

16.1 Rate Expression & Reaction Mechanism

Contents

- ***** 16.1.1 Rate Equations
- ***** 16.1.2 Rate Constants
- ***** 16.1.3 Graphical Representations
- **** 16.1.4 Rate Determining Step
- ***** 16.1.5 Mechanism Problems

16.1.1 Rate Equations

Rate Equations

- The rate of reaction can be found by:
	- Measuring the decrease in the concentration of a reactant overtime
	- Measuring the increase in the concentration of a product overtime
	- The units for rate of reaction are mol dm⁻³ s⁻¹

Rate of Reaction

The following general reaction will be used as an example to study the rate of reaction

$D (aq) \rightarrow E (aq) + F (g)$

The rate ofreaction at different concentrations of Dis measured and tabulated

Rate of reactions table

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A directly proportional relationship between the rate of reaction and concentration of D is observed when the results are plotted on a graph:

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Page 3 of 27

- **Products and catalysts may feature in rate equations**
- **Intermediates do not feature in rate equations**

Order of reaction

- The order of a reactant shows how the concentration of a chemical, typically a reactant, affects the rate ofreaction
- \blacksquare It is the power to which the concentration of that reactant is raised in the rate equation
- The order can a positive, negative or fractional value
	- **•** Orders that are a fraction suggest that the reaction involves multiple steps

Q Examiner Tip

- In an exam, you may be presented with information about a reaction that uses orders that are fractions. Examples include:
	- The decomposition of ethanal Rate = $k[\text{CH}_3\text{CHO}]^{3/2}$
	- The reaction of hydrogen and bromine Rate = $k[H_2][Br_2]^{1/2}$
	- The reaction between carbon monoxide and chlorine Rate = $k[\mathrm{CO}]^2[\mathrm{C}l_2]^{1/2}$
- However,for calculations only values of 0,1 or 2 need to be considered
- \blacksquare When the order of reaction with respect to a chemical is 0
	- Changing the concentration of the chemical has no effect on the rate of the reaction
	- \blacksquare Therefore, it is not included in the rate equation
- When the order of reaction with respect to a chemical is 1
	- The concentration ofthe chemical is directly proportionalto the rate ofreaction, e.g. doubling the concentration of the chemical doubles the rate of reaction
	- The chemical is included in the rate equation
- \blacksquare When the order of reaction with respect to a chemical is 2
	- The rate is directly proportional to the square of the concentration of that chemical, e.g. doubling the concentration of the chemical increases the rate of reaction by a factor of four
	- The chemical is included in the rate equation (appearing as a squared term)
- The overall order of reaction is the sum of the powers of the reactants in a rate equation

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The chemical equation forthe thermal decomposition of dinitrogen pentoxide is:

$$
2N_2O_5(g) \rightarrow 4NO_2(g) + O_2(g)
$$

The rate equation forthis reaction is:

$$
Rate = k[N_2O_5(g)]
$$

- 1. State the order of the reaction with respect to dinitrogen pentoxide
- 2. Deduce the effect on the rate of reaction if the concentration of dinitrogen pentoxide is tripled

Answers

Answer 1:

- \blacksquare Dinitrogen pentoxide features in the rate equation, therefore, it cannot be order zero / 0
- The dinitrogen pentoxide is notraised to a power, which means thatit cannot be order 2 / second order
- Therefore, the order with respect to dinitrogen pentoxide must be order 1 / first order

Answer 2:

- Since the reaction is first order, the concentration of dinitrogen pentoxide is directly proportional to the rate
- This means that if the concentration of the dinitrogen pentoxide is tripled, then the rate of reaction will also triple

Worked example

The following equation represents the oxidation of bromide ions in acidic solution

 BrO_3^- (aq) + 5Br⁻ (aq) + 6H⁺ (aq) \rightarrow 3Br₂ (l) + 3H₂O (l)

The rate equation forthis reaction is:

$$
Rate = k[BrO3^-(aq)][Br^-(aq)][H^+(aq)]
$$

- 1. State the overall order of the reaction
- 2. Deduce the effect on the rate of reaction if the concentration of bromate ions is doubled and the concentration of bromide ions is halved

Answers

Answer 1:

Page 5 of 27

- Allthree reactants feature in the rate equation butthey are notraised to a power,this means that the order with respect to each reactant is order 1/ first order.
- The overall order of the reaction is $1+1+1=3$ or third order.

Answer 2:

- \blacksquare Since each reactant is first order, the concentration of each reactant is directly proportional to the effect that it has on rate
- If the concentration of the bromate ion is doubled, then the rate of reaction will also double
- \blacksquare If the concentration of the bromide ion is halved then the rate will also halve
- Therefore, there is no overall effect on the rate of reaction one change doubles the rate and the other change halves it

Solving Rate Equation Problems

Deducing Rate Equations

The following reaction will be used to deduce the rate equation, using experimental data

 $(CH₃)₃CBr + OH⁻ \rightarrow (CH₃)₃COH + Br⁻$

Table to show the experimental data of the above reaction

- \blacksquare To derive the rate equation for a reaction, you must first determine all of the orders with respect to each of the reactants
- **This can be done using the tabulated data provided**
- \blacksquare Take the reactants one at a time and find the order with respect to each reactant individually
- 1. Identify two experiments where the concentration of one reactant changes, butthe concentrations of all other reactants are constant
- 2. Calculate what happens to the concentration
- 3. Calculate what happens to the rate ofreaction
- 4. Deduce the order of reaction with respect to that chemical
- 5. Repeatthis for all ofthe reactants, one at a time, until you have determined the order with respectto all reactants

Order with respect to $[(CH₃)₃CBr]$

- 1. In experiments 1 and 2, the concentration of $(CH_3)_3$ CBr changes while the concentration of OH⁻ remains constant
- 2. The $[(CH₃)₃CBr]$ has doubled
- 3. The rate of the reaction has also doubled
- 4. Therefore, the order with respect to $[(CH₃)₃CBr]$ is 1 (first order)
	- [Change in concentration]^{order} = change in rate
		- $[2]$ ^{order} = 2
	- $[2]¹ = 2$

Page 7 of 27

Order with respect to [OH⁻]

- 1. In experiments 1 and 3, the concentration of OH⁻ changes while the concentration of (CH₃)₃CBr remains constant
- 2. The $[OH^-]$ has doubled
- 3. The rate of the reaction has increased by a factor of 4
- 4. Therefore, the order with respect to [$\mathsf{OH}^\mathsf{-1}$ is 2 / second order
	- [Change in concentration]^{order} = change in rate
	- $[2]$ ^{order} = 4
	- $[2]² = 2$

Building the rate equation

- **Once the order with respect to all of the reactants is known the rate equation can be constructed**
	- E Zero order reactants are not included in the rate equation
	- First order reactants are included in the rate equation they do notrequire a power
	- Second order reactants are included in the rate equation they are raised to the power of 2
- So,forthis reaction the rate equation will be:

Rate = k [(CH₃)₃CBr] [OH⁻]²

Q Examiner Tip

Examiners will often give concentration and rate data in standard form to test your mathematical skills!Take your time because it is easy to make a mistake - the most common one is failing to notice a factor of ten, e.g. one rate value is x10⁻⁴ while the rest are x10⁻³

16.1.2 Rate Constants

Rate Constants

 \blacksquare The rate constant, (k) , of a reaction can be calculated using the initial rates and the rate equation

Calculating the rate constant from the initial rate

- The reaction of sodium carbonate with chloride ions (from hydrochloric acid) to form sodium chloride will be used as an example to calculate the rate constant from the initial rate and initial concentrations
- **The reaction and rate equation are as follows:**

 Na_2CO_3 (aq) + 2Cl⁻ (aq) + 2H⁺ (aq) \rightarrow 2NaCl (aq) + CO₂ (g) + H₂O (l)

$$
RATE = k[Na,CO,][Cl^{-}]
$$

THIS REARRANGES TO GIVE:

The progress of the reaction can be followed by measuring the initial rates of the reaction using various initial concentrations of each reactant

Experimental results of concentrations & initial rates table

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 \blacksquare To find the rate constant (k):

Substitute the values of one of the experiments to find k (for example measurement 1)

Page 9 of 27

$$
k = \frac{4.38 \times 10^{-6}}{(0.0250) \times (0.0125)}
$$

$$
k = 1.40 \times 10^{-2}
$$

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- \blacksquare The values of **measurement 2** or 3 could also have been used to find k
	- They all give the same result of 1.40×10^{-2}

Calculating units

- \blacksquare When you are asked to calculate the rate constant, k , for a reaction you must also be able to deduce the units
- This is done by replacing the values in the rearranged rate equation with the units of that value
- The units can then be combined or cancelled as required
- For example, to calculate the units for the above reaction:

UNITS OF
$$
k = \frac{(mol\ dm^{-3} s^{-1})}{(mol\ dm^{-3})}
$$

\n
$$
= \frac{s^{-1}}{mol\ dm^{-3}}
$$

\n
$$
= mol^{-1}dm^{3} s^{-1}
$$

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Temperature and the rate constant, k

The following general reaction and rate equation will be used to discuss the effect of temperature on the rate constant, k:

$$
\mathsf{A} + \mathsf{B} \mathbin{\rightarrow} \mathsf{C} + \mathsf{D}
$$

Rate of reaction = $k[A][B]$

- \blacksquare The rate equation shows that rate of reaction depends on the rate constant, k , and the concentration ofthe reactants
	- As the rate ofreaction increases the rate constant will increase
- Increasing the temperature of a reaction increases the rate of a chemical reaction
	- **Remember: this does not necessarily increase the yield of a chemical reaction depending on** whether a reaction is endothermic or exothermic according to Le Châtelier's principle
- \blacksquare Therefore, increasing the temperature also increases the value of the rate constant, k , assuming that the concentration of the reactants remains unchanged
- An exponential relationship between the rate ofreaction and temperature is observed when seen on a graph:

Page 10 of 27

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Your notes

Relationship between temperature and rate constant, k

- The graph shows that the rate of reaction roughly doubles with an increase of 10 $^{\circ}$ C $\,$
	- This general relationship does not apply to all reactions
	- Also, it is not necessarily every 10 $^{\circ}$ C, the rate may double every 9 $^{\circ}$ C or 11 $^{\circ}$ C $\,$
	- The number of degrees needed to double the rate also changes gradually as temperature increases

16.1.3 Graphical Representations

Graphical Representations of Reaction Order

Reaction Order Using Concentration-Time Graphs

- In a zero-order reaction, the concentration of the reactant is inversely proportional to time
	- \blacksquare This means that the reactant concentration decreases as time increases
	- The graph is a straight line going down as shown:

Concentration-time graph of a zero-order reaction

- The gradient of the line is the rate of reaction
	- Calculating the gradient at different points on the graph, will give a constant value forthe rate of reaction
- \blacksquare When the order with respect to a reactant is 0, a change in the concentration of the reactant has no effect on the rate of the reaction
- Therefore:

 $Rate = k$

- \blacksquare This equation means that the gradient of the graph is the rate of reaction as well as the rate constant, k
- In a first-order reaction, the concentration of the reactant decreases with time
	- The graph is a curve going downwards and eventually plateaus:

Page 12 of 27

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Concentration-time graph of a first-order reaction

- n a second-order reaction, the concentration of the reactant decreases more steeply with time
	- The concentration of reactant decreases more with increasing time compared to a first-order reaction
	- **The graph is a steeper curve going downwards:**

Page 13 of 27

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Concentration-time graph of a second-orderreaction

Order of reaction from half-life

- The order of a reaction can also be deduced from its half-life $(t_{1/2})$
- \blacksquare For a zero-order reaction the successive half-lives decrease with time
	- \blacksquare This means that it would take less time for the concentration of reactant to halve as the reaction progresses
- \blacksquare The half-life of a first-order reaction remains constant throughout the reaction
	- \blacksquare The amount of time required for the concentration of reactants to halve will be the same during the entire reaction
- For a second-order reaction, the half-life increases with time
	- This means that as the reaction is taking place, ittakes more time forthe concentration of reactants to halve

Half-lives of zero, first and second-orderreactions

Q Examiner Tip

Make sure that you know the correct shapes forthe concentration-time graphs. It can be easy to confuse some concentration-time graphs with the following rate-concentration graphs, particularly;

- The straight line of a zero-order concentration-time graph with the straight line of a first-order rate-concentration graph.
- The curve of a first-order concentration-time graph with the curve of a second-order rateconcentration graph.

Reaction order using rate-concentration graphs

- In a zero-order reaction, the rate doesn't depend on the concentration of the reactant
	- \blacksquare The rate of the reaction therefore remains constant throughout the reaction

Page 14 of 27

- The graph is a horizontal line
- The rate equation is rate = k

Rate-concentration graph of a zero-orderreaction

- In a first-order reaction, the rate is directly proportional to the concentration of a reactant
	- \blacksquare The rate of the reaction increases as the concentration of the reactant increases
	- \blacksquare This means that the rate of the reaction decreases as the concentration of the reactant decreases when it gets used up during the reaction
	- \blacksquare The graph is a straight line
	- The rate equation is rate = $k[A]$

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reactant

Page 16 of 27

16.1.4 Rate Determining Step

Molecularity & Rate Determining Step

Rate-determining step & intermediates

- A chemical reaction can only go as fast as the slowest part of the reaction
	- So, the rate-determining step is the slowest step in the reaction
- If a reactant appears in the rate-determining step, then the concentration of that reactant will also appear in the rate equation
- For example, the rate equation for the reaction below is rate = $k[\mathrm{CH}_3\mathrm{Br}][\mathrm{OH}^-]$

$CH₃Br + OH⁻ \rightarrow CH₃OH + Br⁻$

- This suggests that both CH₃Br and OH⁻ take part in the slow rate-determining step
- Molecularity is the number of reactant particles that participate in the rate-determining step
- **Fig.** This reaction is a bimolecular reaction
	- **Unimolecular**: one species involved in the rate-determining step
	- **Bimolecular:** two species involved in the rate-determining step
- \blacksquare The intermediate is derived from substances that react together to form it in the rate-determining step
	- For example, for the reaction above the intermediate would consist of CH_3Br and OH $^{\text{-}}$

The intermediate formed from the species that are involved in the rate-determining step (and thus appearin the rate equation)

Q Examiner Tip

Intermediates do not feature in rate equations. Instead, the chemicals required to make the intermediate feature in the rate equation

Identifying the rate-determining step

- The rate-determining step can be identified from a rate equation given that the reaction mechanism is known
- For example, propane (CH₃CH₂CH₃) undergoes bromination under alkaline solutions

Page 18 of 27

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The overall reaction is:

$CH_3CH_2CH_3 + Br_2 + OH^- \rightarrow CH_3CH_2CH_2Br + H_2O + Br^-$

The reaction mechanism is:

Reaction mechanism for the bromination of propane under alkaline conditions

The rate equation is:

$Rate = k[CH_3CH_2CH_3][OH^-]$

- From the rate equation, it can be deduced that only $\mathrm{CH_3CH_2CH_3}$ and OH⁻ are involved in the ratedetermining step and not bromine (Br $_2$)
- $CH_3CH_2CH_3$ and OH⁻ are only involved in the first step of the reaction mechanism, therefore the ratedetermining step is:
	- $CH_3CH_2CH_3 + OH^- \rightarrow CH_3CH_2CH_2^- + H_2O$

Potential Energy Level Profiles

Single-step reactions

- When any reacting molecules collide with bond breaking and bond formation occurring, the interacting molecules will be in an unstable, high-energy state temporarily
	- This transition state will be of a higher energy than either the reactants or products and corresponds to the activation energy
- The exothermic reaction of hydrogen and iodine to form hydrogen iodide will be used to discuss how energy level diagrams relate to the rate-determining step

The potential energy level diagram for the exothermic reaction of hydrogen and iodine

- As the reaction proceeds, covalent bonds startto form between the hydrogen and iodine atoms from the hydrogen and iodine molecules
- At the same time, the covalent bonds within the hydrogen and iodine molecules grow longer and become weaker
- **This results in the transition state complex shown**

Page 20 of 27

The transition state complex for the reaction of hydrogen and iodine

- \blacksquare From this transition state, the bonds between the hydrogen and iodine atoms can continue to grow shorter and stronger resulting in the formation of hydrogen iodide
- I Alternatively, the bonds within the hydrogen and iodine molecules can grow shorter and stronger which would result in the formation of the reactants
- Hydrogen iodide will only form if the hydrogen and iodine molecules collide with kinetic energy greater than or equal to the activation energy
	- The molecules will also need to collide in the correct orientations
- The reaction, or elementary, step with the greatest activation energy will be the rate-determining step and can be used to determine the rate equation

Multi-step reactions

The exothermic reaction of nitrogen dioxide and fluorine to form nitryl fluoride (NO_2 F) will be used to relate rate equations and rate-determining steps to the energy level diagram of a multi-step reaction:

$2NO₂(g) + F₂(g) \rightarrow 2NO₂F(g)$

- This reaction is unlikely to occur in a single step as that would require three molecules to collide in the correct orientation and with sufficient kinetic energy
	- \blacksquare This is even less likely to occur as all three molecules are gaseous
- \blacksquare Experimental data shows that the rate equation for this reaction is:

$Rate = k[NO₂][F₂]$

- One proposed reaction mechanism forthis reaction involves the following elementary steps:
	- Step 1: $NO₂ + F₂ \rightarrow NO₂F + F$
	- Step 2: $NO₂ + F \rightarrow NO₂F$
- \blacksquare Step 1 must be the rate-determining step as it is the only step that has reactants matching the rate equation
	- **Therefore, on a potential energy level diagram the activation energy for step 1 will be greater than** the activation energy for step 2
- The transition state forthis multi-step reaction must be:

$NO₂ + NO₂F + F$

This can be deduced using the the equation for elementary step 1 and the overall equation

Page 21 of 27

- The overall equation states that two $NO₂$ react with one $F₂$
- Elementary step 1 uses one $NO₂$ to form $NO₂F + F$
- This leaves one $NO₂$ along with $NO₂F + F$
- This leads to the following potential energy level diagram:

Potential energy level diagram for the formation of nitryl fluoride

- Key points from the potential energy level diagram are:
	- The overall reaction is exothermic, as stated
	- The rate-determining step is the step that has the greatest activation energy
	- **There is a labelled energy level for the transition state**

Q Examiner Tip

Multi-step reactions where the first elementary step is not the rate-determining step prove more challenging in terms of potential energy level diagrams. Remember that the rate-determining step must have the greatest activation energy. Do not confuse greatest with the highest! When people talk about the highest they are normally referring to where the activation energy peak is, not the actual value itrepresents

Catalysts & Mechanisms

Identifying intermediates & catalyst

- When a rate equation includes a species that is not part of the chemical reaction equation then this species is a catalyst
- For example, the halogenation of butanone under acidic conditions Ē.
- **The overall equation for this reaction is:**

$$
\text{CH}_{3}\text{CH}_{2}\text{COCH}_{3} + I_{2} \xrightarrow{\text{H}^{+}} \text{CH}_{3}\text{CH}_{2}\text{COCH}_{2}I + \text{H}^{+}
$$

Die commonly accepted reaction mechanism is:

Reaction mechanism for the halogenation of butanone under acidic conditions

- The rate-determining step is the slowest step, identified as step 2
- However, the CH₃CH₂C(OH+)CH₃ cannot be used in the rate equation as it is an intermediate, not a reactant

Page 23 of 27

Reaction to form the intermediate found in the rate-determining step

Therefore, the $CH_3CH_2COCH_3$ and H+ that form the intermediate feature in the rate equation:

$Rate = k[CH_3CH_2COCH_3][H^+]$

- The H⁺ is **not** a reactant in the overall chemical equation but it does appear in the rate equation
	- H⁺ must, therefore, be a **catalyst**
	- This is also supported by the H⁺ being used in step 1 and forming as a product in step 2 $\,$

16.1.5 Mechanism Problems

Mechanism Problems

- **EXEC** Chemical kinetics can only suggest a reaction mechanism, they cannot prove it
	- **However, they can be used to disprove a proposed mechanism**
- **Elementary steps are the steps involved in a reaction mechanism**
	- **For example, in the following general reaction:**

 $A + B \rightarrow C + D$

The elementary steps could involve the formation of an intermediate:

Elementary step 1:
$$
A \rightarrow R + D
$$

Elementary step 2: $R + B \rightarrow C$

- It is important that the elementary steps for a proposed mechanism agree with the overall stoichiometric equation
	- **For example, combining the 2 elementary steps above gives the overall stoichiometric equation**

 $A + R + B \rightarrow R + C + D$

 $A + B \rightarrow C + D$

Worked example

Sulfur dioxide reacts with oxygen to form sulfurtrioxide

- 1. Propose a one step mechanism forthe above reaction
- 2. The above reaction is catalysed by the formation of nitrogen dioxide from nitrogen monoxide. Propose a two step mechanism for this reaction.

Answers

Answer 1:

- A one step reaction mechanism is simply the overall stoichiometric equation
- Therefore, the correct answer is $2SO_2 + O_2 \rightarrow 2SO_3$

Answer 2:

One of the two elementary steps for this two step mechanism can be taken from the question:

Elementary step 1: $2NO + O_2 \rightarrow 2NO_2$

Page 25 of 27

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The second elementary step must involve the reaction of the nitrogen dioxide formed with sulfur dioxide:

Elementary step 2: $NO_2 + SO_2 \rightarrow NO + SO_3$ (or $2NO_2 + 2SO_2 \rightarrow 2NO + 2SO_3$)

Q Examiner Tip

It is important that you check that the equations you are proposing for a reaction mechanism. They must add up to the overall stoichiometric equation, otherwise the proposed mechanism is wrong.

Predicting the reaction mechanism

- **The overall reaction equation and rate equation can be used to predict a possible reaction mechanism** of a reaction
	- This shows the individual reaction steps which are taking place
- For example, nitrogen dioxide ($NO₂$) and carbon monoxide (CO) react to form nitrogen monoxide (NO) and carbon dioxide $(CO₂)$
	- The overall reaction equation is:

$$
NO2(g) + CO(g) \rightarrow NO(g) + CO2(g)
$$

• The rate equation is:

Rate = $k \,[\text{NO}_2]^2$

- From the rate equation, it can be concluded that the reaction is zero-order with respect to CO (g) and second-order with respect to $NO₂(g)$
- This means that there are two molecules of $NO₂(g)$ involved in the rate-determining step and zero molecules of CO (g)
	- This means that in terms of molecularity, the rate determining step is bimolecular
- A possible reaction mechanism could therefore be:

Step 1:

 $2NO₂(g) \rightarrow NO(g) + NO₃(g)$ slow (rate-determining step)

Step 2:

 $NO₃(g) + CO(g) \rightarrow NO₂(g) + CO₂(g)$ fast

Overall:

 $2NO₂(g) + NO₃(g) + CO(g) \rightarrow NO(g) + NO₃(g) + NO₂(g) + CO₂(g)$

 $= NO₂(g) + CO(g) \rightarrow NO(g) + CO₂(g)$

Page 26 of 27

Q Examiner Tip

It is important that the elementary steps for a proposed mechanism also agree with the experimentally determined rate equation. The rate equation and the overall reaction must be related, i.e. the correct chemical species involved. Remember: There is no direct link between the orders in the rate equation and the stoichiometry of the overall equation. However, the rate equation can be derived directly from the rate determining step and its stoichiometry

Predicting the reaction order & deducing the rate equation

- The order of a reactant and thus the rate equation can be deduced from a reaction mechanism if the rate-determining step is known
- For example, the reaction of nitrogen oxide (NO) with hydrogen (H₂) to form nitrogen (N₂) and water

 $2NO(g) + 2H_2(g) \rightarrow N_2(g) + 2H_2O(l)$

The reaction mechanism for this reaction is:

Step 1:

 $NO(g) + NO(g) \rightarrow N_2O_2(g)$ fast

Step 2:

 $N_2O_2(g) + H_2(g) \rightarrow H_2O (l) + N_2O (g)$ slow (rate-determining step)

Step 3:

 $N_2O(g) + H_2(g) \rightarrow N_2(g) + H_2O(l)$ fast

- The second step in this reaction mechanism is the rate-determining step
- **The rate-determining step consists of:**
	- N_2O_2 which is formed from the reaction of two NO molecules
	- One H_2 molecule
- The reaction is, therefore, **second order** with respect to NO and fir<mark>st order</mark> with respect to H₂
- So, the rate equation becomes:

 $Rate = k [NO]^2 [H_2]$

The reaction is, therefore, third order overall

Q Examiner Tip

Intermediates in the mechanism cannot appear as substances in the rate equationThis is why you substitute the N_2O_2 in the above example. Step 1 shows that 2NO molecules are required to form the n ecessary N_2O_2

Page 27 of 27

