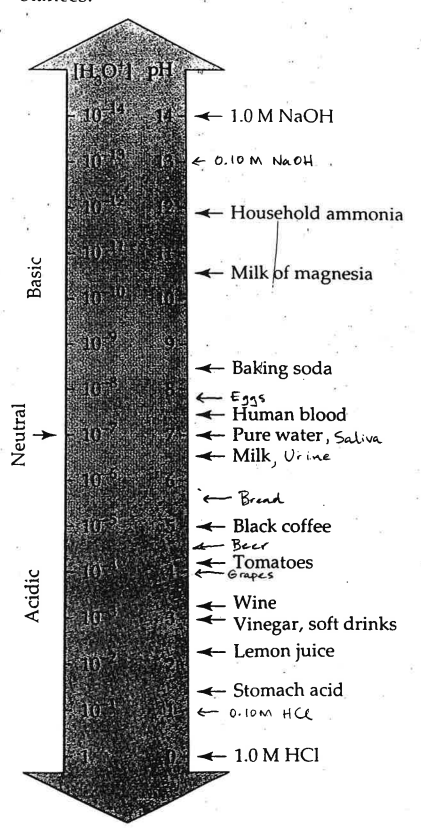
Chemistry 12

Unit IV – Acid/Base I

FIGURE 15.2 The pH scale and pH values for some common substances.



I) Properties of Acids and Bases <u>Acids</u> 1... 2. 3. 4. 5. 6. <u>Bases</u> 1. 2. 3. 4.

5.

6.

2

II) Arrhenius Acids and Bases (1903)

Arrhenius Acid:

Arrhenius Base:

Salt: the neutralization product that results when a strong acid and a strong base react – made up of a _____ and a _____.

e.g.
$$HCl_{(aq)} + NaOH_{(aq)} \Rightarrow H_2O_{(l)} + NaCl_{(aq)}$$

 $str.acid + str.base = water + salt$

e.g.
$$H_2SO_{4(aq)} + KOH_{(aq)} \Rightarrow$$

str. acid + str. base =

Net Ionic Equation for a *strong acid/strong base* neutralization reaction: (net ionic equations display the true chemistry that is occurring – *spectator ions* are not included)

Assignment 1: Hebden p.110-112, Qs #1-4
Read pgs. 112-114 (Think about Qs 5-9 p.114)

III) Bronsted-Lowry Acids and Bases (1923)

Bronsted-Lowry Acid:

Bronsted-Lowry Base:

Assignment 2: Label the acid reactant and the base reactant

- 1. $HCl + H₂O \Rightarrow H₃O⁺ + Cl⁻$
- 2. $NH_3 + H_2O \Leftrightarrow NH_4^+ + OH^-$
- 3. $CO_3^{2-} + H_2O \Leftrightarrow HCO_3^{-} + OH^{-}$

$$\begin{cases} 4. \text{ HPO}_{4^{2^{-}}} + \text{HBr} \Rightarrow \text{H}_{2}\text{PO}_{4^{-}} + \text{Br} \\ \\ 5. \text{ HPO}_{4^{2^{-}}} + \text{HS}^{-} \Leftrightarrow \text{PO}_{4^{3^{-}}} + \text{H}_{2}\text{S} \end{cases}$$

- 6. $H_2PO_4^- + HF \Leftrightarrow H_3PO_4 + F^-$
- 7. $H_2PO_4^- + H_2O \iff HPO_4^{2-} + H_3O^+$

Also do Hebden p.117 Q #11

IV) The Acid/Base Table

- acids are listed on the left from top (strongest) to bottom (weakest)
- bases are listed on the right from bottom (strongest) to top (weakest)
- the separate region at the top represents the **strong** acids (there are SIX), meaning they dissociate 100% (one-way arrow no *equilibrium*)
- the separate region at the bottom represents the **strong** bases, meaning they dissociate 100% as well.
- Alkali metal hydroxides (eg. NaOH) and Alkaline Earth metal hydroxides (eg. Ca(OH)₂) are also strong bases, they are simply not listed on the table.
- Alkali metal hydroxides are very soluble salts (bases), whereas Alkaline Earth metal hydroxides are low soluble (except for Sr(OH)2). Thus, even though Alkaline Earth metal hydroxides are deemed to be strong bases, their low solubility disallows them from generating a relatively high pH.
- Transition metal hydroxides, along with Aluminum hydroxide, are **weak** bases (Fe³⁺, Cr³⁺, and Al³⁺ shown on the table in the middle section).

- the section in the middle of the table represents the weak acids (left) and the weak bases (right). These substances form an equilibrium with water as their dissociation is defined as being <50% (most weak acids/bases, in the Chemistry 12 context, dissociate < 5%).
- therefore, weak acids are poorer conductors of electricity than strong acids (ie. they create less ions in solution than strong acids).

*same with weak bases.

V) H⁺ and H₃O⁺

H⁺ is simply a proton. Why?

When H⁺ is created in water (due to the addition of acid), it immediately reacts with a water molecule to form the ______ ion. Remember, a proton is an extremely concentrated positive charge and water's oxygen atom is partially negative in charge.

$$H^+ + H_2O \Rightarrow$$

proton water hydronium ion

*thus, _____ do NOT exist in water!

Because this reaction is instantaneous, H⁺ and H₃O⁺ are used interchangeably (H⁺ is used primarily for shorthand). The dissociation of acid in water can be shown with or without water in the chemical equation:

i)
$$HCl_{(aq)} \Rightarrow H^+ + Cl^-$$

ii)
$$HCl + H2O(1) \Rightarrow H3O+ + Cl-$$

All of the reactions on the acid/base table are written in shorthand form. Each can be converted to type ii above by adding $H_2O_{(1)}$ to the left side and changing H^+ to H_3O^+ on the right side.

Change three equations from your table to include water.						
An ACID can only act as an acid if it is in the presence of a						
A BASE can only act as a base if it is in the presence of an						
Water can act as either an acid or a base (it is amphiprotic in its nature).						
VI) Further Definitions within Bronsted-Lowry Theory						
Monoprotic Acid ():						
Diprotic Acid ():						
Polyprotic Acid ():						
J I (———————————————————————————————————						
Amphiprotic substance:						
Ampriliprotic substance.						
Look back at questions #4 and #5 in Assignment 2. What do you notice						
about HPO ₄ ²⁻ and its behaviour in either question?						
What can be concluded, then, about HPO ₄ ² -?						

H₂O is amphiprotic as well, but it's a special case; see #3 of Assignment 3.

Generally, amphiprotic substances (except for H₂O) are polyatomic groups that contain at least one proton and are negatively charged.

Exceptions to this rule?

How can you tell if a substance is amphiprotic using the acid/base table?

RULE: In a reaction between two amphiprotic substances, the stronger acid (higher on the left side of the table) will act as the acid and be the proton donor.

Assignment 3: Determine the products

2.
$$HPO_4^{2-} + HSO_4^{-} \Leftrightarrow$$

3.
$$H_2O + HSO_3$$
 \Leftrightarrow

#3: Water is a special-case amphiprotic as it is the _____ weak acid <u>AND</u> the ____ weak base. Thus, any other amphiprotic species will *hydrolyze* both acidically AND basically in water, but one will predominate.

Also do Q12 p.117 and Qs 13-14 p.119.

VII) Conjugate Acid-Base Pairs

Conjugate acid/base pairs are molecules or ions that differ only by a

A base has one ______ proton than its conjugate acid, and an acid has one _____ proton than its conjugate base.

Remember to **adjust the charge** when writing the conjugate. (Gain proton => less - / more +; Lose proton => more - / less +)

What is the conjugate base of each of the following acids?

HClO₄ / PH₄+ /
CH₃COOH / HNO₂ /
H₂CO₃ / HC₂O₄- /

What is the conjugate acid of each of the following bases?

CN- / H2PO3- / F- / NH3 / HCOO- /

Acid/Base Equations

Complete the following equation and identify the conjugate acid-base pairs.

$$HCO_3^- + H_2PO_4^- \Leftrightarrow$$

Assignment 4: Complete the equation and identify conjugate acid-base pairs

1.
$$HSO_4$$
 + $H_2O \Leftrightarrow$

2.
$$H_2O + HNO_2 \Leftrightarrow$$

3.
$$HIO_3 + NH_3 \Leftrightarrow$$

5.
$$HS^- + H_3PO_4 \Leftrightarrow$$

6.
$$HCO_3^- + CN^- \Leftrightarrow$$

7.
$$H_3BO_3 + HO_2$$
 \Leftrightarrow

8.
$$C_2O_4^{2-} + H_2O \Leftrightarrow$$

9.
$$H_2O + H_2SO_3 \Leftrightarrow$$

Assignment 5: Hebden p. 121 #16-19

VIII) Strong and Weak Acids and Bases

Strong acids and bases are ______ ionized (dissociated) in solution. Thus, they react to completion and do not form an equilibrium (see table).

e.g.
$$HCl + H_2O_{(i)} \Rightarrow H_3O^+ + Cl^-$$

Weak acids and bases are not 100% ionized in solution (in fact, by definition, they dissociate <50% -- in Chem 12, the general assumption is that they dissociate <5%). They create an equilibrium with reactants heavily favoured (ie. $[R] >> [P] \rightarrow$ therefore, $K_{eq} _ 1$)

e.g.
$$HF + H_2O \Leftrightarrow H_3O^+ + F^-$$

Only weak acids and bases are in equilibrium with their conjugate pairs. Strong acids (such as HCl) have a conjugate base (Cl-), but they are unable to form an equilibrium because conjugate bases of strong acids cannot act as a base (see table). The same idea pertains to strong bases and their non-functional conjugate acids.

*It is important to mention that the terms **strong** and **weak** differ from the terms **concentrated** and **dilute**.

6M KOH is	and	·
0.0001M KOH is	and	
6M CH₃COOH is	and	·
0.0001M CH₃COOH is	and	
So, to summarize:	<i>y</i>	
In order to determine if an aci	id/base is strong or weak, one mu	ıst determine
In order to determine if the sa must know the	ame chemical is concentrated or d	lilute, one

Leveling Effect

All of the strong acids listed at the top of the acid-base table ionize (dissociate) 100% in solution. Therefore, each of the six are **level** in strength. The same goes for strong bases.

Thus, H_3O^+ is the 'strongest' acid that can exist in water (aqueous solution) because H_3O^+ is the result of adding a STRONG acid to water...similarly with bases \rightarrow OH $^-$ is the 'strongest' base that can exist in water (aqueous solution) because OH $^-$ is the result of adding a STRONG base to water.

Question: Why are 1M HCl and 1M H₂SO₄ *level* in strength if one species is monoprotic and the other is diprotic?

IX) Determining Whether Reactants or Products are Favoured in an Acid/Base Reaction

Note: All Bronsted-Lowry reactions involve the transfer of 1 proton unless one or both species are ______.

$$HCO_{3}^{-} + HF \iff H_{2}CO_{3} + F^{-}$$

There is a 'competition' between the two acids HF and H₂CO₃ to donate a proton. The stronger acid 'wins out' and will give a proton to a greater extent. Since _______ is the stronger acid in this equilibrium, the ______ reaction is favoured (in terms of possessing a lower ______) and therefore ______ are favoured (ie. at equilibrium, [P] ____ [R]).

RULE: The side of the reaction with the _____ acid is always favoured.

REASON: The stronger acid donates a proton more readily, causing the other side (weaker side) to be greater in concentration.

So...if products are favoured: $K_{eq} > 1$; if reactants favoured: $K_{eq} < 1$.

Assignment 6: Label conjugate acid-base pairs and state whether reactants or products are favoured.

- 1. $NH_4^+ + H_2O \Leftrightarrow NH_3 + H_3O^+$
- 2. $H_2S + NH_3 \Leftrightarrow HS^- + NH_4^+$
- 3. $H_2PO_4^- + HS^- \Leftrightarrow HPO_4^{2-} + H_2S$
- 4. $H_2O_2 + SO_3^{2-} \Leftrightarrow HO_{2^-} + HSO_{3^-}$
- 4. $CH_3COOH + PO_4^{3-} \Leftrightarrow CH_3COO^- + HPO_4^{2-}$
- 6. $H_2PO_4^- + C_2O_4^{2-} \Leftrightarrow HPO_4^{2-} + HC_2O_4^{-}$
- 7. $H_2SO_3 + SO_4^{2-} \Leftrightarrow HSO_3^{-} + HSO_4^{-}$

Assignment 7: Review Questions

- 1. Will the K_{eq} be greater or less than 1 for the following equilibrium? Why? $HSO_4^- + NH_3 \Leftrightarrow SO_4^{2-} + NH_4^+$
- 2. Which acid has the higher [H₃O⁺] when reacting with water, HCN or CH₃COOH? Why?

- 3. Will a reaction occur between NH₂ and C₂O₄²-? Explain why or why not.
- 4. Write an equation to show the reaction between NH₂- and water to form a basic solution (containing OH-), and explain why products are favoured.

Assignment 8: Hebden p. 125-126 #21-27, p. 133 #38-46

Approximately 1 in every 10 million water molecules dissociates at 25° C, or approx. 2 in every 20 million water molecules react with each other which is what <u>really</u> happens!

Shorthand water dissociation reaction equation:

Longhand (actual) version:

$[H_3O^+]$ = $[OH^-]$ because every reaction between two water molecules
produces one hydronium ion and one hydroxide ion. In any solution
where $[H_3O^+] = [OH^-]$, the solution is considered to be in pH.
Water reacting with an acid will produce moreions, thereby
creating an solution. Water reacting with a base will
produce moreions, thereby creating a
solution.
Regardless of the chemicals, if water is the 'arena' (ie. reaction takes place in
aqueous solution), then the Kw remains constant at 1.0 x 10 ⁻¹⁴ (since
is the only stress that can alter a Keq value).
Therefore, if one of the hydronium ion or hydroxide ion increases, the
other must decrease in concentration (by the same proportion) in order to
satisfy the Kw equation.
Water dissociation equation: $2H_2O_{(1)} + 59kJ \Leftrightarrow H_3O^+_{(aq)} + OH^{(aq)}$
Scenarios:
ADD ACID (incr. [H ₃ O ⁺]):
1071
In acid: [H ₃ O ⁺] 1.0 x 10 ⁻⁷ M and [OH ⁻] 1.0 x 10 ⁻⁷ M
* Still OH- present, since Kw equilibrium still exists even though it shifted.
(**there will NEVER be 0 M hydronium or hydroxide in an aqueous sol'n).
ADD BASE (incr. [OH-]):
T 1 TH OH 10-10-7M on 1 TOTAL 10-7M
In base: $[H_3O^+]$ 1.0 x 10 ⁻⁷ M and $[OH^-]$ 1.0 x 10 ⁻⁷ M
* Still H ₃ O ⁺ present, since K _w equilibrium still exists even though it shifted.

Assignment 9: Kw Exercises

- 1. Calculate the $[OH^-]$ in a solution in which $[H_3O^+]$ is $1.0 \times 10^{-12} M$. Is the solution neutral, acidic, or basic?
- 2. Calculate the $[H_3O^+]$ in a solution in which $[OH^-]$ is $1.0 \times 10^{-8}M$.
- 3. Calculate the [OH-] for solutions with the given [H₃O+]. Is each solution acidic, basic, or neutral?

a.
$$[H_3O^+] = 1.0 \times 10^{-3}M$$

b.
$$[H_3O^+] = 2.6 \times 10^{-10}M$$

c.
$$[H_3O^+] = 8.7 \times 10^{-7}M$$

4. Calculate the [H₃O⁺] for solutions with the given [OH⁻]. Is each solution acidic, basic, or neutral?

a.
$$[OH^{-}] = 1.0 \times 10^{-2}M$$

b.
$$[OH^-] = 3.4 \times 10^{-6} M$$

c.
$$[OH^{-}] = 9.2 \times 10^{-9} M$$

- 5. What is the $[H_3O^+]$ and $[OH^-]$ in 0.0010M HCl?
- 6. What is the $[H_3O^+]$ and $[OH^-]$ in 0.00345M NaOH?
- 7. Calculate the [H₃O⁺] and [OH⁻] in

c.
$$6.00 \times 10^{-3} M Ca(OH)_2$$

Effect of Temperature on $K_{\rm w}$

$$2H_2O_{(l)} + \ 59kJ \Leftrightarrow \ H_3O^+{}_{(aq)} \ + \ OH^-{}_{(aq)}$$

This reaction is	in the forward direction and
	in the reverse direction. Increasing the
	e equilibrium to the This
_	in $[H_3O^+]$ and $[OH^-]$, and an
	in the value of K _w . Similarly, decreasing the
temperature causes a shi	ft to the, thereby
	the $[H_3O^+]$ and $[OH^-]$, and causing K_w to
	[OH-] change due to a temperature change, they ther, so water at any temperature is still
Assignment 10: Hebden	Read pp. 126-127; Do Qs 28-29.
XI) pH (Potential Hydro	gen Ion)
pH is a measure of the [I	H^{+}] or [H_3O^{+}] in a solution using logarithms (log).
The pH scale is 0-14 at 25 possible.	5°C, however pH values below 0 and above 14 are

[H ₃ O ⁺]	>100	100	10-1	10-2	10-3	10-4	10-5	10-6	10-7	10-8	10-9	10-10	10-11	10-12	10-	10-14	<10-14
(M)															13		
pН	<0	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14	>14

Sigfigs??? See p.16 (booklet) or Hebden p.139 for explanation...

Essentially, only digits AFTER the decimal are significant in pH values.

Each unit change in pH represents a 10x change in [H₃O+] and [OH-].

Remember that $[H_3O^+]$ and $[OH^-]$ are inversely related (as one goes up, the other goes down), in order to maintain K_w . Thus, a high $[H_3O^+]$ in a solution corresponds to a low $[OH^-]$, since their product must always equal 1.0×10^{-14} at 25° C.

** When doing pH problems, assume 25° C unless otherwise stated. When $[H_3O^+] = 1.0 \times 10^{-7} M$, what is the pH and what is $[OH^-]$?

Conclusion:

If $[H_3O^+] > 1.0 \times 10^{-7}M$, such as $1.0 \times 10^{-4}M$, what is the pH? What is $[OH^-]$?

Conclusion: A low pH (less than 7) corresponds to an ______solution.

If $[H_3O^+] < 1.0 \times 10^{-7}M$, such as $1.0 \times 10^{-11}M$, what is the pH and the $[OH^-]$?

Conclusion: A high pH (greater than 7) corresponds to a ______solution.

What if the concentration values are 'in between' those that are given in the table above (ie. coefficients other than 1, 1.0, 1.00, etc...)?

pH can be calculated from [H₃O⁺] by the following: $pH = -\log [H₃O⁺]$

[H₃O⁺] can be calculated from pH by the following: [H₃O⁺] = inv log (-pH) = $10^{(-pH)}$

Fill in the blanks in the following table (use proper sigfigs!):

[H ₃ O ⁺] (M)		9 x 10 ⁻⁹		5.09×10^{-13}	
рН	4.562		13.2		7.32

pOH

pOH is a measure of hydroxide ion concentration using logarithms.

$$pOH = - log [OH^-]$$

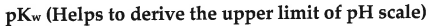
$$[OH^{-}]$$
 = inv log (-pOH)

If pOH < 7; solution is basic

If pOH > 7; solution is acidic.

Assignment 11: pH/pOH Exercises

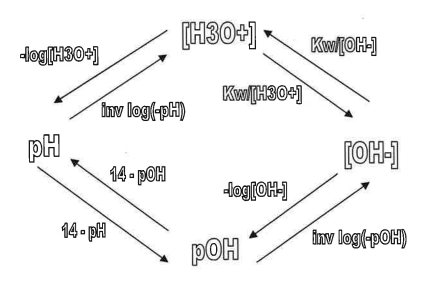
- 1. Find the pH of
 - a) 1.0 x 10⁻⁵M H₃O⁺.
 - b) 2.6 x 10⁻⁷M H₃O⁺.
 - c) 6.7 x 10⁻¹²M H₃O⁺.
- 2. Find the [H₃O+] for each
 - a) pH = 2.35
 - b) pH = 6.456
 - c) pH = 10.76
- 3. Find the pOH of each in #1. Hint: use $K_w = [H_3O^+][OH^-]$
- 4. Find the pOH of each in #2.
- 5. Find the [OH-] for each
 - a) pOH = 2.34
 - b) pOH = 12.59
 - c) pOH = 7.10
- 6. Add the pH and pOH for each of a, b, and c in #1 and 3. What do you notice?



$$pK_w = -\log K_w \text{ at } 25^{\circ}C =$$

Since
$$K_w = [H_3O^+][OH^-] = 1.0 \times 10^{-14}$$
,
Then $pK_w = pH + pOH$ and $14 = pH + pOH$ (at 25°C)

Conversion "Road-Map":



Assignment 12: More pH/pOH Exercises:

- 1. Calculate pH and pOH of (with proper sigfigs)
- a) 1.2 x 10⁻²M H₃O⁺
- b) 3.54 x 10⁻⁸M OH⁻
- c) $8.77 \times 10^{-10} M H_3 O^+$
- d) 6.5 x 10⁻¹M OH⁻

- 2. Calculate $[H_3O^+]$ and $[OH^-]$ for
- (with proper sigfigs)

- a) pH = 3.45
- b) pH = 12.54
- c) pOH = 2.335
- d) pOH = 13.22

Outside the pH scale

At 25°C, very concentrated acidic solutions may have pH values less than 0 and very concentrated basic solutions may have pH values greater than 14. Examples: Find the pH of

- a) 2.0M HBr
- b) 10.0M KOH
- c) 1.5M Sr(OH)2
- d) 1.5 moles Mg(OH)₂ placed into 1L of water *teehee...

Assignment 13: Hebden p. 139 #49, 50, 53 & p. 141 #55, 57

MORE EXAMPLES:

Eg 1: Find the pH of a 0.0020M solution of HNO₃.

Eg 2: Calculate the pH of a 0.010M NaOH solution.

Eg 3: If the pH is decreased from 5.0 to 2.0, what happens to the $[H_3O^+]$ and $[OH^-]$?

Eg 4: If pH is increased from 7.25 to 8.93, what happens to the $[H_3O^+]$?

Eg 5: Calculate the pH of the final solution if 100.0mL of a strong acid at pH = 4.500 is diluted by adding 50.0mL of water.

Eg 6: By how many pH units does the pH change if 80.0mL of 0.0200M HCl is diluted to a final volume of 160.0mL?

Assignment 14: More pH Exercises

- 0. Hebden p.141 #56.
- 1. Calculate the pH, pOH and [OH-] of a 0.00100M solution of HNO₃.
- 2. Calculate the pOH, pH, and $[H_3O^+]$ of a 2.34 x $10^{-4}M$ solution of Ca(OH)₂. Assume 2.34 x 10^4 M is low enough to allow Ca(OH)₂ to fully dissociate
- 3. If the pH is increased from 1 to 6, what happens to the $[H_3O^+]$ and $[OH^-]$?
- 4. If the pH decreases from 9.3 to 6.5, what happens to the $[H_3O^+]$?
- 5. What is the pH of the final solution if 35.00mL of a strong acid at pH 3.56 is diluted by adding 100.0mL of water?
- 6. You have 50.00mL of a 0.00345M solution of HClO₄. How does the pH change if you dilute the solution to a final volume of 175.0mL?

Temperature and pH

At 25°C: $2H_2O_{(1)} + 59kJ \Leftrightarrow H_3O^+_{(aq)} + OH^-_{(aq)}$

 $K_{\rm w}$ = [H₃O⁺] [OH⁻] = 1.0 x 10⁻¹⁴, so p $K_{\rm w}$ = 14

remember: $(pK_w = upper limit of pH scale)$

Therefore, the pH scale (at 25°C) is 0-14

If the temperature is increased, what happens to the above equilibrium and the resulting K_w ? What will happen to the pH scale? Will the pH be acidic, basic, or neutral?

Answer the same questions for when the temperature is decreased.

An increase in temperature to 50° C results in a K_w of 5.48×10^{-14} . Calculate the pH, pOH, [H₃O⁺], and [OH⁻] in water.

A sample of distilled, pure water has a pH of 7.50. Is the temperature greater than or less than 25°C? Explain.

Assignment 15: Hebden p. 139 #51, 52, + question 1 below

- 1. Water at a certain temperature has a K_w of 4.4×10^{-15} .
 - a) Is the water at a temperature above or below 25° C?
 - b) What is the pK_w?
 - c) What would the pH scale be at this temperature?
 - d) Find the $[H_3O^+]$ and $[OH^-]$.
 - e) Find the pH and pOH.
 - f) Is water at this temperature acidic, basic, or neutral?

Mixing an acid with a base produces a solution that can be ______, or ______ depending on the *moles* of H₃O⁺ compared to the *moles* of OH⁻ mixed. Hebden uses a *molarity* method; I teach a *moles* method. Either may be used, but sigfigs could differ. The moles method allows one to better understand the 'true' chemistry at work.

1. Calculate the pH of a solution obtained by adding 50.0mL of 0.10M HCl to 80.0mL of 0.15M NaOH.

Some important information:

- monoprotic acids dissociate into hydronium at a one-to-one ratio;
- diprotic acids dissociate into hydronium at a one-to-two ratio;
- triprotic acids dissociate into hydronium at a one-to-three ratio;
- etc...

When in the presence of an in-excess, strong counterpart.

- 'Diprotic' (or 'polyprotic') Arrhenius bases that are low soluble will dissociate to their fullest extent when in the presence of an in-excess strong acid.

2. Calculate the pH of a solution obtained by adding 30.00g of LiOH to 650.0 mL of 0.400 M H₂SO₄.

3. What mass of NaOH would have to be added to 500.0mL of 0.100M HClO4 in order to produce a solution with a pH of 3.200.

- 4. How many moles of HCl must be added to 40.0 mL of 0.180 M Ca(OH)² to produce a solution having a pH of 7.00? (Assume that there is NO change in volume when the HCl is added).
- *some trivial, but useful, information: most pure acids are gases (HCl included); pure sulfuric acid is a liquid. Acids become aqueous when they are mixed into water. Thus, there is no change in volume in #4 because the HCl must have been added in gaseous form.

Assignment 16: Hebden p. 143 # 58-68 (Change question 60 to 400.0 mL of HCl – new answer: pH of 2.197)

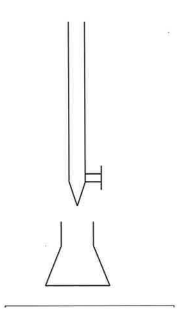
XIII) Titrations

Scenario: You are a public school teacher and you enter the chemical room looking for some HCl of a particular molarity. You see a container marked HCl but the concentration has been rubbed off the bottle. Since money is tight, you want to avoid dumping the solution, but in order to use the acid productively, you'll need to know the concentration. A ______ can be performed in order to determine the unknown concentration.

What is required?

- A measured volume of a _____ (a reactant of known molarity).
- A small, but known volume of the unknown molarity chemical (in our case for this scenario, HCl).
- An acid/base indicator.
- *if the unknown molarity solution is an acid (like our scenario), then the standardized solution will be a base; if the unknown molarity solution is a base, then the standardized solution will be an acid.

 The SET-UP:



Suppose the flask contains the HCl of unknown concentration and the burette contains a standardized base. At some point during the titration,
the moles of OH- added from the burette will equal the moles of H ₃ O+ in
the flask. This is called the
(aka the).
Equivalence Point: mol H ₃ O ⁺ = mol OH ⁻
Strong Acid vs. Strong Base Titration → pH at equiv. pt. is 7.
An is used to help visually determine when the equivalence point has been reached (due to a colour change). The volume of standardized base is determined using the gradations on the burette, and is used to calculate the concentration of the acid.
A 10.00mL sample of HCl of unknown concentration is titrated using 23.62mL of 0.100M NaOH (the standardized solution). Determine [HCl].

Primary Standards (pp. 164-165)

Pure and dry acids or bases are very uncommon as the majority of acids and bases (such as NaOH) absorb water vapour very easily; they are **hygroscopic** (ie. they become hydrates). In order to prepare a standardized solution of, say, NaOH, NaOH would have to be weighed on a scale. However, an unknown percentage of the mass would be due to the absorption of water, and therefore calculating an accurate molarity of the resulting solution would not be possible by simple quantitative methods. A primary standard acid would be required to standardize the standardized solution (ie. 'standardize the standard').

Acids and bases that are non-hygroscopic are known as

________. An accurate molarity
(concentration) of primary standard solutions can be found by direct
calculation, since none of their mass can be attributed in any way to water
vapour acquisition.

Example of a Primary Standard Base: sodium carbonate (Na₂CO₃) – solid.

Examples of Primary Standard Acids: potassium hydrogen phthalate (C₈H₅KO₄) – solid, and oxalic acid (H₂C₂O₄) – solid.

Once a primary standard of known concentration is prepared, it can be used to **standardize** any other acid or base solution. For example, oxalic acid is used to standardize NaOH through titration. Then, that same NaOH, whose concentration is now accurately known, can be used to titrate our unknown concentration of HCl. Likewise, if we have an unknown concentration of base in our lab, we can use sodium carbonate as a primary standard to standardize a hygroscopic acid like HBr, so that we may use the HBr as a standardized solution to titrate our unknown base.

Why wouldn't a chemist simply use primary standards as standardized solutions and, in the process, save time and energy?

Titration	Calcui	lations
IIIIauvii	Calcu	iauviis

Remember: A STRONG base will fully neutralize (dissociate) a weak and/or polyprotic acid and vice versa.

37.86mL of 0.250M NaOH was required to neutralize a 20.0mL sample of HF. Calculate the [HF].

A 15.0mL sample of unknown [KOH] was titrated using 18.56mL of 0.350M HNO₃. Find [KOH].

50.0mL of 0.100M Sr(OH)₂ was required to neutralize a 30.0mL sample of HBr. Determine the [HBr].

Assignment 17: Titration Exercises

1. Find the concentration of an HCl solution if 25.00mL is titrated with 28.46mL of a 0.105M standardized solution of NaOH.

2. You titrated a 30.0mL solution of HNO₃ with 23.75mL of a 0.25M standardized solution of KOH. What is the [HNO₃]?

3. A 35.00mL unknown solution of LiOH is titrated with 17.67mL of 0.200M HI. What is the [LiOH]?

4. A 24.00mL sample of H₂SO₄ is titrated with 32.43mL of 0.150M NaOH solution. Find [H₂SO₄].

5. A 40.00mL sample of Ca(OH)₂ is titrated with 16.55mL of 0.100M HCl. Find [Ca(OH)₂].

6. A 20.00mL sample of H_3PO_4 is titrated with 25.76 mL of a 0.100M $Sr(OH)_2$ solution. Find $[H_3PO_4]$.

Calculating Volumes

What volume of 0.0350M Ba(OH)₂ will be required to neutralize 50.0mL of 0.0275M HCl?

Determining Percent Purity

A 0.74g sample of impure NaCH $_3$ COO is diluted in a volume of 25.0mL. The solution is titrated using 30.0mL of 0.100M HCl. Calculate the % purity of the NaCH $_3$ COO.

Molar Mass Determination

A 0.49g sample of unknown monoprotic acid HA is diluted to 40.0mL. The solution is titrated using 12.0mL of 0.100M NaOH. Calculate the molar mass of the acid.

Chloride Ion (Cl⁻) Titrations – Sol'n Chem. -- Read pp. 99-100 as a class. Main point – Equivalence point is reached when moles Ag⁺ added = moles Cl⁻ in flask. Like acid/base titrations, this is a one-to-one ratio, making the math easier than if the ratio was not one-to-one.

eg1: In order to find the [Cl-] in a sea water sample, a 25.0 mL sample was titrated with a 0.500M AgNO₃ solution, using sodium chromate as an indicator. At the equivalence point, 26.8 mL of AgNO₃ solution had been added. What was the [Cl-] in the sea water?

eg2: What volume of 0.125M silver nitrate will be required to titrate 50.0 mL of 0.0550 M Cl⁻ solution, using a sodium chromate indicator?

eg3. A 5.29g sample of impure NaCl was dissolved and diluted to a total volume of 250.0 mL. If 25.0 mL of the NaCl solution required 28.5 mL of 0.300M silver nitrate solution to reach the equivalence point, using chromate indicator, what was the percentage purity of the original sodium chloride sample?

Assignment 18: Hebden p. 158 #94-97, 102-103, 105, 106; p. 165 #121-122. Also read pp. 99-101; do #70-72, 74-75 (Cl⁻ titrations)