

Key Terms and Definitions

Term	Definition
Entropy (ΔS)	A measure of the disorder in a system. Units: $\text{J K}^{-1} \text{mol}^{-1}$
Standard Entropy Change (ΔS°)	$\Delta S^\circ = \sum S^\circ(\text{products}) - \sum S^\circ(\text{reactants})$
Spontaneous Process	A process where $\Delta G < 0$; can occur without external input.
Gibbs Free Energy (ΔG)	The energy available to do work: $\Delta G = \Delta H - T\Delta S$
Lattice Enthalpy of Formation ($\Delta H^\circ_{\text{LE}}$)	Enthalpy change when 1 mole of an ionic solid forms from its gaseous ions (exothermic).
Lattice Enthalpy of Dissociation	Enthalpy change when 1 mole of ionic solid dissociates into gaseous ions (endothermic).
Atomisation Enthalpy ($\Delta H^\circ_{\text{at}}$)	Enthalpy change to form 1 mole of gaseous atoms from the element.
Electron Affinity ($\Delta H^\circ_{\text{ea}}$)	Enthalpy change when 1 mol of electrons is added to 1 mole of gaseous atoms to form 1^- ions.
Hydration Enthalpy ($\Delta H^\circ_{\text{hyd}}$)	Enthalpy change when gaseous ions dissolve in water to form aqueous ions.

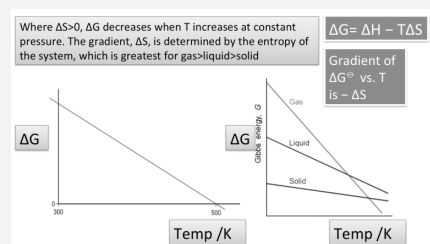
Key Terms and Definitions (cont)

Solution Enthalpy ($\Delta H^\circ_{\text{sol}}$)	Enthalpy change when 1 mol of solute dissolves to infinite dilution.
Perfect Ionic Model	Assumes ions are spherical and charge is evenly distributed with no covalent character.

Entropy Effect

Change	Entropy Effect
Melting / Boiling	Entropy increases significantly
Dissolving ionic lattice	Entropy increases
Increase in gas moles in products	Entropy increases
Cooling a gas	Entropy decreases
Formation of solid from gas	Entropy decreases significantly

Gibbs Free Energy vs. Temperature



Equation: $\Delta G = -\Delta S \cdot T + \Delta H$ ($y = mx + c$)

Slope = $-\Delta S$

Y-intercept = ΔH



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Key Formulas

- ✓ Overall Entropy of a system $\Delta S_{\text{total}} = \Delta S_{\text{system}} + \Delta S_{\text{surroundings}}$
- ✓ Entropy of surroundings $\Delta S_{\text{surroundings}} = -\Delta H/T$
- ✓ Entropy Change of a reaction within a system
 $\Delta S^{\circ}_{\text{system}} = \sum S^{\circ}_{\text{products}} - \sum S^{\circ}_{\text{reactants}}$
- If entropy change is +ve, products more disordered than reactants (natural direction of change)
- & if entropy change is -ve, reactants more disordered than products.
- ✓ Gibbs Free Energy $\Delta G = \Delta H - T\Delta S_{\text{system}}$
- ΔG : Gibbs free energy ($\text{kJ}\cdot\text{mol}^{-1}$)
- ΔH : Enthalpy change ($\text{kJ}\cdot\text{mol}^{-1}$)
- T: Temperature (K)
- ΔS : Entropy change ($\text{J}\cdot\text{K}^{-1}\cdot\text{mol}^{-1}$; convert to kJ by dividing by 1000)
- ✓ Feasibility Conditions
- A reaction is feasible when: $\Delta G < 0$
- To find minimum temperature for spontaneity: $\Delta G = 0 \Rightarrow T = \Delta H/\Delta S_{\text{system}}$
- ✓ Born-Haber Cycle Equation
- For lattice formation enthalpy:
 $\Delta H^{\circ}_{\text{LE}} = \sum \text{enthalpies (atomisation, ionisation, electron affinity, etc.)} + \Delta H^{\circ}_{\text{LE}}$
- ✓ Hydration & Solution Enthalpy (Hess's Cycle) $\Delta H^{\circ}_{\text{sol}} = \sum \Delta H^{\circ}_{\text{f}} - \Delta H^{\circ}_{\text{LE}}$

Kinetics vs. Thermodynamics

A reaction may be thermodynamically feasible ($\Delta G < 0$) but may not occur due to:

- High activation energy
- Slow reaction rate due to kinetic barriers

Factors Affecting Lattice & Hydration Enthalpies

Factor	Effect on ΔH (more negative)
Smaller ionic radius	Increases attraction → More negative ΔH
Higher ionic charge	Increases attraction → More negative ΔH
Higher ionic charge → More negative ΔH Greater hydration energy	Stronger interactions with water → More negative ΔH



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