Shocking revelations: a conductometric titration of barium hydroxide with sulfuric(VI) acid

Strengthen students’ mental models of the particles reacting in solution with this conductometric titration of barium hydroxide with sulfuric(VI) acid

Kit
- 25 cm³ of 0.1 mol dm⁻³ sulfuric(VI) acid
- 50 cm³ of 0.1 mol dm⁻³ barium hydroxide (irritating to skin and eyes)
- Approx 225 cm³ of deionised water
- Magnetic stirrer
- Clamp and stand
- Burette and burette clamp
- 500 cm³ beaker
- Conductivity probe (see tips below)

Preparation
Wear eye protection. Load the sulfuric(VI) acid into the beaker and dilute to approx 250 cm³ with deionised water, add a stirrer bead and place on the magnetic stirrer. Clamp a conductivity probe to hold the tip in the middle of the solution and place a burette loaded with the barium hydroxide solution above, ensuring the probe isn’t directly below the jet.

You might be tempted to alter the concentrations or miss out the diluting step. Don’t be.

In front of the class
You may wish to begin by demonstrating the conductivity of deionised water and each of the reacting solutions. Warned about electrocution hazards, students often believe that pure water is a good conductor. However, the LED of CLEAPSS conductivity indicator will not light, a conductivity probe should give a value below 10 μS/cm and a multimeter’s resistance reading is likely to give an error because the value is too great. Meanwhile 0.1 M acids and bases will give readings in the tens of thousands of μS/cm. If needed, this step will allow you to choose an appropriate range setting for your probe.

Having introduced the chemistry behind the reaction, simply open the tap and watch for the next two minutes as the burette empties into the solution below. For optimum results with quantitative data, let the tap run half open, as there is a lag of...
a few seconds between the addition of solution and the completion of precipitation. The conductivity will decrease until half the barium hydroxide has been delivered and where the reading will be almost zero, before it increases until the burette is emptied.

**Teaching goal**

The point of this demonstration is to strengthen students’ mental models of the particles reacting in solution and how this can be represented in chemical equations. A common misconception among students is that ionic substances behave as molecules leading to difficulties understanding dissociation in solution and later ionic equations.

This demonstration emphasises the ionic and molecular nature of one of the first ionic equations students are likely to meet: that of a neutralisation reaction in which hydroxide ions react with hydrogen ions (equation 1) or perhaps later, more accurately, hydronium ions (equation 2) to produce neutral water molecules.

**Equation 1**: \( \text{H}^+ (aq) + \text{OH}^- (aq) \rightarrow \text{H}_2\text{O} (l) \)

**Equation 2**: \( \text{H}_3\text{O}^+ (aq) + \text{OH}^- (aq) \rightarrow 2\text{H}_2\text{O} (l) \)

The choice of barium hydroxide and sulfuric(VI) acid allows the change in conductivity of the neutralisation reaction to unfold without confusion from other ions which precipitate out of solution (equation 3).

**Equation 3**: \( \text{Ba}^{2+} (aq) + \text{SO}_4^{2-} (aq) \rightarrow \text{BaSO}_4 (s) \)

Students may be familiar with the insolubility of barium sulfate(VI) as it is the product of the sulfate(VI) anion test with acidified barium chloride.
If you simply run 50 cm$^3$ of one 0.1 M solution into 25 cm$^3$ of the other at the same concentration, you will see a curve dipping below a linear trendline before the equivalence point, and then deviating above it afterwards. This is due to dilution factors – the final volume is three times the volume of the initial even if it should have the same number of ions within. These factors are minimised in the demo as described by the initial dilution of the acid.

The second half of the curve is more stretched as the pressure at the burette jet decreases as the contents above empty out. There is also a lag in the rate of increase of conductivity associated with the delay in precipitation of barium and sulfate(VI) ions out of solution.

While it would be too time-consuming for a classroom demonstration, keen students could be invited to collect values from a more dilute reaction and plot these against volume rather than time. This gives a fantastic linear relationship (Figure 2), but one question remains: why is the graph still not symmetric? The answer to this will have to wait until the next issue when we introduce some other ions to the mix.

**Disposal**

Filter the barium sulfate(VI) and dispose of the residue in the normal refuse. You can rinse filtrate down the sink.