<u>Chapter Notes</u> **Subject: Chemistry** Class: XI Chapter: Thermodynamics

Top concepts

1. The branch of science which deals with study of different forms of energy and their interconversion is called thermodynamics.

2. A system in thermodynamics refers to that part of universe in which observations are made.

3. The remaining portion of universe which is not part of system constitutes the surroundings. The surroundings include everything other than the system.

4. The wall (real or imaginary) that separates the system from the surroundings is called boundary.

5. Types of the System

Types of the System	Exchange of	Exchange of matter
	energy	
Open	Yes	Yes
Closed	Yes	No
Isolated	No	No

6. The state of a thermodynamic system is described by its measurable or macroscopic (bulk) properties. The state of the surroundings can never be completely specified.

7. State variables are the measurable properties of system required to describe the state of the system. Examples are temperature, pressure, volume etc.

8. Various types of processes

Type of process	Definition
Isothermal	Process in which temperature of system
	remains constant
Adiabatic Process in which there is no transfer o	
	between the system and surroundings
Isobaric	Process in which pressure of system

9. Every substance possesses definite amount of energy which depends on factors such as chemical nature, temperature and pressure etc. Internal energy, U of the system represents the total energy of the system

10. Work done in a adiabatic process and sign convention

Work done (w _{ad})	Sign of w
By the system	-
On the system	+

11.Heat changes and sign convention

Heat(q)transferred	Sign of q
From surroundings to	+
the system	
From system to the	-
surroundings	

12. First Law of Thermodynamics: $\Delta U = q + w$

13. First law of thermodynamics is also called as *law of conservation of energy i.e. energy can neither be created nor destroyed. It also states that the energy of an isolated system is constant.*

14. Absolute value of the internal energy can never be determined only change in internal energy can be determined.

15. A process or change is said to be reversible, if a change is brought out in such a way that the process could, at any moment, be reversed by an infinitesimal change.

16. Expansion of a gas in vacuum (pex = 0) is called free expansion.

No work is done during free expansion of an ideal gas whether the process is reversible or irreversible

17.For adiabatic change, q = 0 and $\Delta U = W_{ad}$

18. The enthalpy of a system may be defined as the sum of the internal energy and the product of its pressure and volume. It is denoted by the symbol H and is given by H = U + PV

19.Change in enthalpy $\Delta H = \Delta U + p \Delta V$

20.Enthalpy change and sign convention

	Type of reaction	Definition	ΔH
	Exothermic reactions	Reactions in	-
•		which heat is	
		evolved during	
		the reaction	
	Endothermic reactions	Reactions in	+
		which heat is	
		absorbed during	
		the reaction	

21.An extensive property is a property whose value depends on the quantity or size of matter present in the system. Examples are mass, volume, internal energy, enthalpy, heat capacity, etc

22.An intensive property is a property whose value does not depend on the quantity or size of matter present in the system. Examples are temperature, density, pressure etc.

23.Specific heat, also called specific heat capacity is the quantity of heat required to raise the temperature of one unit mass of a substance by one degree Celsius (or one Kelvin).

24. The enthalpy change accompanying a reaction is called the reaction enthalpy ΔH_r

 ΔH_r = (sum of enthalpies of products) - (sum of enthalpies of reactants)

25. The standard enthalpy of reaction is the enthalpy change for a reaction when all the participating substances are in their standard states. The standard state of a substance at a specified temperature is its pure form at pressure of 1 bar.

26.A balanced chemical equation together with the value of its is called a thermochemical equation $\Delta_r H^{\circ}$

27.Hess's Law of Constant Heat Summation: If a reaction takes place in several steps then its standard reaction enthalpy is the sum of the standard enthalpies of the intermediate reactions into which the overall reaction may be divided at the same temperature

28.The lattice enthalpy of an ionic compound is the enthalpy change which occurs when one mole of an ionic compound dissociates into its ions in gaseous state

29.
$$\Delta_{\rm sol} H^{\theta} = \Delta_{\rm lattice} H^{\theta} + \Delta_{\rm hyd} H^{\theta}$$

30.A process that has natural tendency of occurrence in a particular direction and is reversible only by application of some external agency is known as spontaneous process.

31.The processes which are forbidden and are made to take place only by supplying energy continuously from outside the system are called non-spontaneous process.

32.Entropy (S) is the measure of randomness of a system. It is a state function.

33.Entropy increases from solid to gas. Entropy is maximum for gases as they have maximum disorder.

34.According to Second law of thermodynamics entropy of the universe always increases during a spontaneous change

35.For reversible process entropy change is given by

$$\Delta S = \frac{q_{rev}}{T}$$
 where

 $\mathbf{q}_{_{\text{rev}}}$ is heat absorbed or released during the reaction T is the temperature of the reaction

36. At constant pressure $q_{rev}=\Delta H$

 $\Delta S = \Delta H/T$

$$\Delta S(\text{total}) = \Delta S(\text{sys}) + \Delta S(\text{surr}) > 0$$

37. For spontaneous process

38.Decrease in enthalpy and increase in entropy of system independently cannot serve the criteria for spontaneity

39.Gibbs function or Gibbs energy is denoted by G.

G=H-TS

40. The criteria for spontaneous reaction in relation to ΔG at constant pressure and constant temperature.

If Δ G < 0, process is spontaneous

If Δ G = 0, process is in equilibrium

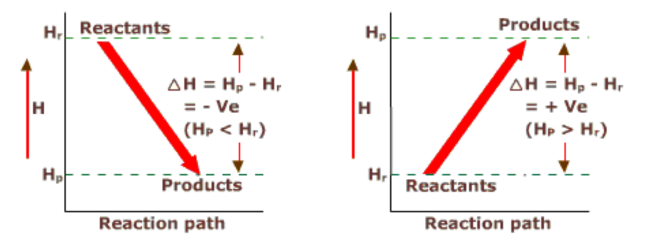
If Δ G > 0, process is non-spontaneous

41.Relation between Gibbs energy change ($\Delta_r G^\circ$) and equilibrium constant (K)

ΔG[°]=-RTInK

42. In case of endothermic reactions enthalpy of reaction ($\Delta_r H^\circ$) is large and positive. In such a case, value of equilibrium constant (K) will be much smaller than 1 and the reaction is unlikely to form much product 43. In case of exothermic reactions enthalpy of reaction ($\Delta_r H^\circ$) is negative and $\Delta_r G^\circ$ is also large and negative. In such cases equilibrium constant (K) is larger than 1 and reactions go to

completion



(a) Enthalpy change during an Exothermic reaction (b) Enthalpy change during Endothermic reaction

44. Different types of enthalpy

No.	Type of	Definition	Symbol
	enthalpy		
1	Standard	The enthalpy change that accompanies	$\Delta_{fus} H^{\theta}$
	Enthalpy of	melting of one mole of a solid	
	fusion	substance in standard state	
	or		
	Molar		
	enthalpy		
	of fusion		
2	Standard	The enthalpy change that accompanies	$\Delta_{vap} H^{\theta}$
	Enthalpy of	vaporizing of one mole of a liquid at	tup.
	vaporization	constant temperature and under	
	or	standard pressure (1bar)	
	Molar		
	enthalpy		
	of		
	vaporization		
I			I

3	Standard	The enthalpy change when one mole of	$\Delta_{mh} H^{\theta}$
	Enthalpy of	a solid substance sublimes at a	∆ _{sub} 11
	sublimation	constant temperature and under	
	sublimation	standard pressure (1bar)	
4	Enthalpy of	The enthalpy change for the formation	$\Delta_f H^{\theta}$
	formation	of one mole of a compound from its	
	or	elements in their most stable states of	
	Standard	aggregation	
	molar		
	enthalpy of		
	formation		
5	Enthalpy of	Enthalpy change when 1 mole of	$\Delta_{comb} H^{\theta}$
	combustion	substance is completely burnt in excess	or
		of oxygen or air	
			$\Delta_{c}H^{\theta}$
7	Enthalpy of	Enthalpy change in breaking one mole	$\Delta_a H^{\theta}$
	atomisation	of bonds of a substance completely into	
		atoms in gaseous state	
8	Bond	Amount of energy required to break	$\Delta_{bond} H^{\theta}$
	dissociation	one mole of bond of a particular type	
	enthalpy	between atoms in gaseous state	

9	Enthalpy of	Enthalpy of solution of a substance is the	$\Delta_{sol}H^{\theta}$
		enthalpy change when one mole of solute dissolves in a specified amount of solvent.	