount of heat required / given off to make 1 mole of a compound from its elemental components. Enthalpy of Formation (also called Heat of Formation)
- Molar Heat of Formation of ALL ELEMENTS is 0 kJ .
- The state of the compound affects the magnitude of $\Delta \mathrm{H}_{\mathrm{f}}$. $\left(\mathrm{H}_{2} \mathrm{O}(\mathrm{g})\right.$ has $\Delta \mathrm{H}_{\mathrm{f}}=-241.8 \mathrm{~kJ} / \mathrm{mol} ; \mathrm{H}_{2} \mathrm{O}$ (l) has $\left.\Delta \mathrm{H}_{\mathrm{f}}=-285.8 \mathrm{~kJ} / \mathrm{mol}\right)$
- Standard Molar Heat of Formation $\left(\Delta \mathrm{H}_{\mathrm{f}}^{\circ}\right)$ is the heat of formation under standard conditions ( 1 atm and $25^{\circ} \mathrm{C}$ ).
- Enthalpies of formation for many chemical substances can be found in tables.

$$
\Delta \mathrm{H}=\sum n \Delta \mathrm{H}_{f_{\text {product }}}^{0}-\sum n \Delta \mathrm{H}_{f_{\text {reactant }}^{0}}^{0}
$$

Example: Calculate the enthalpy of reaction for decomposition of hydrogen peroxide into oxygen and water and determine if the reaction is exothermic or endothermic.
$2 \mathrm{H}_{2} \mathrm{O}_{2}(\mathrm{I}) \rightarrow \mathrm{O}_{2}(\mathrm{~g})+2 \mathrm{H}_{2} \mathrm{O}(\mathrm{g})$
$\Delta \mathrm{H}_{\mathrm{f}}$ of reactants:
$\Delta \mathrm{H}_{\mathrm{f}} \mathrm{H}_{2} \mathrm{O}_{2}(\mathrm{I})=-187.8 \mathrm{~kJ} / \mathrm{mol}$
$\Delta \mathrm{H}_{\mathrm{f}}$ of products:
$\Delta \mathrm{H}_{\mathrm{f}} \mathrm{O}_{2}(\mathrm{~g})=0.0 \mathrm{~kJ} / \mathrm{mol}$
$\Delta \mathrm{H}_{\mathrm{f}} \mathrm{H}_{2} \mathrm{O}(\mathrm{g})=-285.8 \mathrm{~kJ} / \mathrm{mol}$
Using the balanced equation:
$\Delta \mathrm{H}=[0.0+2 \mathrm{~mol}(-285.8 \mathrm{~kJ} / \mathrm{mol})]-[2 \mathrm{~mol}(-187.8 \mathrm{~kJ} / \mathrm{mol})]=-196 \mathrm{~kJ} / \mathrm{mol} \quad$ Because $\Delta \mathrm{H}=-196 \mathrm{~kJ} / \mathrm{mol}$ is negative, the reaction is exothermic

## 11D Calorimetry

## Units of Energy:

calorie--amount of heat needed to raise the temp. of 1.00 gram of water $1.00^{\circ} \mathrm{C}$
kilocalorie-1cal $==1000 \mathrm{Kcal}$ the food calorie with a capital C.
joule--SI unit of energy; $1 \mathrm{cal}=4.184 \mathrm{~J}$
Specific heat,(s) also known as specific heat capacity (Cp) (sometimes referred to as just heat capacity), is defined as the amount of heat necessary to raise the temperature of 1 g of a substance by one degree. joules per gram per degree, $\mathrm{J} / \mathrm{g}^{\circ} \mathrm{C}$ or $\mathrm{J} / \mathrm{g} \mathrm{K}$. The change in temperature $\Delta \mathrm{t}=\mathrm{t}_{\text {final }}-\mathrm{t}_{\text {tinitia }}$ is the same whether you use ${ }^{\circ} \mathrm{C}$ or K .

## $\mathrm{Q}=(\mathrm{mass})(\mathrm{Cp})(\Delta \mathrm{t})$

Specific heat of water (liquid state) $=4.184 \mathrm{~J} / \mathrm{g}^{\circ} \mathrm{C}$ ( or $1.00 \mathrm{cal} / \mathrm{g}^{\circ} \mathrm{C}$ )
Heat(enthalpy) of fusion is the amount of heat required to change a specified amount of a solid to a liquid or released to change a given amount of a liquid to a solid at the melting or freezing point of the substance.
Heat(enthalpy) of vaporization is the amount of heat to change a given amount of substance form an liquid to a gas or released to change a given amount of a gas to a liquid at the substance boiling point.
Calorimetry: is a method of measuring heat flow between systems and surroundings.
The device used to measure heat flow between systems and surroundings is called a calorimeter
$\mathrm{Q}=(\mathrm{mass})(\mathrm{Cp})(\Delta \mathrm{t})$ change in temperature
Example 1: How much heat is required to raise 80.0 g of aluminum from $12^{\circ} \mathrm{C}$ to $46^{\circ}$ ? The specific heat capacity of aluminum is $0.90 \mathrm{~kJ} / \mathrm{kg}^{\circ} \mathrm{C}$. $Q$ is heat so we are solving for $Q$. we can just plug into: $Q=(\mathrm{m})(\mathrm{C})(\Delta \mathrm{t})$, If we use $0.90 \mathrm{~kJ} / \mathrm{kg}{ }^{\circ} \mathrm{C}$ we have to use kg
$(80 \mathrm{~g}) / 1 \times(1 \mathrm{~kg}) / 1000 \mathrm{~g}=0.080 \mathrm{~kg}$.
$\mathrm{Q}=(0.080 \mathrm{~kg})\left(0.90 \mathrm{~kJ} / \mathrm{kg}^{\circ} \mathrm{C}\right)\left(46^{\circ} \mathrm{C}-12^{\circ} \mathrm{C}\right)=\mathbf{2 . 4 8 8} \mathrm{kJ}$


## Example 2: How much water at $25^{\circ} \mathrm{C}$ must be added to 15 kg of water at $10^{\circ} \mathrm{C}$ so that the mixture becomes $20^{\circ} \mathrm{C}$ ? $\mathrm{Qp}=(\mathrm{m})(\mathrm{C})(\Delta \mathrm{t})$

Q hotter water = Q of cooler water LCE: heat will be transferred from the hotter water to cooler water until they are both at the same temperature.
Absolute change in temperature for the hotter water is $25^{\circ} \mathrm{C}-20^{\circ} \mathrm{C}=5^{\circ} \mathrm{C}$
Absolute change in temperature for the cooler water is $=20^{\circ} \mathrm{C}-10^{\circ} \mathrm{C}=10^{\circ} \mathrm{C}$
Q hotter = Q cooler
$(\mathrm{m})(\mathrm{cp})(\Delta \mathrm{t})=(\mathrm{m})(\mathrm{C})(\Delta \mathrm{t})$
(m) $\left(4.184 \mathrm{~J} / \mathrm{g}^{\circ} \mathrm{C}\right)\left(5^{\circ} \mathrm{C}\right)=(15 \mathrm{~kg})\left(4.184 \mathrm{~J} / \mathrm{g}^{\circ} \mathrm{C}\right)\left(10^{\circ} \mathrm{C}\right)$

$$
m=\frac{(15 \mathrm{~kg})\left(4.184 / / g^{\circ} \mathrm{C}\right)\left(10^{\circ} \mathrm{C}\right)}{\left(4.184 / / g^{\circ} \mathrm{C}\right)\left(5^{\circ} \mathrm{C}\right)}
$$



Example 3: We are going to determine the specific heat of a metal using experimental data. A piece of hot metal is placed in cooler water. The heat from the hotter metal will move to the cooler water until both the metal and the water have the final temperature. In this experiment, we used a "coffee cup" calorimeter and gathered the following data:

| Mass of empty cup | 2.31 g |
| :--- | :--- |
| Mass of cup + water | 80.89 g |
| Mass of cup + water + metal | 780.89 g |
| Initial temperature of water | $17.0^{\circ} \mathrm{C}$ |
| Initial temperature of metal | $52.0^{\circ} \mathrm{C}$ |
| Final temperature of system | $27.0^{\circ} \mathrm{C}$ |

The key thermochemistry equation for solving this problem is:
Q metal $=\mathrm{Q}$ water
(mass) ( $\Delta \mathrm{t})(\mathrm{C})=($ mass $)(\Delta \mathrm{t})(\mathrm{C})$
Then, by substitution, we have (metal values on the left, water values on the right):
We need to use the data table to get what we need to substitute into the above equation.
mass of water: 180.98 g cup + water -2.31 g cup $=178.58 \mathrm{~g}$ water
mass of metal: 780.89 g cup + water + metal -180.89 g cup + water $=600.0 \mathrm{~g}$ metal
change in water temperature: $27.0^{\circ} \mathrm{C}-17.0^{\circ} \mathrm{C}=10.0^{\circ} \mathrm{C}$
change in metal temperature: $52.0^{\circ} \mathrm{C}-17.0^{\circ} \mathrm{C}=25.0^{\circ} \mathrm{C}$
Putting the numbers in place gives us:
$(600.0 \mathrm{~g})\left(25.0^{\circ} \mathrm{C}\right)(\mathrm{C})=(178.58 \mathrm{~g})\left(10.0^{\circ} \mathrm{C}\right)\left(4.184 \mathrm{~J} / \mathrm{g}^{\circ} \mathrm{C}\right)$
$C=\frac{(178.58 \mathrm{~g})\left(10.0{ }^{\circ} \mathrm{C}\right)\left(4.184 \mathrm{~J} / \mathrm{g}^{\circ} \mathrm{C}\right)}{(600.0 \mathrm{~g})\left(25.0^{\circ} \mathrm{C}\right)(\mathrm{C})}=\mathbf{0 . 4 9 8} \frac{\mathbf{J}}{\mathbf{g}^{\circ} \mathrm{C}}$


Heat Added (each division $=4 \mathrm{~kJ}$ )

11E $\Delta H_{\mathrm{rxn}} \quad \Delta H_{\mathrm{rxn}}$ values are derived from experimental data. Experiments that measure heat exchange are referred to collectively as calorimetry experiments, and the measurement device is called a calorimeter. There are two important types of calorimetry experiments: constant-pressure calorimetry (also known as coffee cup calorimetry) and constant-volume calorimetry (also known as bomb calorimetry).

## Measuring Heat of Reaction Experimentally

i. A known quantity of reactant is placed in a well-insulated vessel (a Styrofoam cup)
ii. The initial temperature of this reactant is recorded, $\mathrm{t}_{\mathrm{i}}$.
iii. A known quantity of the second reactant is added, the vessel is sealed with a lid and the reaction mixture stirred.
iv. The final temperature of the reaction mixture is recorded, $\mathrm{t}_{\mathrm{f}}$.
$v$. The the heat released or absorbed, in joules, for the reaction is calculated:

## $\mathrm{Q}=($ mass $)(\Delta \mathrm{t})(\mathrm{C}) \Delta \mathrm{t}=\mathrm{t}_{\mathrm{f}}-\mathrm{t}_{\mathrm{i}}$

vi. The enthalpy change in kJ per mole of a given reactant for the reaction is calculated:

$$
\Delta \mathrm{H}=\frac{\mathrm{kJ}}{\mathrm{~mole}}
$$

## Common assumptions for reaction mixtures made up of aqueous solutions:

- density of aqueous solution assumed to be the same as for water, $1 \mathrm{~g} / \mathrm{mL}, 100 \mathrm{~mL}$ of solution is said to have a mass of 100 g
- additivity of volumes of reactants is assumed, 100 mL of reactant $a+200 \mathrm{~mL}$ of reactant $b=300 \mathrm{~mL}$ of reaction mixture
- specific heat capacity of the reaction mixture assumed to be the same as water, specific heat capacity $=4.184 \mathrm{~J} / \mathrm{g}^{\circ} \mathrm{C}$

Heat is not lost to, or absorbed by, the surroundings.

## Example Heat (Enthalpy) of Solution

Ammonium chloride is very soluble in water. When $4.50 \mathrm{~g} \mathrm{NH}_{4} \mathrm{Cl}$ is dissolved in 53.00 g of water, the temperature of the solution decreases from $20.40{ }^{\circ} \mathrm{C}$ to $15.20^{\circ} \mathrm{C}$. Calculate the enthalpy of dissolution of $\mathrm{NH}_{4} \mathrm{Cl}$ (in kJ/mol). (or enthalpy or heat of solution) You are asked to calculate the enthalpy change for the dissolution of ammonium chloride. You are given the masses of the solid and water, and the temperature change that occurs when the two are combined. First calculate the energy change for the surroundings ( Q solution) in the coffee cup calorimeter.
$Q$ solution $=m($ solution $) \times C$ solution $\times \Delta T$
$\mathrm{C}_{\mathrm{p}}$ solution $=4.18 \mathrm{~J} / \mathrm{g} \cdot{ }^{\circ} \mathrm{C}$ ( always assume same as water for coffee cup calorimetry)
m solution $=4.50 \mathrm{~g}+53.00 \mathrm{~g}=57.50 \mathrm{~g}$
$\Delta \mathrm{T}=\mathrm{T}$ final -T initial $=15.20^{\circ} \mathrm{C}-20.40^{\circ} \mathrm{C}=-5.20^{\circ} \mathrm{C}$
$Q$ solution $=m$ (solution) $\times C$ solution $\times \Delta T$
$\mathrm{Q}=(57.50 \mathrm{~g})\left(4.18 \mathrm{~J} / \mathrm{g} \times{ }^{\circ} \mathrm{C}\right)\left(-5.20{ }^{\circ} \mathrm{C}\right)=\mathbf{- 1 2 5 0} \mathrm{J}$ The surroundings lost energy is negative so the system gained energy and the reaction is endothermic and $\Delta H$ is positive
Q dissolution = 1250 J
$\triangle H$ has the units $\mathrm{KJ} /$ mole so we will convert 1250 J to kJ and $4.50 \mathrm{~g} \mathrm{NH}_{4} \mathrm{Cl}$ to moles to determine $\Delta H$
$\frac{1250 \mathrm{f}}{1} \times \frac{1 \mathrm{~kJ}}{1000 \mathrm{f}}=1.250 \mathrm{~kJ} \quad \frac{4.50 \mathrm{~g} \mathrm{NH}_{4} \mathrm{Cl}}{1} \times \frac{1 \mathrm{~mole} \mathrm{NH}_{4} \mathrm{Cl}}{53.492 \mathrm{~g} \mathrm{NH}_{4} \mathrm{Cl}}=\mathbf{0 . 0 8 4 1} \mathrm{mole} \mathrm{NH}_{4} \mathbf{C l}$
$\triangle \mathrm{H}=\frac{1.250 \mathrm{~kJ}}{0.0841 \mathrm{~mol} \mathrm{NH}_{4} \mathrm{Cl}}=14.86 \mathrm{~kJ} / \mathrm{mol}$


## Examples of Particles

| beta particle (electron) | ${ }_{-1}^{0} \mathrm{e}$ |
| :--- | :---: |
| positron | ${ }_{+1}^{0} \mathrm{e}$ |
| alpha particle | ${ }_{2}^{4} \mathrm{He}$ |
| proton | ${ }_{1}^{1} \mathrm{H}$ |
| neutron | ${ }_{0}^{1} n$ |
| gamma ray | ${ }_{0}^{0} \gamma$ |

## 12B Nuclear Reactions

Writing nuclear reactions requires that the atomic number, $Z$, and the mass number, $A$, add up to the same value on each side. In earlier chapters, $Z$ stood for the number of protons. The mass number, $A$, is the sum of protons and neutrons. The mass of a proton or a neutron is about 1 amu .
The alpha decay of radon-198 ${ }_{86}^{198} \mathrm{Rn} \rightarrow{ }_{2}^{4} \mathrm{He}+{ }_{84}^{194} \mathrm{Po}$ The beta decay of uranium -237 ${ }_{92}^{237} \mathrm{U} \rightarrow{ }_{-1}^{0} e+{ }_{93}^{237} \mathrm{~Np}$
Nuclear Transformation = converting one element into another by force.

* bombarding with alpha particles:
$14 / 7 \mathrm{~N}+4 / 2 \mathrm{He}$
1/1 1 H7/80 +
* bombarding with neutrons:
238/92U + 1/1n $\quad \square 239 / 92 \mathrm{U}$

One way to identify the missing component of a nuclear reaction is to use the requirement that the total atomic number, Z , mass number, $A$, (the sum of the protons and neutrons', and so also called the nucleon number) and total charge $Q$ must balance between reactants and products.

## Example Nuclear Transformation

${ }^{249} \mathrm{Cf}+{ }^{18} \mathrm{O} \rightarrow ?+4 n$. The first thing to do is to write each component of the reaction in the form $\frac{\text { atomic number }}{\text { mass number }} \mathrm{X}^{\text {charge }}, 9_{8}^{249} \mathrm{Cf}^{98+}+{ }_{8}^{18} \mathrm{O}^{8+} \rightarrow{ }_{a}^{b} \mathrm{X}^{c}+4{ }_{0}^{1} \mathrm{n}^{0}$, where for now the unknown component is labeled X .
Next, we consider in turn the balance of atomic number (number of protons), mass number (number of protons plus neutrons), and total charge.

- The atomic number balance means:
$98+8 \rightarrow Z+4+0$ and so the number of protons in the unknown species is $Z=98+8-0=106$. From the atomic number we can immediate identify that the unknown nucleus is that of transuranium element with $Z=106$, known as seaborgium, Sg .
- The mass number balance means:
$249+18 \rightarrow A+4 \mu 1$ and so the mass number of the unknown species is $\mathrm{A}=249+18-4=263$. This means the Sg nucleus contains 106 protons and $263-106=157$ neutrons.

| NUCLEAR FISSION | NUCLEAR FUSION |
| :--- | :--- |
| A heavy nucleus breaks up to form <br> two lighter nuclei. | Two light nuclei combine to form <br> a heavy nucleus. |
| It involves a chain reaction. | Chain reaction is not involved. |
| The heavy nucleus is bombarded <br> with neutrons. | Light nuclei are heated to an <br> extremely high temperature. |
| We have proper mechanisms to <br> control fission reaction for <br> generating electricity. | Proper mechanisms to control <br> fusion reaction are yet to be <br> developed. |
| Disposal of nuclear waste is a <br> great environmental problem. | Disposal of nuclear waste is not <br> involved. |
| Raw material is not easily available <br> and is costly. | Raw material is comparatively <br> cheap and easily available. |

## 12C Fission vs. Fusion

There are two main types of nuclear reactions that can be discussed:
Fission: The process of causing a larger atom to split into multiple smaller atoms, releasing energy in the process.

- This is the system that we use in nuclear power plants.
- It is relatively easy to do, but also leaves us with lots of nuclear waste that must be stored safely for thousands of years.
Fusion: The process of causing smaller atoms to fuse together into a larger atom, in the process releasing energy.
- This is the process that drives our sun, and all other suns.
- We can do it under the right conditions in a lab, but we end up putting in more energy than we get out.
- The left over products of these reactions are safe, which is why a lot of research is going into developing fusion reactors.


|  |  |  | Mnemonic |  | prefix | symbol | number | power of 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | King |  | kilo | k | 1000 | $10^{3}$ |
|  |  |  | Henry |  | hecto | h | 100 | $10^{2}$ |
|  |  |  | died |  | deka | da | 10 | $10^{1}$ |
|  |  |  | by |  | base unit | b | 1 | $10^{0}$ |
|  |  |  | drinki |  | deci | d | 0.1 | $10^{-1}$ |
|  |  |  | choco | late | centi | C | 0.01 | $10^{-2}$ |
|  |  |  | milk |  | milli | m | 0.001 | $10^{-3}$ |
|  |  |  | Expanded List of Metric Prefixes |  |  |  |  |  |
|  |  |  | prefix | symbol |  | number |  | power <br> of 10 |
|  |  |  | yotta | Y |  |  |  | $10^{24}$ |
|  |  |  | zetta | Z |  |  |  | $10^{21}$ |
|  |  |  | exa | E | 10000000000 | 000 |  | $10^{18}$ |
|  |  |  | peta | P | 10000000 | 000 |  | $10^{15}$ |
|  |  |  | tera | T | 1000 | 000 |  | $10^{12}$ |
|  |  |  | giga | G |  | 000 |  | $10^{9}$ |
|  |  |  | mega | M |  | 000 |  | $10^{6}$ |
| TABLE 1.4 SI Base Units |  |  | kilo | k | 1000 |  |  | $10^{3}$ |
|  |  |  | hecto | h | 100 |  |  | $10^{2}$ |
| Physical Quantity | Name of Unit | Abbreviation | deka <br> base | da | 10 |  |  | $10^{1}$ |
| Mass |  |  | base unit <br> deci | b | 1 |  |  | $10^{0}$ |
| Length | Meter | kg m | deci | c | 0.01 |  |  | $\frac{10}{10^{-2}}$ |
| Time | Second | $\mathrm{s}^{\text {a }}$ | milli | m | 0.001 |  |  | $10^{-3}$ |
| Temperature | Kelvin | K | micro | $\mu$ | 0.000001 |  |  | $10^{-6}$ |
| Amount of substance | Mole | mol | nano | n | 0.000000001 |  |  | $10^{-9}$ |
| Electric current | Ampere | $\mathrm{A}$ | pico | p | 0.000000000001 |  |  | $10^{-12}$ |
| Luminous intensity | Candela | cd | femto <br> atto | f | 0.000000000000001 |  |  | $10^{-15}$ |
| ${ }^{*}$ The abbreviation sec is frequently used. |  |  | zepto | z | 0.000000000000000000001 |  |  | $10^{-21}$ |
|  |  |  | yocto | y | 0.000000000000000000000001 |  |  | $1{ }^{1} 10^{-24}$ |

