

Edexcel Chemistry A-level

Topic 12: Acid-Base Equilibria Detailed Notes

This work by PMT Education is licensed under CC BY-NC-ND 4.0







Brønsted-Lowry Acids and Bases

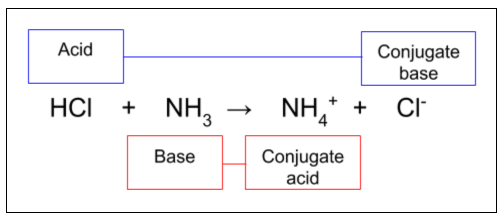
Acid-base equilibria involve the **transfer of protons** between substances. Therefore, substances can be classified as acids or bases depending on their interaction with protons.

A Brønsted-Lowry **acid** is a **proton donor**. For example, ammonium ions (NH_4^+) . A Brønsted-Lowry **base** is a **proton acceptor**. For example, hydroxide ions (OH⁻).

Brønsted–Lowry Conjugate Acid-Base Pairs

A **conjugate acid** is the species formed when a **base accepts a proton**. A **conjugate base** is the species formed when an **acid donates a proton**. These form conjugate acid-base pairs.

Example:

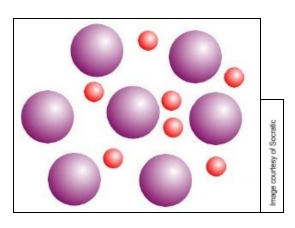


Acid and Base Strength

Acid strength doesn't refer to the concentration of a solution. A strong acid is defined as being:

An acid that completely dissociates to ions when in solution with pH 0-1.

Example:



www.pmt.education

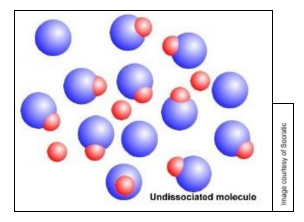




In comparison, a weak acid is defined as being:

An acid that only slightly dissociates when in solution with pH between 3 and 7

Example:



The same definitions are true for **strong** and **weak bases**. Strong bases have pH between 12-14 and weak bases have pH between 7-11.

Enthalpy Change of Neutralisation

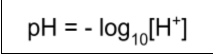
When strong acids and bases react together they will produce very similar enthalpies of neutralisation. This is because the solutions completely dissociate, so the same, simple acid-base reaction occurs between H⁺ and OH⁻ ions to produce water in each case. The other dissociated ions present are simply spectator ions and do not affect the reaction.

However, in reactions of **weak** acids and bases, the ions only slightly dissociate so **other enthalpy changes** also occur within the solution. As a result, the enthalpies of neutralisation can **vary** quite a lot.

Determining pH

pH is a measure of **acidity and alkalinity**. It is a **logarithmic** scale from 0 to 14 that gives the concentration of H^+ ions in a solution. 0 is an **acidic** solution with a high concentration of H^+ ions whereas 14 is a **basic** solution with a low concentration of H^+ ions.

The pH can be calculated using the concentration of hydrogen ions, $[H^+]$, as follows:





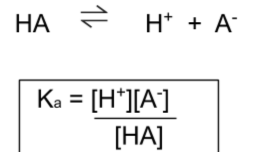


This equation also allows the concentration of H^+ ions to be determined if the pH is known. When using these equations below, the concentration of H^+ ions is given in **mol dm**⁻³.

This concentration of H⁺ ions is equivalent to the **concentration of a strong acid** as it **completely dissociates** to ions in solution.

Weak Acids and Bases

Weak acids and bases only **slightly dissociate** in solution to form an **equilibrium** mixture. Therefore, the reaction has an equilibrium dissociation constant, **K**_a.



In a similar way to $[H^+]$, this constant can be found using pKa:

$$pK_a = -log_{10}K_a$$
 $K_a = 10^{-pK_a}$

The value pKa is a logarithmic acid dissociation constant, representing how acidic something is. A **low value** indicates a **strong acid**.

The relationships of Ka, pKa and $[H^+]$ can be used to find the **pH of weak acids and bases**. Depending on the reaction and the relative concentrations, a different method may have to be used:

HA in excess - Use [HA] and $[A^-]$ along with Ka to find $[H^+]$, then pH.

 $A^{\scriptscriptstyle -}$ in excess - Use Kw to find [H^+], then pH.

 $HA = A^{-}$ - In this case, pKa is equal to pH, therefore find pKa.





Diluting Acids

If you dilute a strong acid **10 times** its pH will increase by **one unit**, because pH is a **logarithmic scale**. Diluting it **100 times** and **1000 times** would, therefore, increase the pH by **two units** and **three units**, respectively.

Weak acids do not behave in the same way. Weak acids are **not fully dissociated in solution**, so diluting them causes the **equilibrium** to **shift** to oppose the change. This means a **10x dilution** of a weak acid would increase the pH by **less than one unit**.

Calculating the Ka of Weak Acids

Weak acids only **partially dissociate** in solution. Therefore, the **equilibrium** of a weak acid has to be taken into account.

- 1. The initial concentrations, change in concentrations and equilibrium concentrations of the reactants and products have to be found.
- 2. The concentration of H^+ ions can then be found using the pH given.
- 3. This value can be used to find the actual equilibrium concentrations.
- 4. Finally, these values can be substituted into the expression for Ka.

Example:

A weak acid, HA, with a concentration of 0.25 M has a pH of 3.5. What is its Ka?

$$HA + H_2O \rightleftharpoons A^- + H_3O^+$$

We assume that 'x' mol of HA dissociates to A^- and H_3O^+ .

| | HA | A- | H₃O⁺ |
|---------|----------|----|------|
| Initial | 0.25 | 0 | 0 |
| Change | -X | +χ | +x |
| End | 0.25 - x | х | х |

Using the pH value give, the $[H^+]$ (= x) can be calculated:

$$[H_3O^+] = [H^+] = 10^{-pH}$$

 $[H^+] = 10^{-3.5} = 3.16 \times 10^{-4} M$
 $x = 3.16 \times 10^{-4} M$





Substituting into the expression for Ka and using the equilibrium concentrations:

Ka =
$$[H^+] [A^-]$$
 = $(x) (x)$
[HA] $(0.25-x)$

We can now substitute in the value of x calculated earlier: $x = 3.16 \times 10^{-4} M$

Ka =
$$(3.16 \times 10^{-4})^2$$

 $(0.25 - (3.16 \times 10^{-4}))$
Ka = 3.99 x 10⁻⁷

Ionic Product of Water

Water **slightly dissociates** to form hydroxide and hydrogen ions as an equilibrium with its own equilibrium constant, **Kw**.

$$Kw = [H^+][OH^-]$$

At **25°C**, room temperature, Kw has a constant value of **1.0 x 10⁻¹⁴**. However, as temperature changes, **this value changes**.

The **forward** reaction in the equilibrium of water is **endothermic** and is therefore favoured when the temperature of the water is increased. As a result, as temperature increases, **more** H⁺ **ions** are produced meaning the water becomes **more acidic**.

$H_2O \Rightarrow H^+ + OH^-$

In the same way that pKa can be calculated from Ka, pKw can be calculated from Kw.

$$\mathsf{pK}_\mathsf{w} = -\log_{10}\mathsf{K}_\mathsf{w}$$

 $K_w = 10^{-pK_w}$





The pH of a strong base can be calculated using pKw or Kw. For a **strong base**, the concentration of OH^{-} will be the **same** as the concentration of the base.

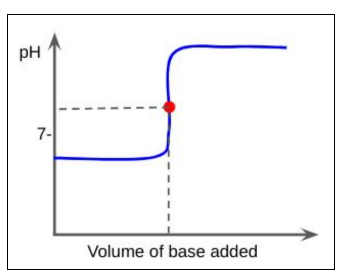
 $\begin{aligned} \mathsf{Kw} &= [\mathsf{OH}^{-}] \, [\mathsf{H}^{+}] \\ \textit{Rearrange Kw to find } [\mathsf{H}^{+}]: \, [\mathsf{H}^{+}] &= \underbrace{\mathsf{Kw}}_{[\mathsf{OH}^{-}]} \\ \textit{Use Kw} &= 10^{-14} \textit{ and the concentration of the base to find } [\mathsf{H}^{+}]. \\ \textit{Finally, calculate pH using: } \mathsf{pH} &= -\log_{10}[\mathsf{H}^{+}] \end{aligned}$

Titration Curves

A pH titration curve shows how the pH of a solution **changes** during an **acid-base reaction**. When an acid and base react, a **neutralisation point** is reached, identified as a large **vertical section** of the graph, through the neutralisation or equivalence point.

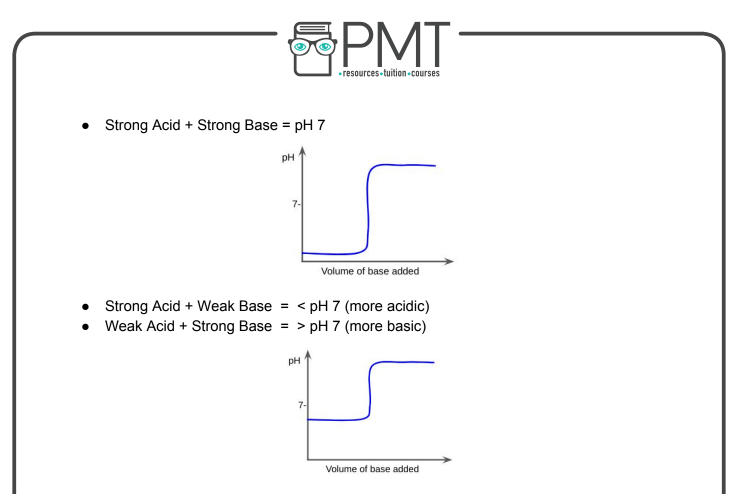
To obtain a pH titration curve, alkali is slowly added to an acid (or vice versa) and the pH is regularly measured with a pH probe. The smaller the added volumes, the more accurate the curve produced.

Example: The red dot indicates the equivalence point



For a strong acid-strong base reaction, this neutralisation point occurs around **pH 7**. Other combinations of strong and weak acids and bases results in a **different neutralisation point**:

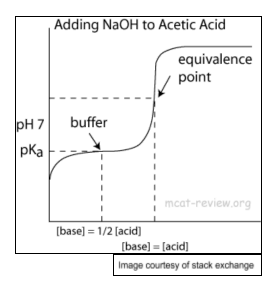




• Weak Acid + Weak Base = normally pH 7 but hard to determine

Calculating Ka from Titration Curves

The vertical region of a titration curve is the **equivalence point**. At **half-equivalence point** (when half the volume of acid/base at the equivalence point is added) the **pH is equal to pKa**.



Therefore, by reading the pH at half the equivalence point, Ka can be easily calculated from pKa.





Indicators

Specific chemical indicators have to be used for specific reactions as they can **only indicate a pH change within a certain range**.

The two most common indicators used at A-Level are methyl orange and phenolphthalein:

Methyl Orange - used for reactions with a **more acidic** neutralisation point. - red in acids and turns **yellow** at the neutralisation point.

Phenolphthalein - used for reaction with a **more basic** neutralisation point. - pink in alkalis and turns **colourless** at the neutralisation point.

| Indicator | pH at colour change | Colour in acid | Colour in base |
|-----------------|---------------------|----------------|----------------|
| Methyl orange | 3-5 | Red | Yellow |
| Phenolphthalein | 8-10 | Colourless | Pink |
| Litmus | 5-8 | Red | Blue |

It is important that, depending on the strength of the titration reactants, the **correct indicator is selected** so that the colour change occurs at neutralisation.

Buffer Action

A buffer solution is formed from a weak acid and its salt or a weak acid and a base. This produces a mixture containing H^+ ions and a large reservoir of OH^- ions which helps to resist any change in pH. Therefore, a buffer solution is defined as:

A solution which is able to resist changes in pH when small volumes of acid or base are added.

The large reservoir of OH- ions allows the **ratio of acid to base** in the mixture to be kept almost **constant**.

Consider the following buffer solution:

 $NH_3 + H_2O \stackrel{\scriptstyle \Rightarrow}{} NH_4^+ + OH^-$

The OH⁻ concentration will increase if a small amount of base is added, making the solution more basic. The extra OH⁻ ions will react with the NH_4^+ ions, to form the original reactants. Therefore, the equilibrium will shift to the left to remove the OH⁻ ions and stop the pH from changing largely.

S www.pmt.education





Buffer Calculations

These are long calculations that use acid-base calculations. There are two types:

Acid + Base - Find the number of moles of each species.

- Calculate their concentration when at equilibrium using the total volume.
- Use Ka to find [H+] and therefore pH.

Acid + Salt - Find the moles of the salt.

- Use Ka to find pH.

Example:

A buffer solution contains 0.35 mol dm⁻³ methanoic acid and 0.67 mol dm⁻³ sodium methanoate. For methanoic acid, Ka = 1.6×10^{-4} mol dm⁻³. Find the pH of this buffer.

We **assume** that the **sodium methanoate completely dissociates** so that the equilibrium concentration of HCOO⁻ is the same as the initial concentration of HCOO⁻Na⁺. Similarly, HCOOH only slightly dissociates so we assume that the **equilibrium concentration** is equal to the **initial concentration**.

1. First, find the expression for Ka for methanoic acid HCOOH \Rightarrow H⁺ + HCOO⁻

$$Ka = [H^+] \times [HCOO^-]$$
$$[HCOOH]$$

2. Rearrange the expression to find $[H^*]$

 $[H^{+}] = \frac{\text{Ka x [HCOOH]}}{[HCOO^{-}]}$ $[H^{+}] = 1.6 \times 10^{-4} \times (0.35/0.67) = 8.4 \times 10^{-5}$

3. Convert $[H^+]$ to pH

$$pH = -log_{10}(8.4 \times 10^{-5}) = 4.08$$





Adding Small Volumes

The pH of a buffer solution doesn't change much but will change in the order of 0.1 or 0.01 units of pH when a small volume of acid or base is added.

Adding small amounts of **acid** (H^+) **increases the concentration** of the acid in the buffer solution meaning the overall solution will get slightly **more acidic**.

Adding small amounts of **base (OH**) **decreases the concentration** of acid in the buffer solution meaning the overall solution will get slightly more **basic**.

Uses of Buffers

Buffer solutions are common in **nature** in order to keep systems regulated. This is important as **enzymes** or reactions in living organisms often require a **specific pH**, which can be maintained using a buffer solution.

Another important buffer in nature is found in the human circulatory system. The **pH of human blood** is maintained in a buffer between **carbonic acid and bicarbonate ions**. These ions **neutralise any acidic substances** that enter the bloodstream, converting them to carbonic acid and water.

