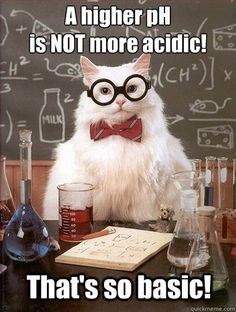
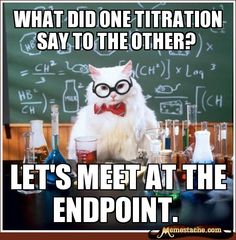
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Acids and Bases (Teacher)





(Discover and save creative ideas n.d.)

|  |  |  |  |
| --- | --- | --- | --- |
| **Week** | **topic** | **unit content** | **assessment** |
| 5 – 10 | Acids and Bases | * models and theories are contested and refined or replaced when new evidence challenges them, or when a new model or theory has greater explanatory scope. * Davy initially proposed that acids were substances that contained replaceable hydrogen (hydrogen that could be partly or totally replaced by metals) and bases were substances that reacted with acids to form salts and water. * Arrhenius model, which includes only soluble acids and bases, identified acids as substances which produce hydrogen ions in solution and bases as substances which produce hydroxide ions in solution * the Brønsted-Lowry model describes acid-base behaviour in terms of proton donors and proton acceptors. This approach includes a wider range of substances and can be more broadly applied. * acids are substances that can act as proton (hydrogen ion) donors and can be classified as monoprotic or polyprotic, depending on the number of protons available for donation * the relationship between acids and bases in equilibrium systems can be explained using the Brønsted-Lowry model and represented using chemical equations that illustrate the transfer of protons between conjugate acid-base pairs * the strength of acids is explained by the degree of ionisation at equilibrium in aqueous solution which can be represented by chemical equations and acidity constants (Ka) * the hydrolysis of salts of weak acids and weak bases can be represented using equations; the Brønsted-Lowry model can be applied to explain the acidic, basic and neutral nature of salts derived from bases and monoprotic and polyprotic acids * water is a weak electrolyte; the self-ionisation of water is represented by Kw = [H+][OH‒] where Kw = 1.0 x 10-14 at 25 °C * Kw can be used to calculate the concentration of hydrogen ions or hydroxide ions in solutions of strong acids or bases * the pH scale is a logarithmic scale and the pH of a solution can be calculated from the concentration of hydrogen ions using the relationship pH = ‒ log10 [H+] * acid-base indicators are weak acids, or weak bases, in which the acidic form is a different colour from the basic form * buffer solutions are conjugate in nature and resist changes in pH when small amounts of strong acid or base are added to the solution; buffering capacity can be explained qualitatively; Le Châtelier’s Principle can be applied to predict how buffers respond to the addition of hydrogen ions and hydroxide ions * volumetric analysis methods involving acid-base reactions rely on the identification of an equivalence point by measuring the associated change in pH, using appropriate acid-base indicators or pH meters, to reveal an observable end point * data obtained from acid-base titrations can be used to calculate the masses of substances and concentrations and volumes of solutions involved | Practical 2  Test 2 |

**Chemical and Physical Properties of Acids and Bases**

Acidic solutions generally:

* conduct an electric current
* turn litmus red
* taste sour

Basic solutions generally:

* also conduct an electric current
* turn litmus blue
* taste bitter
* have a slippery feel

Fill in the chemical equation and the net ionic equation for the reactions below illustrating some of the **chemical properties** of acids and bases.

1. Acidic solutions dissolve most metals forming a salt and hydrogen gas. Au, Ag, Pt and Cu are unaffected by dilute acids.

**ACID + METAL → SALT + H2(g)**

(Eg: Some hydrochloric acid solution is poured onto a strip of nickel metal)

**2HCl(aq) + Ni(s) → NiCl2(aq) + H2(g)**

**2H+(aq) + Ni(s) → Ni2+(aq) + H2(g)**

1. **ACID + METAL HYDROXIDE → SALT + H2O**

(Eg: Calcium hydroxide solution is added to a solution of nitric acid)

**2HNO3(aq) + Ca(OH)2(aq) → Ca(NO3)2(aq) + 2H­2O(l)**

**H+(aq) + OH−(aq) → H2O(l)**

1. **ACID + METAL OXIDE → SALT + H2O**

(Eg: Some solid copper (II) oxide is added to a solution of sulfuric acid)

**H2SO4(aq) + CuO(s) → CuSO4(aq) + H­2O(l)**

**2H+(aq) + CuO(s) → Cu2+(aq) + H2O(l)**

1. **ACID + CARBONATE → SALT + H2O + CO2(g)**

(Eg: Hydrochloric acid solution is added to nickel carbonate solid)

**2HCl(aq) + NiCO3(s) → NiCl2(aq) + H2O(l) + CO2(g)**

**2H+(aq) + NiCO3(s) → Ni2+(aq) + H2O(l) + CO2(g)**

1. **ACID + HYDROGENCARBONATE → SALT + H2O + CO2(g)**

(Eg: Ethanoic acid solution is added to solid zinc hydrogencarbonate)

**2CH3COOH(aq) + Zn(HCO3)2(s) → Zn2+(aq) + 2CH3COO-(aq) + 2H2O(l) +2CO2(g)**

1. **ACID + METAL SULFITE → SALT + H2O + SO2(g)**

(Eg: Hydrochloric acid solution and sodium sulphite powder are mixed)

**2HCl(aq) + Na2SO3(s) → 2NaCl(aq) + H2O(l) + SO2(g)**

**2H+(aq) + Na2SO3(s) → 2Na+ (aq) + H2O(l) + SO2(g)**

1. **BASE + AMMONIUM SALT → SALT + H2O + NH3(g)**

(Eg: A potassium hydroxide solution is mixed with an ammonium chloride solution)

**KOH(aq) + NH4Cl(aq) → KCl(aq) + H2O(l) + NH3(g)**

**NH4+(aq) + OH-(aq) → H2O(l) + NH3(g)**

1. **BASE + NON-METAL OXIDE → SALT + H2O**

(Eg: Sulfur dioxide gas is bubbled through a potassium hydroxide solution)

**SO2(g) + 2KOH(aq) → K2SO3(aq) + H2O(l)**

**SO2(g) + 2OH- (aq) → SO32-(aq) + H2O(l)**

***Complete Lucarelli Set 3***

###### **Theories of Acids and Bases**

###### **Davy Theory**

###### Acids: Have replaceable H (hydrogen could be partly or totally replaced by metals).

###### eg hydrochloric acid:

###### 2HCl(aq) + Mg(s) → MgCl2(aq) + H2(g)

###### Bases: Reacted with acids to form salt and water.

###### eg sodium hydroxide:

###### NaOH(s) + HNO3(aq) → NaNO3(aq) + H2O(l)

###### **Arrhenius Theory**

Acids: Have H in their formula and produce hydrogen ions (H+) when dissolved in water.

eg hydrochloric acid: HC*l*(g) → H+(aq) + C*l*−(aq)

eg ethanoic acid: CH3COOH(s) → H+(aq) + CH3COO−(aq)

Bases: Have OH in their formula and produce hydroxide ions (OH−) when dissolved in water.

eg sodium hydroxide: NaOH(s) → Na+(aq) + OH−(aq)

eg strontium hydroxide: Sr(OH)2(s) → Sr2+(aq) + 2OH−(aq)

Neutralisation: an acid plus a base produces a neutral solution (ie H+(aq) + OH−(aq) → H2O(l))

Problems with the theory:

* Some bases produce OH− ions in solution yet do not have OH in their formula (eg NH3 and CO32-).
* Restricted to aqueous solutions.
* Not all salts are neutral.
* Does not allow for the existence of hydronium ions (H3O+).

***Brønsted-Lowry Theory***

Acids are proton (H+) donors.

Strong acid: **HC*l*(aq) + H2O(l) → H3O+(aq) + C*l*−(aq)**

HCl is donating a proton and acting as an acid. The H2O is accepting a proton and acting as a base.

Weak acid: CH3COOH(aq) + H2O(l)  H3O+(aq) + CH3COO−(aq)

**CH3COOH(aq) is donating a proton and acting as an acid. The H2O is accepting a proton and acting as a base**.

Bases are proton (H+) acceptors.

**eg ammonia: NH3(aq) + H2O(l) NH4+(aq) + OH−(aq)**

**NH3(aq) is accepting a proton and acting as a base. The H2O is donating a proton and acting as an acid.**

eg hydrogencarbonate ion: HCO3− (aq) + H2O(l)  H2CO3(aq) + OH−(aq)

HCO3-(aq) is accepting a proton and acting as a base. The H2O is donating a proton and acting as an acid.

Neutralisation: reaction between a proton donor and a proton acceptor.

eg carbonate ion plus water: CO32-(aq) + H2O(l) HCO3−(aq) + OH−(aq

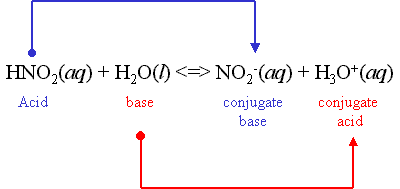
**Conjugate acid- base pairs**

Brønsted-Lowry theory recognises the reversibility of proton transfer reactions.

* **Once an acid has donated a proton it has the potential to act as a base. We say it has formed its conjugate base.**
* **Once a base has accepted a proton it has the potential to act as an acid. We say it has formed its conjugate acid.**
* This means both the forward and the reverse reactions are acid-base reactions.

eg NH3(g) + H2O(l)  NH4+(aq) + OH−(aq) H2O(l) + H2O(l)  H3O+(aq) + OH−(aq)

* The acid and the base it forms (or the base and the acid it forms) are called a conjugate acid-base pair.
* In any conjugate acid- base pair the formula of the acid is always greater by the equivalent of H+.
* The stronger the acid the weaker its conjugate base.



(Blaber, 2000)

**Acidity constant (Ka) and acid strength**

**An acid- base transfer reaction is an example of an equilibrium process. The extent to which an acid ionises in aqueous solutions can be determined from the equilibrium constant Ka also known as the acidity constant. It is a measure to which the proton transfer goes to completion.**

* The larger the Ka the greater the tendency of the acid to donate a proton to water ie the stronger the acid is the greater the degree of its ionisation. (Strong acids fully ionise in water.)
* Eg the proton transfer equation for ethanoic acid is:

CH3COOH(aq) + H2O(l)  H3O+(aq) + CH3COO−(aq)

its Ka expression is: Ka = 

The Ka value at 25°C is 1.8 x 10-5. This indicates the reaction only proceeds to a very limited extent and only about 1% of ethanoic acid molecules are ionised.

* **The larger the value of Ka the more the equilibrium position favours products, hence the greater the tendency of the acid to form hydrogen ions and so the stronger the acid is.**
* The Ka value also indicates the strength of the acid’s conjugate base to accept a proton back from water so, if Ka is large (strong acid) its conjugate base is weak. As Ka is progressively smaller, the acid is progressively weaker and the conjugate base becomes progressively stronger. (**The stronger the acid is the weaker its conjugate-base.)**
* Acid-base reactions tend to occur in the direction in which the stronger acid and stronger base react to form a weaker acid and weaker base.

**Eg NH3(aq) + H2O(l) NH4+(aq) + OH–(aq)Equilibrium lies to the left**

**HCN(aq) + H2O(l) H3O+(aq) + CN-(aq) Equilibrium lies to the left**

###### **Polyprotic Acids**

**Monoprotic acids can donate only one proton (hydrogen ion) per molecule. Eg HCl; CH3COOH; HNO3**

While weak monoprotic acids produce fewer hydrogen ions in solution than strong acids, they react to the same extent with bases, providing that the concentration is the same.

**HC*l*(aq) + H2O(l) → H3O+(aq) + C*l*−(aq)**

**Diprotic acids contain two acidic protons per molecule. Eg H2SO4; H2CO3**

Explain (using equations) why one mole of H2SO4 yields between one and two moles of H3O+ ions in solution.

H2SO4(aq) + H2O(l) → HSO4−(aq) + H3O+(aq)

HSO4−(aq) + H2O(l) SO42−(aq) + H3O+(aq)

This process of becoming ions over multiple steps is called **successive ionisation.**

Polyprotic acids are acids which contain two or more acidic protons. Use equations to show the successive ionisation of phosphoric acid.

**H3PO4(aq) + H2O(l) H2PO4−(aq) + H3O+(aq)**

**H2PO4−(aq) + H2O(l) HPO42−(aq) + H3O+(aq)**

**HPO42−(aq) + H2O(l) PO43−(aq) + H3O+(aq)**

Acid-Base Properties of Salts

**When an acid reacts with a base a salt and water are produced. Although this is called a neutralisation reaction, the salt formed in the reaction is not always neutral. The acid-base properties of the salt can be understood in terms of Bronsted-Lowry theory. If one of the ions present in the salt reacts with water (and the ion acts as a proton donor) then its solution will be acidic. If one of the ions present reacts with water (and its ion acts as a proton acceptor) then its solution is basic. The reaction of the ions with water is also known as hydrolysis.**

Eg 1

An ammonium chloride solution (NH4Cl(aq)) is acidic. The ammonium ion (NH4+) is a weak acid in water as acts as a proton donor and hydrolyses to produce hydronium ions (H3O+). The chloride ion (Cl-) is neutral.

**NH4+(aq) + H2O(l)  NH3(aq) + H3O+(aq)**

Eg 2

A sodium ethanoate solution (NaCH3COO(aq)) is basic. The ethanoate ion (CH3COO-) is a weak base in water as it acts as a proton acceptor and hydrolyses to produce hydroxide ions (OH-).The sodium ion (Na+) is neutral.

CH3COO-(aq) + H2O(l)  CH3COOH(aq) + OH-(aq)

# Summary

The table below classifies common ions according to whether they cause aqueous solutions to be acidic, basic or neutral.

|  |  |  |  |
| --- | --- | --- | --- |
| **Type of Ion** | **Acidic** | **Basic** | **Neutral** |
| Anions | Anions produced in first stage of ionisation of polyprotic acids: HSO4−, H2PO4− | Anions from  weak acids:  F−, CO32-, HS−, S2-, HPO42-, PO43-, HCO3−, CH3COO− , SO42- | Anions from  strong acids:  Cl−, Br−, I−, NO3− |
| Cations | NH4+, Fe3+, Al3+, Cr3+  Sn4+ | Not common | Li+, Na+, K+, Mg2+, Ca2+, Ba2+ |

Predict whether aqueous solutions of the following salts will be acidic, basic or neutral and justify your choice by writing suitable chemical equations.

1. Ammonium nitrate
2. Calcium hydrogencarbonate
3. Potassium ethanoate
4. Magnesium bromide
5. Calcium hydrogensulfate
6. **Acidic**

**NO3­- ions are neutral**

**NH4+(aq) + H2O(l)  NH3(aq) + H3O+(aq)**

1. **Basic**

**Ca2+ ions are neutral**

**HCO3- (aq)  + H2O(l)  H2CO3(aq) +  OH-(aq)**

1. **Basic**

**K+ ions are neutral**

**CH3COO-(aq) + H2O(l)  CH3COOH(aq) + OH-(aq)**

1. **Neutral**

**Mg2+ and Br- ions are both neutral**

1. **Acidic**

**Ca2+ ions are neutral**

**HSO4-(aq) + H2O(l) SO42-(aq) + H3O+(aq)**

***Complete Essential Chemistry Set 4***

### Self- Ionisation of Water, Kw and pH

**Water is an extremely weak electrolyte. The ionisation of water molecules into hydronium and hydroxide ions is very small. This is known as self-ionisation. It is given by:**

**2H2O(l)  H3O+(aq) + OH−(aq)**

At hh, the equilibrium constant for the ionisation of water (Kw) equals 1.00 ×10-14. It is given by: Kw = [H3O+][OH–]

In water or any solution where H3O+ and OH- have the same concentration, the solution is said to be neutral. [H3O+] = [OH–] = 1.00 ×10-7 mol L-1 at 25 °C.

From this, an acidic solution must have [H3O+] > [OH–] and a basic solution must have [H3O+] < [OH–]. Also, since [H3O+] × [OH–] = 1.00 ×10-14 for all solutions at 25 °C, the concentration of one of these ions can be calculated given the concentration of the other.

Eg. Calculate the concentration of hydrogen ions in a 0.001 mol L-1 NaOH solution.

**[OH–] = [NaOH] = 1 x 10-3 mol L-1**

**[H3O+] × [OH–] = 1.00 ×10-14 mol L-1**

**[H3O+] = 1.00 ×10-14/ 1 x 10-3**

**= 1.00 x 10 -11 mol L-1**

Eg. Calculate the concentration of hydroxide and hydrogen ions in a solution made by dissolving 2.34 g of Ba(OH)2 solid in 543 mL of water.

n Ba(OH)2 = 2.34/171.316 = 0.01366 mol

n(OH-) = 2x n Ba(OH)2

= 2 x 0.01366 mol = 0.02732 mol

[OH–] = n/v = 0.02732/ 0.543 = **5.03 x 10-2** mol L-1

[H+] × [OH–] = 1.00 ×10-14 mol L-1

[H+] = 1.00 ×10-14/ [OH–]

= **1.99 x 10-13** mol L-1

**[OH–] = 5.03 x 10-2**

**[H+] = 1.99 x 10-13**

**pH**

**pH is a number indicating how acidic or basic a solution is. It was developed by Søren Sørenson.**

**It is related to [H+] / [H3O+] through the expression: pH = –log10[H+] / [H3O+].**

**A pH value below 7 is acidic, while a pH value above 7 is basic and pH 7 is neutral.**

Eg. Calculate the pH of the following solutions:

1. A 0.002 mol L-1 solution of HC*l*.

pH = 2.7

1. A 0.007 mol L-1 solution of H2SO4 (assuming complete ionisation)

H2SO4(aq) → 2H+(aq) + SO42-(aq)

Complete ionisation produces **2 x** 0.007 mol L-1 H+ ions = 0.014 mol L-1

pH = - log [0.014] = 1.85

1. A 0.005 mol L-1 solution of NaOH.

NaOH(aq) → Na+(aq) + OH-(aq)

Complete ionisation produces 0.005 mol L-1 OH- ions

[H+] = 1.00 x 10-14/0.005 = 2 x 10-12 mol L-1

pH = - log [2 x 10-12 ] = 11.70

1. Calculate the final pH of a solution made up by adding 5.67 g of HC*l* solid to 23.5 mL of a 3.96 ×10-1 mol L-1 Ba(OH)2 solution (assume no change in volume).

-0.76

1. Calculate [H+] of HNO3 solution with a pH of 1.38.

0.0417 mol L-1

1. Calculate [OH-] and [H3O+] of a Mg(OH)2 solution with a pH of 12.68.

[H3O+] = 2.09 x 10-13

[OH-] = 4.79 x 10-2

pH can be measured using an indicator solution (a specific weak acid that changes colour over a range of pH), indicator paper or a pH meter (which is much more precise, if used correctly). A pH meter measures the voltage difference across two electrodes placed in the solution.

* The self-ionisation of water is an **endothermic process**. An increase in temperature will favour the endothermic (forward) reaction which would result in an increased equilibrium constant. (eg at 40 °C, Kw = 2.916 ×10-14).
* Similarly, at lower temperatures, the exothermic (reverse) reaction is favoured and so the equilibrium constant is reduced (eg at 10 °C, Kw = 2.93 ×10-15).

It is important to remember that even though pure water at temperatures above 40 °C has a pH less than 7, it is not acidic (this is because a neutral pH of 7 only applies at 25 °C). Pure water at, whatever temperature, is always neutral.

***Complete Essential Chemistry Set 5***

**Buffers**

**A buffer is a solution that resists changes in its pH when small amounts of an acid or base are added. It is made up of a mixture of a weak acid and its conjugate base or a weak base and its conjugate acid. They are often prepared by adding the salt of the conjugate acid or base to the weak base or acid (respectively).**

Eg Mixing 2 mol of ethanoic acid, CH3COOH, with 2 mol sodium ethanoate NaCH3COO and making the solution volume up to 1.00 L. This solution contains a high concentration of weak acid, CH3COOH, and its conjugate weak base, CH3COO- (from NaCH3COO).

The equilibrium set up is given as: CH3COOH(s) + H2O(l) H3O+(aq) + CH3COO−(aq)

If acid is added, the system adjusts (according to Le Châtelier’s Principle) by shifting the equilibrium to the left. In this case, the acetate ions will react with the excess H3O+ to keep the overall [H3O+] relatively constant. Similarly, if a base is added, the system adjusts by shifting the equilibrium to the right to replace the H3O+ which reacted with the OH–.

In both these cases, the acid or base can be consumed without a great alteration to the [H3O+] in the original mixture. As a result, the pH of the mixture does not alter appreciably. Of course, buffer solutions have their limits and eventually if enough acid or base is added, the pH will change significantly.

Explain buffering using Collision Theory

Using the ethanoic/ sodium ethanoate buffer system:

If **acid** **is added** to the system this increases the H+/ H3O+ concentration. This increases the frequency of collisions between H3O+ and CH3COO‒ and so increases the rate of the reverse reaction relative to the forward reaction and the equilibrium shifts left. This will consume H3O+ until new equilibrium is established where H3O+ concentration is very close to what it was originally (maintaining pH).

If **base is added** to the system this decreases the H+/ H3O+ concentration. This reduces the frequency of collisions between H3O+ and CH3COO‒ and so reduces the rate of the reverse reaction relative to the forward reaction and the equilibrium shifts right. This will increase H3O+ until new equilibrium is established where H3O+ concentration is very close to what it was originally (maintaining pH).

The buffer capacity of a solution depends on two factors:

* The relative concentrations of the weak acid (base) and its conjugate base (acid). Equal concentrations of these two offers the best buffering capacity.
* The concentration of the weak acid (base) and its conjugate base (acid). The greater the concentrations (in general) the greater the buffering capacity.

Blood has a system of buffers present in it to maintain a fairly consistent pH range between 7.35 and 7.45. This is essential to maintain the body systems functioning.

One of the buffer systems to help blood maintain this narrow pH range is the **carbonic acid** and **hydrogen carbonate ion** buffer system. Carbonic acid is the weak acid and the hydrogen carbonate ion is the weak conjugate base.

H2CO3(aq) + H2O(l)  HCO3−(aq) + H3O+(aq)

**If blood becomes too acidic, [H3O+]becomes too high,equilibrium shifts to the left as the HCO3‒ reacts with the excess H3O+. If blood becomes too basic, [H3O+]falls too low,equilibrium shifts to the right producing more H3O+.**

An enzyme found in blood called carbonic anhydrase catalyses the formation of carbonic acid from carbon dioxide which ensures the equilibrium adjusts quickly to changes in H2CO3(aq) or CO2(aq).

The H2CO3/HCO3–/CO2 buffer system is particularly useful because it allows the excess H3O+ to exit the body through the lungs as CO2.

Another blood buffer system is the dihydrogenphosphate and hydrogenphosphate, H2PO4–/HPO42- buffer system.

***Complete Essential Chemistry Set 6***

**Acid-Base Titrations**

Acid- base titrations (also called volumetric analysis) are an analytical procedure often used to determine the concentration of a solution or the amount of a particular substance present. The concentration of one solution must be accurately known. This is called the **standard solution.** The purpose of the titration is to determine the concentration of the acid or base in the unknown solution.

Five main pieces of equipment are used in titrations:

* Pipette – used to accurately deliver a known volume (aliquot) of liquid. It is rinsed with the liquid to be delivered from it prior to use.
* Burette – used to accurately deliver a variable volume (titre) of liquid.

It is rinsed with the liquid to be delivered from it prior to use.

* Volumetric flask – used to prepare an accurately known volume of solution.

It is rinsed with de-ionised water prior to use.

* Analytical balance – used to weigh out an accurately known mass.
* Conical flask – used to hold the solutions during the titration. It is rinsed with de-ionised water prior to use.

In the titration process, a carefully measured volume of one solution, called an aliquot, is added to a conical flask. A variable volume of the other solution, called the titre, is then carefully added from a burette into the conical flask until the reaction between the acid and base is complete. This theoretical point in the titration when neither acid nor base remain is called the equivalence point.

**Indicators** are used to determine when stoichiometrically equivalent amounts of the two reactants have reacted. This point is called the equivalence point. The point at which the indicator **changes colour** is called the end point. To be a useful indicator, the end point should match the equivalency point.

For an indicator (which is a weak acid) in solution, the following reaction takes place:

HInd(aq) H+(aq) + Ind–(aq)

The colour changes are due to the difference in the colours of HInd and Ind–.

## A standard solution has an accurately known concentration. A primary standard solution is used to determine the concentration of a secondary standard solution. A primary standard solution should possess the following characteristics:

* **Has a high degree of purity and a known formula.**
* **The reactions it is involved in are known.**
* **It must be stable. That is, it should not decompose in air, react with oxygen, carbon dioxide or other gases in the air**
* **It must not absorb water from the atmosphere. That is, it is not hygroscopic (absorbs water from the atmosphere) or deliquescent (absorbs so much water that it eventually dissolves in the water it absorbs).**
* **It should have a relatively high formula mass to minimize weighing errors (percentage uncertainty is minimized).**

Common primary standards used in acid-base titrations include **anhydrous sodium carbonate, oxalic acid (H2C2O4)** and **potassium hydrogenphthalate (KHC8H4O4).**

When performing titrations, a practice run is always performed and at least three trials which should agree to within ±0.3 mL (at least two of them). The practice run is never used in the calculating of the concentration.

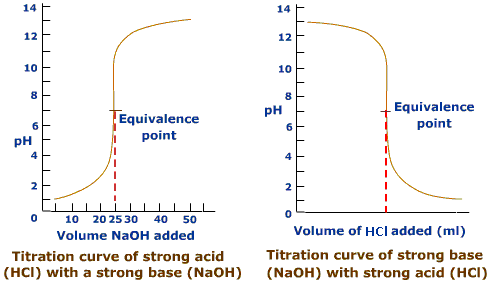
## The Products of Neutralisation Reactions

Not all equivalency points of acid-base reactions will be neutral. Only the reaction between a strong acid and a strong base will produce a neutral solution. As such, the choice of indicator is very important. Below is a list of common indicators and their useful pH range.

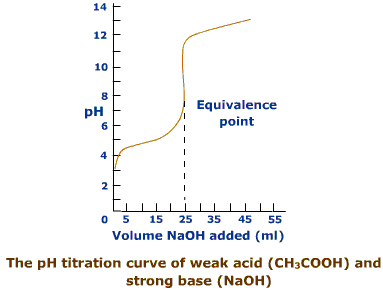
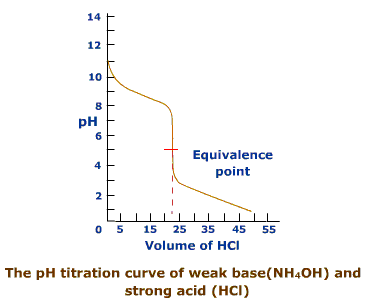
|  |  |  |  |
| --- | --- | --- | --- |
| Indicator | Colour in Acid | Colour in Base | pH Range |
| Universal indicator | Red | Purple | 0.0-14.0 |
| Methyl orange\* | Red | Yellow | 3.1-4.4 |
| Bromophenol blue\* | Yellow | Purple | 3.0-4.6 |
| Methyl red | Red | Yellow | 4.4-6.2 |
| Litmus | Red | Blue | 4.5-8.3 |
| Bromothymol blue | Yellow | Blue | 6.0-7.6 |
| Phenolphthalein\* | Colourless | Pink | 8.3-10.0 |
| Alizarine yellow R | Yellow | Red | 10.2-12.0 |

\*denotes more commonly used (Although Universal Indicator is never used as an acid/base titration, you must be familiar with its colours in various pH ranges i.e. red 0-2, orange 3-4, yellow 5-6, green 7, blue 8-10, violet 11-14)

During a titration, the pH does not change in a linear fashion. Below are typical titration curves for various reactions. Note that the equivalence point occurs at the turning point of the curve.



### 



(NSC. Pearson 2016)

### Reactions Between a Weak Acid and a Strong Base

Write an equation for the reaction between acetic acid and sodium hydroxide in aqueous solution. Which salt is produced? Are solutions of this salt acidic, basic or neutral? Which indicator should be used?

* **CH3COOH(aq) + OH–(aq) → CH3COO–(aq) + H2O(l) then**
* **CH3COO–(aq) + H2O(l)  CH3COOH(aq) + OH–(aq)**
* **Result is a basic solution. Indicator to use is phenolphthalein.**

Write an equation for the reaction between hypochlorous acid (HC*l*O) and potassium hydroxide in aqueous solution. Which salt is produced? Are solutions of this salt acidic, basic or neutral? Which indicator should be used?

* **HC*l*O(aq) + OH–(aq) → C*l*O–(aq) + H2O(l) then**
* **C*l*O- (aq) + H2O(l) HC*l*O(aq) + OH- (aq)**
* **Result is a basic solution. Indicator to use is phenolphthalein.**

Summing up, the end point of titrations between a weak acid and a strong base in aqueous solution are basic and the indicator to be used is phenolphthalein.

### Reactions Between a Strong Acid and a Weak Base

Which salt is formed when an aqueous solution of ammonia is added to a hydrochloric acid solution? Are solutions of this salt acidic, basic or neutral? Which indicator should be used?

* **NH3(aq) + H+(aq) → NH4+(aq) followed by NH4+(aq) + H2O(l)  NH3(aq) + H3O+(aq)**
* **Result is an acidic solution. Indicator to use is methyl orange or bromophenol blue.**

Which salt is formed when an aqueous solution of sodium hydrogencarbonate is added to nitric acid. Are solutions of this salt acidic, basic or neutral? Which indicator should be used?

* **HCO3–(aq) + H+(aq) → CO2(g) + H2O(l)**
* **followed by CO2(g) + H2O(l)  H2CO3(aq)**
* **then H2CO3(aq) + H2O(l)  HCO3−(aq) + H3O+(aq)**
* **Result is an acidic solution. Indicator to use is methyl orange or bromophenol blue.**

Summing up, the end point of titrations between a strong acid and a weak base in aqueous solution are acidic and the indicator to be used is methyl orange or bromophenol blue.

### Reaction Between a Weak Acid and A Weak Base

The reaction between acetic acid and ammonia can be represented by the equation:

* CH3COOH(aq) + NH3(aq) → CH3COO–(aq) + NH4+(aq)

The pH of solutions of salts such as ammonium acetate depend on the relative strengths of the anion as a base and the cation as an acid. Ammonium acetate solutions are very close to neutral. As such, a pH meter is definitely the more preferred choice to monitor this type of acid-base titration as the pH change around the equivalence point is too gradual.

**The titration calculation**

Eg 20.00 mL aliquots of hydrochloric acid solution are titrated with 9.039 x 10-2 mol L-1sodium carbonate solution. Several titrations were performed and the volumes of sodium carbonate solution used were 19.45 mL, 18.90 mL, 18.77 mL and 18.85 mL. What is the concentration of the hydrochloric acid in the solution?

C(HCl)  = 1.703 x 10-1 mol L-1

In a **back titration**, a known volume of an excess reactant is added to the reactant being analysed. A titration is then carried out to determine the amount of excess reactant remaining after the reaction.

Eg Magnesium oxide is not very soluble in water, and is difficult to titrate directly. Its purity can be determined by use of the back titration method. 4.06 g of impure magnesium oxide was completely dissolved in 100.0 mL of hydrochloric acid, of concentration 2.00 mol L-1 (in excess). The excess acid required 19.70 mL of sodium hydroxide (0.200 mol L-1) for neutralisation. Determine the percentage purity of the magnesium oxide in the original sample.

Purity MgO = 97.3%

Sometimes the solution to be analysed is too concentrated for the standard solution and would require a titre too large for the burette. In this case the solution of high concentration is **diluted**.

The following equation will be used in these calculations: c1V1 = c2V2 where

c1 = original undiluted solution concentration in mol L-1

V1 = original undiluted solution volume in litres

c2 = new diluted solution concentration in mol L-1

V2 = new diluted solution volume in litres

Eg. A laboratory technician needs to accurately confirm the concentration of a batch of commercial hydrochloric acid. In order to do so a 15.00 mL sample is transferred to a 2.500 L volumetric flask and made up to the mark with distilled water. 20.00 mL aliquots require on average 16.65 mL of 9.08 x 10-2 mol L-1 NaOH(aq) for equivalence. Determine the concentration of HCl(aq) in the commercial solution.

**n(NaOH) = cV**

**= 9.08 x 10-2 x 16.65 x 10-3**

**=1.51 x 10-3 mol**

**HCl(aq) + NaOH(aq) → NaCl(aq) + H2O(l)**

**Thus n(HCl) = n(NaOH) = 1.51 x 10-3 mol**

**c (HCl diluted) = =  = 7.56 x 10-2 mol L-1**

**c1V1 = c2V2**

**c1 = =  = 12.6 mol L­-1 HCl(aq)**

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**cf: Ethanoic acid solution is added to zinc hydrogencarbonate solution**

**CH3COOH(aq) + HCO3-(aq) → CH3COO-(aq) + H2O(l) +CO2(g)**

**cf: Hydrochloric acid is added to solid zinc hydrogencarbonate**

**2H+ (aq) + Zn(HCO3)2(s) → Zn2+(aq) + 2H2O(l) +2CO2(g)**