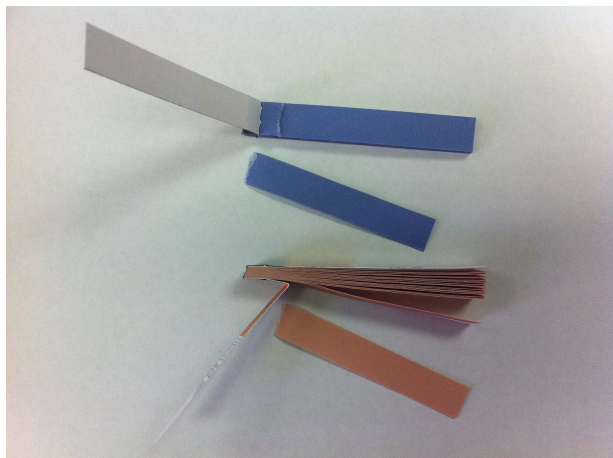


# Acids and Bases

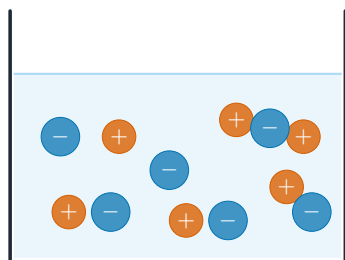
## AP Chemistry

### Introduction to Acids and Bases



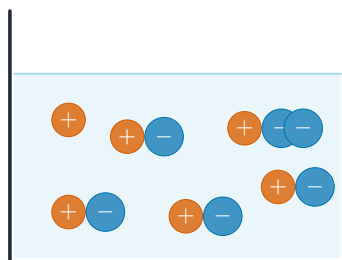
*Litmus paper: acid-base indicators change colour with  $H^+$  concentration (pH)*

By the **Brønsted–Lowry** definition, an **acid** 酸 is a proton ( $H^+$ ) **donor** and a **base** 碱 is a proton **acceptor**. When an acid donates a proton it becomes its **conjugate base** 共轭碱; a base gaining a proton becomes its **conjugate acid** 共轭酸. Water is **amphoteric** – it can act as either.



**strong acid**

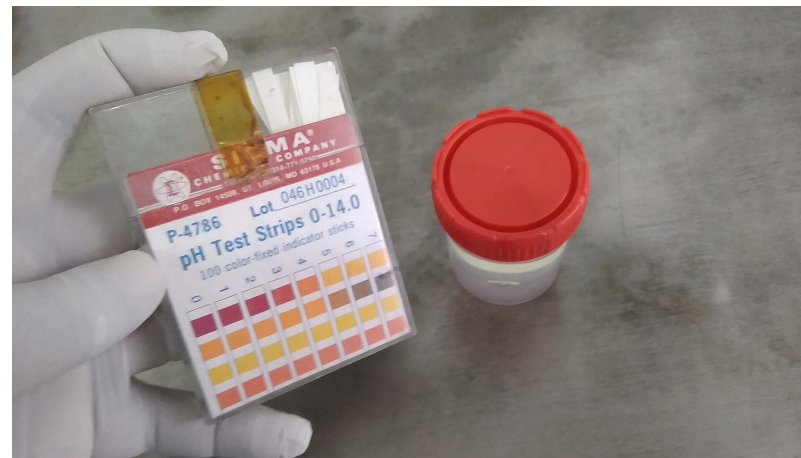
fully dissociated: many ions



**weak acid**

mostly molecules: few ions

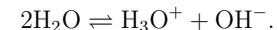
*A strong acid is fully dissociated; a weak acid is only partly dissociated*



*pH test strips: acid-base strength and concentration set the pH you measure*

### The Autoionization of Water

Even pure water conducts a tiny current, because water molecules react with each other in a reaction called **autoionization** 自偶电离:



One water molecule passes a proton to another, making a **hydronium ion** 水合氢离子 ( $H_3O^+$ , the ion we loosely write as  $H^+$ ) and a hydroxide ion. This equilibrium has its own constant, the **ion-product constant of water** 水的离子积:

$$K_w = [H_3O^+][OH^-] = 1.0 \times 10^{-14}$$

at 25 °C. Defining  $pOH = -\log[OH^-]$  and taking  $-\log$  of the  $K_w$  expression gives the rule the rest of this topic leans on:

$$pH + pOH = pK_w = 14.0$$

(again at 25 °C). Know one of pH or pOH and you get the other by subtracting from 14.

In pure water the two ions form in equal numbers, so  $[H_3O^+] = [OH^-]$  and  $pH = pOH = 7.0$ . This is what **neutral** 中性 means. Acidic means  $[H_3O^+] > [OH^-]$  (pH below 7); basic is the reverse.

**Worked example.** A solution has  $[H_3O^+] = 2.0 \times 10^{-3}$  M. Find  $[OH^-]$ . Because  $K_w$  always holds in water,

$$[OH^-] = \frac{K_w}{[H_3O^+]} = \frac{1.0 \times 10^{-14}}{2.0 \times 10^{-3}} = 5.0 \times 10^{-12} \text{ M.}$$

The hydroxide concentration is far smaller than the hydronium, confirming the solution is acidic.

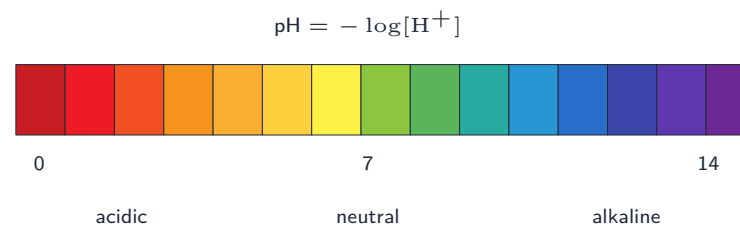
$K_w$  is **temperature dependent**: autoionization is endothermic, so heating water shifts it forward and raises  $K_w$ . At 50 °C,  $K_w > 1.0 \times 10^{-14}$ , so neutral water has  $\text{pH} = \text{pOH} < 7.0$ . It is still neutral, because the ion concentrations are still equal – neutral means equal ions, not a pH of exactly 7.



*pH strips: water autoionizes so pure water is pH 7; acids and bases shift  $H^+$  and  $OH^-$ .*

## pH and pOH of Strong Acids and Bases

The **pH** pH 值 scale measures acidity:  $\text{pH} = -\log[\text{H}^+]$ , with  $\text{pH} + \text{pOH} = 14$  at 25 °C. A **strong acid** 强酸 or **strong base** 强碱 dissociates **completely**, so its ion concentration equals its concentration – read pH directly. Lower pH means more acidic.



The *pH* scale: *pH* is the negative log of the hydrogen-ion concentration

**Worked example.** Find the pH of 0.010 M HCl. Because HCl is a strong acid it is fully dissociated, so  $[\text{H}^+] = 0.010 \text{ M}$  and

$$\text{pH} = -\log(0.010) = 2.0.$$

For a strong **base** like 0.010 M NaOH,  $\text{pOH} = 2.0$ , so  $\text{pH} = 14 - 2.0 = 12.0$ .

## Weak Acid and Base Equilibria

A **weak acid** 弱酸 only **partly** dissociates, described by an equilibrium constant  $K_a$  (larger  $K_a$  = stronger weak acid); a weak base has  $K_b$ . For any conjugate acid–base pair the two are linked through water,  $K_a \times K_b = K_w$ , so  $\text{p}K_a + \text{p}K_b = 14$  – a stronger acid always has a weaker conjugate base. Because dissociation is small, find  $[\text{H}^+]$  with an ICE table and the  $K_a$  expression, often using the small- $x$  approximation.

**Worked example.** Find the pH of 0.10 M acetic acid,  $K_a = 1.8 \times 10^{-5}$ . Let  $x = [\text{H}^+]$  at equilibrium; the ICE table gives  $K_a = \frac{x^2}{0.10 - x} \approx \frac{x^2}{0.10}$  (small- $x$  approximation), so

$$x = \sqrt{K_a \times 0.10} = \sqrt{1.8 \times 10^{-6}} = 1.3 \times 10^{-3} \text{ M} \Rightarrow \text{pH} = -\log(1.3 \times 10^{-3}) = 2.9.$$

The weak acid is far less acidic than a strong acid of the same concentration (which would be pH 1.0).

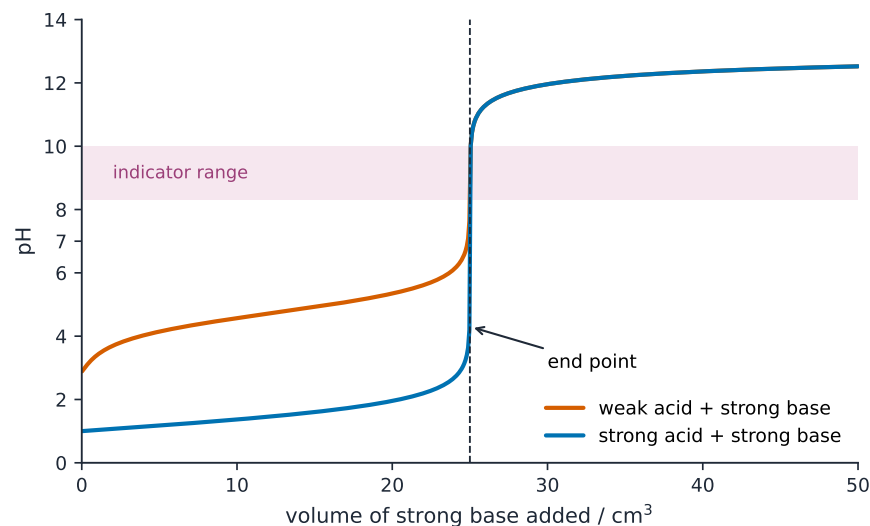
## Acid-Base Reactions and Buffers

A **buffer** 缓冲溶液 resists pH change. It contains a weak acid **and** its conjugate base (or a weak base and its conjugate acid) in comparable amounts. Added acid is neutralized by the conjugate base, and added base by the weak acid, so pH barely moves.

## Acid-Base Titrations

A **titration curve** 滴定曲线 plots pH as base (or acid) is added. Key points: the **equivalence point** 等当点 (moles of acid = moles of base – a steep jump), and the **half-equivalence point**, where  $\text{pH} = \text{p}K_a$  (half the weak acid is converted, so a buffer is at its center).

An **acid-base indicator** 酸碱指示剂 is itself a weak acid whose protonated and deprotonated forms have **different colours**, so its colour responds to pH. You pick an indicator whose colour change happens near the titration's equivalence-point pH, so the colour flips just as the reaction completes.



*A titration curve has a steep jump near the equivalence point*

## Molecular Structure of Acids and Bases

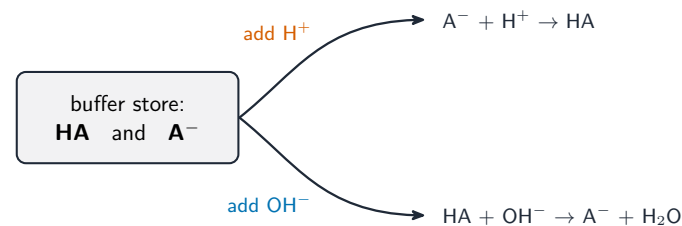
Strength has structural roots. An acid is stronger when its **conjugate base is more stable** – for example, more electronegative atoms or resonance spreading the negative charge stabilize it. For oxoacids, more oxygen atoms on the central atom means a stronger acid.

## pH and pK<sub>a</sub>

$\text{p}K_a = -\log K_a$ : a **smaller**  $\text{p}K_a$  means a **stronger** acid. Comparing pH to  $\text{p}K_a$  tells you the dominant form: below  $\text{p}K_a$  the acid form dominates; above it, the conjugate base form dominates.

## Properties of Buffers

A buffer works best when the weak acid and conjugate base concentrations are **similar** (pH near  $\text{p}K_a$ ). Choose a buffer whose  $\text{p}K_a$  is close to the target pH. Diluting a buffer barely changes its pH, because the acid-to-base **ratio** stays the same.



either way, the pH barely changes

*A buffer resists pH change by mopping up added acid or base*

## The Henderson-Hasselbalch Equation

For a buffer, the pH follows from the acid-to-base ratio:

$$\text{pH} = \text{p}K_a + \log \frac{[\text{A}^-]}{[\text{HA}]}$$

When the concentrations are equal, the log is zero and  $\text{pH} = \text{p}K_a$ . Use it to design a buffer or find its pH quickly.

**Worked example.** A buffer holds 0.20 M acetic acid ( $pK_a = 4.74$ ) and 0.30 M acetate. Its pH is

$$\text{pH} = 4.74 + \log \frac{0.30}{0.20} = 4.74 + \log(1.5) = 4.74 + 0.18 = 4.92.$$

A little more conjugate base than acid pushes the pH just above  $pK_a$ , as the equation predicts.

## Buffer Capacity

**Buffer capacity** 缓冲容量 is how much acid or base a buffer can absorb before its pH changes sharply. It is greatest when the components are **concentrated** and in roughly **equal** amounts. Once one component is used up, the buffer fails.

## pH and Solubility

The solubility of a salt with a basic anion **rises** in acidic solution: the added  $\text{H}^+$  reacts with the anion, removing it from the solubility equilibrium and pulling more solid to dissolve (Le Chatelier applied to  $K_{sp}$ ). So pH can control whether an ionic solid dissolves.

## Exam tips

- **Neutral means equal ions, not pH 7.** In pure water  $[\text{H}_3\text{O}^+] = [\text{OH}^-]$ ; since  $K_w$  rises with temperature, neutral pH drifts below 7 above 25 °C. Use  $K_w = [\text{H}_3\text{O}^+][\text{OH}^-] = 1.0 \times 10^{-14}$  (25 °C) to convert between  $[\text{OH}^-]$  and  $[\text{H}_3\text{O}^+]$ .
- $\text{pH} = -\log[\text{H}^+]$  is **logarithmic** — each unit is a ten-fold change in  $[\text{H}^+]$ .
- A **strong** acid/base dissociates completely (so  $[\text{H}^+] = \text{concentration}$ ); a **weak** one only partly, needing  $K_a$ .
- A **buffer** contains a weak acid **and** its conjugate base together; it resists pH change by mopping up added acid or base.
- On a titration curve the **equivalence point** is the steep jump; the **half-equivalence point** has  $\text{pH} = pK_a$ .
- A smaller  $pK_a$  means a stronger acid.