Elements – (Metals)

Properties of Metals

Properties:
1) Electrical and thermal conduction
2) Luster
3) Deform under stress without cleaving
4) Form positive ions

Why:
1) Electrical and thermal conduction
   Electrical and thermal conduction because of movement of free electrons while metals cations remain fixed

2) Luster
   Electrons are excited to higher energy state by light all wavelengths. Electrons fall back to lower levels and re-emit light so metals have shiny surface.

3) Deform under stress without cleaving
   Held together by mobile electrons
   Ductile – drawn into wire
   Malleable – pounded into plate

4) Form positive ions
   Metals readily lose one or more electrons since electron configuration is such that many inner electrons shield full charge of nucleus and so outer electrons are not as strongly held. For example K forms K⁺ but Cl is apt to gain electron to become Cl⁻. Notice that both K⁺ and Cl⁻ are isoelectronic (same electronic configuration as) with noble gas Ar.

High Density because of close packed arrangement
   many metals have 12 nearest neighbors

Group 1 and 2 metals exceptions to above - they are soft metals
   They have largest radii and smallest masses

Most metals have high M.P. and B.P. which shows the strength of the metallic bond
Occurrence

Some pure metals – native metals example - gold (not reactive)

Most metals obtain from oxides example - iron oxide

or from carbonates or sulfides converted to oxides

Group 1 (IA) metals- alkali metals

<table>
<thead>
<tr>
<th>Element</th>
<th>Symbol</th>
<th>Outermost e</th>
<th>Electron Configuration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li</td>
<td>Lithium</td>
<td>2s(^1)</td>
<td>s(^2) p(^1)</td>
</tr>
<tr>
<td>Na</td>
<td>Sodium</td>
<td>3s(^1)</td>
<td>s(^2) p(^4)</td>
</tr>
<tr>
<td>K</td>
<td>Potassium</td>
<td>4s(^1)</td>
<td>s(^2) p(^6) d(^1)</td>
</tr>
<tr>
<td>Rb</td>
<td>Rubidium</td>
<td>5s(^1)</td>
<td>s(^2) p(^6) d(^5)</td>
</tr>
<tr>
<td>Cs</td>
<td>Cesium</td>
<td>6s(^1)</td>
<td>s(^2) p(^6) d(^5)</td>
</tr>
<tr>
<td>Fr</td>
<td>Francium (rare)</td>
<td>7s(^1)</td>
<td>s(^2) p(^6) d(^6)</td>
</tr>
</tbody>
</table>

Properties

Compared to other metals

Low density

Low melting point, low boiling point (Cs M.P. = 29 °C)

Soft (can cut with steel knife)

Have one electron beyond noble gas so form +1 ions

easy to get off first electron, difficult to get off 2\(^{nd}\) electron

Good reducing agents because they will undergo oxidation

\( \text{Li} \rightarrow \text{Li}^+ + e^- \) occurs readily

Reaction examples

With chlorine gas

\( 2\text{K}(s) + \text{Cl}_2(g) \rightarrow 2\text{KCl}(s) \) where made of K\(^+\) cations and Cl\(^-\) anions

Can have explosive reaction with water - more reactive as go down group

\( 2\text{K} + 2\text{H}_2\text{O} \rightarrow 2\text{KOH(aq)} + \text{H}_2(g) + \text{energy} \)

Less reactive metals will react only if water and acid (not just water)

\( \text{Zn (s) + 2 H}^+(aq) \rightarrow \text{Zn}^{2+}(aq) + \text{H}_2(g) \)

Group 1 metals have strong tendency to oxidize \(+ E_{ox}^\circ\)

The larger the \(+ E_{ox}^\circ\) the more the readily reaction occurs

\( \text{Na} \rightarrow \text{Na}^+ + e^- \rightarrow +2.71 \text{ V} \) more reactive than Zn

\( \text{Zn} \rightarrow \text{Zn}^{2+} + 2e^- \rightarrow +.76 \text{ V} \)
**Group 2 (IIA) Metals – alkaline earth metals**

**List**

<table>
<thead>
<tr>
<th>Be</th>
<th>Beryllium</th>
<th>2s²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mg</td>
<td>Magnesium</td>
<td>3s²</td>
</tr>
<tr>
<td>Ca</td>
<td>Calcium</td>
<td>4s²</td>
</tr>
<tr>
<td>Sr</td>
<td>Strontium</td>
<td>5s²</td>
</tr>
<tr>
<td>Ba</td>
<td>Barium</td>
<td>6s²</td>
</tr>
</tbody>
</table>

**Properties and Reactions**

Second most reactive metals  
Not found free in nature  
Produced by electrolysis of molten chlorides  

M.P. much greater than Group 1 metals  
(extreme example: Sr M.P. = 800°C, Cs M.P. = 29°C)

Tend to form +2 ions  

**Reaction examples**

\[
\text{Mg} + \text{X}_2 \rightarrow \text{MgX}_2 \quad \text{X} = \text{halogen F, Cl, Br, I}
\]

\[
2\text{Ca} + \text{O}_2 \rightarrow 2\text{CaO}
\]
Group 13 (IIIA) Metals

List

<table>
<thead>
<tr>
<th>Element</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>Aluminum</td>
</tr>
<tr>
<td>Ga</td>
<td>Gallium</td>
</tr>
<tr>
<td>In</td>
<td>Indium</td>
</tr>
<tr>
<td>Tl</td>
<td>Thallium</td>
</tr>
</tbody>
</table>

Properties

Al is most abundant metal of earth’s crust (8%)  
other Group 13 metals found in trace amounts

Al important in containers and light weight parts

Al tends to form 3+ ions
**Group 14 (IVA) metals**

**List**

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Ge</td>
<td>Germanium</td>
</tr>
<tr>
<td>Sn</td>
<td>Tin</td>
</tr>
<tr>
<td>Pb</td>
<td>Lead</td>
</tr>
</tbody>
</table>

**Properties**

- Form +4, +2 oxidations states
- But not +4 ions, +4 ox nu is associated with covalent compounds

Generally form covalent compounds rather than ions - example PbCl₄ is actually covalent and not ionic because metal is more like nonmetal and less like metal in some ways.
Group 3 – 12 (IIIB – IIB)- Transition Metals

Properties

High M.P. and B.P.  (Group 3 exception Zn, Cd, Hg)
   Extreme example Hg liquid at room temp.

Good conductors of electricity and heat
   (Group 11 are outstanding conductors Cu, Ag, Au)

Electronic configuration:
   Filling of d orbitals
   Variety of oxidation states possible for transition metals
   Be aware of range of possibilities

   +1 important only for Cu⁺, Ag⁺, Au⁺, Hg₂²⁺
   +2 important for 4th period (s orbital e⁻ come off first)
   Maximum ox. nu. in middle of d period

Unpaired electrons cause many transition metal compounds to be:
   colored solids (most pure organic compounds are white solids)
   and paramagnetic (attracted to a magnetic field –
       unpaired electrons are like tiny magnets in atoms)

Relative Reactivity

Observe that gold does not react in hydrochloric or nitric acid (a classic test for gold) while zinc reacts with both and copper with only nitric.

<table>
<thead>
<tr>
<th></th>
<th>HCl(aq)</th>
<th>HNO₃(aq)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Cu</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Au</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

So therefore the relative reactivity is:  Zn > Cu > Au

Air oxidation of metals causes corrosion that
   destroys metal and changes properties

Rust costs billions of dollars a year.

Gold is prized for jewelry because it does not react.
Lanthanides (inner transition elements)

List
Elements number 51 through 71 (see Periodic Table)
Filling of 4f orbitals

Properties and Reactions

Lanthanide contraction - atomic ionic radii decrease with increasing atomic number because 4f orbitals are not outermost

Also because 4f are not outermost, addition of electrons does not cause much variation in properties

Chemistry of lanthanides are all very similar

All form +3 oxidation state through loss of 2 s e\(^{-}\) and 1 f e\(^{-}\)

Elements are very difficult to separate and are frequently found as mixtures of lanthanide elements in nature

Very Reactive:

\[
2 \text{M} + 3 \text{X}_2 \rightarrow 2 \text{MX}_3 \\
2 \text{M} + 3 \text{O}_2 \rightarrow 2 \text{M}_2\text{O}_3
\]
Extra Topics in Chemistry 1120 Metals

Metallic Bond and Conduction

Array of positive ions surrounded by outer electrons

Because of low electronegativities, ionization energies of outer e- are loosely held and free to move

Energy band within crystal

Each atom brings to the crystal certain orbitals and the orbitals may be partially or completely filled

Bands = energy level
(in diagram below energy increases going down
usually higher energy level as you go up in diagram)

N orbitals form band with N levels
And can contain 2N electrons since can have 2 e in orbital

http://www.chem.queensu.ca/people/faculty/mombourquette/FirstYrChem/Molecular/bands/index.asp
Valence and Conduction Bands

Below return to normal view of higher energy as you go up

Valence band - band formed from outer electrons

Conduction band - empty sites available for electrons to move in

Forbidden energy zone (band gap) - gap between energy bands
Temperature Effects

Electronic Conductivity of semiconductor increases with increase in temperature so the electrons move up to the conduction band.

Electronic conductivity of metals decreases with increase in temperature because the vibration of atoms limits flow of electrons.

Extrinsic Semiconductor-
   semiconductor in which impurities have been added

Si, Ge     diamond lattice     4 valence e-

Low conductivity

Add B (low concentration) with 1B/ million Si atoms will change conductivity by factor of 100,000

P-type semiconductor
Add IIIA (group 3) element
B three valence e- to bulk Si create a hole vacancy neutral charge
   but e- can move

N-type semiconductor
Add VA (group 5) element
P five valence e- to bulk Si creates an extra electron

http://www.chembio.uoguelph.ca/educmat/chm729/band/psemi.htm
Group 1 (IA) metals- alkali metals (more)

Reactions

\[ 2M + X_2 \rightarrow 2 MX \]  
\[ \text{Halogens } \rightarrow \text{ Halide} \]

\[ 2M + H_2 \rightarrow 2 MH \]  
\[ \text{Hydrogen } \rightarrow \text{ Hydride} \]

\[ 2M + S \rightarrow M_2S \]  
\[ \text{Sulfur } \rightarrow \text{ Sulfide} \]

\[ 2M + H^+ (aq) \rightarrow 2 M^+ + H_2 \]  
\[ \text{Violent reaction} \]

\[ 2M + 2 \text{H}_2\text{O} \rightarrow 2 \text{MOH} (aq) + \text{H}_2(g) \]  
\( \text{Can be violent} \)

Production – Laboratory

Very Reactive metals – react with water

Group 1  
Li, Na, K, Rb  
explosive reaction

Group 2  
Ca, Sr, Ba  
vigorous

\[ 2 \text{Na} (s) + 2 \text{H}_2\text{O} \rightarrow 2 \text{Na}^+ (aq) + 2 \text{OH}^- + \text{H}_2 \]

Oxidation: \( 2 \text{Na} \rightarrow 2 \text{Na}^+ + 2 \text{e}^- \)

Reduction: \( 2 \text{e}^- + 2 \text{H}_2\text{O} \rightarrow 2 \text{OH}^- + \text{H}_2 \)

Metals have strong tendency to oxidize + \( E_{\text{ox}}^0 \)

The larger the +\( E_{\text{ox}}^0 \) the more the energetic the reaction

Less reactive metals will react with acid in water

\[ \text{Zn} (s) + 2 \text{H}^+ (aq) \rightarrow \text{Zn}^{2+} (aq) + \text{H}_2 (g) \]

\[ \text{Na} \rightarrow \text{Na}^+ + \text{e}^- +2.71 \text{V} \]  
\( \text{more reactive than Zn} \)

\[ \text{Zn} \rightarrow \text{Zn}^{2+} + 2 \text{e}^- +.76 \text{V} \]
Reactions - metals

Because of the high bond energy, most reactions with H₂ require high temperature to form metal hydrides

Salt-like Hydrides (M⁺ H⁻ like ionic solids)
Group 1 and heavy Group 2 (Ca, Sr, Ba)

\[
\begin{align*}
2 \text{Na (s)} &+ \text{H}_2 (g) \rightarrow 2 \text{NaH (s)} \\
\text{Ca (s)} &+ \text{H}_2 (g) \rightarrow \text{CaH}_2 (s)
\end{align*}
\]

Interstitial Hydrides

Pt, Pd, Ni
Hydrogen absorbed H-H bond broken, atomic hydrogen is found between layers of metal

Complex Hydrides

Lithium Aluminum hydride

\[
\text{Li}^+ \quad \text{H}^- \quad \text{Al}^- \quad \text{H}^+
\]

http://en.wikipedia.org/wiki/LiAlH₄

with metal oxides to produce water and metal

\[
\text{WO}_3 (s) + 3 \text{H}_2 (g) \rightarrow \text{W (s)} + 3 \text{H}_2\text{O (g)}
\]

Commercial production of tungsten is an expensive process

Reactions – Non metals

Oxygen

\[
2 \text{H}_2 \quad + \quad \text{O}_2 \rightarrow 2 \text{H}_2\text{O}
\]

with water

High T to start, oxyhydrogen torch 2800°C, space shuttle engines
Metals

Halogens

\[ \text{H}_2 + \text{F}_2 \rightarrow 2 \text{HF} \]

Hydrogen fluoride

F₂, Cl₂ room temp
Br₂, I₂ 300°C

Nitrogen

\[ 3 \text{H}_2 (g) + \text{N}_2 (g) \rightarrow 2 \text{NH}_3 (g) \]

Fe catalyst
500 atm, 500°C

Haber process used to make ammonia

Sulfur

\[ \text{H}_2 (g) + \text{S} (g) \rightarrow \text{H}_2\text{S} (g) \]

Hydrogen sulfide

High Temp

Group 2 (IIA) Metals – alkaline earth metals (more)

Reactions

\[ \text{M} + \text{X}_2 \rightarrow \text{MX}_2 \quad \text{X} = \text{halogen F, Cl, Br, I} \]

\[ 2 \text{M} + \text{O}_2 \rightarrow 2 \text{MO} \]

\[ \text{M} + \text{S} \rightarrow \text{MS} \]

\[ \text{M} + \text{H}_2 \rightarrow \text{MH}_2 \]

\[ \text{M} + 2 \text{H}_2\text{O} \rightarrow \text{M(OH)}_2 + \text{H}_2 \quad \text{M} = \text{Ca, Sr, Ba} \]

(Be not react w/ H₂O)

\[ \text{M} + 2 \text{H}^+ \rightarrow \text{M}^{2+} + \text{H}_2 \]
Solubility

Compare solubility of Group 1 and Group 2 salts
Most group 1 metal salts are very soluble
A number of group 2 metal salts are low solubility

<table>
<thead>
<tr>
<th>Anion</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>OH⁻</td>
<td>Hydroxide</td>
</tr>
<tr>
<td>SO₄²⁻</td>
<td>Sulfate</td>
</tr>
<tr>
<td>CO₃²⁻</td>
<td>Carbonate</td>
</tr>
<tr>
<td>C₂O₄²⁻</td>
<td>Oxalates</td>
</tr>
<tr>
<td>F⁻</td>
<td>Fluorides</td>
</tr>
<tr>
<td>CrO₄²⁻</td>
<td>Chromates</td>
</tr>
</tbody>
</table>

Metals Reducing Agents

Metals have low ionization energy, small electron affinity, and easily lose electrons to become positive

\[
M \rightarrow M^{+} + e^- \\
M \rightarrow M^{2+} + 2 e^- \\
Zn + 2 H^+ (aq) \rightarrow Zn^{2+} (aq) + H_2 (g) \\
Ca + 2 NO_3^- + 4 H^+ \rightarrow Ca^{2+} + 2 NO_2 + 2 H_2O
\]

Ease of oxidation is same as reactivity with H⁺

Group 1 is most reactive (lowest ionization energy) and Group 2 is less reactive

Compare ability of one metal to react by oxidizing another

\[
Zn (s) + Cu^{2+} (aq) \rightarrow Zn^{2+} (aq) + Cu (s)
\]

So Zn is more reactive than Cu because it is more easily oxidized or causes Cu to be reduced

Can quantify relative reactivity in electrochemistry chapter
Nonmetals as Oxidizing Agents

Strongest oxidizing agents are found going up and across to the right of the periodic table

\[ F \rightarrow F^- \]
\[ Cl \rightarrow Cl^- \]

F will go to an ion more than Cl

\[ Cl_2 + 2 Br^- \rightarrow 2 Cl^- + Br_2 \]

Cl is reduced (gain e)
Br is oxidized (lose e)

Reactions with Oxygen:

Metals-

\[ 2 Fe + 3/2 O_2 + x H_2O (l) \rightarrow Fe_2O_3xH_2O \]

Iron oxide crystal includes water molecules

\[ Cu + O_2 (g) \rightarrow CuO (s) \text{ Slow at room T} \]
\[ 2 Mg (s) + O_2 (g) \rightarrow 2 MgO (s) \text{ Fast at high T} \]

Can predict product if unique ox nu of metal

Nonmetals-

\[ 2 H_2 (g) + O_2 (g) \rightarrow 2 H_2O \]
\[ C (s) + O_2 (g) \rightarrow CO_2 (g) \]
\[ S (s) + O_2 (g) \rightarrow SO_2 (g) \]

\[ N_2 + O_2 \rightarrow 2 NO \text{ Only at very high T, otherwise not in atmosphere} \]
\[ CH_4 + 2 O_2 \rightarrow CO_2 + 2 H_2O \]

More complicated molecules react to form would as direct elements
Lewis Acids and Bases

Lewis acid - electron pair acceptor
Lewis base – electron pair donor

Lewis Acid


Lewis Base

http://bilbo.chm.uri.edu/CHM112/lectures/Image1110.gif

Bronsted Lowry acid and base also fits Lewis definition but there is also a more general definition

<table>
<thead>
<tr>
<th>Acid</th>
<th>Base</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{H}_3\text{O}^+$</td>
<td>$\text{OH}^-$</td>
<td>$\text{2 H}_2\text{O}$</td>
</tr>
<tr>
<td>$\text{Ag}^+$</td>
<td>$\text{Cl}^-$</td>
<td>$\text{AgCl}$</td>
</tr>
</tbody>
</table>
Ore

Naturally occurring material from which pure metal may be extracted

Ores when mined include other unwanted materials such as silica, clay, granite, and gangue (worthless rocks in which minerals are found)

Metallurgy

extraction of metals from ores and converting into usable form

Three stages
1. preliminary treatment – desired part of ore concentrated impurities removed or mineral converted to another form
2. reduction – metal compound reduced to free metal
3. refining – metal purified and if needed other substances added to give desired product

Preliminary Treatment

Concentration of ore removal of gangue
Physical separation
Mineral particles may be separated by floatation

Oil – mineral particles go into oil
Water – used to remove gangue

Chemical separation

Example make steel from Fe$_3$O$_4$

Iron containing ore crushed
Magnetic Fe$_3$O$_4$ separated from gangue by attracting it with electromagnets
Reduction

Reduction in blast furnace

\[
\begin{align*}
\text{Heat} & \quad \text{Fe}_3\text{O}_4 (s) + \text{CO (g)} \rightarrow 2 \text{FeO (s)} + \text{CO}_2 (g) \\
\text{Heat} & \quad \text{FeO (s)} + \text{CO (g)} \rightarrow \text{Fe (l)} + \text{CO}_2 (g)
\end{align*}
\]

Hot air reacts with carbon to form CO
Gives off heat
CO reacts with iron oxide

Refining

Pig iron contains up to 40% C, 2% Si, P, S
These impurities are removed by direct stream of high pressure oxygen into pig iron
Oxides formed as gases and removed by stream of oxygen

Steel- other metals added as needed to improve properties