1.9 Rate Equations

Rate Equations

The rate equation relates mathematically the rate of reaction to the concentration of the reactants.





Continuous Monitoring

When we follow one experiment over time recording the change in concentration we call it a continuous rate method.

The gradient represents the rate of reaction. The reaction is fastest at the start where the gradient is steepest. The rate drops as the reactants start to get used up and their concentration drops. The graph will eventual become horizontal and the gradient becomes zero which represents the reaction having stopped.

Measurement of the change in volume of a gas

This works if there is a change in the number of moles of gas in the reaction. Using a gas syringe is a common way of following this. It works quite well for measuring continuous rate but a typical gas syringe only measures 100ml of gas so you don't want at reaction to produce more than this volume. Quantities of reactants need to be calculated carefully.

Mg + HCl \rightarrow MgCl₂ +H₂

The initial rate is the rate at the start of the reaction, where it is fastest. It is obtained by taking the gradient of a continuous monitoring conc vs time graph at time = zero. A measure of initial rate is preferable as we know the concentrations at the start of the reaction





Typical Method

- Measure 50 cm³ of the 1.0 mol dm⁻³ hydrochloric acid and add to conical flask.
- Set up the gas syringe in the stand
- Weigh 0.20 g of magnesium.
- Add the magnesium ribbon to the conical flask, place the bung firmly into the top of the flask and start the timer.
- Record the volume of hydrogen gas collected every 15 seconds for 3 minutes.

Large Excess of reactants

In reactions where there are several reactants, if the concentration of one of the reactant is kept in a large excess then that reactant will appear not to affect rate and will be pseudo-zero order. This is because its concentration stays virtually constant and does not affect rate.

Comparing continuous rate curves



measured.

A Common Clock Reaction (no need to learn details)

graph at time = zero

Hydrogen peroxide reacts with iodide ions to form iodine. The thiosulfate ion then immediately reacts with iodine formed in the second reaction as shown below.

 $\mathsf{H}_2\mathsf{O}_2(\mathsf{aq}) + 2\mathsf{H}^+(\mathsf{aq}) + 2\mathsf{I}^-(\mathsf{aq}) \rightarrow \mathsf{I}_2(\mathsf{aq}) + 2\mathsf{H}_2\mathsf{O}(\mathsf{I})$

 $2S_2O_3^{2-}(aq) + I_2(aq) \rightarrow 2I^{-}(aq) + S_4O_6^{2-}(aq)$

When the I_2 produced has reacted with all of the limited amount of thiosulfate ions present, excess I_2 remains in solution. Reaction with the starch then suddenly forms a dark blue-black colour.

A series of experiments is carried out, in which the concentration of iodide ions is varied, while keeping the concentrations of all of the other reagents the same. In each experiment the time taken (t) for the reaction mixture to turn blue is measured.

In clock reactions there are often two successive reactions and an end point is achieved when one limited reactant runs out, resulting in a sudden colour change

By repeating the experiment several times, varying the concentration of a reactant e.g. I⁻, (keeping the other reactants at constant concentration)you can determine the order of reaction with respect to that reactant

The initial rate of the reaction can be represented as (1/t)

Experiment	Sulfuric acid (H⁺)	Starch	Water	Potassium iodide(I ⁻)	Sodium Thiosulfate S ₂ O ₃ ²⁻
1	25	1	20	5	5
2	25	1	15	10	5
3	25	1	10	15	5
4	25	1	5	20	5
5	25	1	0	25	5

Working out orders from experimental initial rate data

Normally to work out the rate equation we do a series of experiments where the initial concentrations of reactants are changed (one at a time) and measure the initial rate each time.

Working out rate order graphically

In an experiment where the concentration of one of the reagents is changed and the reaction rate measured it is possible to calculate the order graphically

Taking rate equation

Rate = $k [Y]^n$

Log both sides of equation Log rate = log k + n log [Y] Y = c + m x

A graph of log rate vs log [Y] will yield a straight line where the gradient is equal to the order n



log [Y]

In this experiment high concentrations with quick times will have the biggest percentage errors.

This initial rate data can also be presented in a table.

Example: work out the rate equation for the following reaction, $A+B+2C \rightarrow D+2E$, using the initial rate data in the table

Experiment	[A] mol dm ⁻ 3	[B] mol dm ⁻³	[C] mol dm ⁻³	Rate mol dm ⁻³ s ⁻¹
1	0.1	0.5	0.25	0.1
2	0.2	0.5	0.25	0.2
3	0.1	1.0	0.25	0.4
4	0.1	0.5	0.5	0.1

In order to calculate the order for a particular reactant it is easiest to compare two experiments where **only that reactant** is being changed

If conc is doubled and rate stays the same: order= 0

If conc is doubled and rate doubles: order= 1

If conc is doubled and rate quadruples : order= 2

For reactant A compare between experiments 1 and 2

For reactant A as the concentration **doubles** (B and C staying constant) so does the rate. Therefore the order with respect to reactant **A is first order**

For reactant B compare between experiments 1 and 3 :

As the concentration of B doubles (A and C staying constant) the rate quadruples.

Therefore the order with respect to B is 2nd order

For reactant C compare between experiments 1 and 4 :

As the concentration of C doubles (A and B staying constant) the rate stays the same.

Therefore the order with respect to C is zero order

The overall rate equation is $r = k [A] [B]^2$

The reaction is 3^{rd} order overall and the unit of the rate constant =mol⁻²dm⁶s⁻¹

Working out orders when two reactant concentrations are changed simultaneously

In most questions it is possible to compare between two experiments where only one reactant has its initial concentration changed. If, however, both reactants are changed then the effect of both individual changes on concentration are multiplied together to give the effect on rate.

In a reaction where the rate equation is $r = k [A] [B]^2$

If the [A] is x2 that rate would $\mathbf{x2}$

If the [B] is x3 that rate would $x3^2 = x9$

If these changes happened at the same time then the rate would x2x9=x 18

Example work out the rate equation for the reaction, between X and Y, using the initial rate data in the table

Experiment	Initial concentration of X/ mol dm ⁻³	Initial concentration of Y/ mol dm ⁻³	Initial rate/ mol dm ⁻³ s ⁻¹
1	0.05	0.1	0.15 x 10 ^{–6}
2	0.10	0.1	0.30 x 10 ^{−6}
3	0.20	0.2	2.40 x 10 ⁻⁶

For reactant X compare between experiments 1 and 2

For reactant X as the concentration **doubles** (Y staying constant) so does the rate. Therefore the order with respect to reactant **X is first order**

Comparing between experiments 2 and 3 :

Both X and Y double and the rate goes up by 8

We know X is first order so that will have doubled rate

The effect of Y, therefore, on rate is to have quadrupled it.

Y must be second order

The overall rate equation is $r = k [X] [Y]^2$

The reaction is 3^{rd} order overall and the unit of the rate constant =mol^{-2}dm^6s^{-1}

Calculating a value for k using initial rate data

Using the above example, choose any one of the experiments and put the values into the rate equation that has been rearranged to give k. Using experiment 3:

 $\mathbf{r} = \mathbf{k} [\mathbf{X}] [\mathbf{Y}]^2 \longrightarrow \mathbf{k} = \frac{\mathbf{r}}{[\mathbf{X}] [\mathbf{Y}]^2}$

 $k = 2.40 \times 10^{-6}$ 0.2×0.2^{2}

k = 3.0 x 10⁻⁴ mol⁻²dm⁶s⁻¹

Remember k is the same for all experiments done at the same temperature. Increasing the temperature increases the value of the rate constant k



Effect of Temperature on Rate Constant: The Arrhenius Equation

Increasing the temperature increases the value of the rate constant **k**

Increasing temperature increases the rate constant k. The relationship is given by the Arrhenius equation $\mathbf{k} = Ae^{-Ea/RT}$ where A is a constant R is gas constant and Ea is activation energy.

The Arrhenius equation can be rearranged

 $\ln k = \text{constant} - \text{Ea}/(\text{RT})$

k is proportional to the rate of reaction so $\ln k$ can be replaced by $\ln(rate)$

From plotting a graph of ln(rate) or ln k against 1/T the activation energy can be calculated from measuring the gradient of the line



k



Rate Equations and Mechanisms

A mechanism is a series of steps through which the reaction progresses, often forming intermediate compounds. If all the steps are added together they will add up to the overall equation for the reaction Each step can have a different rate of reaction. The slowest step will control the overall rate of reaction. **The slowest step is called the rate-determining step**.

The molecularity (number of moles of each substance) of the molecules in the slowest step will be the same as the order of reaction for each substance. e.g. 0 moles of A in slow step would mean A is zero order. 1 mole of A in the slow step would mean A is first order

