

## 1 – Collision theory

<b>Collision theory</b>	For a reaction to occur, particles must <b>collide</b> with <b>sufficient energy</b> .
<b>Activation energy</b>	<b>Minimum</b> amount of <b>energy</b> that reactant particles must have when they <b>collide</b> in order to <b>react</b> .

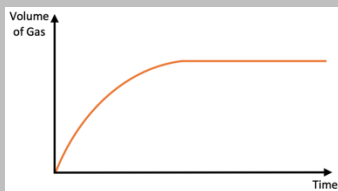
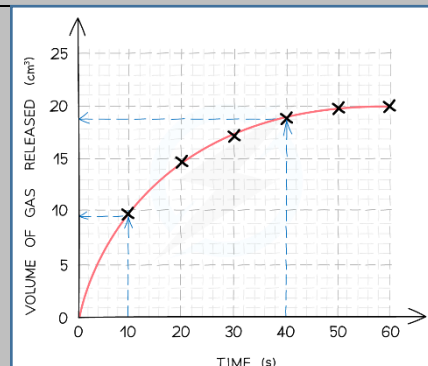
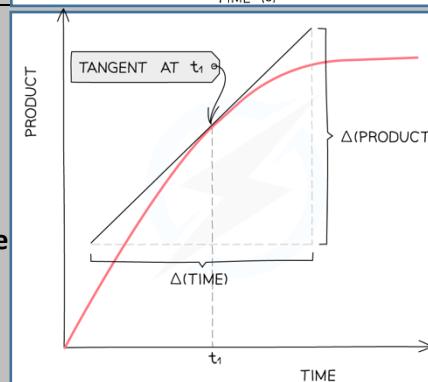
## 2 – Factors affecting rate of reaction

<b>Temperature</b>	<b>Increase</b> temperature → particles have <b>more energy</b> so move <b>faster</b> → collide more <b>frequently</b> → more collisions have <b>activation energy</b> → greater chance of <b>successful collisions</b> .	
<b>Concentration/ Pressure</b>	<b>Increase concentration</b> of solution → more particles in same <b>volume</b> → particles collide more <b>frequently</b> .	<b>Increase pressure</b> of gas → same number of particles <b>occupy smaller space</b> → particles are <b>closer</b> → collide more <b>frequently</b> .
<b>Surface area</b>	Breaking a solid into <b>smaller pieces</b> <b>increases surface area to volume ratio</b> → particles have more <b>area to collide with</b> → <b>collisions</b> occur more <b>frequently</b> .	
<b>Catalyst</b>	A substance that <b>speeds up rate</b> of reaction <b>without being used</b> up in the reaction. They work by <b>decreasing</b> the <b>activation energy</b> needed by providing an <b>alternative reaction pathway</b> .	

## 3 – Measuring rates of reaction

<b>Calculation</b>	$\text{mean rate} = \frac{\text{quantity of reactant used}}{\text{time taken}} \text{ units = g/s}$ $\text{mean rate} = \frac{\text{quantity of product formed}}{\text{time taken}} \text{ units = g/s or cm}^3/\text{s}$
<b>Measuring mass</b>	Measure mass at the <b>start</b> and <b>end</b> of a reaction to measure <b>mass of gas</b> produced. Not suitable for <b>hydrogen</b> due to small mass.
<b>Measuring volume</b>	<b>Volume of gas produced</b> can be measured using a <b>gas syringe</b> or an <b>upside-down measuring cylinder</b> filled with water.
<b>Precipitation</b>	Can be used when <b>initial solution</b> is <b>transparent</b> , and the <b>product</b> is a <b>precipitate</b> , so the solution becomes <b>opaque</b> . The <b>cloudiness</b> of the solution is called the ' <b>turbidity</b> '. Observe a <b>mark</b> through the solution and measure <b>time</b> taken to <b>disappear</b> .

## 4 – Rate of reaction graphs

<b>Shape of graphs</b>	The <b>steeper</b> the line, the <b>faster</b> the rate. As the line gets <b>less steep</b> , the rate is <b>slowing down</b> . A <b>horizontal line</b> means the reaction is <b>finished</b> .	
<b>Calculating mean rate</b>	$\text{mean rate} = \frac{\text{change in } y}{\text{change in } x}$ e.g $y = 19 - 8.5 = 10.5$ $x = 40 - 10 = 30$ $10.5/30 = 0.35 \text{ cm}^3/\text{s}$	
<b>Calculating instantaneous rate</b>	A <b>tangent</b> need to be drawn to calculate <b>change in x and y</b> then the rate of reaction at a <b>particular point</b> .  1. Draw a <b>tangent</b> at the <b>time</b> the rate needs to be <b>calculated</b> . 2. Calculate the <b>gradient</b> of the tangent: <b>change in product ÷ change in time</b> 3. Gradient = rate of reaction.	

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Chemistry C6 – Rate and extent of chemical change

## 5 – Required practical methods

<b>Observing colour change</b>	1. 10cm <sup>3</sup> of <b>sodium thiosulfate</b> and 40cm <sup>3</sup> of <b>water</b> into <b>conical flask</b> (diluted to 8g/dm <sup>3</sup> ). 2. Place flask on a <b>white tile</b> with <b>black cross</b> . 3. Add 10cm <sup>3</sup> of dilute <b>HCl</b> and start <b>timer</b> . 4. Gently <b>swirl</b> the flask and stop timer when the <b>cross is no longer visible</b> . 5. <b>Repeat</b> using <b>different concentrations</b> of sodium thiosulfate. 6. <b>Repeat</b> whole investigation twice more and <b>calculate a mean</b> for each concentration.
<b>Measuring volume of gas produced</b>	1. Add 50cm <sup>3</sup> of <b>1.0mol/dm<sup>3</sup> HCl</b> to a <b>conical flask</b> . 2. Fit a <b>bung</b> and <b>delivery tube</b> to the conical flask. 3. Half fill a <b>trough</b> with <b>water</b> . 4. Fill a <b>measuring cylinder</b> with <b>water</b> , ensure it stays full and <b>invert</b> it into the <b>water trough</b> , with the <b>delivery tube</b> positioned into it. 5. Add a <b>3cm</b> strip of <b>magnesium</b> to the <b>conical flask</b> and <b>replace</b> the bung, then start the <b>timer</b> . 6. Record the <b>volume</b> of hydrogen <b>gas</b> given off in <b>10 second</b> intervals. 7. <b>Repeat</b> using <b>1.5mol/dm<sup>3</sup> acid</b> .

## 6 - Reversible reactions and equilibrium

<b>Reversible reaction</b>	A reaction where the <b>products</b> can react to produce the <b>original reactants</b> . Shown by: $\rightleftharpoons$	
<b>Equilibrium</b>	The point at which the <b>rates</b> of the <b>forward</b> and <b>backward</b> reactions in a <b>reversible</b> reaction are the <b>same</b> . The amounts of <b>reactants</b> and <b>products</b> in the reaction <b>container don't change</b> .	
<b>Positions of equilibrium</b>	If the <b>equilibrium</b> lies to the <b>right</b> : the <b>concentration</b> of <b>products</b> is <b>higher</b> than the <b>reactants</b> .	If the <b>equilibrium</b> lies to the <b>left</b> : the <b>concentration</b> of <b>reactants</b> is <b>greater</b> than the <b>products</b> .
<b>Changing the direction</b>	<b>Heating</b> a <b>reversible</b> reaction moves it in the <b>endothermic</b> direction. <b>Cooling</b> a <b>reversible</b> reaction moves it in the <b>exothermic</b> direction.	

## 7 – Le Chatelier's principle (HT)

<b>Le Chatelier's principle</b>	If you <b>change</b> the <b>conditions</b> of a <b>reversible</b> reaction, at <b>equilibrium</b> , the system will try and <b>counteract</b> the <b>change</b> .
<b>Changing temperature</b>	<b>Decrease</b> temperature → reaction moves in <b>exothermic</b> direction. <b>Increase</b> temperature → reaction moves in <b>endothermic</b> direction.
<b>Changing concentration</b>	<b>Increase concentration</b> of <b>reactants</b> → system makes more products (the <b>forward reaction increases</b> ). <b>Decrease concentration</b> of <b>products</b> → system tries to make more products by reducing reactants ( <b>forward reaction increases</b> ).
<b>Changing pressure</b>	Only affects <b>equilibrium</b> involving <b>gases</b> . <b>Increase</b> pressure → equilibrium tries to reduce it and moves to the side with <b>fewer molecules</b> . <b>Decrease</b> pressure → equilibrium tries to increase it and moves to the side with <b>more molecules</b> .

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### Chemistry C6 – Rate and extent of chemical reactions.

