

Edexcel Chemistry A-level

Topic 8: Energetics I

Detailed Notes

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Enthalpy Change (AH°)

Enthalpy change is heat energy change and is represented by the symbol ΔH° . Enthalpy is measured under standard conditions of **100 kPa pressure** and a specified temperature, generally **298 K**.

In a reaction, bonds are broken and then bonds are made. For bonds to be **broken**, energy is **taken in** from the surroundings and when bonds are **formed**, energy is **given out**. The overall energy change of the reaction depends on how much energy is transferred in these processes.

When energy is taken in from the surroundings, the **enthalpy change is positive**. When energy is released to the surroundings, the **enthalpy change is negative**.

Overall enthalpy change (ΔH) can be calculated as follows:



Enthalpy changes in a reaction can be shown on an **energy level diagram**. These also indicate if the reaction is endothermic or exothermic.

Example:



In **endothermic** reactions, more energy is needed to break bonds than make new ones. Therefore, the overall ΔH is positive and heat is taken in from the surroundings.

In **exothermic** reactions, the opposite is true, with more energy being needed to make new bonds than break existing ones. Therefore **heat is given out** and ΔH is negative.





Measuring Enthalpy Change

Enthalpy Change of Reaction (ΔH°)

This is defined as:

The enthalpy change when quantities of substances in standard states react completely under standard conditions.

Enthalpy Change of Formation (A_fH°)

This is defined as:

The enthalpy change when one mole of a substance is produced from its elements under standard conditions.

Enthalpy Change of Combustion ($\Delta_c H^o$)

This is defined as:

The enthalpy change when one mole of a substance is burned completely in oxygen under standard conditions.

Enthalpy Change of Neutralisation ($\Delta_n H^o$)

This is defined as:

The enthalpy change when solutions of acid and alkali react together under standard conditions to produce one mole of water.

Calorimetry

Calorimetry is an **experimental method** for finding enthalpy change by measuring **temperature change over time**. When recorded and plotted on a graph, data can be **extrapolated** to give an accurate value for the change in temperature.

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This measured change in temperature, ΔT , is **proportional** to the energy change:

(where q = energy change (J), m = mass (g), c = specific heat capacity (J $g^{-1} \circ C^{-1}$), ΔT = temperature change (°C))

This equation allows the energy change of a substance of known mass to be calculated. It depends on its **specific heat capacity**, defined as:

The energy required to raise 1g of substance by 1K without a change of state.

Using this energy value (q), **enthalpy change per mole** (J mol⁻¹) of a substance can be calculated:

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Sources of Error

 Δ H values found using calorimetry are **never completely accurate** as energy is easily lost from the system. Heat loss can occur due to **conduction**, **convection** or inaccuracies in measuring temperatures. The heat loss to the surroundings can be reduced by putting a **lid** on the calorimeter and **insulating** the outsides of the calorimeter using an insulator like polystyrene.

Also, the **specific heat capacity** of the solution is taken to be 4.18 kJ mol⁻¹, which is the value for water and not the **actual solution**. The specific heat capacity of the calorimeter is not taken into account so this leads to inaccuracies in the calculation.

Hess's Law

Energy in a reaction system must be **conserved**, as energy cannot be created or destroyed. Therefore the **overall enthalpy change** for a reaction is **the same**, regardless of the **route taken**.

This is **Hess's Law** and it is used to determine the enthalpy changes for reactions that cannot be found directly using an experimental method. To do this, a **triangular cycle method** is used with an intermediate product. The direction of the arrows indicates whether the values should be added or taken away. They can be treated like vectors:

Example:







Enthalpies of Formation

Hess's law can be used to calculate an enthalpy change using given enthalpies of formation. When setting up the triangular diagram, the **arrows point out from the central product C** as both A and B are formed from the elements at C.

Example:



Note: ΔH^1 *is subtracted as the reaction goes in the opposite direction to the arrow.*

Enthalpies of Combustion

Hess's law can also be used to calculate an enthalpy change using given enthalpies of combustion. When setting up the triangular diagram, the **arrows point towards the central product (which is always H_2O and CO_2)** as both A and B burn to form the products at C.

Example:



Note: ΔH^2 is subtracted as the reaction goes in the opposite direction to the arrow.





Bond Enthalpies

Bond enthalpy data is an averaged value representing:

The energy required to break one mole of the stated bond in a gaseous state, under standard conditions.

Different covalent bonds require different amounts of energy to be broken. Values can be found experimentally using **calorimetry** methods. The bond enthalpy values calculated in this way often differ from the **databook values** as they are **not exact** and **vary in each situation**. This means the databook values are **averaged values**.

Mean bond enthalpy values tell you how much energy is required to break a particular bond, averaged out across the range of compounds containing that bond. These values relate to how **strong** a bond is, so they can be used to infer which bonds might **break first** in a chemical reaction. Bonds with **lower** bond enthalpy values would be expected to break first.

If a reaction requires the breaking of many strong bonds or the reaction is endothermic overall, then it is likely that the reaction will occur **slowly** at **room temperature**.

Bond Enthalpy Calculations

Mean bond enthalpy values can be used to calculate the **overall enthalpy change** for a reaction.

$$\Delta H_{reaction} = \Sigma H_{bonds \ broken} - \Sigma H_{bonds \ formed}$$

This means that the enthalpy change for a reaction is equal to the **sum** of the bond enthalpies for the bonds **broken** in the reaction, minus the **sum** of the bond enthalpies for the bonds **formed** in the reaction.

Mean bond enthalpy values are often **tabulated**. You need to identify the bonds broken and formed during the reaction in order to calculate the overall enthalpy change.

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Example:

Calculate the enthalpy change when methane undergoes complete combustion in oxygen using the bond enthalpies given.

Bond	Bond enthalpy (kJ mol ⁻¹)
С-Н	413
0=0	498
О-Н	464
C=O	799

Chemical equation: $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O$

Bonds formed

C=O x 2 = 799 x 2 = 1598 H-O x 4 = 464 x 4 = 1856

Bonds broken

C-H x 4 = 413 x 4 = 1652 O=O x 2 = 498 x 2 = 996

Enthalpy change = (1652+996) - (1856+1598)= 2648 - 3454 = -806 kJ mol⁻¹

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