### **Chapter 5**

# Introduction to Reactions in Aqueous Solutions

### **Aqueous Solutions**

Solvent

liquid water

### • Solute

substance being dissolved

### • Electrolyte

substance that, when dissolved in a polar solvent like water, produces a solution that can conduct electricity

(because it produces ions, which are charged)

### **Electrolytes**

### **Strong** electrolyte:

When dissolved in water, dissociates almost completely into ions

- <u>Highly water-soluble ionic compounds</u> (e.g. NaCl, K<sub>2</sub>SO<sub>4</sub>)
- <u>Strong acids</u> (e.g. HCl, H<sub>2</sub>SO<sub>4</sub>)
- <u>Strong bases (e.g. KOH)</u>

In this course, we will assume that all ionic compounds dissolve as separated cations and anions

### Weak electrolyte:

When dissolved in water, only some of it produces ions

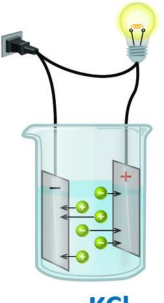
- <u>Weak acids</u> (e.g.  $HC_2H_3O_2$ )
- <u>Weak bases</u> (e.g. NH<sub>3</sub>)

### Non-electrolyte:

Does not dissociate into ions at all (e.g. sugars, alcohols, oils)

### **Electrolyte solutions conduct electricity**

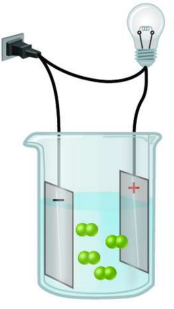
### ions can carry the electric charge though the solution



### KCl High conductivity

Strong Electrolytes conduct current efficiently (bulb shines brightly at high concentration) Acetic Acid Low conductivity

### Weak Electrolytes conduct only a small current (bulb glows dimly even at high concentration)



### Sugar No conductivity

### Nonelectrolytes

conduct no current (bulb remains unlit).

### **Be aware:**

The terms "strong electrolyte" or "weak electrolyte" refers to the **<u>substance</u>** dissolved (i.e. solute)

• A concentrated solution of a strong electrolyte is a good conductor and can make the bulb glow bright.

### But ...

- A <u>dilute</u> solution of a strong electrolyte won't conduct much electricity, and the bulb will <u>not</u> glow bright.
  - Even if the dissolved "strong electrolyte" fully dissociated, if there isn't much of it, it will still have low conductivity

On the other hand,

Even a concentrated solution of a weak electrolyte is not a good conductor and will not make the bulb glow bright. A molecular substance like hydrogen chloride (HCI), which is <u>not</u> ionic, but fully dissociates into ions when dissolved in water, is <u>also</u> a strong electrolyte.

Because it's a strong acid, fully dissociating into H<sup>+</sup>(aq) and Cl<sup>-</sup>(aq) ions

Note: Acids produce H<sup>+</sup> ions in water, along with an anion

Strong acids are strong electrolytes

-- fully ionize in water

Weak acids are weak electrolytes

-- only partially ionize in water

Almost all the bases we see in this course are **metal hydroxides** like NaOH or KOH, and are strong electrolytes (<u>because they are ionic compounds</u>).

Remember: we will assume that all ionic compounds fully dissociate into their ions when dissolved

Note: Bases produce  $OH^-$  ions in water, along with a cation NaOH(aq)  $\rightarrow$  Na<sup>+</sup>(aq) + OH<sup>-</sup>(aq) One weak base we encounter is ammonia: NH<sub>3</sub>

• A <u>molecular</u> compound (as opposed to the usual metal hydroxide like NaOH)

-- no guarantee it will fully ionize

-- indeed it doesn't

- A small portion reacts with water to form NH<sub>4</sub><sup>+</sup> and OH<sup>-</sup> Produced by reacting with water
- Not fully converted to ions
  - -- so it is a **weak electrolyte**

### **Types of Chemical Reactions we will look at:**

- Precipitation Acid–Base

Double-displacement reactions
Also called: Double-replacement Exchange **Methathesis** 

Oxidation–Reduction

A wide variety of reactions

### **Precipitation Reactions**

A **<u>double displacement</u>** reaction in which a <u>solid</u> forms and separates from the solution.

Precipitate (*verb*): to drop out of solution Precipitate (*noun*): the solid that forms

We will consider precipitation of <u>ionic compounds</u>

Remember: When ionic compounds dissolve in water, the resulting solution contains the **separated ions**.

If ions from two soluble ionic compounds

<u>cation</u> from one, <u>anion</u> from the other correspond to an <u>insoluble</u> compound

### they combine and precipitate

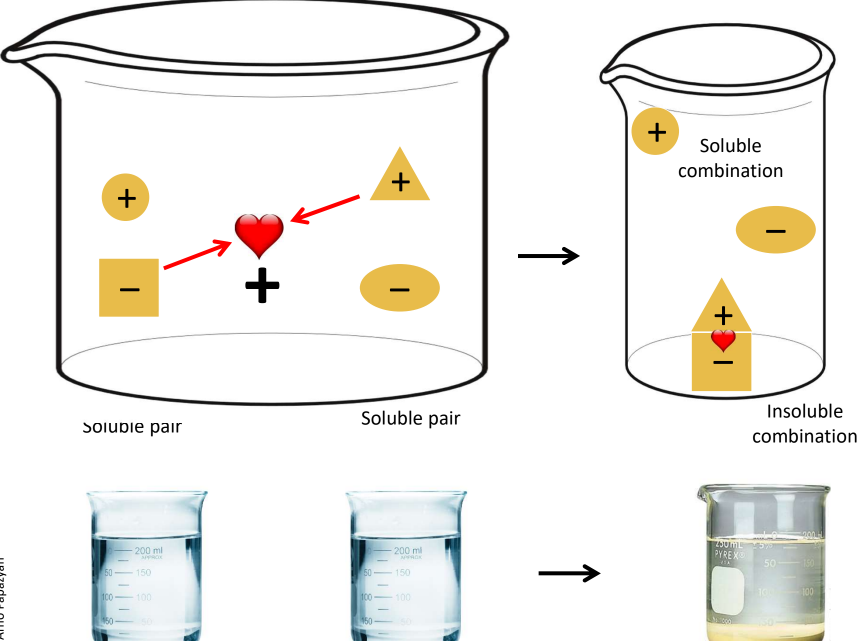
if they are in the same solution

it doesn't matter if there are soluble combinations

### insoluble combination wins

- Because these ions are "happier" together in solid form
- That's why that combination is insoluble

#### **Precipitation Reactions**



### Terms related to precipitation

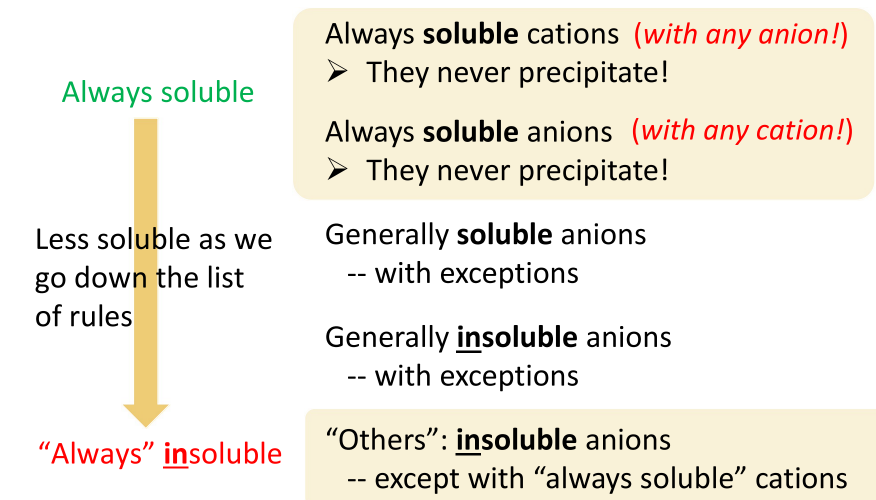
- *Soluble:* solid dissolves in solution; (*aq*) is used in the reaction equation.
- *Insoluble:* solid does not dissolve in solution; (*s*) is used in reaction equation.
- Insoluble and slightly soluble are often used interchangeably. We will do that too.

What's called "soluble" or "insoluble" may differ slightly, depending on the industry, discipline, or preferences of an instructor

Marginally soluble: considered "soluble", but barely
 -- cannot have concentrated solutions

### **Solubility Rules for cation-anion pairs**

Before you get scared by a bunch of rules to memorize, observe that the solubility rules are ordered in decreasing solubility:



### **Solubility Rules for cation-anion pairs**

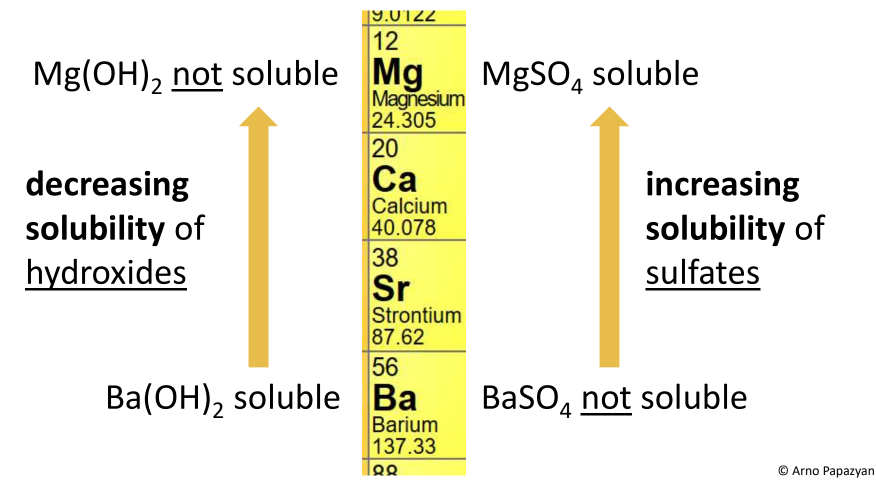
- 1. Alkali metal (Group 1) cations and NH<sub>4</sub><sup>+</sup> are <u>always soluble</u>
- 2. Nitrate (& perchlorate, chlorate, acetate) salts are <u>always soluble</u>.
- 3. Most Cl<sup>-</sup>, Br<sup>-</sup>, and l<sup>-</sup> salts are soluble (except with Ag<sup>+</sup>, Pb<sup>2+</sup>, Hg<sub>2</sub><sup>2+</sup>). halides
- 4. Most **sulfates** are soluble (except with Sr<sup>2+</sup>, Ba<sup>2+</sup>, Ca<sup>2+</sup>, Hg<sub>2</sub><sup>2+</sup>, Pb<sup>2+</sup>; and Ag<sub>2</sub>SO<sub>4</sub> is only "marginally soluble").
- Most hydroxides are insoluble or only slightly soluble. Exceptions: see Rule 1, and Ba(OH)<sub>2</sub>, Sr(OH)<sub>2</sub>, Ca(OH)<sub>2</sub> are somewhat soluble
- 6. Salts of **"other"\*** anions (CO<sub>3</sub><sup>2–</sup>, CrO<sub>4</sub><sup>2–</sup>, PO<sub>4</sub><sup>3–</sup>, C<sub>2</sub>O<sub>4</sub><sup>2–</sup> etc. we see in General Chemistry\*\*) are insoluble (exceptions: see Rule 1)

*\** anions <u>not mentioned explicitly</u> in the rules

\*\* **not literally** all other anions known to humankind!

## Additional observations about alkaline earth ionic compound solubilities

For Group 2 metals, hydroxide & sulfate solubility trends are opposite of each other



### **On sulfides:**

- Sulfide ion, S<sup>2-</sup>, is generally insoluble (except, as always, with alkali metals and ammonium)
- Sulfides of Group 2 metals (alkaline earth) are soluble.

But:

- Sulfide ion in water is unstable in the long term.
- It slowly reacts with water to produce H<sub>2</sub>S gas.

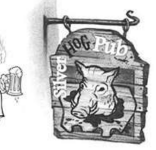
Because its solutions are unstable, I don't include sulfide in the "normal" rules of solubility.

### A silly way to remember the Solubility Rules

- -- mnemonics are about the exceptions
- -- exceptions are insoluble in rules 3 & 4, and soluble in later rules

### Some silly mnemonics for the Solubility Rules

- 1. Group I and <u>am</u>monia **"I am" always soluble**
- 2. perchlorate, chlorate, acetate, nitrate "pecan" always soluble
- 3. Halides (Cl<sup>-</sup>, Br<sup>-</sup>, I<sup>-</sup>) generally soluble
  - except with Ag<sup>+</sup>, Hg<sub>2</sub><sup>2+</sup>, Pb<sup>2+</sup>
     "He lost his halo at the <u>Silver Hog Pub</u>"
- 4. Sulfate (SO<sub>4</sub><sup>2-</sup>) generally soluble
  - except with Sr<sup>2+</sup>, Ba<sup>2+</sup>, Ca<sup>2+</sup>, Ag<sup>+</sup>, Hg<sub>2</sub><sup>2+</sup>, Pb<sup>2+</sup>
     "So 4 his fun, Sr. (Señor) Baca goes to the Silver Hog Pub"
- 5. Hydroxide (OH<sup>-</sup>) generally insoluble
  - except with Sr<sup>2+</sup>, Ba<sup>2+</sup>, Ca<sup>2+</sup>
     "Oh, except for <u>Sr. Baca!"</u>
- Compounds of other anions are insoluble (exception: Rule 1)
   <u>Others</u> are powders, except where <u>I am</u>



### Which of the following ions forms a compound with Pb<sup>2+</sup> that is soluble in water?

- a) S<sup>2-</sup>
- b) Cl<sup>-</sup>
- c)  $NO_3^-$
- d) SO<sub>4</sub><sup>2-</sup>
- e) Na<sup>+</sup>

Precipitation reactions can be represented in three different forms:

- Molecular equation
- Complete ionic equation
- Net ionic equation

### "Molecular" Equation

A better term would have been "Formula Equation", or maybe "Chemical Equation" since there are no molecules around.

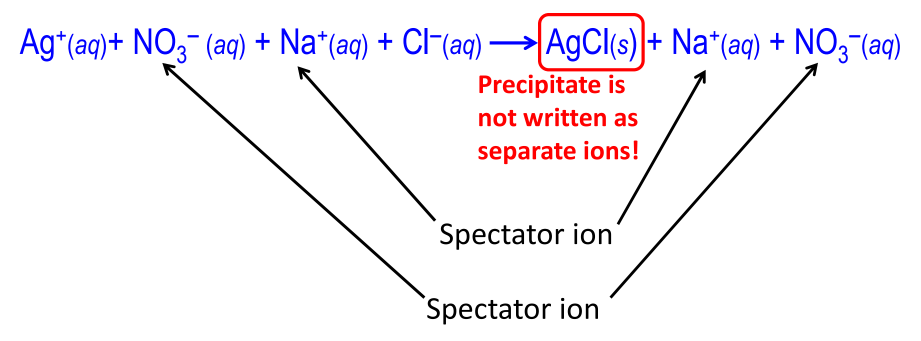
- > All ions or ionic compounds
- Reactants and products shown as compounds
   -- instead of the dissociated ions
- Use solubility rules to determine which compounds are aqueous and which compounds are solids.

AgNO<sub>3</sub>(aq) + NaCl(aq) 
$$\rightarrow$$
 AgCl(s) + NaNO<sub>3</sub>(aq)  
Ag<sup>+</sup> and Cl<sup>-</sup> are not soluble together

### **Complete Ionic Equation**

All <u>soluble</u> substances that are ionized in aqueous solution are represented as ions.

### Only the precipitate is written as a compound



Spectator ions don't participate in the reaction.

They appear on both sides of the reaction in the same state.

### **Net Ionic Equation**

Includes only the ions that do something.
 i.e. the ions that precipitated

### $Ag^+(aq) + Cl^-(aq) \rightarrow AgCl(s)$

• No spectator ions

Na<sup>+</sup> and NO<sub>3</sub><sup>-</sup> are spectator ions. They remain in solution. So they don't show up in the net ionic equation

### **Concept Practice**

Write the correct "<u>molecular</u>" equation, <u>complete ionic</u> <u>equation</u>, and <u>net ionic equation</u> for the reaction between cobalt(II) chloride and sodium hydroxide.

Formula (Molecular) Equation:

 $CoCl_2(aq) + 2NaOH(aq) \rightarrow Co(OH)_2(s) + 2NaCl(aq)$ 

<u>Complete Ionic Equation</u>:

 $Co^{2+}(aq) + 2CI^{-}(aq) + 2Na^{+}(aq) + 2OH^{-}(aq) \rightarrow Co(OH)_{2}(s) + 2Na^{+}(aq) + 2CI^{-}(aq)$ 

Net Ionic Equation:

 $Co^{2+}(aq) + 2OH^{-}(aq) \rightarrow Co(OH)_{2}(s)$ 

### Acids, Bases, and Their Reactions

Acid-base "neutralization" reactions are also "double-displacement"

We touched on acids and bases earlier

- > When we discussed inorganic nomenclature
- And earlier in this chapter when we considered electrolytes

We will now look at them at more depth

### **Acids & bases as defined by Arrhenius**

- Oldest acid-base concept
- Applies in aqueous solutions
- Acids provide H<sup>+</sup> in aqueous solution
- Bases provide OH<sup>-</sup> in aqueous solution
- When an acid and a base "neutralize" each other
   >H<sup>+</sup> and OH<sup>-</sup> combine to form H<sub>2</sub>O

Remaining parts of the acid and the base form a "salt"

### Acids as defined by Arrhenius ("Arrhenius Acids")

### Acid: provides H<sup>+</sup>

 $H\{Anion\}(aq) \rightarrow H^+(aq) + \{Anion\}(aq)$ 

$$HNO_3$$
 (aq)  $\rightarrow H^+$  (aq) +  $NO_3^-$  (aq)

<u>"Acidic" hydrogens</u> are attached to an anion

 $H^+(aq)$  produced actually attaches to a water molecule, forming  $H_3O^+$ 

- In the formula of an Arrhenius acid, the acidic hydrogen is written first.
- Any other hydrogens later in the formula are not acidic.
   HC<sub>2</sub>H<sub>3</sub>O<sub>2</sub>
   acidic not acidic
- An acid with one acidic hydrogen is "<u>monoprotic</u>" HNO<sub>3</sub>
- An acid with two acidic hydrogens is "<u>di</u>protic" H<sub>2</sub>SO<sub>4</sub>
- An acid with three acidic hydrogens is "<u>tri</u>protic" H<sub>3</sub>PO<sub>4</sub>

### Weak acids:

Only some of the weak acid molecules dissociate into ions

 $HC_2H_3O_2(aq) \rightarrow$ 

Most acetic acid molecules remain undissociated

 $H^+(aq) + C_2H_3O_2^-(aq)$ Only some of the molecules dissociate into ions

The format of a reaction equation gives the impression that all reactions are always complete. That's not true

### **Strong acids:**

<u>Practically all</u> of the strong acid molecules dissociate into H<sup>+</sup> and the anion of the acid.

HBr  $(aq) \rightarrow H^+(aq) + Br^-(aq)$  All of the acid molecules dissociate into ions

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### List of "Strong Acids" we need to know:

- $HI(aq) \longrightarrow H^+(aq) + I(aq)$
- HBr(aq)  $\rightarrow$  H<sup>+</sup>(aq) + Br<sup>-</sup>(aq)  $\succ$  compounds of
- $HCl(aq) \rightarrow H^+(aq) + Cl^-(aq)$
- Binary hydrogen compounds of halogens, except HF(aq)
- $HClO_4(aq) \rightarrow H^+(aq) + ClO_4^-(aq)$
- $HNO_3(aq) \rightarrow H^+(aq) + NO_3^-(aq)$

 $H_2SO_4(aq) \rightarrow H^+(aq) + HSO_4^-(aq)$ 

Even for a strong acid like  $H_2SO_4$ , if there is more than one acidic H, only the 1<sup>st</sup> dissociation is complete. The 2<sup>nd</sup> H doesn't dissociate easily.

### "Weak Acids" we need to know:

- Any acid not listed among the "strong"
  - Some may be fairly strong, but just don't dissociate fully
- Organic acids such as acetic acid  $(HC_2H_3O_2)$
- **HF**(*aq*)
- H<sub>3</sub>PO<sub>4</sub>
- The anion produced after diprotic or triprotic acids, even if they are strong in releasing their first acidic H

 $HSO_4^-(aq) \rightarrow H^+(aq) + SO_4^{2-}(aq)$ Incomplete dissociation

### Extracurricular knowledge, if you enjoy obscure info The strange case of Hydrogen Fluoride (HF)



- HF is normally regarded as a weak acid.
- In dilute aqueous solution, after HF dissociates fully, F<sup>-</sup> forms a tight ion pair with the H<sub>3</sub>O<sup>+</sup> just created
   ➤ makes H<sub>3</sub>O<sup>+</sup> unavailable to function as free H<sup>+</sup>(aq)
- Actually when there isn't much water around, HF is a stronger acid than our usual "strong" acids like H<sub>2</sub>SO<sub>4</sub>, HCl, etc.

### **Nevertheless:**

In the world of General Chemistry where aqueous (and not very concentrated) solutions rule, <u>we will regard HF</u> <u>as a "weak acid"</u>

### **Bases as defined by Arrhenius ("Arrhenius bases")**

**Base: provides OH<sup>-</sup>** 

{Cation}OH (aq)  $\rightarrow$  {Cation}<sup>+</sup> (aq) + OH<sup>-</sup> (aq)

NaOH (aq)  $\rightarrow$  Na<sup>+</sup> (aq) + OH<sup>-</sup> (aq)

- Bases in the Arrhenius definition act by producing OH<sup>-</sup> ions
- Most inorganic bases are metal hydroxides, such as NaOH, KOH, Ca(OH)<sub>2</sub>.
- In General Chemistry it's common to assume all metal hydroxides to be strong bases.
- Ammonia  $(NH_3)$  is the only weak base we will need to know in this course.

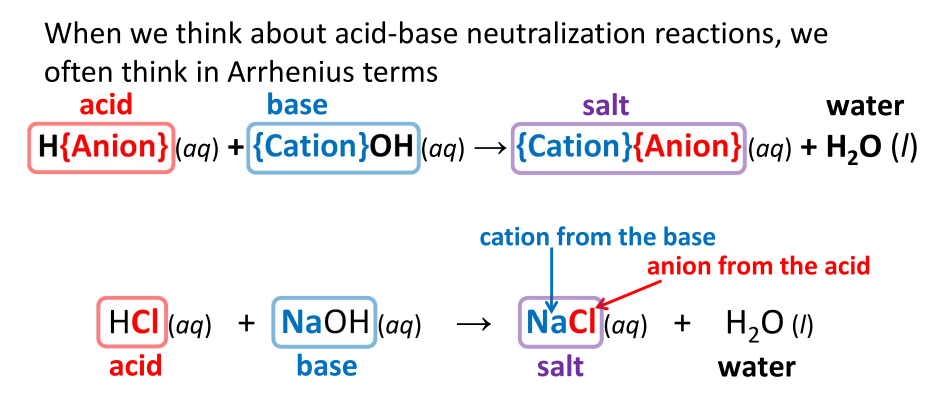
 $NH_3$ , a molecular compound and a weak base, creates the hydroxide ion by reacting with water:

 $NH_3(aq) + H_2O(l) \rightarrow NH_4^+(aq) + OH(aq)$ 

Since it is a weak base, the reaction is not complete, and most of the ammonia molecules dissolved in water remain as NH<sub>3</sub> molecules.

When you see a reagent bottle labeled "NH<sub>4</sub>OH" or "ammonium hydroxide", it is mostly dissolved ammonia rather than the "compound" NH<sub>4</sub>OH.

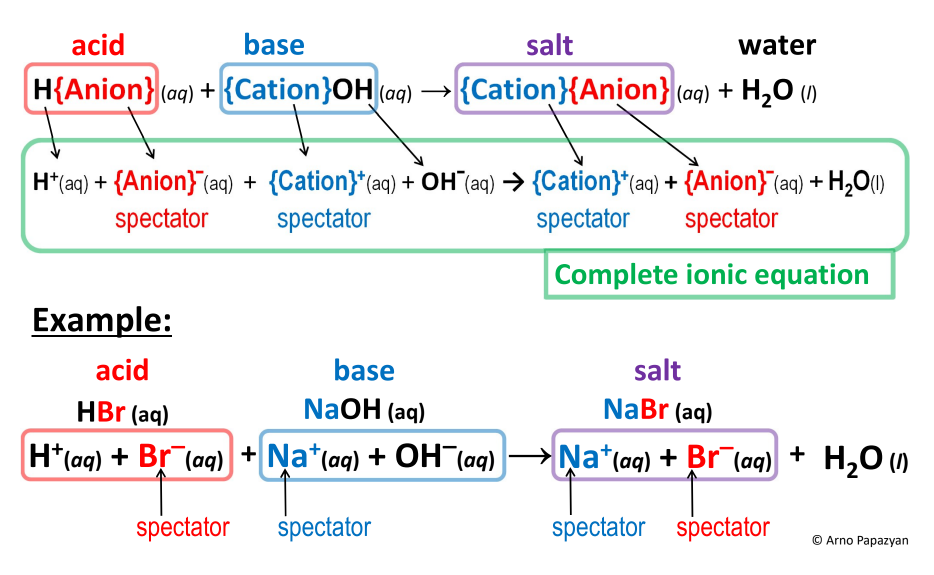
#### **Arrhenius acid-base reactions (Neutralization)**



- Ions are shown only as part of compounds above.
- So it is the "molecular equation" representation of an acidbase reaction.

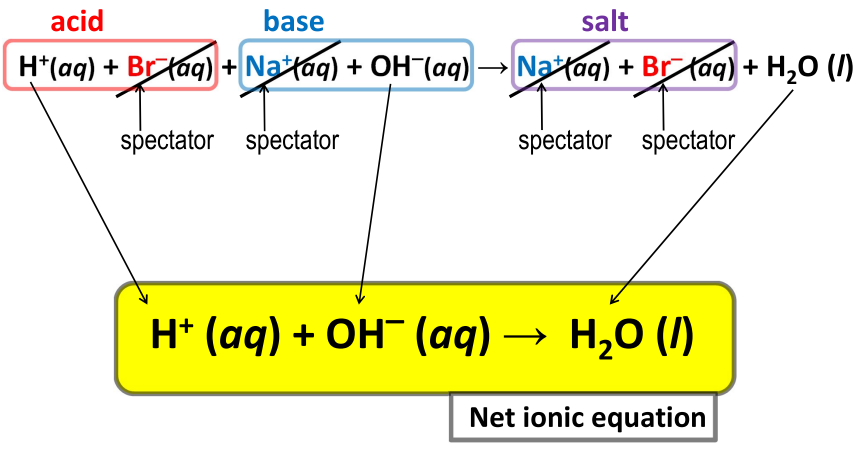
#### **<u>Complete ionic</u>** equation for Arrhenius acid-base reactions

Acid, base, and the produced salt written as ions



#### **<u>Net</u>** ionic equation for an Arrhenius acid-base reaction

#### Obtained when we eliminate spectator ions from the complete ionic equation



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It is part of general chemistry folklore to write ionic equations of acid-base reactions of weak acids with an undissociated weak acid molecule because their ionization is incomplete when alone.

For a weak acid like acetic acid  $(HC_2H_3O_2)$ :

"Traditional" Complete Ionic Equation  $HC_{2}H_{3}O_{2}(aq) + Na^{+}(aq) + OH^{-}(aq) \rightarrow Na^{+}(aq) + C_{2}H_{3}O_{2}^{-}(aq) + H_{2}O(I)$ 

"Traditional" Net Ionic Equation  $HC_{2}H_{3}O_{2}(aq) + OH^{-}(aq) \rightarrow C_{2}H_{3}O_{2}(aq) + H_{2}O(I)$  Similarly, for a <u>weak base</u>, the "traditional" way to write the ionic equations is:

"Traditional" Complete Ionic equation  $H^+(aq) + Br^-(aq) + NH_3(aq) \longrightarrow NH_4^+(aq) + Br^-(aq)$ Weak base

"Traditional" Net Ionic equation  $H^+(aq) + NH_3(aq) \longrightarrow NH_4^+(aq)$ Weak base That reasoning is probably not right, because:

- H<sup>+</sup> cation diffuses drastically faster than the undissociated acid molecule, and will generally reach the OH<sup>-</sup> first
- H<sup>+</sup> reacts much more readily than the undissociated acid molecule because H<sup>+</sup> is the strongest possible acid in aqueous solution (the weaker the acid, the more drastic this difference becomes)

Undissociated acid concentration being higher than the H<sup>+</sup> concentration is unlikely to justify the traditional way the ionic equations are written.

A similar argument can be made for most weak bases as well.

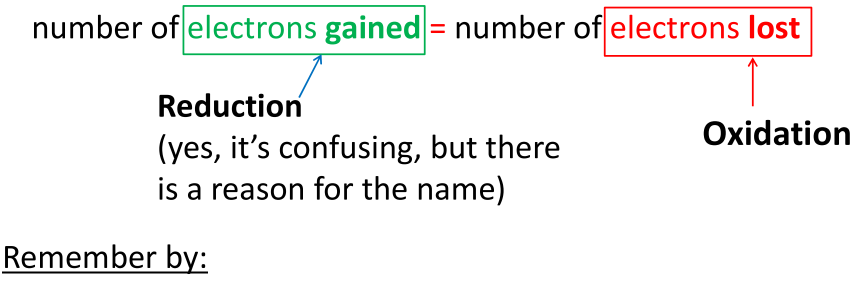
#### However:

In future courses, you might be expected to "know" to write the ionic equations for weak acid and bases in the "traditional" way.

- I will not ask you to write weak acid/base ionic equations
- But it would be wise to be aware of this tradition

#### **Redox Reactions** Oxidation-Reduction reactions

- One or more electrons are transferred.
- Electrons are not created or destroyed in chemical reactions:



Oxidation Is Loss of e<sup>-</sup>

**R**eduction Is Gain of e<sup>-</sup>

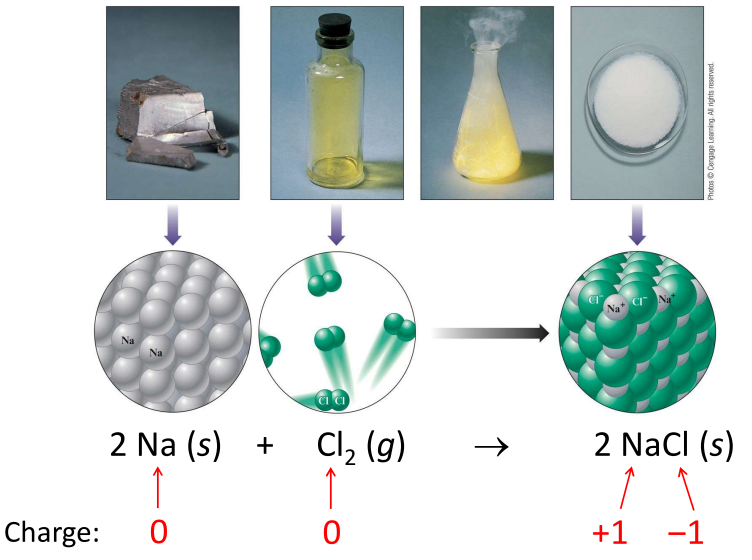
#### OIL RIG

## Originally "oxidation" referred literally to combining with oxygen

- The more oxygens combined with an atom, the more "oxidized" it was
- Oxygen was thought of as "oxide" in its compounds (with its –2 charge)
- So, the more oxygens bonded to an atom, the more positive it was
  - To have a neutral compound
- Oxidation was then generalized to mean an increase in "charge" (and not necessarily bonding to oxygen)
- And charge itself was generalized to the more abstract "oxidation state" (which is often not an actual charge)

**Redox Reactions** 

#### A simple redox reaction: Reaction of Sodium and Chlorine



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## We keep track of electron gain and loss using Oxidation numbers (Oxidation states)

- They are like charges, but not exactly
- Oxidation numbers are the charges that would exist if the atoms always formed ionic bonds, with the electron ending up on the atom that likes them more.
   e.g. oxygen likes electrons more than hydrogen
- Oxidation numbers are often not actual charges
   -- when the bonds the atom makes is not ionic
  - -- But charges on ions are real
- Even when they are not actual charges, the net change in oxidation states does determine the net number of electrons transferred

#### **Hierarchy of Rules for Assigning Oxidation States**

(1) The oxidation state of an atom in a free element is 0.

(2) The oxidation state of a monoatomic ion is equal to its charge.

(3) The sum of the oxidation states of all atoms in

(3a) a neutral molecule or formula unit is 0

(3b) a polyatomic ion is equal to the charge of the ion

Sum of oxidation states in an entity = net charge

(4) In their compounds,

- (4a) **Group I** metals have an oxidation state of +1
- (4b) **Group II** metals have an oxidation state of +2
- (5) In their compounds, nonmetals are assigned oxidation states according to a hierarchical table. Elements higher in the table have priority over lower entries.

"Oxygen is -2" can be overridden by a "higher" rule like "Fluorine is -1"

Fluorine	-1
Hydrogen	+1
Oxygen	-2
Group 7A	-1
Group 6A	-2
Group 5A	-3

#### **Examples on monatomic species**

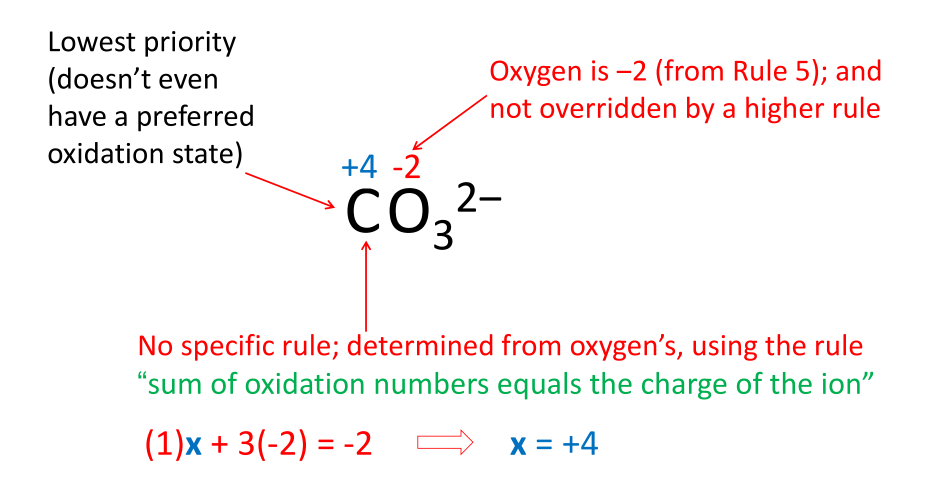
- Cu(s) Oxidation state = 0
- $H_2(g)$  Oxidation state = 0
- $H_2(I)$  Oxidation state = 0
- H (g) Oxidation state = 0
- H<sup>+</sup>(aq) Oxidation state = +1
- $S_8(s)$  Oxidation state = 0
- $S^{2-}(s)$  Oxidation state = -2
- Fe<sup>3+</sup>(aq) Oxidation state = +3

## Finding the oxidation states of elements in a compound or polyatomic ion

- 1. If you recognize a polyatomic ion, treat it separately first
- 2. Find the lowest priority element in compound or ion
  - Deal with it last
  - Its oxidation state will be determined by higher-priority rules
- 3. Assign other elements their preferred oxidations states
- Determine the oxidation state of the last (lowest priority) element using the oxidation states of other elements and Rule 3
  - It might not agree with its "default" oxidation state (e.g. oxygen might end up with -1 or +2 instead of the "default" -2). That's completely ok.

**Redox Reactions** 

Find the oxidation states for each of the elements in:



Find the oxidation states for each of the elements in  $K_2Cr_2O_7$ Suppose we didn't recognize the  $Cr_2O_7$  part as the  $Cr_2O_7^{2-}$  ion and we looked at the entire formula

The oxidation state of a monatomic ion is equal to its charge (Rule 2); Group 1A metals are always +1 in compounds

Lowest priority (doesn't even have a preferred oxidation state) - Oxygen is -2 (from Rule 5); and not overridden by a higher rule

No specific rule; determined from others, using the rule "sum of oxidation numbers in a compound equals zero"  $(2)(+1) + (2)x + 7(-2) = 0 \implies x = +6$ 

We got lucky because K has a high-priority oxidation state

Oxygen is -2

Find the oxidation states for each of the elements in  $K_2Cr_2O_7$ Suppose we recognized the  $Cr_2O_7$  part as the  $Cr_2O_7^{2-}$  ion

Lowest priority (doesn't even have a preferred oxidation state) determined from oxygen and"sum of oxidation numbers inan ion equals ion charge"+6 -2 -2 (from Rule 5);and notoverridden by ahigher rule

 $(2)x + 7(-2) = -2 \implies x = +6$ 

Then we would have determined the oxidation state of the metal as we always did. Its charge needs to neutralize the anion(s) in the formula. But we already know K is always +1.

$$^{+1} + ^{+6} - ^{-2} K_2 Cr_2 O_7$$

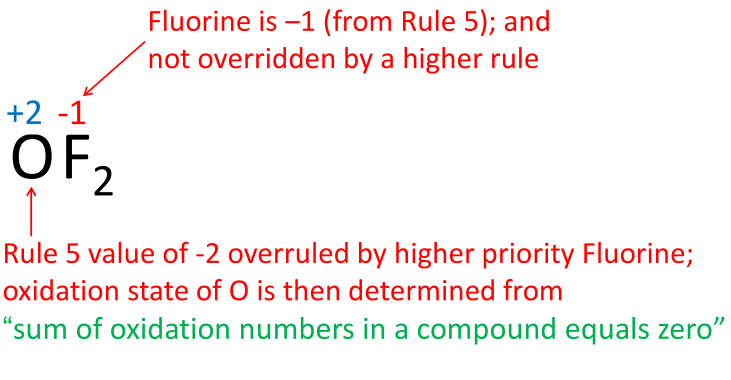
#### Find the oxidation states for each of the elements in CuSO<sub>4</sub>

Here we must recognize  $SO_4$  as  $SO_4^{2-}$ . Otherwise we either can't determine both Cu and S, or worse, we assign -2 to S as "higher priority" than Cu (since there is a rule for S, but there isn't one for Cu), and end up assigning +10 to Cu!

"sum of oxidation numbers in an ion equals ion charge"  $\mathbf{x} + 4(-2) = -2 \implies \mathbf{x} = +6$  Oxygen = -2 as highest priority element here

Because it needs to neutralize the 2- charge of the sulfate. We could do this before learning about oxidation numbers! +2 +6 -2 Cu SO<sub>4</sub>

#### Find the oxidation states for each of the elements in:



 $(1)x + 2(-1) = 0 \implies x = +2$ 

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Find the oxidation states for each of the elements in:

Group 1 metals: +1 in their compounds (Part of Rule 4) +1 -1  $LiH \leftarrow$  Lower priority; its preference is part of Rule 5

Rule 5 value of +1 overruled by higher priority Rule 4 and Rule 2: "sum of oxidation numbers in a compound equals zero"

$$(1)(+1) + (1)x = 0 \implies x = -1$$

## Re-stating "oxidation" and "reduction" in terms of oxidation state (instead of electrons)

**Oxidation**: **increase** in oxidation state (loss of electrons)

**Reduction**: decrease in oxidation state (gain of electrons)

#### **Oxidation and reduction are coupled**

- Electrons lost by one atom must be gained by another.
- Electrons are not lost; they are transferred

Number of electrons lost = Number of electrons gained

- If there is an atom getting oxidized, there must be at least one other atom getting reduced
- If there is an atom getting reduced, there must be at least one other atom getting oxidized

#### **Oxidation and reduction are coupled**

#### **Oxidized** substance: reducing agent

If something is oxidized, it must be reducing something else

#### Reduced substance: oxidizing agent

If something is reduced, it must be oxidizing something else

#### And remember:

- "oxidizing agent" and "reducing agent" are always on the reactant side!
- They cannot be products!

#### An oxidizing agent (oxidant)

- causes another substance to be oxidized
- contains an element whose oxidation state *decreases* in a redox reaction
- gains electrons (electrons are found on the left side of its half-equation)
- is reduced

#### A reducing agent (reductant)

- causes another substance to be reduced
- contains an element whose oxidation state *increases* in a redox reaction
- Ioses electrons (electrons are found on the right side of its half-equation)
- > is oxidized

In a redox reaction:

- What we said about electrons applies equally to oxidation states.
- An increase in the oxidation state of an atom is coupled with a decrease in the oxidation state of another atom

#### It's a redox reaction if ...

## The oxidation state of any element changes in the reaction

Remember: If there is an element that changed its oxidation state, there must be another that also changed in the opposite direction

## It's <u>probably</u> a redox reaction if there is an elemental substance such as $O_2(g)$ , Cu(s), etc., anywhere in the reaction

- Oxidation state is zero for an elemental substance
- In a <u>compound</u> atoms normally have nonzero oxidation states.
- If the elemental substance like  $O_2(g)$ ,  $H_2(g)$ ,  $S_8(s)$ , Fe(s) appears on either side of a reaction, it must occur in a compound on the other side of the reaction.

There are exceptions to:

"It's <u>probably</u> a redox reaction if there is an elemental substance such as  $O_2(g)$ , Cu(s), etc., anywhere in the reaction"

*Exception*:

When the element is in an "elemental state" on both sides of the reaction. For example, there is no redox here:  $O_2(g) + O(g) \rightarrow O_3(g)$ 

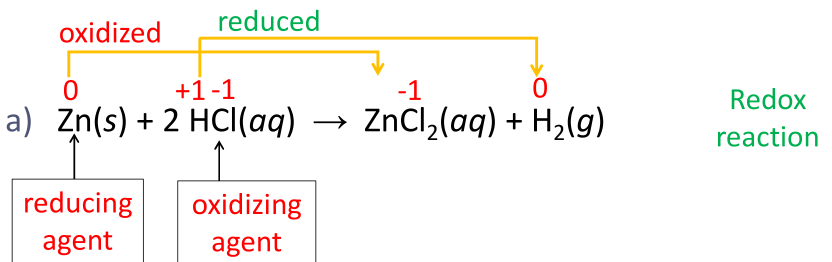
#### *Exception*:

When the element happens to have an average of zero oxidation state in a compound. For example, there is no redox here:

 $C_6H_{12}O_6(s) \rightarrow C(s) + H_2O(g)$ 

#### Example

Which of the following are oxidation-reduction reactions? Identify the oxidizing agent and the reducing agent. Remember: they are always <u>reactants</u>, <u>never products</u>



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#### **Recap:**

#### Oxidation

- O.S. of some element *increases* in the reaction.
- Electrons are on the right of the equation

#### Reduction

- O.S. of some element *decreases* in the reaction.
- Electrons are on the left of the equation.

#### **Balancing Oxidation-Reduction Equations**

- We can't count on "balancing by inspection" or even the more general procedure we learned earlier.
- We need to use a method specialized to redox reactions.
- Especially because we will be expected to bring in  $H^+$ ,  $OH^-$ , and  $H_2O$  into the reaction equation even if they are not initially in the given equation, as we will normally consider redox reactions in aqueous solution.

#### **Balancing Oxidation-Reduction Equations**

The following equations are all balanced in terms of atoms and charges, but only one is balanced in terms of the number of electrons lost and gained.

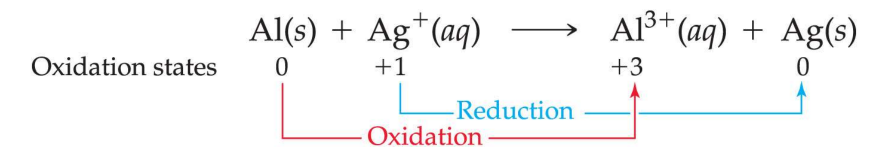
# $2 \operatorname{MnO_4}^- + \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 3 \operatorname{O_2} + 4 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 3 \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 4 \operatorname{O_2} + 6 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 5 \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 5 \operatorname{O_2} + 8 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 7 \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 7 \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 7 \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 7 \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 7 \operatorname{H_2O_2} + 6 \operatorname{H^+} \longrightarrow 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 7 \operatorname{H_2O_2} \times 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{MnO_4}^- + 7 \operatorname{H_2O_2} \times 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{Mn^{2+}} \times 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} + 10 \operatorname{H_2O} \times 2 \operatorname{Mn^{2+}} \times 2 \operatorname{Mn^{2+}} + 6 \operatorname{O_2} \times 2 \operatorname{Mn^{2+}} \times 2$

#### A General Procedure for Balancing Redox Reactions Using the Half-Reaction Method

- 1. Assign oxidation states.
- 2. Separate the overall reaction into two half-reactions.
- 3. Balance each half-reaction with respect to number of atoms of each element.
- 4. Balance each half-reaction with respect to charge by adding electrons.
- 5. Make the number of electrons lost/gained in both halfreactions equal by multiplying.
- 6. Add the two half-reactions together.
- 7. Verify that the reaction is balanced.

#### **Balancing Redox Equations**

 Assign oxidation numbers to all atoms to determine what is being oxidized and what is being reduced.



• Divide the reaction into two half-reactions, one for oxidation and one for reduction.

Oxidation:  $Al(s) \longrightarrow Al^{3+}(aq)$ Reduction:  $Ag^+(aq) \longrightarrow Ag(s)$ 

- In this case, the half-reactions are already balanced with respect to mass—the number of each type of atom on both sides of each half-reaction is the same.
- However, the equations are not balanced with respect to charge.
- We balance the charge of each half-reaction individually by adding the appropriate number of electrons to make the charges on both sides equal.

$$Al(s) \longrightarrow Al^{3+}(aq) + 3e^{-}$$
$$1e^{-} + Ag^{+}(aq) \longrightarrow Ag(s)$$

(zero charge on both sides) (zero charge on both sides)

- Since these half-reactions must occur together, the number of electrons lost in the oxidation halfreaction must equal the number gained in the reduction half-reaction.
- We equalize these by multiplying one or both halfreactions by appropriate whole numbers to equalize the electrons lost and gained.

$$Al(s) \longrightarrow Al^{3+}(aq) + 3e^{-}$$
$$3 \times [1e^{-} + Ag^{+}(aq) \longrightarrow Ag(s)]$$

• Add the half-reactions together, canceling electrons and other species as necessary.

$$Al(s) \longrightarrow Al^{3+}(aq) + 3e^{-}$$
$$3e^{-} + 3Ag^{+}(aq) \longrightarrow 3Ag(s)$$
$$Al(s) + 3Ag^{+}(aq) \longrightarrow Al^{3+}(aq) + 3Ag(s)$$

• Verify that the equation is balanced, with respect to both mass and charge.

$$Al(s) \longrightarrow Al^{3+}(aq) + 3e^{-}$$

$$3e^{-} + 3Ag^{+}(aq) \longrightarrow 3Ag(s)$$

$$Al(s) + 3Ag^{+}(aq) \longrightarrow Al^{3+}(aq) + 3Ag(s)$$

Reactants	Products
1 Al	1 Al
3 Ag	3 Ag
+3 charge	+3 charge

 Notice that the charge need not be zero on both sides of the equation—it just has to be *equal* on both sides. The equation is balanced.

# Balancing Redox Reactions in Aqueous Solutions

Balancing redox reactions <u>in aqueous solution</u> might look like alchemy where the initial equation might be missing an element or two (or more) on one side of the equation or the other!

You might feel like you are doing things we are not supposed to do in balancing, such as adding new reactants or products to the equation. That's because aqueous-solution redox reactions are often given in "bare-bones" form.

- We are expected to "fill in the blanks" and create a complete equation while balancing it.
- Actually, it's easier to start balancing without the species that are not involved in redox and then add them later. We just need to get used to it.

Balancing Redox Reactions in Aqueous Solutions -- "ionic" or "net ionic" equations

The "bare-bones" (i.e. skeletal) reactions are usually given in "net ionic" form

Some authors seem to use the term "ionic equation" to mean "net ionic equation"; so beware!

"ionic equation" and "net ionic equation" are analogous to those for double-displacement (metathesis) reactions:

- Ionic equation: include the "counter-ions" that would allow you to combine ions into neutral chemical formulas
- Net-ionic equation: do not include spectator ions (those "counter-ions" of ions that are actually involved in redox

# Balancing Redox Reactions in <u>Acidic</u> Conditions

- Write both half reactions (oxidation & reduction)
- Balance all atoms other than O and H
- Balance O by adding  $H_2O$
- Balance H by adding H<sup>+</sup>
- Balance charge by adding e<sup>-</sup>
- Multiply both half reactions by a factor to equate no of  $e^{\scriptscriptstyle -}$
- Add the two half-reactions and cancel out the e<sup>-'</sup>s and any substances appearing on both side
- Double check: ✓ All elements balanced

✓ Charges balanced

#### **EXAMPLE / PRACTICE**

Balance the redox reaction  $Fe^{2+}(aq) + MnO_4^{-}(aq) \rightarrow Fe^{3+}(aq) + Mn^{2+}(aq)$ in acidic solution, using the half-reaction method

**1.Assign oxidation states** to all atoms and identify the substances being oxidized and reduced. 2. Separate the overall reaction into two OXIDATION  $Fe^{2+}(aq) \rightarrow Fe^{3+}(aq)$ half-reactions, one for oxidation and one for reduction. **REDUCTION**  $MnO_4^{-}(aq) \rightarrow Mn^{2+}(aq)$ **3.**Balance each half-reaction with respect to **mass** in the following order: All elements other than hydrogen and oxygen are • Balance all elements other than H and O. balanced, so you can proceed to the next step.  $Fe^{2+}(aq) \rightarrow Fe^{3+}(aq)$ • Balance O by adding H<sub>2</sub>O.  $MnO_4^{-}(aq) \rightarrow Mn^{2+}(aq) + 4 H_2O(l)$ 

Balance the redox reaction  $Fe^{2+}(aq) + MnO_4^{-}(aq) \rightarrow Fe^{3+}(aq) + Mn^{2+}(aq)$ in acidic solution, using the half-reaction method (continued)

**4.Balance each half-reaction with respect to charge** by adding electrons to the right side of the oxidation halfreaction and the left side of the re-(The sum of the charges on both s should be equal.)

5. Make the number of electrons in both half-reactions equal by multiplying one or both half- reactions by a small whole number.

6.Add the two half-reactions together, canceling electrons and other species as necessary.

• Balance H by adding H<sup>+</sup>.

 $5 \operatorname{Fe}^{2+}(aq) \longrightarrow 5 \operatorname{Fe}^{3+}(aq) + 5e^{2}$  $5e^{-} + 8 \operatorname{H}^{+}(aq) + \operatorname{MnO}_{4}^{-}(aq) \longrightarrow \operatorname{Mn}^{2+}(aq) + 4 \operatorname{H}_{2}O(l)$  $5 \text{ Fe}^{2+}(aq) + 8 \text{ H}^{+}(aq) + \text{MnO}_{4}^{-}(aq) \longrightarrow 5 \text{ Fe}^{3+}(aq) + \text{Mn}^{2+}(aq) + 4 \text{ H}_{2}O(l)$ 

 $5 \times [Fe^{2+}(aq) \rightarrow Fe^{3+}(aq) + le^{-}]$  $5e^{-} + 8 H^{+}(aq) + MnO_{4}^{-}(aq) \rightarrow Mn^{2+}(aq) + 4 H_{2}O(l)$ 

$$Fe^{2+}(aq) \rightarrow Fe^{3+}(aq) + 1e^{-}$$
  
5e<sup>-</sup> + 8 H<sup>+</sup>(aq) + MnO<sub>4</sub><sup>-</sup>(aq)  $\rightarrow$  Mn<sup>2+</sup>(aq) + 4 H<sub>2</sub>O(*l*)

$$\operatorname{Fe}^{2+}(aq) \longrightarrow \operatorname{Fe}^{3+}(aq) + \mathbf{1e}^{-}$$

 $Fe^{2+}(aq) \rightarrow Fe^{3+}(aq)$ 

8  $H^+(aq)$  + MnO<sub>4</sub><sup>-(aq)</sup>  $\rightarrow$  Mn<sup>2+(aq)</sup> + 4  $H_2O(l)$ 

Balance the redox reaction  $Fe^{2+}(aq) + MnO_4^{-}(aq) \rightarrow Fe^{3+}(aq) + Mn^{2+}(aq)$ in acidic solution, using the half-reaction method (*continued*)

7.Verify that the reaction is balanced
with respect to both mass* and charge.

Reactants	Products	
5 Fe	5 Fe	
8 H	8 H	
1 Mn	1 Mn	
4 O	4 O	
+17 charge	+17 charge	

\* Not literally "mass". It just means the number of atoms for each element

# Balancing Redox Reactions in **Basic** Conditions

- Write both half reactions
- Balance all atoms other than O and H
- Balance O by adding H<sub>2</sub>O
- Balance H by adding H<sup>+</sup>
- Balance charge by adding e<sup>-</sup>
- Multiply both half reactions by a factor to equate no of e<sup>-</sup>
- Add the two half-reactions and cancel out the electrons
- "Neutralize" H<sup>+</sup> by adding same no. of OH<sup>-</sup> to both sides
- Combine H<sup>+</sup> and OH<sup>-</sup> on the same side to make H<sub>2</sub>O
- Cancel appropriate no. of  $H_2O$  (and any other species) appearing on both sides, so that the remainder appears on only one side
- Double check: ✓ All elements balanced

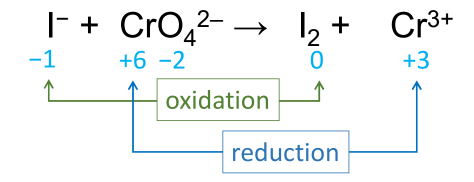
✓ Charges balanced

## Start like acidic

## Practice – Balancing redox reactions, Basic Conditions

Balance the reaction equation  $I^- + CrO_4^{2-} \rightarrow I_2 + Cr^{3+}$ 

assign oxidation states and determine element oxidized and element reduced



separate into oxidation & reduction half-reactions

oxidation:  $I^- \rightarrow I_2$ reduction:  $CrO_4^{2-} \rightarrow Cr^{3+}$  Balance the reaction equation in basic solution  $I^- + CrO_4^{2-} \rightarrow I_2 + Cr^{3+}$  (continued)

•

 balance atoms other than O and H oxidation:  $I^- \rightarrow I_2$  $2 \mid \rightarrow \mid_2$ reduction:  $CrO_4^{2-} \rightarrow Cr^{3+}$  $CrO_4^{2-} \rightarrow Cr^{3+} + 4H_2O$  balance O by adding H<sub>2</sub>O to side that lacks O balance H by adding H<sup>+</sup> to side that lacks H  $CrO_4^{2-} + 8H^+ \rightarrow Cr^{3+} + 4H_2O$  balance each half-reaction with respect to charge by adding electrons on product side for  $2I^- \rightarrow I_2 + 2e^$ oxidation adding electrons on reactant side for  $CrO_{a}^{2-} + 8H^{+} + 3e^{-} \rightarrow Cr^{3+} + 4H_{2}O$ reduction

Balance the reaction equation in basic solution  $I^- + CrO_4^{2-} \rightarrow I_2 + Cr^{3+}$  (continued)

Multiply, if needed, each halfreaction by an appropriate factor so that both involve the same number of electrons (and they are the smallest possible integers)

$$(2I^{-} \rightarrow I_{2} + 2e^{-}) \times 3$$
  
 $(CrO_{4}^{2-} + 8H^{+} + 3e^{-} \rightarrow Cr^{3+} + 4H_{2}O) \times 2$   
 $\downarrow$   
 $6I^{-} \rightarrow 3I_{2} + 6e^{-}$   
 $2CrO_{4}^{2-} + 16H^{+} + 6e^{-} \rightarrow 2Cr^{3+} + 8H_{2}O$ 

Add the half reactions and cancel electrons

$$2CrO_4^{2-} + 16H^+ + 6I^- + 6e^- \rightarrow 2Cr^{3+} + 8H_2O + 3I_2 + 6e^-$$

$$2CrO_4^{2-} + 16H^+ + 6I^- \rightarrow 2Cr^{3+} + 8H_2O + 3I_2$$

Balance the reaction equation in basic solution  $I^- + CrO_a^{2-} \rightarrow I_2 + Cr^{3+}$ (continued)

So far it has been the same as for balancing in acidic solution

Now add enough OH<sup>-</sup> to neutralize H<sup>+</sup> to produce  $H_2O$ , and the same number of OH<sup>-</sup> to the other side

Cancel  $H_2O$  (and any other species appearing on both sides at this point)

 $2CrO_4^{2-} + 16H^+ + 16OH^- + 6I^- \rightarrow 2Cr^{3+} + 8H_2O + 3I_2 + 16OH^ 16H_{2}O$  $2CrO_4^{2-} + 16H_2O + 6I^- \rightarrow 2Cr^{3+} + 8H_2O 3I_2 + 16OH^-$ 

 $8^{2}$  2CrO<sub>4</sub><sup>2-</sup> + 16H<sub>2</sub>O + 6l<sup>-</sup>  $\rightarrow$  2Cr<sup>3+</sup> + 8H<sub>2</sub>O + 3l<sub>2</sub> + 16OH<sup>-</sup>

**Balanced in basic solution**  $2CrO_4^{2-} + 8H_2O + 6I^- \rightarrow 2Cr^{3+} + 3I_2 + 16OH^-$ 

Balance the reaction equation in basic solution  $I^- + CrO_4^{2-} \rightarrow I_2 + Cr^{3+}$  (continued)

Finally, make sure that all the elements as well as charges are balanced:

```
2CrO_4^{2-} + 8H_2O + 6I^- \rightarrow 2Cr^{3+} + 3I_2 + 16OH^-
```

reactant side	Element	product side
2	Cr	2
6	I	6
16	0	16
16	Н	16
-10	charge	-10

#### **Solving Stoichiometry Problems for Reactions in Solution**

- 1. Identify the species present in the combined solution, and determine what reaction occurs.
- 2. Write the balanced equation (net ionic or "molecular", depending on which seems more convenient) for the reaction.
- 3. Calculate the moles of reactants.
- 4. Determine which reactant is limiting.
- 5. Calculate the moles of product(s), as required.
- 6. Convert to grams or other units, as required.

Not really different from general stoichiometry questions!

We just use volumes and concentrations to find moles (or vice versa)

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Any reaction that can occur in solution can be the subject of solution stoichiometry, such as:

Acid-base neutralization reactions

**Redox reactions** 

**Precipitation reactions** 

To do anything useful, we must know:

- The nature of the reaction.
- The amounts of chemicals present in the solutions.
   And to know the amount of a chemical present in solution, we need to know the volume of the solution and its concentration

#### **Performing Calculations for Acid–Base Reactions**

- No different from other reactions, really.
- Often we do "titrations"
- In a titration there is no limiting reactant, by definition
- Balancing is easy because H<sup>+</sup> and OH<sup>-</sup> must come in equal moles
  - we can even do the stoichiometry without writing the reaction
  - but don't feel you must be able to

## **Acid–Base Titrations**

**Titration:** delivery of a measured volume of a solution of known concentration (the titrant) into a solution containing the substance being analyzed (the analyte).

*Equivalence point:* exactly enough titrant added to react completely with the analyte.

> The ideal we try to achieve at the end of titration

**Endpoint:** the indicator changes color so you can tell the equivalence point has been reached. It's an experimental approximation of the equivalence point.

What we can achieve in practice, with our indicator

## There is no limiting or excess reactant in titrations -- by design

#### Practice

For the titration of sulfuric acid  $(H_2SO_4)$  with sodium hydroxide (NaOH), how many mL of 0.1054 *M* sodium hydroxide would be required to react with 1.00 L of 0.500 *M* sulfuric acid to reach the endpoint?

$$H_2SO_4(aq) + 2 NaOH(aq) \rightarrow Na_2SO_4(aq) + 2 H_2O(aq)$$

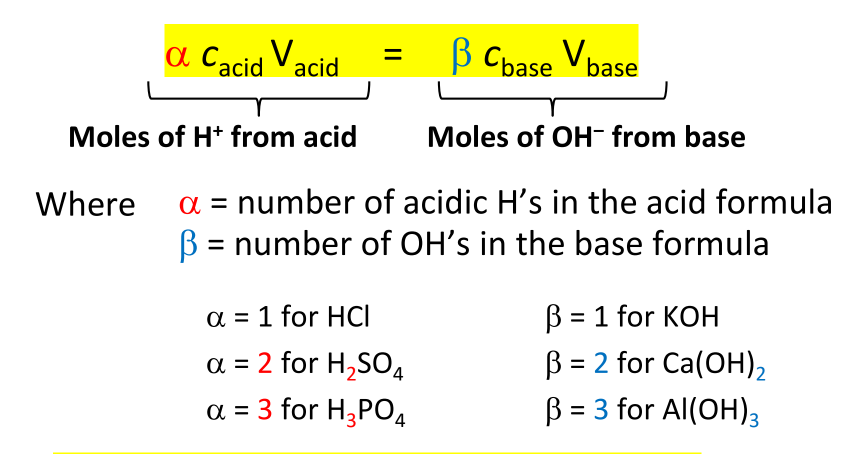
$$n_{H_2SO_4} = c_{H_2SO_4} V_{H_2SO_4} = (0.500)(1.00) = 0.500 \text{ mol } H_2SO_4$$
$$n_{NaOH} = 0.500 \text{ mol } H_2SO_4 \frac{2 \text{ mol } NaOH}{1 \text{ mol } H_2SO_4} = 1.00 \text{ mol } NaOH$$

 $\mathbf{n}_{_{\mathrm{NaOH}}} = c_{_{\mathrm{NaOH}}} \mathbf{V}_{_{\mathrm{NaOH}}}$ 

 $1.00 \text{ mol} = (0.1054 \text{ mol/L}) V_{NaOH}$ 

$$V_{NaOH} = 9.49 L = 9.49 x 10^3 m L$$

When the acid and base are used in stoichiometric amounts, such as during titrations, you can use the following formula:



Don't confuse it with the dilution formula!

Then the previous example:

For the titration of sulfuric acid  $(H_2SO_4)$  with sodium hydroxide (NaOH), how many mL of 0.1054 *M* sodium hydroxide would be required to react with 1.00 L of 0.500 *M* sulfuric acid to reach the endpoint?

could be solved this way:

 $\alpha c_{acid} V_{acid} = \beta c_{base} V_{base}$ (2) (0.500) (1.00) = (1) (0.1054) V\_{NaOH}

 $V_{NaOH} = 9.49 L = 9.49 x 10^3 mL$ 

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Acid-base neutralization and redox reactions are often used in titrations where, by design, there is no limiting or excess reactant.

Titration with a redox reaction (or any reaction) is in principle similar to that with acid-base reaction.

With precipitation reactions, we often have a limiting reactant – therefore also excess reactant(s).

We don't often see precipitation reactions in the context of titration.

## **Practice**

10.0 mL of a 0.30 *M* sodium phosphate solution reacts with 20.0 mL of a 0.20 *M* lead(II) nitrate solution (assume volumes add up perfectly). Will there be a reaction? If so, what is the product?

With two solutions of ionic compounds, typically we think of a double-displacement reaction of the "precipitation" kind.

Will there be a precipitate? **If not**, the ions would still be floating around just as before the two solutions were mixed, and there would be **no reaction**.

sodium phosphate: Na<sub>3</sub>PO<sub>4</sub> (aq) exists as 
$$3 \text{ Na}^+(aq) + PO_4^{3-}(aq)$$
  
insoluble together  
lead (II) nitrate: Pb(NO<sub>3</sub>)<sub>2</sub> (aq) exists as  $Pb^{2+}(aq) + 2 \text{ NO}_3^-(aq)$   
"Molecular" (or "formula") equation:  
 $2 \text{ Na_3}PO_4(aq) + 3 \text{ Pb}(\text{NO}_3)_2(aq) \rightarrow Pb_3(PO_4)_2(s) + 6 \text{ NaNO}_3(aq)$ 

precipitate

## Practice

10.0 mL of a 0.30 *M* sodium phosphate solution reacts with 20.0 mL of a 0.20 *M* lead(II) nitrate solution (assume volumes add up perfectly). What mass of precipitate will form?

