5 Chemical energetics

Enthalpy changes and entropy changes accompany chemical reactions. This topic demonstrates why some reactions and processes are spontaneous and others are not.

- 5.1 Lattice energy
- 5.2 ionic solid, its aqueous solution and Born-Haber cycles

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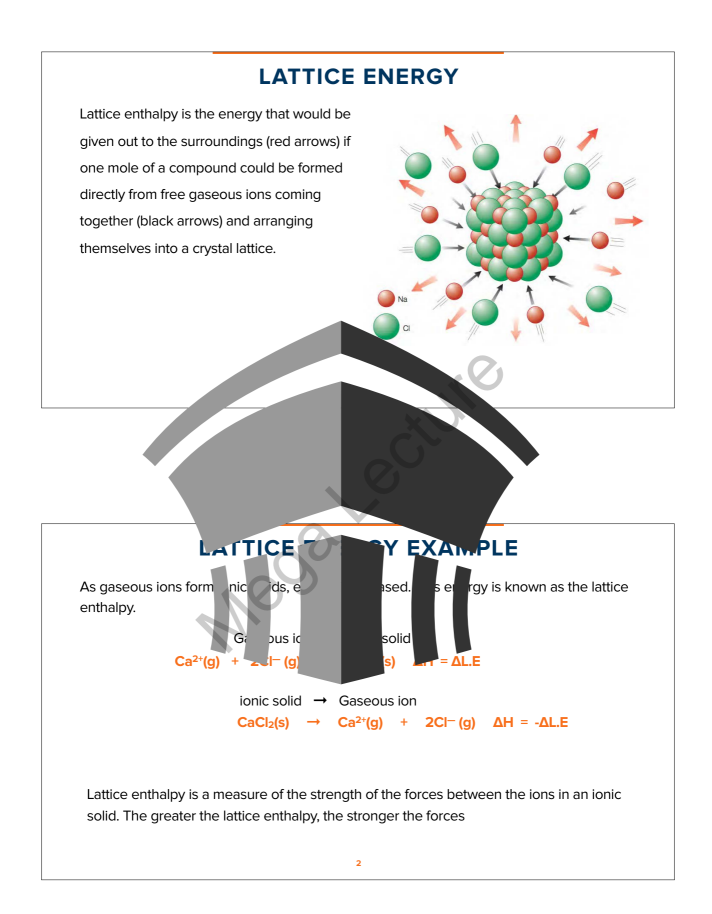


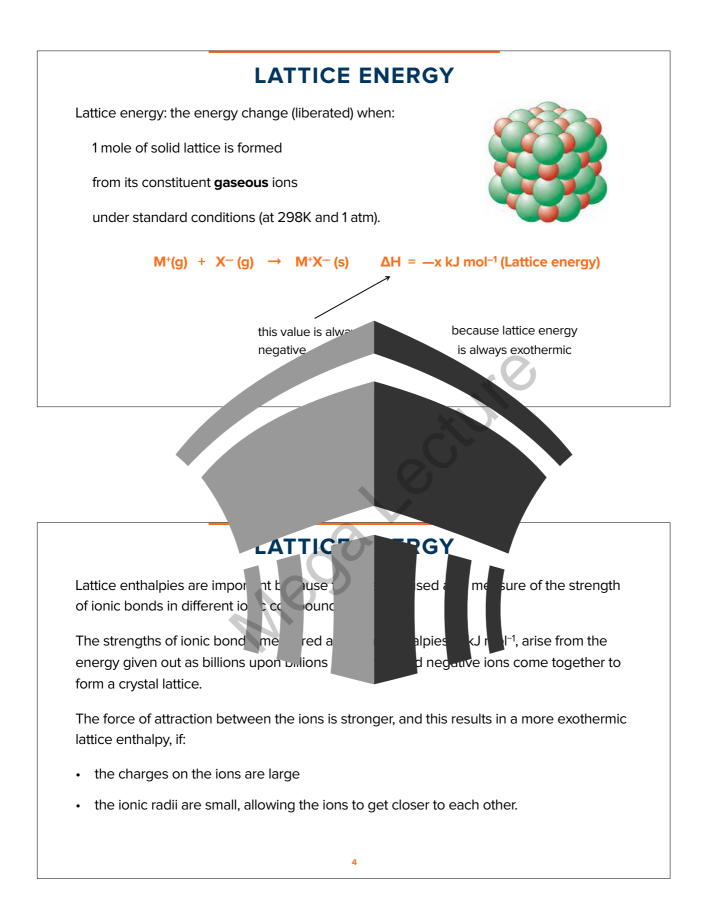
ENERGETICS

5 Chemical energetics

Enthalpy changes and entropy changes accompany chemical reactions. This topic demonstrates why some reactions and processes are spontaneous and others are not.

	Learning outcomes Candidates should be able to:
5.1 Enthalpy change, ΔH	a) explain that chemical reactions are accompanied by energy changes, principally in the form of heat energy; the energy changes can be exothermic (ΔH is negative) or endothermic (ΔH is positive)
	b) explain and use the terms:
	 (i) enthalpy change of reaction and standard conditions, with particular reference to: formation, combustion, hydration, solution neutralisation, atomisation
	(ii) bond energy (ΔH positive, i.e. bond breaking)
	(iii) <i>lattice energy (</i> Prative , i.e. gaseous ions to solid lattice)
	c) calculate entry r_{s} opropriate experimental results, including set of the relation $mc\Delta T$
	r charge and of ioni rad ¹ lattic ergy
5.2 Hess' Law, including Born-Haber cycles	and carry out gy terms, with
	 change enthalpy changes use found by direct experiment, exactly baloy change of mation from enthalpy changes of mation paraway diagram, in terms of the enthalpy change of mation and of the activation energy

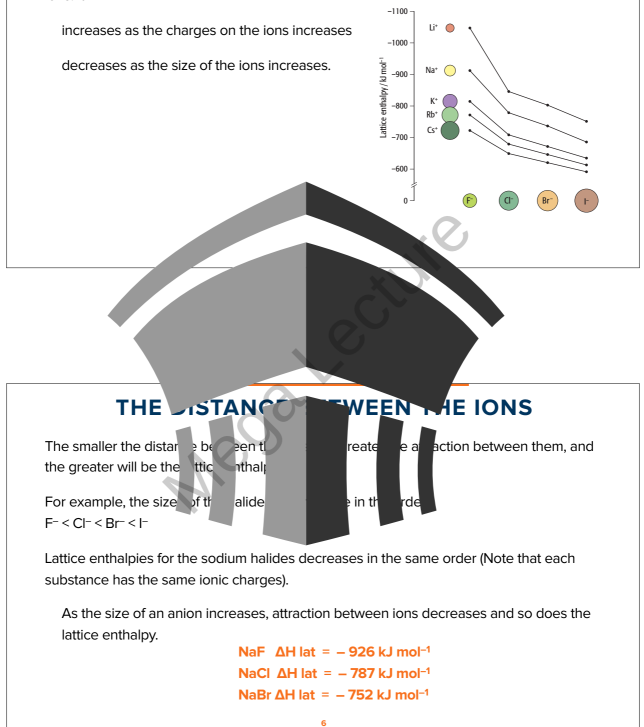


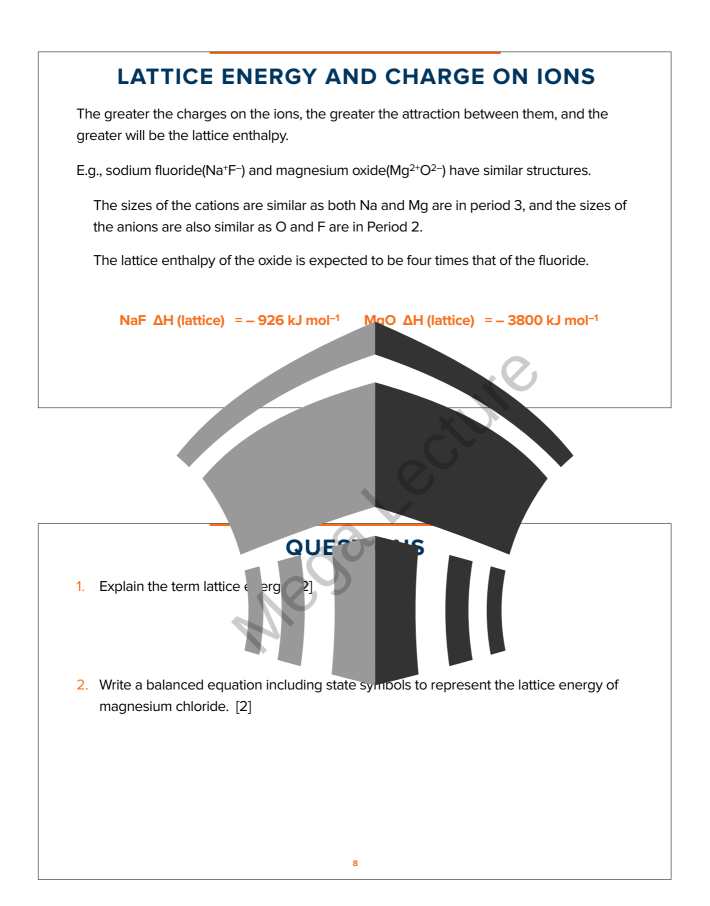


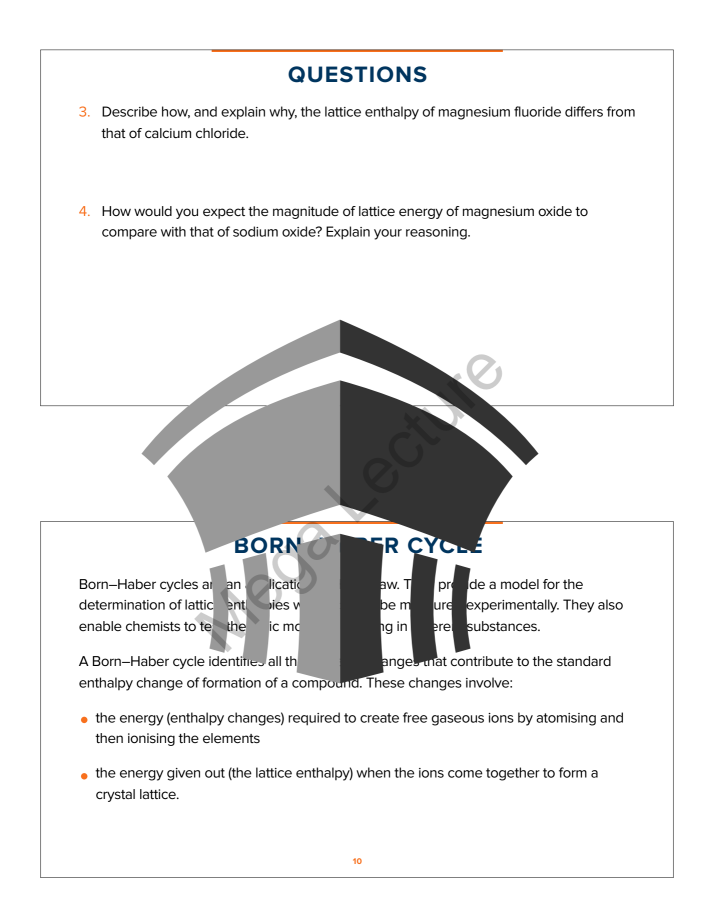
LATTICE ENERGY

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The magnitude of Lattice energy depends on the the electrostatic attraction between the ions. It



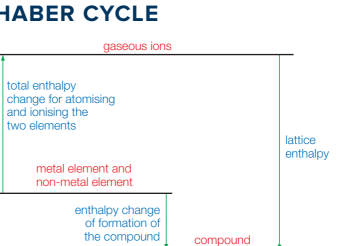




BORN-HABER CYCLE

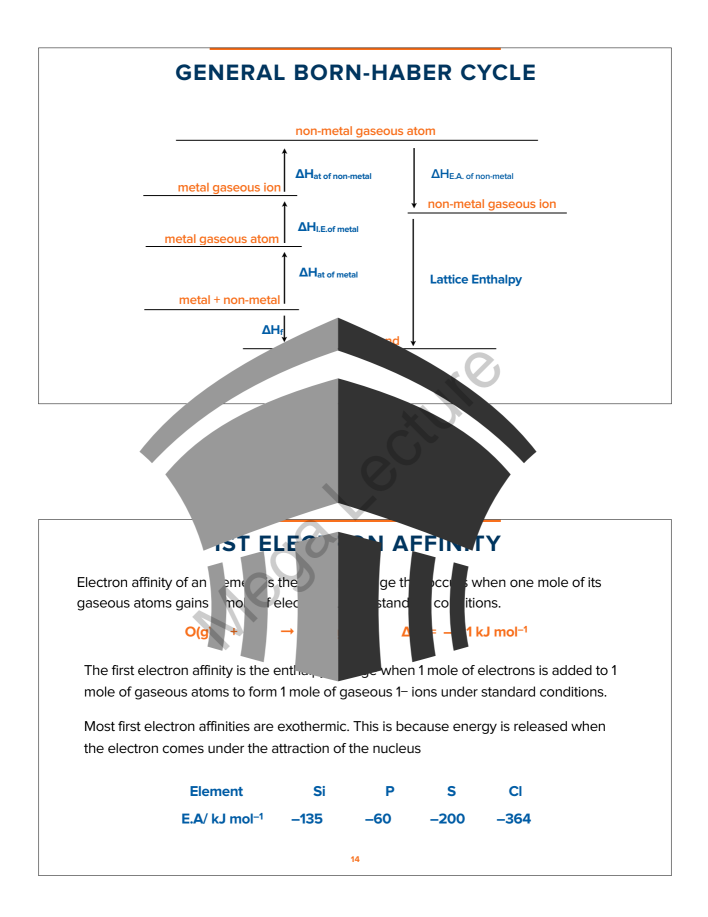
A Born–Haber cycle is usually set out like an enthalpy profile diagram, with enthalpy changes one after another.

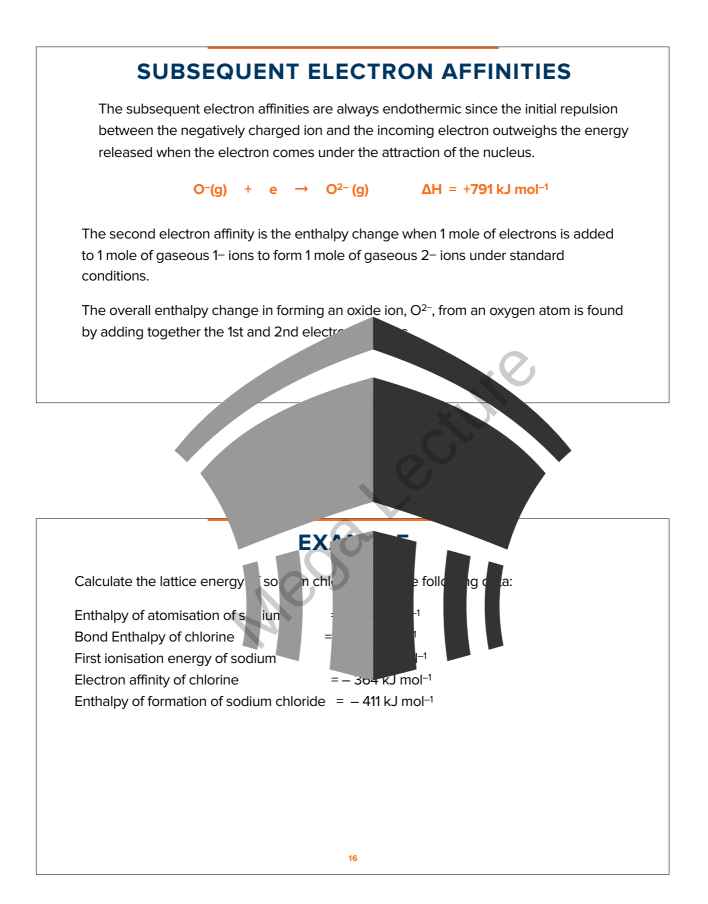
All the processes in the cycle can be measured experimentally except the lattice enthalpy. So, by using Hess's law it is possible to calculate the lattice enthalpy.

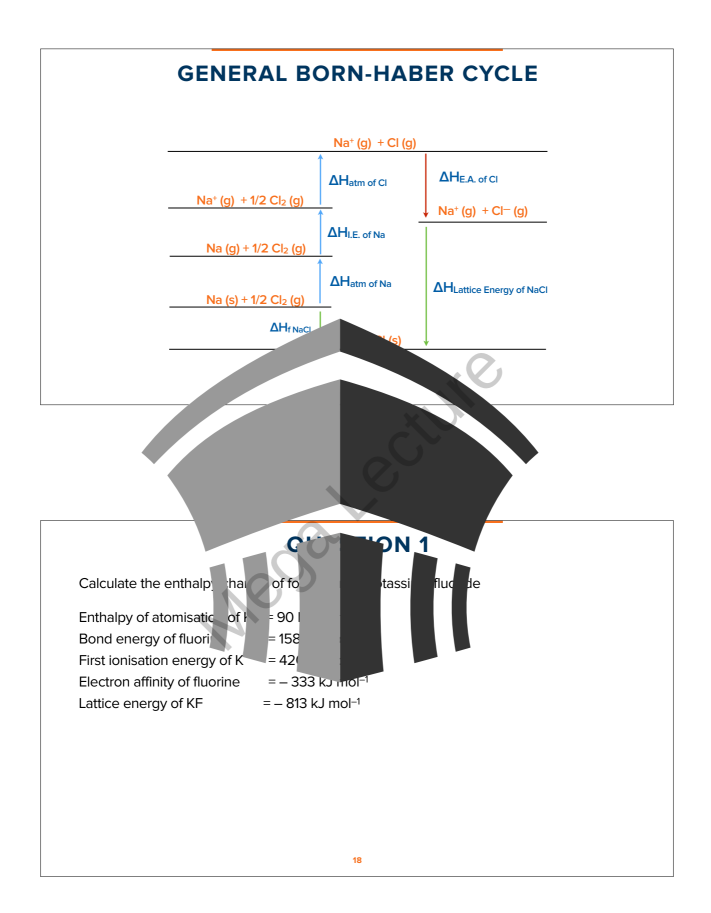


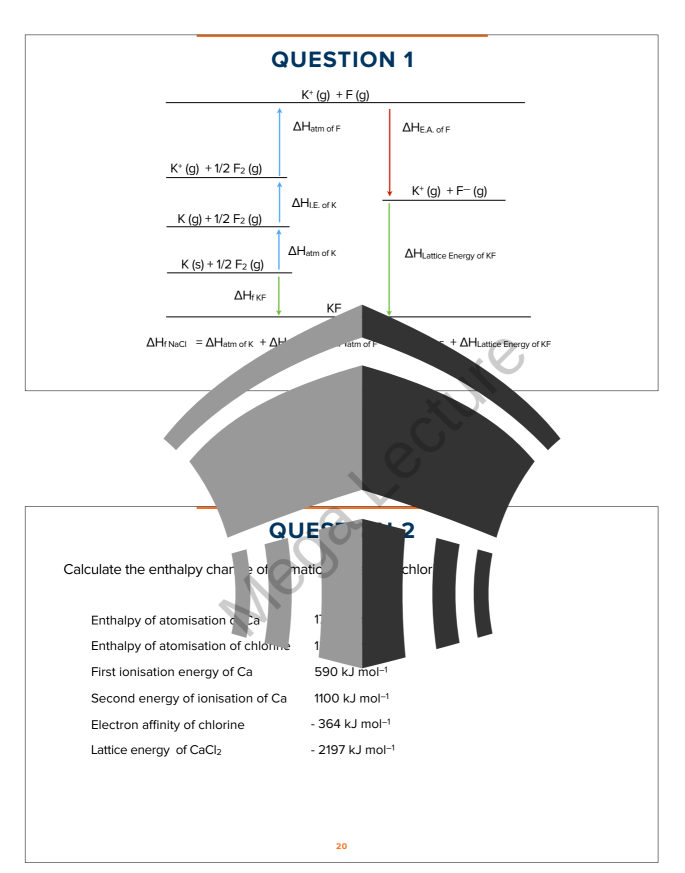
E JRN-H

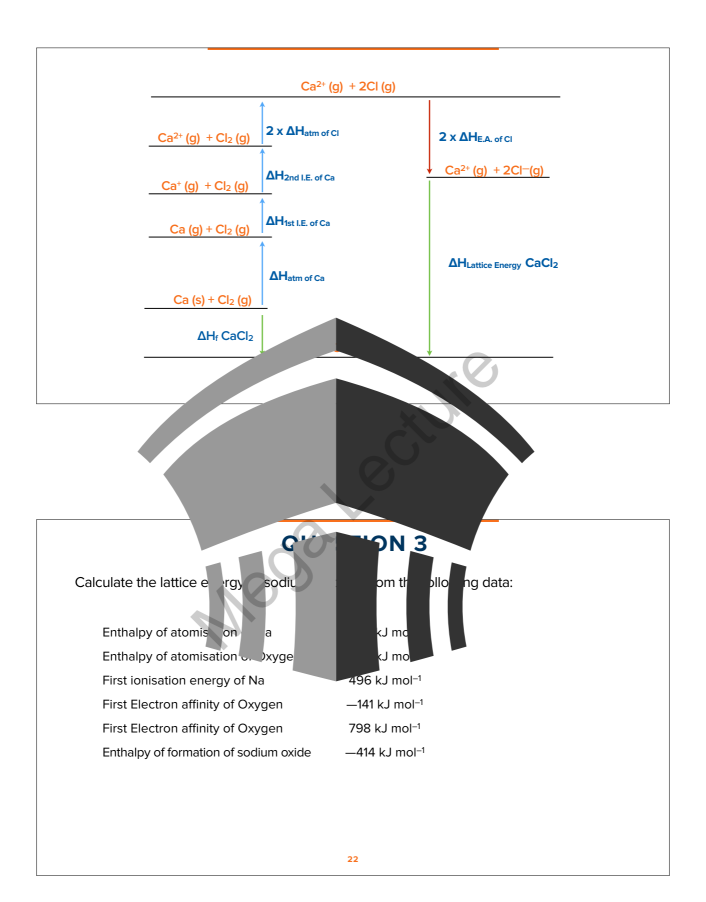
A method for analysis of the	noc nical	d ba: on	ess' law.		
With the calculation of lattice	entr ies tr	ed a			
atomisation energies for tr	ie e ents,				
ionisation energies for the metal component, ΔH_{ie} ,					
electron affinities for the non-metal component, $\Delta H_{\mbox{\tiny ea}}$, and					
the enthalpy change of formation, ΔH_f , for the compound.					
		etal + Ionisation Energy ron affinity of non-meta			
12					



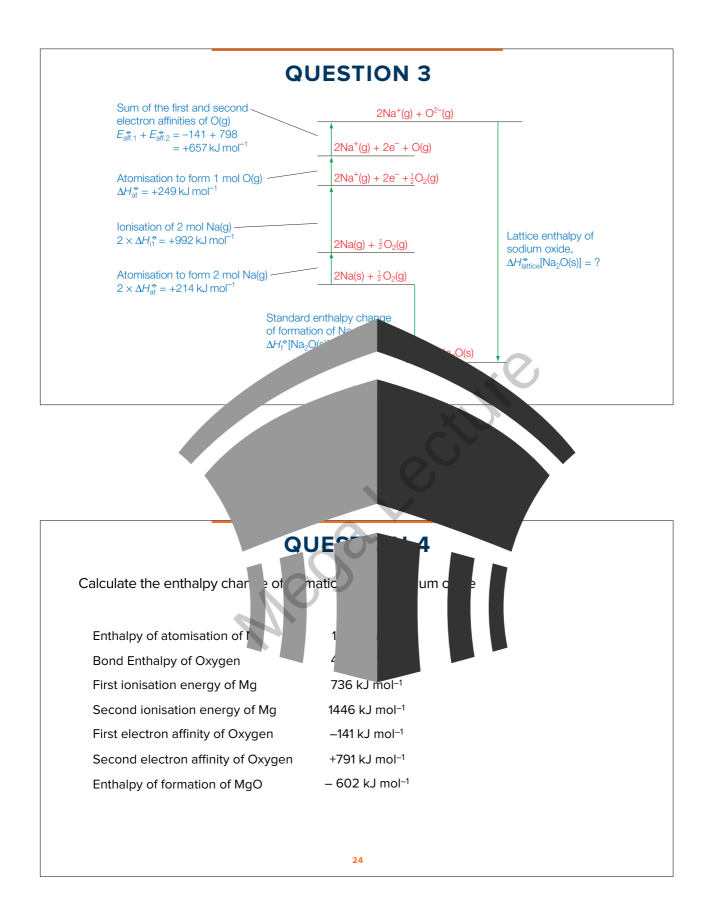


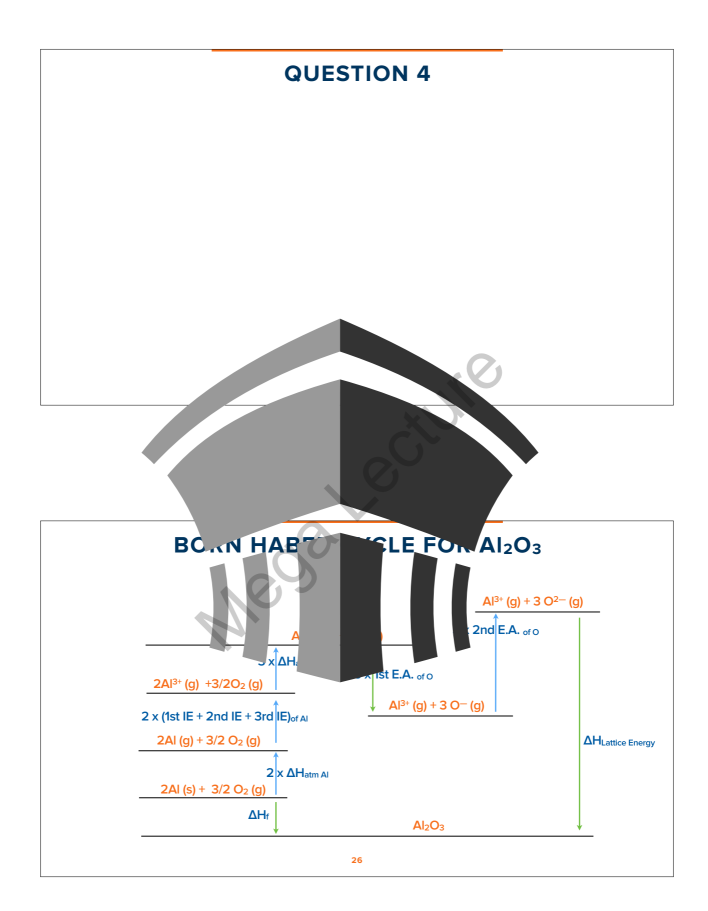






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BORN-HABER CYCLES

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If the cycles are examined it will be seen that:

the largest single contributions in each cycle are made by lattice energy and ionization energy and these two are always opposite in sign

the magnitude of the enthalpy of formation depends on the relative values of the lattice energy and the ionisation energy

if ionisation energy is too high to be supplied by lattice energy then such a compound will not form ionically

ENTHALPY JANGE

Why do ionic crystals dissol in er, er attracted to each other?

An ionic compound such as odi chlor hexane, but it will dissolve in a port solve

ING DISSOLVING

ns in lat e are strongly disso in r n-polar solvents like

When one mole of sodium chloride

dissolves in excess water to produce a very angle solution under standard conditions, there is an enthalpy change of +3.8 kJ mol⁻¹. This enthalpy change is described as the standard enthalpy change of solution, ΔH solution, of sodium chloride. The process can be summarised by the equation:

NaCl(s) + aq \rightarrow Na⁺(aq) + Cl⁻ (aq) ΔH solution = +3.8 kJ mol⁻¹

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ENTHALPY CHANGES DURING DISSOLVING When ionic compounds, like NaCl, dissolve in water the crystal lattice should be broken down and this requires input of energy. This energy is supplied through the energy of hydration. bond between bond forming ions weakens new bonds formed lattice of ions ENTHAL, CHAI URING DISSOLVING First of all, Na+ and C rom t boli NaCI crystals to form wellion. ust be the spaced ions in the ga ous ate, N . This verse of the lattice enthalpy (+787 kJ mo In the second stage, gaseou, Na⁺(g) , drated by polar water molecules s are forming a solution of sodium chloride, ING and Cl-(aq). Under standard conditions, this process is the sum of the standard enthalpy changes of hydration of Na⁺(g) and Cl⁻(g). 30

D

ENTHALPY CHANGES DURING DISSOLVING

Enthalpy of Hydration:

The enthalpy change of hydration, ΔH hyd, is the enthalpy change when 1 mole of a specified gaseous ion dissolves in sufficient water to form a very dilute solution.

