	414 merniouynumics und mernioenemistry	
9.	Heat produced in calories by the combustion of one gram of carbon is called	20
	(a) Heat of combustion of carbon	-0
	(b) Heat of formation of carbon	
	(c) Calorific value of carbon	21
	(d) Heat of production of carbon	
10.	Conditions of standard state used in thermochemistry is	
	(a) $0^{\circ}C$ and 1 atm (b) $20^{\circ}C$ and 1 atm	22
	(c) $25^{\circ}C$ and 1 atm (d) $0K$ and 1 atm	
11.	The temperature of the system decreases in an [KCET 2005]	23
	(a) Adiabatic compression(b) Isothermal compression(c) Isothermal expansion(d) Adiabatic expansion	-3
12.	For the isothermal expansion of an ideal gas	
12,	(a) E and H increases	
	(b) E increases but H decreases	24
	(c) H increases but E decreases	
	(d) E and H are unaltered	
3.	If a refrigerator's door is opened, then we get	25
0	[CPMT 1980]	0
	(a) Room heated	
	(b) Room cooled	26
	(c) More amount of heat is passed out	
	(d) No effect on room	
4.	The cooling in refrigerator is due to [KCET 2005]	
	(a) Reaction of the refrigerator gas	27
	(b) Expansion of ice	-/
	(c) The expansion of the gas in the refrigerator	
	(d) The work of the compressor	- 0
5.	Point out the wrong statement in relation to enthalpy	28
	(a) It is a state function	
	(b) It is an intensive property	_
	(c) It is independent of the path followed for the change	_
	(d) Its value depends upon the amount of substance in the system	_
6.	the system Which of the following is zero for an isochoric process	1.
0.	(a) dP (b) dV	
	(a) dF (b) dV (c) dT (d) dE	
7	Mark the <i>correct</i> statement [MP PET 1997]	
17.	(a) For a chemical reaction to be feasible, ΔG should be	
	(a) For a chemical reaction to be leasible, ΔG should be zero	2.
	(b) Entropy is a measure of order in a system	
	(c) For a chemical reaction to be feasible, ΔG should be	
	positive	
	(d) The total energy of an isolated system is constant	
18.	In an isochoric process the increase in internal energy is	
	[MP PMT 1997]	
	(a) Equal to the heat absorbed	
	(b) Equal to the heat evolved	
	(c) Equal to the work done	3.
	(d) Equal to the sum of the heat absorbed and work done	J .
9.	It is general principle that the less energy a system	
	contains, it is [MH CET 1999]	
	(a) Loss stable (b) More stable	

(b) More stable (a) Less stable

		(c) Unstable	(d) More unstable
	20.	Internal energy is an examp	ple of [Pb. PMT 2000]
		(a) Path function	(b) State function
		(c) Both (a) and (b)	(d) None of these
	21.	The process, in which no he	eat enters or leaves the system,
		is termed as [Pb. PMT 1999); Kerala (Med.) 2002; J & K 2005]
		(a) Isochoric	(b) Isobaric
		(c) Isothermal	(d) Adiabatic
	22.	The intensive property amo	ong these quantities is
			[KCET 2002]
		(a) Enthalpy	(b) Mass/volume
]		(c) Mass	(d) Volume
	23.	In thermodynamics which	one of the following is not an
		intensive property	[Kerala (Med.) 2002]
		(a) Pressure	(b) Density
		(c) Volume	(d) Temperature
	24.	If in a container neither	mass and nor heat exchange
		occurs then it constitutes a	[MP PMT 1993; AFMC 2003]
		(a) Closed system	(b) Open system
		(c) Isolated system	(d) Imaginary system
	25.	Which of the following is no	ot a state function [DCE 2002]
		(a) ΔS	(b) ΔG
		(c) ΔH	(d) ΔQ
	26.	Which of the following is tr	· · ~
	20.		2002; MP PET 1995, 98, 2004;
			MT 1990; MP PMT 1998, 2002]
		(a) $\Delta H = 0$	(b) $\Delta W = 0$
		(c) $\Delta Q = 0$	(d) $\Delta V = 0$
	27.	Which of the following is no	
	Z /•	which of the following is no	[MH CET 2004; AIIMS 2001]
		(a) Internal energy	(b) Enthalpy
		(c) Work	(d) Entropy
	28.	Among them intensive prop	· · · · · · · · · · · · · · · · · · ·
	_0,	(a) Mass	(b) Volume
		(c) Surface tension	
			(a) Entimpy
	_		

First law of thermodynamics and Hess law

- The first law of thermodynamics is only
 - (a) The law of conservation of energy
 - (b) The law of conservation of mass
 - (c) The law of conservation of momentum
 - (d) Both (a) and (b)
- A mixture of two moles of carbon monoxide and one mole of oxygen, in a closed vessel is ignited to convert the carbon monoxide to carbon dioxide. If ΔH is the enthalpy change and ΔE is the change in internal energy, then [KCET 2005]
 - (a) $\Delta H > \Delta E$
 - (b) $\Delta H < \Delta E$
 - (c) $\Delta H = \Delta E$

>>>

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(d) The relationship depends on the capacity of the vessel

Which of the following is always negative for exothermic reaction? [BCECE 2005]

- (a) ΔH (b) ΔS (c) ΔG
 - (d) None of these

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- **4.** The relation between ΔE and ΔH is
 - [MP PET 1992; MP PMT 1996; MP PET/PMT 1998]

(a)
$$\Delta H = \Delta E - P \Delta V$$
 (b) $\Delta H = \Delta E + P \Delta V$

(c)
$$\Delta E = \Delta V + \Delta H$$
 (d) $\Delta E = \Delta H + P \Delta V$

5. At constant *T* and *P*, which one of the following statements is correct for the reaction, $CO(g) + \frac{1}{2}O_2(g) \rightarrow CO_2(g)$

[AIIMS 1982, 83; KCET 1988; BHU 1995; MP PET 1997, 99]

- (a) ΔH is independent of the physical state of the reactants of that compound
- (b) $\Delta H > \Delta E$
- (c) $\Delta H < \Delta E$
- (d) $\Delta H = \Delta E$
- 6. For the reaction of one mole of zinc dust with one mole of sulphuric acid in a bomb calorimeter, ΔU and w correspond to [AIIMS 2005]
 (a) ΔU < 0, w = 0 (b) ΔU = 0, w < 0
 - (c) $\Delta U > 0, w = 0$ (d) $\Delta U < 0, w > 0$
- 7. Which is not the correct relation between enthalpy (ΔH) and intrinsic energy (ΔE) [EAMCET 1992]
 - (a) $\Delta H = \Delta E + P \Delta V$ (b) $\Delta H = \Delta E + n RT$
 - (c) $\Delta H = \Delta E P \Delta V$ (d) $\Delta E = \Delta H P \Delta V$
- 8. The law of Lavoisier and Laplace illustrates

[KCET 1989]

- (a) The principle of conservation of energy
- (b) Equivalence of mechanical and thermal energy
- (c) The principle of conservation of matter
- (d) Equivalence of mechanical and chemical energy
- **9.** For the reaction $N_2 + 3H_2 = 2NH_3$; $\Delta H =$

[Roorkee 2000; CBSE PMT 1991, 2002]

(a) $\Delta E - RT$ (b) $\Delta E - 2R$

- (c) $\Delta E + RT$ (d) $\Delta E + 2RT$
- **10.** If ΔH is the change in enthalpy and ΔE the change in internal energy accompanying a gaseous reaction

[KCET 1989; CBSE PMT 1990]

- (a) ΔH is always greater than ΔE
- (b) $\Delta H < \Delta E$ only if the number of moles of the products is greater than the number of the reactants
- (c) ΔH is always less than ΔE
- (d) $\Delta H < \Delta E$ only if the number of moles of the products is less than the number of moles of the reactants
- "The resultant heat change in a reaction is the same whether it takes place in one or several stages." This statement is called [MP PMT/PET 1988; MP PMT 1989]
 - (a) Lavoisier and Laplace law
 - (b) Hess's law

- (c) Joule's law
- (d) Le-chatelier's principle
- **12.** Hess's law of constant heat summation in based on

[MP PET 2001]

- (a) $E = mc^2$
- (b) Conservation of mass
- (c) First law of thermodynamics
- (d) None of the above
- **13.** Consider the reaction : $N_2 + 3H_2 \rightarrow 2NH_3$ carried out at constant temperature and pressure. If ΔH and ΔU are the enthalpy and internal energy changes for the reaction, which of the following expression is true **[AIEEE 2005]**
 - (a) $\Delta H = 0$ (b) $\Delta H = \Delta U$
 - (c) $\Delta H < \Delta U$ (d) $\Delta H > \Delta U$
- 14. Which of the following is the correct equation
 - [CBSE PMT 1996]
 - (a) $\Delta U = \Delta Q W$ (b) $\Delta W = \Delta U + \Delta Q$
 - (c) $\Delta U = \Delta W + \Delta Q$ (d) None of these
- **15.** Hess law is applicable for the determination of heat of

[AIIMS 1998; Pb. PET/PMT 1999]

(a) Reaction	(b) Formation
(c) Transition	(d) All of these

- **16.** Enthalpy for the reaction $C + O_2 \rightarrow CO_2$ is
 - [DPMT 1987, 91]
 - (a) Positive (b) Negative
 - (c) Zero (d) None
- **17.** During an isothermal expansion of an ideal gas its

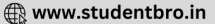
[CBSE PMT 1991]

- (a) Internal energy increases(b) Enthalpy decreases
- (c) Enthalpy remains unaffected
- (d) Enthalpy reduces to zero
- 18. The work done in ergs for the reversible expansion of one mole of an ideal gas from a volume of 10 litres to 20 litres at 25 ° C is [CMC Vellore 1991]
 - (a) $2.303 \times 298 \times 0.082 \log 2$
 - (b) $298 \times 10^7 \times 8.31 \times 2.303 \log 2$
 - (c) $2.303 \times 298 \times 0.082 \log 0.5$
 - (d) $8.31 \times 10^7 \times 298 2.303 \log 0.5$
 - (e) $2.303 \times 298 \times 2\log 2$
- 19. The law of conservation of energy states that

[NCERT 1984]

- (a) The internal energy of a system is constant
- (b) The heat content of a system is constant
- (c) Energy is neither created nor destroyed
- (d) There is an equivalence between energy and mass





- The enthalpies of the elements in their standard states are 20. assumed to be (a) Zero at 298 K (b) Unit at 298 K (c) Zero at all temperatures (d) Zero at 273 K The heat *Q* for a reaction at constant volume is equal to 21. (b) $H_{R} - H_{P}$ (a) $H_P - H_R$ (d) None of these (c) $E_P - E_R$ Which of the following expressions represents the first 22. law of thermodynamics [MP PET 1996, 2000; AFMC 1997; BHU 1999; AMU 2000; KCET (Med.) 2000, 01; CBSE PMT 2000; MP PMT 2002] (a) $\Delta E = -q + W$ (b) $\Delta E = q - W$ (d) $\Delta E = -q - W$ (c) $\Delta E = q + W$ In a reversible isothermal process, the change in internal 23. energy is (a) Zero (b) Positive (c) Negative (d) None of these 24. A schematic plot of lnK_{eq} versus inverse of temperature for a reaction is shown below 6.0 Keqц 2.0 1.5×10-3 $1/T(K^{-1})$ 2.0×10⁻³ The reaction must be [AIEEE 2005] (a) Exothermic (b) Endothermic (c) One with negligible enthalpy change (d) Highly spontaneous at ordinary temperature Hess law of heat summation includes [AFMC 1992] 25. (a) Initial reactants only
 - (b) Initial reactants and final products
 - (c) Final products only
 - (d) Intermediates only
- 26. An ideal gas at constant temperature and pressure expands, then its [BHU 1998]
 - (a) Internal energy remains same
 - (b) Internal energy decreases
 - (c) Internal energy increases

27.

- (d) Entropy first increases and then decreases
- The internal energy of a substance

[KCET 1998; AFMC 2001; AIIMS 2001]

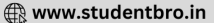
- (a) Increases with increase in temperature
- (b) Decreases with increase in temperature
- (c) Can be calculated by the relation $E = mc^2$
- (d) Remains unaffected with change in temperature
- The relation between change in internal energy (ΔE) , 28. change in enthalpy (ΔH) and work done (W) is represented as [AFMC 1999] (a) $\Delta H = \Delta E + W$ (b) $W = \Delta E - \Delta H$ (c) $\Delta E = W - \Delta H$ (d) $\Delta E = \Delta H + W$ The enthalpy of neutralization of which of the following 29. acids and bases is nearly -13.6 Kcal [Roorkee 1999] (a) HCN and NaOH (b) HCl and KOH (c) HCl and NaOH (d) HCl and NH₄OH 30. Work done during isothermal expansion of one mole of an ideal gas from 10 atm to 1 atm at 300 K is (Gas constant = [AIIMS 2000] 2) (a) 938.8 cal. (b) 1138.8 cal. (c) 1381.8 cal. (d) 1581.8 cal. Joule-Thomson expansion is [JIPMER 2000] 31. (a) Isobaric (b) Isoenthalpic (c) Isothermal (d) None of these In which of the following $\Delta E = \Delta H$ [MP PMT 2001] 32. (a) $N_2O_4(g) \rightleftharpoons 2NO_2(g)$ (b) $2SO_2(g) + O_2(g) \Rightarrow 2SO_3(g)$ (c) $H_2(g) + I_2(g) \Rightarrow 2HI(g)$ (d) $H_2(g) + \frac{1}{2}O_2(g) \Rightarrow H_2O(l)$ In an adiabatic expansion of an ideal gas 33. [KCET (Med.) 2001; MH CET 2000] (a) $W = -\Delta E$ (b) $W = \Delta E$ (c) $\Delta E = 0$ (d) W = 034. For the reaction $CH_{3}COOH(l) + 2O_{2}(g) = 2CO_{2}(g) + 2H_{2}O(l)$ at $25^{\circ}C$ and 1 *atm*. pressure, $\Delta H = -874 \ kJ$. Then the change in internal energy (ΔE) is [Orissa JEE 2002] (a) -874 kJ(b) -871.53 kJ(c) $-876.47 \, kJ$ (d) + 874 kJIn a closed insulated container, a liquid is stirred with a 35. paddle to increase its temperature. In this process, which [CBSE PMT 2002] of the following is true (a) $\Delta E = W = Q = 0$ (b) $\Delta E \neq 0, Q = W = 0$ (d) $\Delta E = Q \neq 0, W = 0$ (c) $\Delta E = W \neq 0, Q = 0$ 36. Hess law is based on [MH CET 2002] (a) Law of conservation of mass (b) Law of conservation of energy (c) Enthalpy is a state function (d) None of these
- **37.** For the reaction,

 $C_3H_8(g) + 5O_2(g) \rightarrow 3CO_2(g) + 4H_2O(l)$

at constant temperature, $\Delta H - \Delta E$ is **[CBSE PMT 2003]**

- (a) -RT (b) +RT
- (c) -3RT (d) +3RT
- **38.** According to Hess's law, the heat of reaction depends upon





- (a) Initial condition of reactants
- (b) Initial and final conditions of reactants
- (c) Intermediate path of the reaction
- (d) End conditions of reactants
- **39.** The value of $\Delta H \Delta E$ for the following reaction at 27 ° C will be $2NH_3(g) \rightarrow N_2(g) + 3H_2(g)$ [Kerala (Med.) 2002]
 - (a) $8.314 \times 273 \times (-2)$ (b) $8.314 \times 300 \times (-2)$

(c)
$$8.314 \times 27 \times (-2)$$
 (d) $8.314 \times 300 \times (2)$

40. For which of the following $\Delta E = \Delta H$ [MP PET 2003]

(a)
$$N_2O_4(g) \rightleftharpoons 2NO_2(g)$$

(b) $2SO_2(g) + O_2(g) = 2SO_3(g)$

(c)
$$H_2(g) + Cl_2(g) \Rightarrow 2HCl(g)$$

(d)
$$H_2(g) + \frac{1}{2}O_2(g) \Rightarrow H_2O(l)$$

41. One mole of an ideal gas is allowed to expand freely and adiabatically into vacuum until its volume has doubled. A statement which is not true concerning this expression is

[Pb. PMT 1998]

[MP PMT 2003]

- (a) $\Delta H = 0$ (b) $\Delta S = 0$ (c) $\Delta E = 0$ (d) W = 0
- **42.** At $27^{\circ}C$, one mole of an ideal gas is compressed isothermally and reversibly from a pressure of 2 *atm* to 10 *atm*. The values of ΔE and *q* are (*R* = 2) [BHU 2001]
 - (a) 0, 965.84 *cal*
 - (b) 965.84 cal, + 965.84 cal
 - (c) + 865.58 *cal*, 865.58 *cal*
 - (d) 865.58 cal, 865.58 cal
- **43.** The work done by a system is 8 *joule*, when 40 *joule* heat is supplied to it. What is the increase in internal energy of system

[BHU 2001]		U U
	(b) 30 <i>J</i>	(a) $25 J$
	(d) 28 J	(c) $32 J$

44. A system absorb 600*J* of heat and work equivalent to 300*J* on its surroundings. The change in internal energy is

							[Pb.	PM	T 20	004]
(a)	300 J				(b)	400 J					
(c)	500 J				(d)	600 J					
* • *	1 1	1		. 1	1		c		1	c	

- 45. Work done during isothermal expansion of one mole of an ideal gas from 10 atom. to 1 atm at 300K is [BHU 2004]
 (a) 4938.8 J
 (b) 4138.8 J
 - (c) 5744.1J (d) 6257.2J
- **46.** If gas, at constant temperature and pressure expands then its

[MH CET 2003]

- (a) Entropy increases and then decreases
- (b) Internal energy increases
- (c) Internal energy remains the same
- (d) Internal energy decreases

For the reaction 47. $PCl_5(g) \rightarrow PCl_3(g) + Cl_2(g)$ [MH CET 2004] (a) $\Delta H = \Delta E$ (b) $\Delta H > \Delta E$ (c) $\Delta H < \Delta E$ (d) None of these 48. Enthalpy (H) is equal to [MH CET 2004] (a) Internal energy (E) (b) Product of pressure (P) and volume (V) of gas (c) Internal energy (E)+PV (d) Work (W) done by a system ΔE° of combustion of isobutylene is $-X k J mol^{-1}$. The 49. value of ΔH^{o} is [DCE 2004] (a) = ΔE^{o} (b) $> \Delta E^{\circ}$ (c) = 0(d) $<\Delta E^{o}$ One mole of an ideal gas is allowed to expand reversibly 50. and adibatically from a temperature of $27^{\circ}C$. If the work done during the process is 3 kJ, then final temperature of the gas is $(C_V = 20 J / K)$ [Pb. CET 2002] (a) 100 K (b) 150 K (c) 195 K (d) 255 K Which of the following is correct regarding the internal 51. energy of a substance [Pb. CET 2002] (a) It is a state function (b) It increases with increase in temperature (c) It can be calculated by experiment (d) It remains unaffected with change in temperature For the gaseous reaction, 52. $N_2O_4 \rightarrow 2NO_2$ [Pb. CET 2003] (a) $\Delta H < \Delta E$ (b) $\Delta H = \Delta E$ (c) $\Delta H = 0$ (d) $\Delta H > \Delta E$ $2C + O_2 \rightarrow 2CO; \Delta H = -220 \, kJ$ 53.

Which of the following statement is correct for this reaction

[BVP 2004]

[AFMC 2004]

- (a) Heat of combustion of carbon is 110 kJ
- (b) Reaction is exothermic
- (c) Reaction needs no initiation
- (d) All of these are correct

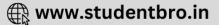
54. An ideal gas expands in volume from $1 \times 10^{-3} m^3$ to $1 \times 10^{-2} m^3$ at 300 K against a constant pressure of $1 \times 10^5 Nm^{-2}$. The work done is **[AIEEE 2004]**

- (a) 270 kJ (b) -900 kJ
- (c) -900 J (d) 900 kJ

55. Internal energy is

- (a) Partly potential and partly kinetic
- (b) Totally kinetic
- (c) Totally potential
- (d) None of these
- **56.** For which of the reaction $\Delta H = \Delta E$ [MP PET 2004]





- (a) $N_2 + 3H_2 \Rightarrow 2NH_3$ (b) $N_2 + O_2 \rightarrow 2NO$
- (c) $2Na + Cl_2 \rightarrow 2NaCl$ (d) $PCl_5 \rightarrow PCl_3 + Cl_2$
- **57.** The work done during the expansion of a gas from a volume of $4dm^3$ to $6dm^3$ against a constant external pressure of 3atm is (1Latm = 101.32 J)[CBSE PMT 2004]
 - (a) +304J (b) -304J
 - (c) -6J (d) -608J

58. For the reaction, $A_{(s)} + 2B_{(g)} \rightarrow 4C_{(s)} + D_{(l)}$. ΔH and ΔU are related as **[Orissa JEE 2004]**

(a) $\Delta H = \Delta U$ (b) $\Delta H = \Delta U + 3RT$

(c) $\Delta H = \Delta U + RT$ (d) $\Delta H = \Delta U - 3RT$

IInd & IIIrd Law of thermodynamics and Entropy

1. For a reversible spontaneous change ΔS is

(a)
$$\frac{\Delta E}{T}$$
 (b) $\frac{P \Delta V}{T}$
(c) $\frac{q}{T}$ (d) $RT \log K$

- **2.** When disorder of a system increases, the change is said to be
 - (a) Exothermic (b) Non-spontaneous
 - (c) Endothermic (d) Spontaneous
- **3.** The spontaneous flow of heat is always
 - (a) From low to high pressure
 - (b) From high to high pressure
 - (c) Unidirectional from lower temperature to higher temperature
 - (d) Unidirectional from the higher to lower temperature
 - Mixing of non-reacting gases is generally accompanied by
 - (a) Decrease in entropy

4.

7.

- (b) Increase in entropy
- (c) Change in enthalpy
- (d) Change in free energy
- **5.** An irreversible process occuring isothermally in an isolated system leads to
 - (a) Zero entropy
 - (b) An increase in the total entropy of the system
 - (c) A decrease in the total entropy of the system
 - (d) None of these
- **6.** The entropy values (in $JK^{-1} mol^{-1}$) of $H_{2(g)} = 130.6$, $Cl_{2(g)} = 223.0$ and $HCl_{(g)} = 186.7$ at 298 *K* and 1 *atm* pressure. Then entropy change for the reaction

 $H_{2(g)} + Cl_{2(g)} \rightarrow 2HCl_{(g)}$ is [BHU 2005]

(a) +540.3 (b) +727.3

(c)
$$-166.9$$
 (d) $+19.8$

- Which of the following is the least random state of water (a) Ice
 - (b) Liquid water
 - (c) Steam
 - (d) All present same random state
- 8. Which one of the following process is non-spontaneous
 - (a) Dissolution of $CuSO_4$ in water
 - (b) Reaction between H_2 and O_2 to form water

- (c) Water flowing down hill
- (d) Flow of electric current from low potential to high potential
- **9.** Which of the following is zero during adiabatic expansion of the gas
 - (a) ΔT (b) ΔS
 - (c) ΔE (d) None of these
- **10.** The entropy of crystalline substances at absolute zero going by the third law of thermodynamics should be taken as

[J & K 2005]

- (b) 50
- (c) Zero

11.

- (d) Different for different substances
- In which state, the matter have highest entropy
- (a) Solid (b) Liquid
- (c) Gas (d) Equal in all
- **12.** Which of the following pairs of a chemical reaction is certain to result in spontaneous reaction ?**[CBSE PMT 2005]**
 - (a) Exothermic and decreasing disorder
 - (b) Endothermic and increasing disorder
 - (c) Exothermic and increasing disorder
 - (d) Endothermic and decreasing disorder
- **13.** When one mole of monoatomic ideal gas at T *K* undergoes adiabatic change under a constant external pressure of 1 atm changes volume from 1 litre to 2 litre. The final temperature in Kelvin would be **[IIT 2005]**

(a)
$$\frac{T}{2^{(2/3)}}$$
 (b) $T + \frac{2}{3 \times 0.0821}$
(c) T (d) $T - \frac{2}{3 \times 0.0821}$

c) T (d)
$$T - \frac{2}{3 \times 0.0821}$$

- 14. 9.0 gm of H_2O is vaporised at 100 °C and 1 atm pressure. If the latent heat of vaporisation of water is x J/gm, then ΔS is given by
 - (a) $\frac{x}{373}$ (b) $\frac{18x}{100}$
 - (c) $\frac{18x}{373}$ (d) $\frac{1}{2} \times \frac{18x}{373}$
- **15.** The ΔS for the vaporisation of 1 mol of water is 88.3 J/mole K. The value of ΔS for the condensation of 1 mol of vapour will be

(a)
$$88.3 J/mol K$$
 (b) $(88.3)^2 J/mol K$

(c)
$$-88.3 J/mol K$$
 (d) $\frac{1}{88.3} J/mol K$

16. For which reaction from the following, ΔS will be maximum

[AlIMS 1982, 83, 94; AMU 2000]

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(a)
$$Ca(s) + \frac{1}{2}O_2(g) \rightarrow CaO(s)$$

- (b) $CaCO_3(s) \rightarrow CaO(s) + CO_2(g)$
- (c) $C(s) + O_2(g) \rightarrow CO_2(g)$

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(d) $N_2(g) + O_2(g) \rightarrow 2NO(g)$

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		The
17.	The occurrence of a reaction is impossible if [AIIMS 1982, 91; MP PET 1997; MP PMT 1999; DPMT 2002]	27
	(a) ΔH is +ve; ΔS is also + ve but $\Delta H < T\Delta S$ (b) ΔH is -ve; ΔS is also - ve but $\Delta H > T\Delta S$	28
18.	(c) ΔH is $-$ ve; ΔS is $+$ ve (d) ΔH is $+$ ve; ΔS is $-$ ve If the enthalpy of vaporization for water is 186.5 <i>kJ</i> mol^{-1} , the entropy of its vaporization will be [CPMT 1988; DCE 2000]	20
	(a) $0.5 JK^{-1}mol^{-1}$ (b) $1.0 JK^{-1}mol^{-1}$	
	(c) $1.5 JK^{-1}mol^{-1}$ (d) $2.0 JK^{-1}mol^{-1}$	
19.	Which of the following statement is true. The entropy of the universe [MP PMT 1993; CET Pune 1998]	30
	(a) Increases and tends towards maximum value(b) Decreases and tends to be zero	
	(c) Remains constant	31
	(d) Decreases and increases with a periodic rate	
20.	When enthalpy and entropy change for a chemical reaction are -2.5×10^{3} cal and 7.4 cal deg ⁻¹ respectively. Predict the reaction at 298 <i>K</i> is	
	[AFMC 1998; MH CET 1999; CBSE PMT 2000]	
	(a) Spontaneous(b) Reversible(c) Irreversible(d) Non-spontaneous	32
21.	The total entropy change for a system and its surroundings increases, if the process is (a) Reversible (b) Irreversible	
	(c) Exothermic (d) Endothermic	
22.	For chemical reactions, the calculation of change in entropy is normally done	
	(a) At constant pressure(b) At constant temperature	
	(c) At constant temperature and pressure both	33
	(d) At constant volume	
23.	When the value of entropy is greater, then the ability of work	
	(a) Is maximum(b) Is minimum(c) Is medium(d) None of these	
24.		34
•	$H_2O(l) \Rightarrow H_2O(g)$ at 100 °C and 1 atmosphere	0
	[KCET 1991; AIIMS 1996]	
	(a) $\Delta E = 0$ (b) $\Delta H = 0$	
25.	(c) $\Delta H = \Delta E$ (d) $\Delta H = T\Delta S$ The enthalpy change for the transition of liquid water to	35
-J.	steam, $\Delta H_{vap} = 37.3 kJ mol^{-1}$ at 373 K. The entropy	
	change for the process is [MP PET 1994]	
	(a) $111.9 Jmol^{-1}K^{-1}$ (b) $37.3 Jmol^{-1}K^{-1}$	
	(c) $100 Jmol^{-1}K^{-1}$ (d) $74.6 Jmol^{-1}K^{-1}$	30
26.	When a solid is converted into liquid, entropy [MP PET 1993, 95; MP PMT 1996]	
	(a) Becomes zero (b) Remains the same	

Inern	nodynamics and Thermochemistry 419
27.	In a spontaneous process, the entropy of the system and its surroundings [MP PET 1996; NCERT 1975]
	(a) Equals zero(b) Decreases(c) Increases(d) Remains constant
28.	The positive value of ΔS indicates that [MP PMT 1997]
20,	(a) The system becomes less disordered
	(b) The system becomes more disordered
	(c) The system is in equilibrium position
	(d) The system tends to reach at equilibrium position
29.	For reaction $2Cl(s) \rightarrow Cl_2(s)$, the signs of ΔH and ΔS
-,-	respectively are [KCET 1996]
	(a) $+,-$ (b) $+,+$
	(c) $-, -$ (d) $-, +$
30.	The enthalpy of water is $386 kJ$. What is entropy of water
0	[BHU 1997]
	(a) $0.5 kJ$ (b) $1.03 kJ$
	(c) $1.5 kJ$ (d) $22.05 kJ$
31.	For which of the processes is ΔS negative [Pb. PMT 1998]
<u> </u>	(a) $H_2(g) \rightarrow 2H(g)$
	(b) $N_2(g)1atm \rightarrow N_2(g)8 atm$
	(c) $2SO_3(g) \rightarrow 2SO_2(g) + O_2(g)$
	(d) $C_{(diamond)} \rightarrow C_{(graphite)}$
32.	Identify the correct statement regarding entropy
J	[CBSE PMT 1998; BHU 2001]
	(a) At $0^{\circ}C$, the entropy of a perfectly crystalline
	substance is taken to be zero
	(b) At absolute zero of temperature, the entropy of a perfectly crystalline substance is + <i>ve</i>
	(c) At absolute zero of temperature, the entropy of all crystalline substances is taken to be zero
	(d) At absolute zero of temperature, the entropy of a perfectly crystalline substance is taken to be zero
33.	One mole of an ideal gas at $300 K$ is expanded isothermally from an initial volume of 1 litre to 10 litres. The change in energy for this process is $(R = 2 cal mol^{-1} K^{-1})$
	[CBSE PMT 1998] (a) 163.7 cal (b) 850.2 cal
	(a) 163.7 cal (b) 850.2 cal (c) 1381.1 cal (d) Zero
0.4	
34.	
	then what is increase in entropy [BHU 1998]
	(a) $43.4 J/mole$ (b) $87.2 J/mole$
e -	(c) 900 <i>J/mole</i> (d) Zero
35.	A container has hydrogen and oxygen mixture in ratio of 4 : 1 by weight, then [BHU 1998]
	(a) Internal energy of the mixture decreases
	(b) Internal energy of the mixture increases
	(c) Entropy of the mixture increases
	(d) Entropy of the mixture decreases

36. If 0.75 *mole* of an ideal gas expands isothermally at 27 ° C from 15 *litres* to 25 *litres*, the maximum work obtained is **[AFMC 1999]**

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(a) 8.40 J	(b) 9.34 <i>J</i>
(c) 10.86 <i>J</i>	(d) 10.43 <i>J</i>

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(d) Increases

(c) Decreases

420 Thermo	dynamics and	d Thermochemistry	
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3 7•	In which of the following conditions a chemical reaction can not occur [AMU 1999]		
	(a) ΔH and ΔS increase and $T\Delta S > \Delta H$	I	
	(b) ΔH and ΔS decrease and $\Delta H > T\Delta S$		
	(c) ΔH increases and ΔS decreases		
	(d) ΔH decreases and ΔS increases		
38.	An engine operating between $150^{\circ}C$ and $25^{\circ}C$ takes	2	
, 0.	500 J heat from a higher temperature reservoir if there are no frictional losses, then work done by engine is	e .	
	[MH CET 1999]		
	(a) $147.7J$ (b) $157.75J$		
~	(c) $165.85J$ (d) $169.95J$		
9.	The entropy of a perfectly crystalline solid at absolute zero is)	
	[AMU 2000]	1	
	(a) Positive (b) Negative		
	(c) Zero (d) Not definite		
).			
	non-spontaneous change [AMU 2000; KCET (Med.) 2000)]	
	(a) Positive ΔH and positive ΔS		
	(b) Negative ΔH and negative ΔS		
	(c) Positive ΔH and negative ΔS		
	(d) Negative ΔH and positive ΔS		
1.	Following data is known about melting of a compound		
	<i>AB</i> . $\Delta H = 9.2 kJ mol^{-1}$, $\Delta S = 0.008 kJ K^{-1} mol^{-1}$. Its		
	melting point is [Pb. PMT 2000; AIIMS 2000]	J	
	(a) 736 <i>K</i> (b) 1050 <i>K</i>		
	(c) 1150 K (d) 1150 ^{o}C		
2.	The volume of a gas decreases from 500 cc to 300 cc		
	when a sample of gas is compressed by an average		
	pressure of 0.6 <i>atm</i> . During this process 10 J of heat is		
	liberated. The change in internal energy is [Pb. PMT 2000)]	
	(a) $-2.16 J$ (b) $12.156 J$		
	(c) $2.16 J$ (d) $101.3 J$		
3.	The standard entropies of $CO_2(g), C(s)$ and $O_2(g)$ are)	
	213.5, 5.690 and 205 JK^{-1} respectively. The standard	ł	
	entropy of formation of $CO_2(g)$ is [CPMT 2001]	1	
	(a) $1.86 JK^{-1}$ (b) $1.96 JK^{-1}$		
	(c) $2.81 JK^{-1}$ (d) $2.86 JK^{-1}$		
4.	Which of the following endothermic processes are spontaneous		

- spontaneous[CBSE PMT 2001](a) Melting of ice(b) Evaporation of water
 - (c) Heat of combustion (d) Both (a) and (b)
- **45.** For the reaction $H_2O(s) = H_2O(l)$ at $0^{\circ}C$ and normal pressure **[KCET 2001]**
 - (a) $\Delta H > T \Delta S$ (b) $\Delta H = T \Delta S$
 - (c) $\Delta H = \Delta G$ (d) $\Delta H < T\Delta S$
- **46.** One mole of an ideal gas is allowed to expand reversibly and adiabatically from a temperature of 27 ° C. If the work done during the process is 3 kJ, the final temperature will be equal to $(C_v = 20 JK^{-1})$

[KCET 2000; AFMC 2000; AIIMS 2001]

- (a) 150 *K* (b) 100 *K*
- (c) $26.85^{\circ}C$ (d) 295K

47. The entropy change, in the conversion of one mole of liquid water at 373 *K* to vapour at the same temperature is (Latent heat of vaporization of water, $\Delta H_{vap} = 2.257 \text{ kJ}/\text{g}$)

[AFMC 2002; KCET 2002]

- (a) $105.9 Jk^{-1}$ (b) $107.9 Jk^{-1}$ (c) $108.9 Jk^{-1}$ (d) $109.9 Jk^{-1}$
- **48.** A heat engine absorbs heat Q_1 at temperature T_1 and heat Q_2 at temperature T_2 . Work done by the engine is $(Q_1 + Q_2)$. This data **[AIEEE 2002]**
 - (a) Violates Ist law of thermodynamics
 - (b) Violates Ist law of thermodynamics if Q_1 is -ve
 - (c) Violates Ist law of thermodynamics if Q_2 is -ve
 - (d) Does not violate Ist law of thermodynamics
- **49.** Equal volumes of monoatomic and diatomic gases at same initial temperature and pressure are mixed. The ratio of specific heats of the mixture (C_p / C_v) will be

[AFMC 2002]

[CBSE PMT 2002]

- (a) 1 (b) 2
- (c) 1.67 (d) 1.5 **50.** The law formulated by Dr. Nernst is [Kerala (Med.) 2002]
 - (a) First law of thermodynamics
 - (b) Second law of thermodynamics
 - (c) Third law of thermodynamics
 - (d) Both (a) and (b)

51. The unit of entropy is

- (a) $J mol^{-1}$ (b) $JK mol^{-1}$
- (c) $J mol^{-1}K^{-1}$ (d) $J^{-1}K^{-1}mol^{-1}$
- **52.** The entropy changed involved in the conversion of 1 *mole* of liquid water at 373 K to vapour at the same temperature will be
 - $[\Delta H_{vap} = 2.257 \, kJ \,/ \, gm]$ [MP PET 2002]
 - (a) 0.119 kJ (b) 0.109 kJ
 - (c) 0.129 kJ (d) 0.120 kJ
- **53.** When a liquid boils, there is
 - (a) An increase in entropy
 - (b) A decrease in entropy
 - (c) An increase in heat of vaporization
 - (d) An increase in free energy

54. Which one of the following has ΔS^{o} greater than zero

[AIIMS 2003]

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[JIPMER 2002]

- (a) $CaO + CO_2(g) \rightleftharpoons CaCO_3(s)$
- (b) $NaCl(aq) \Rightarrow NaCl(s)$

- (c) $NaNO_3(s) \Rightarrow Na^+(aq) + NO_3^-(aq)$
- (d) $N_2(g) + 3H_2(g) = 2NH_3(g)$
- **55.** The molar heat capacity of water at constant pressure is $75 \ JK^{-1}mol^{-1}$. When 1.0 *kJ* of heat is supplied to 100 *g* of water which is free to expand the increase in temperature of water is **[CBSE PMT 2003]** (a) 6.6 K (b) 1.2 K
 - (c) 2.4 K (d) 4.8 K

			Therm	modynamics and Thermochemistry 421	
56.	joules. 128 joule energy is	ct a gas in a cylinder, is 462 evolved in the process. What change in the process [MP PMT :	67.	pressure. Then [KCET	re and [2004]
	(a) $+ 590$ joules	(b) -334 joules	20031	(a) $\Delta H = \Delta E$ (b) $\Delta H = T\Delta S$	
	(c) $+ 334 joules$	(d) -590 joules		(c) $\Delta H = 0$ (d) $\Delta S = 0$	1
5 7.	Highest entropy is in	[BVP 2003]	68.	The enthalpy of vapourization of a liquid is $30 k$.	$J mol^{-1}$
	(a) Hydrogen	(b) Water		and entropy of vapourization is 75 $J mol^{-1}K$. The	-
	(c) Graphite	(d) Mercury		point of the liquid at 1 <i>atm</i> is [IIT JEE (Screening)	2004]
58.	$H^+(aq) + OH^-(aq) - S^o(298 K)K^{-1}mol^{-1} - 10.7$	$\longrightarrow H_2O(l)$ ₊₇₀		(a) 250K (b) 400K (c) 450K (d) 600K	
	Standard entropy change for	or the above reaction is [DPMT 2004]		Heat of reaction	
	(a) $60.3 JK^{-1}mol^{-1}$	(b) 80.7 $JK^{-1}mol^{-1}$			
	(c) $-70 \ JK^{-1}mol^{-1}$	(d) $+10.7 JK^{-1}mol^{-1}$	1.	If $C + O_2 \rightarrow CO_2 + 94.2 kcal$	
59 .	One mole of water at 100	°C is converted into steam at		$H_2 + \frac{1}{2}O_2 \rightarrow H_2O + 68.3 kcal$	
		sure of 1 atm. The change in vaporisation of water at		$CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O + 210.8 \ kcal$	
	$100^{\circ}C = 540 \ cal \ / \ gm$	[Pb. PMT 2004]		then the possible heat of methane will be [MP PE]	Г 1990]
	(a) 8.74	(b) 18.76		(a) 47.3 <i>kcal</i> (b) 20.0 <i>kcal</i>	
	(c) 24.06	(d) 26.06		(c) $45.9 \ kcal$ (d) $-47.3 \ kcal$	
60.	For a carnot engine, the so	urce is at 500 K and the sink at	2.	Molar heat of vaporisation of a liquid is $6kJ mol^{-1}$. If the
	300 K. What is efficiency of (a) 0.2			entropy change is $16 J mol^{-1} K^{-1}$, the boiling point	t of the
	(c) 0.6	(d) 0.3		liquid is [KCE]	[2005]
61.		e temperature of 1 mol of a		(a) $375^{\circ}C$ (b) $375K$	
	substance by $1^{\circ}C$ is called			(c) $273 K$ (d) $102^{\circ} C$	
	(a) Specific heat	(b) Molar heat capacity	3.	The enthalpy of fusion of ice per mole	
	(c) Water equivalent	(d) Specific gravity		(a) $18 kJ$ (b) $8 kJ$	
62.	Maximum entropy will be i	n which of the following		(c) $80 kJ$ (d) $6 kJ$	
		[DCE 2004]	4.	In which of the following neutralisation reactio	ns, the
	(a) Ice	(b) Liquid water		heat of neutralisation will be highest	~ 1
	(c) Snow	(d) Water vapours		[MP PMT 1989, 91; AIIM5	5 1999]
63.		f $CO_2(g), C(s)$ and $O_2(g)$ are		(a) NH_4OH and CH_3COOH	
		respectively. The standard		(b) NH_4OH and HCl	
	entropy of formation of CC	<i>P</i> ₂ is [Pb. CET 2001]		(c) <i>NaOH</i> and CH_3COOH	
	(a) 2.76 JK^{-1}	(b) 2.12 JK^{-1}		(d) NaOH and HCl	
	(c) $1.12 JK^{-1}$	(d) 1.40 JK^{-1}	5.	If $S + O_2 \rightarrow SO_2$; ($\Delta H = -298.2$)	
64.	Which law of thermody entropy at different temper	anamics help in calculating ratures [Pb. CET 2003]		$SO_2 + \frac{1}{2}O_2 \rightarrow SO_3; (\Delta H = -98.2)$	
	(a) First law	(b) Second law		$SO_3 + H_2O \rightarrow H_2SO_4; (\Delta H = -130.2)$	
	(c) Third law	(d) Zeroth law			
65.	Entropy is maximum in cas			$H_2 + \frac{1}{2}O_2 \rightarrow H_2O; (\Delta H = -287.3)$	
-	(a) Steam	(b) Water at $0^{\circ}C$		then the enthalpy of formation of H_2SO_4 at 298K s	will be
	(c) Water at $4^{\circ}C$	(d) Ice		[DPMT 1983; Orissa JEB	
66.		s a thermodynamic parameter,		(a) $-433.7 kJ$ (b) $-650.3 kJ$	
	the criterion for the sponta	neity of any process is		(c) $+320.5 kJ$ (d) $-813.9 kJ$	
		[CBSE PMT 2004]	6.	From Kirchhoff's equation which factor affects the	heat of

ss is taking place at constant temperature and

[MP PMT 1989, 91; AIIMS 1999]

- OH and CH_3COOH
- OH and HCl
- OH and CH₃COOH

$$SO_2 + \frac{1}{2}O_2 \rightarrow SO_3; (\Delta H = -98.2)$$
$$SO_3 + H_2O \rightarrow H_2SO_4; (\Delta H = -130.2)$$

$$H_2 + \frac{1}{2}O_2 \to H_2O; (\Delta H = -287.3)$$

		[DPMT 1983; Orissa JEE 2005]
(a)	–433.7 kJ	(b) $-650.3 kJ$
(c)	+320.5 kJ	(d) $-813.9 kJ$

6. From Kirchhoff's equation which factor affects the heat of reaction [MP PMT 1990]

- (b) Temperature (a) Pressure
- (d) Molecularity (c) Volume

(b) $\Delta S_{surroundin gs} > 0$ only (c) $\Delta S_{system} + \Delta S_{surroundin gs} > 0$

(a) $\Delta S_{system} > 0$ only

(d) $\Delta S_{system} - \Delta S_{surroundin\,gs} > 0$

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- The molar neutralization heat for KOH and HNO3 as 7. compared to molar neutralization heat of NaOH and HCl
 - (a) Less
 - (c) Equal
- (b) More (d) Depends on pressure

[MP PMT 1989]

- 8. An exothermic reaction is one in which the reacting [CPMT 1974, 79; Bihar MEE 1982; substances KCET 1992; JIPMER 2001]
 - (a) Have more energy than the products
 - (b) Have less energy than the products
 - (c) Are at a higher temperature than the product
 - (d) None of the above
- The heat evolved in the combustion of benzene is given by 9.

$$C_6H_6 + 7\frac{1}{2}O_2 \rightarrow 6CO_2(g) + 3H_2O(l); \Delta H = -3264.6 kJ$$

Which of the following quantities of heat energy will be evolved when $39 g C_6 H_6$ are burnt

[NCERT 1978; MP PET 1990; JIPMER 2001]

- (a) 816.15 kJ (b) 1632.3 kJ
- (c) 6528.2 kJ (d) 2448.45 kJ

Thermochemical reactions 10.

$$C(\text{graphite}) + \frac{1}{2}O_2(g) \rightarrow CO(g); \Delta H = -110.5 \, kJ$$
$$CO(g) + \frac{1}{2}O_2(g) \rightarrow CO_2(g); \Delta H = -283.2 \, kJ$$

From the above reaction, the heat of reaction for $C(\text{graphite}) + O_2(g) \rightarrow CO_2(g)$ will be

[CPMT 1988; MP PMT 1989]

(b) $+ 393.7 \, kJ$ (a) $- 393.7 \, kJ$ (c) -172.7 kJ(d) + 172.7 kJ

- The following is (are) endothermic reaction [IIT JEE 1999] 11.
 - (a) Combustion of methane
 - (b) Decomposition of water
 - (c) Dehydrogenation of ethane to ethylene
 - (d) Conversion of graphite to diamond
- Evaporation of water is 12.

[CPMT 1973; DPMT 1982; MP PMT 1989; MP PET 1999] (a) An endothermic change

- (b) An exothermic change
- (c) A process where no heat change occurs
- (d) A process accompanied by chemical reaction
- An exothermic reaction is one which 13.

[NCERT 1977; MP PMT 1990]

- (a) Takes place only on heating (b) Is accompanied by a flame
- (c) Is accompanied by a absorption of heat
- (d) Is accompanied by evolution of heat
- An endothermic reaction is one in which 14.

[MNR 1980; NCERT 1976]

- (a) Heat is converted into electricity (b) Heat is absorbed
- (c) Heat is evolved
- (d) Heat is converted into mechanical work
- Which of the following statement is correct
- 15. [NCERT 1978]
 - (a) ΔH is positive for exothermic reaction
 - ΔH is negative for endothermic reaction (b)
 - The heat of neutralization of strong acid and strong (c) base is always the same

(d) The enthalpy of fusion is negative

- The absolute enthalphy of neutralisation of the reaction 16. $MgO(s) + 2HCl(aq) \rightarrow MgCl_2(aq) + H_2O(l)$ will be
 - [CBSE PMT 2005]
 - (a) Less than $-57.33 \ kJ \ mol^{-1}$
 - (b) $-57.33 \text{ kJ mol}^{-1}$
 - (c) Greater than $-57.33 \text{ kJ mol}^{-1}$
 - (d) 57.33 kJ mol⁻¹
- ΔH_{f}° (298 K) of methanol is given by the chemical 17. equation [AIIMS 2005]
 - (a) $CH_4(g) + 1/2O_2(g) \rightarrow CH_3OH(g)$
 - (b) C (graphite) $+1/2O_2(g) + 2H_2(g) \rightarrow CH_3OH(l)$
 - (c) C (diamond) $+1/2O_2(g) + 2H_2(g) \rightarrow CH_3OH(l)$
 - (d) $CO(g) + 2H_2(g) \rightarrow CH_3OH(l)$
- 18. If ΔH_f^o for H_2O_2 and H_2O are $-188 \, kJ/mole$ and -286 kJ/mole. What will be the enthalpy change of the reaction $2H_2O_2(l) \rightarrow 2H_2O(l) + O_2(g)$ [MP PMT 1992]
 - (a) $-196 \, kJ / mole$ (b) 146 kJ / mole

(c)
$$-494 \, kJ / mole$$
 (d) $-98 \, kJ / mole$

The heat of transition (ΔH_t) of graphite into diamond 19. would be, where

 $C(\text{graphite}) + O_2(g) \rightarrow CO_2(g); \Delta H = x kJ$

 $C(\text{diamond }) + C_2(g) \rightarrow CO_2(g); \Delta H = y kJ$ [Pb. PET 1985]

(a)
$$(x+y)kJmol^{-1}$$
 (b) $(x-y)kJmol^{-1}$

(c)
$$(y-x)kJmol^{-1}$$
 (d) None of these

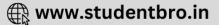
20. Correct relationship between heat of fusion (ΔH_{fus}) , heat of vaporization (ΔH_{vap}) and heat of sublimation (ΔH_{sub}) is

- (a) $\Delta H_{fus} = \Delta H_{vap} + \Delta H_{sub}$
- (b) $\Delta H_{vap} = \Delta H_{fus} + \Delta H_{sub}$
- (c) $\Delta H_{sub} = \Delta H_{vap} + \Delta H_{fus}$
- (d) $\Delta H_{sub} = \Delta H_{vap} \Delta H_{fus}$
- Which of the following is an example of endothermic 21. reaction [MP PMT 1980]
 - (a) $C_2H_2 + 2H_2 \rightarrow C_2H_6; \Delta E = -314.0 kJ$
 - (b) $C + O_2 \rightarrow CO_2$; $\Delta E = -393.5 \, kJ$
 - (c) $N_2 + O_2 \rightarrow 2NO; \Delta E 180.5 kJ$
 - (d) $2H_2 + O_2 \rightarrow 2H_2O; \Delta E + 571.8 kJ$
- If the enthalpy of B is greater than of A, the reaction 22. $A \rightarrow B$ is [MP PMT 1997]
 - (a) Endothermic (b) Exothermic
 - (c) Instantaneous (d) Spontaneous
- 23. Given that
 - $2C(s) + 2O_2(g) \rightarrow 2CO_2(g); \Delta H = -787 kJ$

$$H_2(g) + \frac{1}{2}O_2(g) \rightarrow H_2O(l); \Delta H = -286 \, kJ$$

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$$C_2H_2(g) + 2\frac{1}{2}O_2(g) \rightarrow 2CO_2(g) + H_2O(l); \Delta H = -1301 kJ$$
Heat of formation of acetylene is
[KCET 1989; Orissa JEE 2005]
(a) -1802 kJ
(b) +1802 kJ
(c) -800 kJ
(d) +228 kJ
24. Enthalpy change for reaction, $\frac{1}{2}H_2 + \frac{1}{2}CI_2 \rightarrow HCI$, is called
(a) Enthalpy of combination
(b) Enthalpy of reaction
(c) Enthalpy of formation
(d) Enthalpy of reaction
(c) Enthalpy of neutralization is about 57.3 kJ for the pair
(a) HCI and NH₄OH
(b) NH₄OH and HNO₃
(c) HCI and NaOH
(d) CH₃COOH and NaOH
25. Conversion of oxygen to ozone represented by the equation $3O_2 \rightarrow 2O_3$ is an endothermic reaction.
Enthalpy change ΔH accompanying the reaction[**KCET 1988**,
(a) Is negative
(b) Is positive
(c) Is zero
(d) Depends
on temperature
27. The mutual heat of neutralisation of 40 gm of NaOH and 60 gm CH₃COOH will be
[MP PET/PMT 1988]
(a) 56.1 kcal
(b) -67.6 kcal
(c) More than 56.1 kcal
(d) $13.7 kcal$
28. The heat of formations of $CO(g)$ and $CO_2(g)$ are -26.4 kcal and -94.0 kcal respectively. The heat of combustion of carbon monoxide will be
[MP PET/PMT 1988; EAMCET 1993]
(a) +26.4 kcal
(b) -67.6 kcal
(c) -120.6 kcal
(c) -120.6 kcal
(c) -120.6 kcal
(c) +121 kJ / mole
(c) -124 kJ / mole
(c) +121 kJ / mole
(c) -124 kJ / mole
(c) +121 kJ / mole
(c) -242 kJ / mole
(d) +242 kJ / mole
30. Heat of neutralisation of H₄O₄H and H₂I is
11. EAMCET 1980; Rowrke 1990; MP PMT 1994]
(a) 13.7 kcal
(b) >13.7 kcal
(c) <13.7 kcal
(c) <13.7 kcal
(d) None of the above
31. Heat of combustion Af Of CH₄, C₂H₆, C₂H₄ and C₂H₂
gases are - 212.8, - 373.0, - 337.0 and - 310.5 kcal
respectively at the same temperature. The best fuel
among these gases is

2

2

2

2

- (b) $C_2 H_6$ (a) CH_4
- (d) $C_2 H_2$ (c) $C_2 H_4$
- Heat of formation of $CO_2(g)$, $H_2O(l)$ and $CH_4(g)$ are 32. - 94.0, - 68.4 and - 17.9 kcal respectively. The heat of combustion of methane is
 - (b) 136.8 kcal (a) - 212.9 kcal
 - (d) 105.2 kcal (c) – 304.3 kcal

- The heat of reaction does not depend upon 33.
 - (a) Temperature of the reaction
 - (b) Physical state of reactants and products
 - (c) Whether the reaction is carried out at constant pressure or at constant volume
 - (d) The method by which the final products are obtained from the reactants
- Heat of neutralisation of a strong acid by a strong base is 34. a constant value because [KCET 1984]
 - (a) Salt formed does not hydrolyse
 - (b) Only H^+ and OH^- ions react in every case
 - (c) The strong base and strong acid react completely
 - (d) The strong base and strong acid react in aqueous solution
- Heat of neutralisation of an acid by a base is highest when 35. [KCET 1985]
 - (a) Both the acid and base are weak
 - (b) Both the acid and base are strong
 - (c) The acid is strong and the base is weak
 - (d) The acid is weak and the base is strong

The standard molar heat of formation of ethane, CO_2 and water (l) are respectively -21.1, -94.1 and -68.3kcal. The standard molar heat of combustion of ethane will be

[IIT JEE 1986; DPMT 2005]

(a) – 372 <i>kcal</i>	(b) 162 <i>kcal</i>
(c) –240 <i>kcal</i>	(d) 183.5 kcal

- Which of the following reaction is endothermic 37.
 - [AFMC 1988]
 - (a) $CaCO_3 \rightarrow CaO + CO_2$
 - (b) $Fe + S \rightarrow FeS$
 - (c) $NaOH + HCl \rightarrow NaCl + H_2O$
 - (d) $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O_2$
- In the reaction 38.

36. 921

39.

$$CO_2(g) + H_2(g) \rightarrow CO(g) + H_2O(g); \Delta H = 80 \, kJ$$

 ΔH is known as

[CPMT 1987; KCET 1993; Bihar CEE 1995]

- (a) Heat of formation (b) Heat of combustion
- (c) Heat of neutralization (d) Heat of reaction
- Heat of combustion of a substance

[CPMT 1987, 96; AFMC 1992]

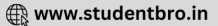
- (a) Is always positive
- (b) Is always negative
- (c) Is equal to heat of formation
- (d) Nothing can be said without reaction

The heat change for the reaction $H_2 + \frac{1}{2}O_2 \rightarrow H_2O$ is 40. called

- (a) Heat of reaction (b) Heat of formation
- (c) Heat of vaporisation (d) None of the above
- 41. The heat change ΔH for the reaction
 - $2CO + O_2 \rightarrow 2CO_2; \Delta H = -135 \ kcal$ is called
 - (a) Heat of formation (b) Heat of reaction
 - (d) Heat of solution (c) Heat of combustion
- The heats of combustion of rhombic and monoclinic 42. sulphur are respectively 70960 and 71030 calories. What

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	will be the heat of conversion of rhombic sulphur to monoclinic			
	[MP PMT/PET 1988]			
	(a) 70960 calories (b) 71030 calories			
	(c) -70 calories (d) $+70$ calories			
43 .	The heat of formation of $H_2O(l)$ is – 68.0 <i>kcal</i> , the heat			
	of formation of $H_2O(g)$ is likely to be			
	(a) -68.0 kcal (b) -67.4 kcal			
	(c) $80.0 \ kcal$ (d) $-58.3 \ kcal$			
44.	Which of the following fuels will have the highest calorific value (kJ/kg) [MP PMT 1990]			
	(a) Charcoal (b) Kerosene			
	(c) Wood (d) Dung			
45.	The heat of combustion of carbon is -94 kcal at 1 atm			
т	pressure. The intrinsic energy of CO_2 is			
	(a) $+94 \ kcal$ (b) $-94 \ kcal$			
	(c) +47 kcal (d) -47 kcal			
46.	The heat of neutralisation of a strong acid and a strong			
alkali is $57.0 kJ mol^{-1}$. The heat released when $0.5 mol$				
	of HNO_3 solution is mixed with 0.2 mole of KOH is			
	[KCET 1991; AIIMS 2002; AMU (Engg.) 2002]			
	(a) $57.0kJ$ (b) $11.4kJ$			
	(c) $28.5 kJ$ (d) $34.9 kJ$			
47.	A solution of 500 ml of 0.2 M KOH and 500 ml of			
	0.2 <i>M HCl</i> is mixed and stirred; the rise in temperature is			
	T_1 . The experiment is repeated using 250 ml each of			
	solution, the temperature raised is T_2 . Which of the			
	following is true[EAMCET 1987; MP PET 1994](1) T T T			
	(a) $T_1 = T_2$ (b) $T_1 = 2T_2$			
	(c) $T_1 = 4T_2$ (d) $T_2 = 9T_1$			
48.	Enthalpy of a compound is equal to its			
	[CMC Vellore 1991]			
	(a) Heat of combustion (b) Heat of formation			
	(c) Heat of reaction (d) Heat of solution			
	(e) Heat of dilution			
49 .	Which is the best definition of "heat of neutralization"			
	[CMC Vellore 1991]			

- (a) The heat set free when one gram molecule of a base is neutralized by one gram molecule of an acid in dilute solution at a stated temperature
- (b) The heat absorbed when one gram molecule of an acid is neutralized by one gram molecule of a base in dilute solution at a stated temperature
- (c) The heat set free or absorbed when one gram atom of an acid is neutralized by one gram atom of a base at a stated temperature
- (d) The heat set free or absorbed when a normal solution containing one gram equivalent of an acid is neutralized by a normal solution containing one gram equivalent of a base at a stated temperature
- (e) The heat set free when one gram equivalent of an acid is neutralized by one gram equivalent of a base in dilute solution at a stated temperature
- **50.** The compound with negative heat of formation are known as [DPMT 1981]

- (a) Endothermic compound
- (b) Exothermic compound
- (c) Heat of formation compound
- (d) None of the above
- **51.** If $H^+ + OH^- \rightarrow H_2O + 13.7 \, kcal$, then the heat of neutralization for complete neutralization of one mole of H_2SO_4 by base will be [MP PMT 1990]
 - (a) 13.7 *kcal* (b) 27.4 *kcal*
 - (c) 6.85 kcal (d) 3.425 kcal
- 52. The lowest value of heat of neutralization is obtained for [KCET 1988; MP PMT 1990]
 - (a) HCl + NaOH
 - (b) $CH_3COOH + NH_4OH$
 - (c) $NH_4OH + HCl$
 - (d) $NaOH + CH_3COOH$
- **53.** Heat of neutralisation for the given reaction $NaOH + HCl \rightarrow NaCl + H_2O$ is $57.1 kJ mol^{-1}$. What will be the heat released when 0.25 mole of NaOH is titrated against 0.25 mole of HCl [CPMT 1990]
 - (a) $22.5 kJ mol^{-1}$ (b) $57.1 kJ mol^{-1}$
 - (c) $14.3 kJ mol^{-1}$ (d) $28.6 kJ mol^{-1}$
- **54.** Reaction, $H_2(g) + I_2(g) \rightarrow 2HI(g) \quad \Delta H = -12.40 \text{ kcal}$. According to this, the heat of formation of *HI* will be [MP PET 1990]

- (c) 6.20 *kcal*(d) 6.20 *kcal*55. All reactions with chemical dissociation are[MP PMT 1990]
 - (a) Reversible
 - (b) Reversible and endothermic
 - (c) Exothermic

(d) Reversible or irriversible and endothermic or exothermic

- **56.** The heat of formation is the change in enthalpy accompanying the formation of a substance from its elements at 298 K and 1 atm pressure. Since the enthalpies of elements are taken to be zero, the heat of formation (ΔH) of compounds[NCERT 1978; KCET 1993]
 - (a) Is always negative
 - (b) Is always positive
 - (c) May be negative or positive
 - (d) Is zero

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- **57.** The heat of combustion of $CH_{4(g)}, C_{(\text{graphite})}$ and $H_{2(g)}$ are 20 *kcal*, -40 *kcal* and -10 *kcal* respectively. The heat of formation of methane is **[EAMCET 1998]**
 - (a) $-4.0 \, kcal$ (b) $+40 \, kcal$
 - (c) $-80 \ kcal$ (d) $+80 \ kcal$
- **58.** If a chemical reaction is accompanied by the evolution of heat, it is [BHU 1979]
 - (a) Catalytic (b) Photochemical
 - (c) Endothermic (d) Exothermic

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59. A reaction that takes place with the absorption of energy is

[EAMCET 1977]

- (a) Burning of a candle (b) Rusting of iron
- (c) Electrolysis of water (d) Digestion of food
- **60.** Which of the following is an endothermic reaction

[EAMCET 1980; MP PMT 1980; IIT JEE 1989; JIPMER 2002]

- (a) $2H_2 + O_2 \rightarrow 2H_2O$
- (b) $N_2 + O_2 \rightarrow 2NO$
- (c) $2NaOH + H_2SO_4 \rightarrow Na_2SO_4 + 2H_2O$
- (d) $3O_2 + C_2H_5OH \rightarrow 2CO_2 + 3H_2O$
- 61. One of the phenomena which cannot be described as combustion is [EAMCET 1979]
 - (a) Oxidation of coal in air
 - (b) Burning of magnesium in nitrogen
 - (c) Reaction of antimony in chlorine
 - (d) Lighting of an electric lamp
- 62. Which of the following statements is correct about heat of combustion [MADT Bihar 1982]
 - (a) It may be exothermic in some cases and endothermic in other cases
 - (b) It is applicable to gaseous substances only
 - (c) It is always an exothermic reaction
 - (d) Its value does not change with temperature

63.
$$H_2 + \frac{1}{2}O_2 \rightarrow H_2O; \Delta H = -68.39 \, kcal$$

$$K + H_2O + \text{Water} \rightarrow KOH(aq) + \frac{1}{2}H_2; \Delta H = -48 \text{ kcal}$$

 $KOH + Water \rightarrow KOH(aq); \Delta H = -14 \ kcal$

The heat of formation of *KOH* is (in *kcal*) [CPMT 1988]

(a) -68.39 + 48 - 14 (b) -68.39 - 48 + 14

(c) 68.39 - 48 + 14 (d) 68.39 + 48 + 14

64. Which of the following reaction can be used to define the heat of formation of $CO_2(g)$

[MP PMT 1989; MH CET 2001]

- (a) $C(\text{graphite}) + O_2(g) = CO_2(g)$
- (b) $CH_4(g) + 2O_2(g) = CO_2(g) + 2H_2O(l)$

(c)
$$CO(g) + \frac{1}{2}O_2(g) = CO_2(g)$$

(d)
$$C_6H_6(l) + 7\frac{1}{2}O_2(g) = 6CO_2 + 3H_2O(l)$$

65. The enthalpy of formation of ammonia is $-46.0 kJ mol^{-1}$. The enthalpy change for the reaction $2NH_3(g) \rightarrow 2N_2(g) + 3H_2(g)$ is

(a) $46.0 \, kJ \, mol^{-1}$ (b) $92.0 \, kJ \, mol^{-1}$

- (c) $-23.0 \ kJ \ mol^{-1}$ (d) $-92.0 \ kJ \ mol^{-1}$
- **66.** The heat of combustion of benzene determined in a bomb calorimeter is $-870 \ kcal \ mol^{-1}$ at $25^{\circ} C$. The value of ΔE for the reaction is

(a) –1740 <i>kcal</i>	(b) –870 <i>kcal</i>
-----------------------	----------------------

- (c) $-872 \, kcal$ (d) $-874 \, kcal$
- **67.** The formation of water from $H_2(g)$ and $O_2(g)$ is an exothermic reaction because [MP PMT/PET 1988]
 - (a) The chemical energy of $H_2(g)$ and $O_2(g)$ is more than that of water
 - (b) The chemical energy of $H_2(g)$ and $O_2(g)$ is less than that of water
 - (c) Not dependent on energy
 - (d) The temperature of $H_2(g)$ and $O_2(g)$ is more than that of water
- **68.** In the reaction for the transition of carbon in the diamond form to carbon in the graphite form, ΔH is -453.5 *cal*. This points out that **[BHU 1981; KCET 1986, 89]**
 - (a) Graphite is chemically different from diamond
 - (b) Graphite is as stable as diamond
 - (c) Graphite is more stable than diamond
 - (d) Diamond is more stable than graphite
- **69.** In the combustion of hydrocarbons, ΔH is **[BHU 1981]** (a) Negative (b) Zero
 - (c) Positive (d) Undeterminate
- **70.** The standard heats of formation in *kcal mol*⁻¹ of $NO_2(g)$ and $N_2O_4(g)$ are 8.0 and 2.0 respectively. The heat of dimerization of NO_2 in *kcal* is

 $2NO_2(g) = N_2O_4(g)$ (a) 10.0 (b) - 6.0 (c) - 12.0 (d) - 14.0

71. The difference between heats of reaction at constant pressure and at constant volume for the reaction $2C_6H_6(l)+15O_2(g) \rightarrow 12CO_2(g)+6H_2O(l)$ at $25^\circ C$ in kJ is

[IIT JEE 1991; Kerala PMT 2004]

[NCERT 1983]

(a) -7.43		(b) $+3.72$
(c) -3.72		(d) + 7.43
~ (]!	1)	

72. C (diamond) $+C_2(g) \rightarrow CO_2(g); \Delta H = -395 kJ$

C (graphite) $+O_2(g) \rightarrow CO_2(g); \Delta H = -393.5 \ kJ$

From the data, the ΔH graphite, is	when diamond is formed from [CBSE PMT 1989; BHU 1987]
(a) $-1.5 kJ$	(b) $+1.5 kJ$
(c) $+3.0 kJ$	(d) $-3.0 kJ$
	walking of boot of formation

- 73. Which of the following values of heat of formation indicates that the product is least stable [MP PMT 1991]
 (a) -94 kcal
 (b) -231.6 kcal
 - (c) +21.4 kcal (d) +64.8 kcal
- **74.** Which of the following equations correctly represents the standard heat of formation (ΔH_f^o) of methane

[IIT JEE (Screening) 1992]

- (a) $C(diamond) + 2H_2(g) = CH_4(g)$
- (b) $C(graphite) + 2H_2(g) = CH_4(l)$
- (c) $C(graphite) + 2H_2(g) = CH_4(g)$

(d)
$$C(graphite) + 4H = CH_4(g)$$

75. In which of the following reactions does the heat change represent the heat of formation of water **[EAMCET 1991]** (a) $2H_2(g) + Q_2(g) \rightarrow 2H_2Q(l): \Delta H = -116 \, kcal$

$$\frac{1}{1}$$

(b)
$$H_2(g) + \frac{1}{2}O_2(g) \to H_2O(l); \Delta H = -58 \ kcal$$

(c) $H^+(aq.) + OH^-(aq.) \rightarrow 2H_2O(l); \Delta H = -13.7 \, kcal$

(d)
$$C_2H_2(g) + 2\frac{1}{2}O_2(g) \rightarrow 2CO_2(g) + H_2O(l);$$

 $\Delta H = -310 \, kcal$

- **76.** Values of heats of formation for SiO_2 and M_gO are -48.4 and -34.7 kJ respectively. The heat of the reaction $2Mg + SiO_2 \rightarrow 2MgO + Si$ is **[KCET 1990]**
 - (a) 21.16 kJ (b) -21.10 kJ

(c)
$$-13.62 kJ$$
 (d) $13.6 kJ$

77. Based on the following thermochemical equations $H_2O(g) + C(s) \rightarrow CO(g) + H_2(g); \Delta H = 131 \, kJ$

$$CO(g) + \frac{1}{2}O_2(g) \rightarrow CO_2(g); \Delta H = -282 \, kJ$$
$$H_2(g) + \frac{1}{2}O_2(g) \rightarrow H_2O(g); \Delta H = -242 \, kJ$$

 $C(s) + O_2(g) \to CO_2(g); \Delta H = X kJ$

 The value of X is
 [CBSE PMT 1992]

 (a) -393 kJ (b) -655 kJ

- (c) +393 kJ (d) +655 kJ
- 78. Heat of transition is the heat evolved or absorbed when a substance is converted from [KCET 1984]
 - (a) Solid to liquid
 - (b) Solid to vapour
 - (c) Liquid to vapour
 - (d) One allotropic form to another allotropic form
- **79.** For the allotropic change represented by equation $C(diamond) \rightarrow C(graphite)$; the enthalpy change is $\Delta H = -1.89 \, kJ$. If $6 \, g$ of diamond and $6 \, g$ of graphite are separately burnt to yield carbon dioxide, the heat liberated in the first case is **[KCET 1988; DPMT 2000]**
 - (a) Less than in the second case by 1.89 kJ
 - (b) More than in the second case by 1.89 kJ
 - (c) Less than in the second case by 11.34 kJ
 - (d) More than in the second case by 0.945 kJ
- **80.** For an exothermic reaction

[MP PET 1994; Manipal MEE 1995]

- (a) H of the products is less than H of the reactants
- (b) H of the products is more than H of the reactants
- (c) H of the products is equal to H of the reactants
- (d) ΔH is always positive
- 81. The energy evolved is highest for which of the following reactions [MP PET 1994]

(a)
$$F + e^- \rightarrow F^{-1}$$
 (b) $Cl + e^- \rightarrow Cl^-$

(c) $S + 2e^- \to S^{2-}$ (d) $O + 2e^- \to O^{2-}$

- 82. In the combustion of 2.0 gm of methane 25 kcal heat is liberated, heat of combustion of methane would be [MP PMT 1994]
 - (a) 100 kcal (b) 200 kcal
 - (c) 300 kcal (d) 400 kcal
- **83.** Complete combustion of CH_4 gives [BHU 1995]

(a)
$$CO_2 + H_2O$$
 (b) $CO_2 + H_2$
(c) $COCI$ (d) $CO + CO_2 + H_2O$

(c)
$$COCl_2$$
 (d) $CO+CO_2+H_2O$

84. If enthalpies of formation of $C_2H_4(g), CO_2(g)$ and $H_2O(l)$ at 25 ° C and 1 *atm* pressure be 52, - 394 and $-286 kJ mol^{-1}$ respectively, the enthalpy of combustion of $C_2H_4(g)$ will be

[CBSE PMT 1995; AIIMS 1998; Pb. PMT 1999]

(a) $+1412 kJ mol^{-1}$ (b) $-1412 kJ mol^{-1}$

(c)
$$+141.2 kJ mol^{-1}$$
 (d) $-141.2 kJ mol^{-1}$

85. Carbon and carbon monoxide burn in oxygen to form carbon dioxide according to the following reactions

$$C + O_2 \rightarrow CO_2; \Delta H = -394 \ kJ \ mol^{-1}$$

 $2CO + O_2 \rightarrow 2CO_2; \Delta H = -569 \text{ kJ mol}^{-1}$

The heat of formation of 1*mol* of carbon monoxide is thus
[MP PET 1995]

(b)
$$-109.5 kJ mol^{-1}$$

(c)
$$-175.0 kJ mol^{-1}$$
 (d) $-87.5 kJ mol^{-1}$

86. Heat of formation in the reaction

(a) $-219.0 kJ mol^{-1}$

- $H_2 + Cl_2 \rightarrow 2HCl + 44 \ kcal$ is [MP PMT 1995] (a) 44 \ kcal (b) 44000 \ kcal
- (c) 22 *kcal* (d) 11 *kcal*
- **87.** When water is added to quick lime, the reaction is
 - [MP PMT 1995] (a) Exothermic (b) Endothermic
 - (c) Explosive (d) None of these
- **88.** In an exothermic reaction ΔH is

(a) Positive

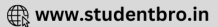
- [MP PMT 1995; CPMT 1999]
- (b) Negative
- (c) Zero (d) Both positive or negative
- **89.** The heat change for the following reaction $C(s)+2S(s) \rightarrow CS_2(l)$ is known as **[MP PMT 1996]**
 - (a) Heat of vaporization (b) Heat of solution
 - (c) Heat of fusion (d) Heat of formation
- **90.** The enthalpy of combustion of benzene from the following data will be

(i) $6C(s) + 3H_2(g) \rightarrow C_6H_6(l); \Delta H = +45.9 kJ$

(ii)
$$H_2(g) + \frac{1}{2}O_2(g) \rightarrow H_2O(l); \Delta H = -285.9 \, kJ$$

(iii)
$$C(s) + O_2(g) \rightarrow CO_2(g); \Delta H = -393.5 \, kJ$$

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	(a) $+ 3172.8 \ kJ$	(b) – 1549.2 <i>kJ</i>		
	(c) $-3172.8 \ kJ$	(d) $-3264.6 kJ$		
91.	Enthalpy of formation of	f <i>HF</i> and <i>HCl</i> are $-161 kJ$ and		
	-92 kJ respectively. Wh	ich of the following statements is		
	incorrect	[KCET 2003]		
	(a) <i>HCl</i> is more stable t	than <i>HF</i>		
	(b) <i>HF</i> and <i>HCl</i> are ex	othermic compounds		
	(c) The affinity of fluor the affinity of chlorir	rine to hydrogen is greater than ne to hydrogen		
	(d) <i>HF</i> is more stable the	nan <i>HCl</i>		
92.	The heat of reaction at co	onstant pressure is given by		
		[MP PMT 1997]		
	(a) $E_P - E_R$	(b) $E_R - E_P$		
	(c) $H_P - H_R$	(d) $H_R - H_P$		
93.	In an endothermic reacti	on the value of ΔH is		
		998; BHU 2000; MP PMT 2000]		
	(a) Zero	(b) Positive		
	(c) Negative	(d) Constant		
94.		yde produced 13.95 kcal of heat		
		on combustion in O_2 . Calculate the heat of combustion of		
	CH ₃ CHO	[Bihar CEE 1995]		
	(a) 279 kcal	(b) 972 kcal		
	(a) 2/9 kcal (c) 27.9 kcal	(d) $\frac{9}{2}$ kcal		
	1			
95.	$C + \frac{1}{2}O_2 \to CO; \Delta H = -42$	2 <i>kJ</i>		
	$CO + \frac{1}{2}O_2 \rightarrow CO_2; \Delta H =$	-24 kJ		
	The heat of formation of	<i>CO</i> ₂ is [CPMT 1996]		
	(a) $-16 kJ$	(b) + 66 kJ		
	(c) $+ 16 kJ$	(d) – 66 <i>kJ</i>		
96.	Standard molar enthalpy	of formation of CO_2 is equal to		
-		[IIT JEE 1997; BHU 2001]		
	(a) Zero			
	(b) The standard mola	ar enthalpy of combustion of		
	gaseous carbon			
	(a) The group of stondard	molor onthelping of formation of		

- (c) The sum of standard molar enthalpies of formation of CO and O_2
- (d) The standard molar enthalpy of combustion of carbon (graphite)
- In the complete combustion of butanol $C_4H_9OH(l)$, if 97. ΔH is enthalpy of combustion and ΔE is the heat of combustion at constant volume, then [EAMCET 1997]
 - (a) $\Delta H < \Delta E$
 - (b) $\Delta H = \Delta E$
 - (c) $\Delta H > \Delta E$
 - (d) $\Delta H, \Delta E$ relation cannot be predicted

 $C + O_2 \rightarrow CO_2; \Delta H = X$ 98.

 $CO + \frac{1}{2}O_2 \rightarrow CO_2; \Delta H = Y$

Then the heat of formation of CO is

		[BHU 1997; DPMT 2002]
(a)	X - Y	(b) $Y - 2X$
(c)	X + Y	(d) $2X - Y$

99. The values of heat of formation of SO_2 and SO_3 are -298.2 kJ and -98.2 kJ. The heat of reaction of the following reaction will be

$SO_2 + \frac{1}{2}O_2 \rightarrow SO_3$	[BHU 1997; CBSE PMT 2000]
(a) $-200 kJ$	(b) $-356.2 kJ$
(c) $+200 kJ$	(d) $-396.2 kJ$

100. Calculate the standard heat of formation of carbon disulphide (l), given that the standard heat of combustion of carbon (s), sulphur (s) and carbon disulphide (l) are

-393.3, -293.72 and $-1108.76 kJ mol^{-1}$ respectively

- -1

.

(a)
$$-128.02 kJ mol^{-1}$$
 (b) $+12.802 kJ mol^{-1}$
(c) $+128.02 kJ mol^{-1}$ (d) $-12.802 kJ mol^{-1}$
101. Fermentation is a reaction called [RPMT 1997]
(a) Endothermic (b) Exothermic
(c) Isotemperature (d) Reversible

102. 8 gm of CH_4 is completely burnt in air. The number of moles of water produced are [Orissa JEE 1997] (a) 0.5 (b) 1 (c) 2 (d) 18

103. For the reaction $A(g) + 2B(g) \rightarrow 2C(g) + 3D(g)$, the value of ΔE at $27^{\circ}C$ is 19.0 kcal. The value of ΔH for the reaction would be $(R = 2.0 cal K^{-1} mol^{-1})$

[Roorkee Qualifying 1998]

(a) 20.8 <i>kcal</i>	(b) 19.8 <i>kcal</i>
(c) 18.8 <i>kcal</i>	(d) 20.2 <i>kcal</i>

104. The enthalpy of solution of $BaCl_2(s)$ and $BaCl_2.2H_2O(s)$ are -20.6 and $8.8 kJ mol^{-1}$ respectively. The enthalpy change for the reaction $BaCl_2(s) + 2H_2O \rightarrow is$

[Roorkee Qualifying 1998]

(a) 29.4 <i>kJ</i>	(b) $-11.8 kJ$
(c) $-20.6 kJ$	(d) $-29.4 kJ$

- **105.** The enthalpy change of a reaction does not depend on [AIIMS 1997]
 - (a) The state of reactants and products
 - (b) Nature of reactants and products
 - (c) Different intermediate reaction
 - (d) Initial and final enthalpy change of a reaction

106.
$$S + \frac{3}{2}O_2 \rightarrow SO_3 + 2x \, kcal$$

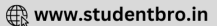
 $SO_2 + \frac{1}{2}O_2 \rightarrow SO_3 + y \, kcal$

Find out the heat of formation of SO_2

[AIIMS 1997; CBSE PMT 1999; KCET 1999]

(a)	2x - y	(b)	2x + y
(c)	x + y	(d)	2x/y

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107	The combustion enthalpies of carbon, hydrogen and	11	
10/1	methane are $-395.5 kJ mol^{-1}$, $-285.8 kJ mol^{-1}$ and $-890.4 kJ mol^{-1}$ respectively at $25^{\circ}C$. The value of		
	standard formation enthalpies of methane at that temperature is [Pb. PMT 1998]		
	(a) $890.4 kJ mol^{-1}$ (b) $-298.8 kJ mol^{-1}$		
	(c) $-74.7 kJ mol^{-1}$ (d) $-107.7 kJ mol^{-1}$	11	
108.	Equal volumes of methanoic acid and sodium hydroxide are mixed. If x is the heat of formation of water, then heat evolved on neutralisation is [BHU 1998]		
	 (a) More than x (b) Equal to x (c) Twice of x (d) Less than x 		
100.	The neutralisation of a strong acid by a strong base		
10 91	liberates an amount of energy per mole of <i>H</i> ⁺ that[BHU 1998 (a) Depends upon which acid and base are involved	8] 11'	
	(b) Depends upon the temperature at which the reaction takes place		
	(c) Depends upon which catalyst is used(d) Is always the same		
110.	When $0.5g$ of sulphur is burnt to $SO_2, 4.6kJ$ of heat is	11	
1101	liberated. What is the enthalpy of formation of sulphur dioxide [KCET 1998; AFMC 2001]	110	
	(a) $+ 147.2 kJ$ (b) $- 147.2 kJ$		
	(c) $-294.4 kJ$ (d) $+294.4 kJ$		
111.	Ozone is prepared by passing silent electric discharge		
	through oxygen. In this reaction[AFMC 1998](a) Energy is given out		
	(b) Energy is absorbed		
	(c) Oxygen is loaded with energy		
	(d) Oxygen is dissociated into atoms		
112.	equation,	11	
	$C_6H_{12}O_6 + 6O_2 \rightarrow 6CO_2 + 6H_2O, \Delta H = -72 kcal$. How		
	much energy will be required for the production of 1.6 g of glucose (Molecular mass of glucose = 180 g) [AFMC 1999]		
	(a) 0.064 kcal (b) 0.64 kcal	12	
	(c) 6.4 kcal (d) 64 kcal		
113.			
	$C(s) + O_2(g) \rightarrow CO_2(g), \Delta H = -394 kJ$		
	$2H_2(g) + O_2(g) \rightarrow 2H_2O(l), \Delta H = 568 kJ$	12	
	$CH_4(g) + 2O_2(g) \rightarrow CO_2(g) + 2H_2O(l) \ \Delta H = -892 \ kJ$		
	Heat of formation of CH_4 is		
	(a) $-70 kJ$ (b) $71.8 kJ$		
	(c) $-244 kJ$ (d) $+782 kJ$		
114.	Calculate the heat of formation of $PCl_5(s)$ from the following data	12	
	$2P(s) + 3Cl_2(g) \rightarrow 2PCl_3(l); \Delta H = -151.8 kcal$		
	$PCl_3(l) + Cl_2(g) \rightarrow PCl_5(s); \Delta H = -32.8 kcal$		
	[JIPMER 1999]		
	(a) $-108.7 kcal$ (b) $+108.7 kcal$	12	
	(c) $-184.6 \ kcal$ (d) $+184.6 \ kcal$		

115.	When $50 cm^3$ of a strong acid is added to $50 cm^3$ of an	
	alkali, the temperature rises by $5^{\circ}C$. If 250 cm ³ of each	
	liquid are mixed, the temperature rise would be	

[KCET 1999]

(a) $5^{\circ}C$ (b) $10^{\circ}C$

(c) $25^{\circ}C$ (d) $20^{\circ}C$

16. $H_2(g) + \frac{1}{2}O_2(g) \to H_2O(l);$

 $\Delta H \text{ at } 298 \ K = -285.8 \ kJ$ The molar enthalpy of vaporisation of water at 1 *atm* and 25 ° C is 44 kJ. The standard enthalpy of formation of 1 *mole* of water vapour at 25 ° C is **[KCET 1999]** (a) -241.8 kJ (b) 241.8 kJ (c) 329.8 kJ (d) -329.8 kJ When 4 g of iron is burnt to ferric oxide at constant

17. When 4 *g* of iron is burnt to ferric oxide at constant pressure, 29.28 *kJ* of heat is evolved. What is the enthalpy of formation of ferric oxide (At. Wt. of Fe = 56)

[AIIMS 1999]

(a) $-81.98 kJ$	(b) – 819.8 <i>kJ</i>
(c) $-40.99 kJ$	(d) + 819.8 kJ

- **118.** When a strong acid, strong base or their salt are dissolved in water, they are completely ionised. If a strong acid is added to a strong base, H^+ ions from the former combine with OH^- ions of the latter forming water. The formation of each water molecule liberates a certain quantity of energy and the reaction is exothermic. The heat liberated when one mole of water is formed by combining hydrochloric acid and sodium hydroxide is 13.7 *kcal*. The heat liberated when one mole of water is formed by combining sulphuric acid and sodium hydroxide is **[AMU (Engg.)** :
 - (a) 25.5 *kcal* (b) 8.5 *kcal*

(c) 13.7 <i>kcal</i> (d) 34	. kcal
-----------------------------	--------

119. 2.1 *g* of *Fe* combines with *S* evolving 3.77 *kJ*. The heat of formation of *FeS* in *kJ/mol* is [AMU (Engg.) 1999]

(a)
$$-1.79$$
 (b) -100.5

(c) -3.77 (d) None of these

120. In the reaction: $H_2 + Cl_2 \rightarrow 2HCl, \Delta H = 194 \ kJ$. Heat of

formation of *HCl* is **[KCET 2000; AFMC 2000; CPMT 2000]**
(a)
$$+ 97 kJ$$
 (b) $+ 194 kJ$
(c) $- 194 kJ$ (d) $- 97 kJ$

- **121.** Enthalpy of neutralisation of acetic acid by *NaOH* is -50.6 kJ/mol. and the heat of neutralisation of a strong
- acid with a strong base is -55.9 kJ/mol. What is the value of ΔH^2 for the ionisation of CH_3COOH [AIIMS 2000]

(a) $+5.3 kJ/mol$	(b) + 6.2 <i>kJ/mol</i>
(c) $+ 8.2 kJ/mol$	(d) + 9.3 kJ/mol

- **122.** Heat of neutralization of the acid-base reaction is 57.32 *kJ* for [JIPMER 2000]
 - (a) $HNO_3 + LiOH$ (b) HCOOH + KOH

(c) $HCl + NH_4OH$ (d) $CH_3COOH + NaOH$

123. The heat change for the following reaction at 298 o K and at constant pressure is +7.3 *kcal*

 $A_2B(s) \rightarrow 2A(s) + 1/2B_2(g), \Delta H = +7.3 kcal$

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	[DCE 2000]
(a) 7.3 <i>kcal</i>	(b) More than 7.3

(c) Zero (d) None of these

The heat change at constant volume would be

The heat evolved in the combustion of benzene is given by 124. the equation

$$C_6H_6(l) + 7\frac{1}{2}O_2(g) \rightarrow 3H_2O(l) + 6CO_2(g)$$

 $\Delta H = 781.0 \, kcal \, mol^{-1}$

Which of the following quantities of heat energy will be evolved when 39 g of benzene is burnt in an open container

[MP PET 2000]

(a) 122.25 kcal mol^{-1} (b) 390.5 kcal mol^{-1}

(c)
$$1562.0 \ kcal \ mol^{-1}$$
 (d) $586.75 \ kcal \ mol^{-1}$

125.
$$H_2(g) + Cl_2(g) \rightarrow 2HCl(g).\Delta H = -44kcal$$

 $2Na(s) + 2HCl(g) \rightarrow 2NaCl(s) + H_2(g), \Delta H = -152 \ kcal$

For the reaction $Na(s) + \frac{1}{2}Cl_2(g) \rightarrow NaCl(s), \Delta H =$

[KCET 2000]

(a) $-108 \ kcal$	(b) – 196 <i>kcal</i>
(c) – 98 <i>kcal</i>	(d) 54 kcal

126. The ΔH_f^o for $CO_2(g), CO(g)$ and $H_2O(g)$ are - 393.5, -110.5 and - 241.8kJmol⁻¹ respectively. The standard enthalpy change (in kJ) for the reaction $CO_2(g) + H_2(g) \rightarrow$ $CO(g) + H_2O(g)$ is [IIT JEE (Screening) 2000]

(a)	524.1	(b) 41.2	
(c)	- 262.5	(d) -41.2	

127. Which of the following compounds will absorb the maximum quantity of heat when dissolved in the same amount of water ? The heats of solution of these compounds at $25^{\circ} C$ in kJ/mole of each solute is given in brackets [AMU (Engg.) 2000]

(b) $KCl(\Delta H = +17.64)$ (a) $HNO_{3}(\Delta H = -33)$

(c) $NH_4 NO_3 (\Delta H = +25.5)$ (d) $HCl(\Delta H = -74.1)$

128. In the reaction $C + 2S \rightarrow CS_2 + \Delta H, \Delta H$ is the

		[AMU (Engg.) 2000]
(a)	Heat of combustion	(b) Heat of neutralisation
(c)	Heat of solution	(d) None of these

- 129. The heat of formation of methane $C(s) + 2H_2(g) \rightarrow CH_4(g)$ at constant pressure is 18500 cal at 25°C. The heat of reaction at constant volume would be [MH CET 2000] (a) 17904 cal (b) 18202 cal
 - (c) 18798 cal (d) 19096 cal
- **130.** The enthalpy of combustion of $C_6 H_{6(l)}$ is -3250 kJ.When 0.39 q of benzene is burnt excess of oxygen in an open vessel, the amount of heat evolved is

[KCET 2000; AFMC 2000; DCE 2000] (b) 16.25 kJ

(a) 16.25 J (c) 32.5 J (d) 32.5 kJ

131.	$C(s) + O_2(g) \rightarrow CO_2; \Delta H = -94 \ kcal$		
	$CO(g) + \frac{1}{2}O_2(g) \rightarrow CO_2; \ \Delta H = 135.2 \ kcal$		
	Then heat of formation of $CO(g)$ is [DCE 2001]		
	(a) – 26.4 <i>kcal</i>	(b) 41.2 <i>kcal</i>	
	(c) 26.4 <i>kcal</i>	(d) – 229.2 <i>kcal</i>	
132.	• The heat of neutralization of <i>HCl</i> and <i>NaOH</i> is		
		[MP PET 2001]	
	(a) Zero	(b) $-57.3 kJ$	
	(c) $+ 57.3 kJ$	(d) None of these	

133. The following thermochemical reactions are given

$$M + \frac{1}{2}O_2 \rightarrow MO + 351.4 \, kJ$$
$$X + \frac{1}{2}O_2 \rightarrow XO + 90.8 \, kJ$$

It follows that the heat of reaction for the following process $M + XO \Rightarrow MO + X$ is given by

(a)
$$422.2 \, kJ$$
 (b) $268.7 \, kJ$
(c) $-442.2 \, kJ$ (d) $260.6 \, kJ$

134. If the heat of combustion of carbon monoxide at constant volume and at $17^{\circ}C$ is - 283.3 kJ, then its heat of combustion at constant pressure $(R = 8.314 \ J \text{ degree}^{-1} mol^{-1})$ [CPMT 2001] (a) -284.5 kJ(b) 284.5 kJ (c) 384.5 kJ (d) $-384.5 \, kJ$

135. Heat of formation of H_2O is -188 kJ/mole and H_2O_2 is -286 kJ/mole. The enthalpy change for the reaction $2H_2O_2 \rightarrow 2H_2O + O_2$ is [BHU 2001] (a) 196 kJ (b) -196 kJ

(c)
$$984 \, kJ$$
 (d) $-984 \, kJ$

136. The values of ΔH for the combustion of ethene and ethyne are - 341.1 and - 310.0kcal respectively. Which of the following is a better fuel [BHU 2001] (a) C_2H_2 (b) $C_2 H_4$

(c) Both of these (d) None of these

137. For exothermic reaction, the equilibrium constant

- (a) Increases with increase of P
- (b) Decreases with increase of P
- (c) Increases with increase of temperature
- (d) Decreases with increase of temperature
- **138.** In order to decompose 9 *q* water 142.5 *kJ* heat is required. Hence the enthalpy of formation of water is [KCET 2001] (a) -142.5 kJ(b) + 142.5 kJ

(c)
$$-285 kJ$$
 (d) $+285 kJ$

139. What is Δn for combustion of 1 mole of benzene, when both the reactants and the products are gas at 298 K [Pb. PMT 2001]

(a) 0 (b)
$$3/2$$

(c) $-3/2$ (d) $1/2$

140. If
$$C + O_2 \rightarrow CO_2 + 94.2 \, kcal$$

$$H_2 + \frac{1}{2}O_2 \rightarrow H_2O + 68.3 \, kcal$$

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 $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O + 210.8 kcal$

(a) 47.3 *kcal* (b) 20.0 *kcal*

(c) $45.9 \ kcal$ (d) $-47.3 \ kcal$

141. Heat of neutralization of strong acid and weak base is **[UPSEAT 2001]**

(a)
$$57.1 \, kJ \, mol^{-1}$$

- (b) $13.7 kJ mol^{-1}$
- (c) Less than $13.7 k cal mol^{-1}$
- (d) More than $13.7 k cal mol^{-1}$
- **142.** A system is changed from state *A* to state *B* by one path and from *B* to *A* another path. If E_1 and E_2 are the corresponding changes in internal energy, then

[Pb. PMT 2001] (a) $E_1 + E_2 = -ve$ (b) $E_1 + E_2 = +ve$

(c) $E_1 + E_2 = 0$ (d) None of these

143. The heat evolved during the combination of 24 *g C* and 128 *g S* following the change is

$C + S_2 \rightarrow CS_2; \Delta H = 22.0 kcal$		[MH CET 2001]
(a) 11 kcal	(b) $22 kcal$	

(a)	11 KCul	(D)	32 KCUI
(c)	44 kcal	(d)	22 kcal

144. When the aqueous solution of 0.5 mole HNO_3 is mixed

with the 0.3 mole of OH^- solution, then what will be the liberated heat (Enthalpy of neutralization is = 57.1 *kJ*)

[Kerala CET 2005]

(a)	28.5 kJ	(b)	17.1 kJ
(a)	$A = = I_{A}T$	(4)	1 = 1.T

(c)	45.7 <i>КЈ</i>	(d) $1.7 kJ$

(e) 2.85 kJ
145. A cylinder of gas is assumed to contain 11.2 kg of butane (C₄H₁₀). If a normal family needs 20000 kJ of energy per day. The cylinder will last: (Given that △H for combustion of butane is - 2658 kJ) [AMU (Engg.) 2002]

(a)	20 days	(h)	25 days
(u)	20 au y 5	(0)	2.) uu yo

(c)	26 days	(d)	24 days

- **146.** Compounds with high heat of formation are less stable because

 [KCET 2002]
 - (a) High temperature is required to synthesise them
 - (b) Molecules of such compounds are distorted
 - (c) It is difficult to synthesis them
 - (d) Energy rich state leads to instability
- **147.** The heat evolved in the combustion of methane is given by the following equations:

$$CH_4(g) + 2O_2(g) \rightarrow CO_2(g) + H_2O(l); \Delta H = -890.3 kJ$$

How many grams of methane would be required to produce 445.15 kJ of heat of combustion

(a)	4 <i>g</i>	(b) 8 <i>g</i>
(c)	12 g	(d) 16 g

148. Heats of combustion (ΔH°) for C(s), $H_2(g)$ and $CH_4(g)$ are -94, -68 and $-213 \ kcal/mol$. respectively. The value

of ΔH^{o} for the reaction,

$C(s) + 2H_2(g) \rightarrow CH_4(g)$ is	[CBSE PMT 2002]
(a) - 85 <i>kcal</i>	(b) – 111 <i>kcal</i>
(c) - 17 <i>kcal</i>	(d) – 170 <i>kcal</i>

- **149.** If the value of ΔH in a reaction is positive, then the reaction is called [BHU 2002]
 - (a) Exothermic (b) Endothermic
 - (c) Polymorphic (d) Polytropic
- **150.** Enthalpy of neutralisation of *NH*₄*OH* and *HCl*, is numerically
 - [JIPMER 2002; Kurukshetra CEE 2002]
 - (a) $57.1 \ kJ \ mol^{-1}$ (b) $< 57.1 \ kJ \ mol^{-1}$
 - (c) $> 57.1 \ kJ \ mol^{-1}$ (d) Zero
- 151. The heat of neutralisation will be highest in [MP PMT 2002](a) NH₄OH and CH₃COOH
 - (b) NH_4OH and HCl
 - (c) *KOH* and CH_3COOH
 - (d) KOH and HCl
- **152.** If a mole of H_2 molecule is heated to high temperature the following reaction takes place [Kerala (Med.) 2002]
 - (a) $H_2(g) + 436 kJ = H(g) + H(g)$
 - (b) $H_2(g) + 820 kJ = 2H_2(g)$
 - (c) $2H_2(g) + 436 J = 2H_2$
 - (d) $H_2 + H_2 = H^+ + H^+$
- **153.** Which of the following reactions is not exothermic [MP PET 2002]
 - (a) $C(s) + O_2(g) \rightarrow CO_2(g)$
 - (b) $C(s) + 2S(s) \rightarrow CS_2(g)$
 - (c) $CH_4(g) + 2O_2(g) \rightarrow CO_2(g) + 2H_2O(l)$
 - (d) $CO(g) + \frac{1}{2}O_2(g) \to CO_2(g)$
- 154. On combustion, carbon forms two oxides CO and CO₂, Heat of formation of CO₂ gas is 94.3 kcal and that of CO is 26.0 kcal. Heat of combustion of carbon is [JIPMER 2002]
 - (a) $26.0 \ kcal$ (b) $-94.3 \ kcal$
 - (c) 68.3 *kcal* (d) 120.3 *kcal*
- 155. 1 mole of conc. HCl requires X moles of dilute NaOH for neutralisation and 1 mole of concentrate H₂SO₄ requires Y moles of small dilute NaOH then which of the following reaction is true [MH CET 2002]

(a)
$$Y = \frac{1}{2}X$$
 (b) $X = \frac{1}{2}Y$

(c) X = 2Y (d) None of these

156. Which of the reaction defines ΔH_f^o

[IIT -JEE (Screening) 2003]

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$$[\mathbf{A}(\mathbf{h}) \mathbf{2}(\mathbf{0})] + O_2(g) \rightarrow CO_2(g)$$

- (b) $\frac{1}{2}H_2(g) + \frac{1}{2}F_2(g) \rightarrow HF(g)$
- (c) $N_2(g) + 3H_2(g) \rightarrow 2NH_3(g)$

(d)
$$CO(g) + \frac{1}{2}O_2(g) \rightarrow CO_2(g)$$

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157. One gram sample of NH_4NO_3 is decomposed in a bomb calorimeter. The temperature of the calorimeter increases

by 6.12 *K* the heat capacity of the system is 1.23 kJ/g/deg. What is the molar heat of decomposition for NH_4NO_3

[AIIMS 2003]

(a) − 7.53 <i>kJ/mol</i>	(b) – 398.1 <i>kJ/mol</i>
(c) $-16.1 kJ/mol$	(d) – 602 <i>kJ/mol</i>

158. For which one of the following equations is $\Delta H_{\text{react}}^{o}$ equal

[CBSE PMT 2003]

(a) $2CO(g) + O_2(g) \rightarrow 2CO_2(g)$

to ΔH_f^o for the product

- (b) $N_2(g) + O_3(g) \rightarrow N_2O_3(g)$
- (c) $CH_4(g) + 2Cl_2(g) \rightarrow CH_2Cl_2(l) + 2HCl(g)$
- (d) $Xe(g) + 2F_2(g) \rightarrow XeF_4(g)$

159. Enthalpy change for a reaction does not depend upon

- [AIEEE 2003]
- (a) The physical states of reactants and products
- (b) Use of different reactants for the same product
- (c) The nature of intermediate reaction steps
- (d) The differences in initial or final temperatures of involved substances

160. $C_{\text{dia}} + O_2 \to CO_2; \ \Delta H = -395.3 \, kJ \,/\, mole$

	f = -393.4 kJ / mole	$C_{gr} + O_2 \rightarrow CO_2; \Delta H$
[BHU 2003]		$C_{gr} \rightarrow C_{dia}; \Delta H =$
	(b) – 1.9	(a) – 3.8
	(d) + 1.9	(c) $+3.8$

161. What is the weight of oxygen that is required for the complete combustion of 2.8 kg of ethylene? [BVP 2003]
(a) 0.6 kg
(b) 06.0 kg

(a)	9.6 <i>KY</i>	(b) $96.0 kg$
<pre></pre>		(1) - 1

- (c) 6.4 kg (d) 2.8 kg
- **162.** For complete neutralization of *HCl* with *NaOH*, the heat of neutralization is

 [MP PET 2003]

(a)
$$+13.70 kJ mol^{-1}$$
 (b) $-13.70 kJ mol^{-1}$

(c)
$$-57.32 kJ mol^{-1}$$
 (d) $+57.32 kJ mol^{-1}$

163. The heat of combustion of carbon to CO_2 is $-393.5 \, kJ / mol$. The heat released upon formation of 35.2 g of CO_2 from carbon and oxygen gas is **[UPSEAT 2004]**

(a)
$$+315 kJ$$
 (b) $-31.5 kJ$
(c) $215 kI$ (d) $+215 kI$

(c)
$$-315 kJ$$
 (d) $+31.5 kJ$

164. Which of the following equations correctly represents the standard heat of formation (ΔH_f^o) of methane

[UPSEAT 2004]

(a)	C (diamond)	$+4H_{(g)} \rightarrow$	$CH_{4(g)}$
• •	- ((8)	+(8)

- (b) C (diamond) $+2H_{2(g)} \rightarrow CH_{4(g)}$
- (c) C (graphite) $+2H_{2(g)} \rightarrow CH_{4(g)}$
- (d) C (graphite) $+4H_{(g)} \rightarrow CH_{4(g)}$

165. If the heat of formation of CO_2 is -393 kJ. The amount of heat evolved in the formation of 0.156 kg of CO_2 is

[MH CET 2004]

(a)	–1357.9 kJ	(b)	-1275.9 kJ

- (c) -1572.0 kJ (d) -1165.5 kJ
- **166.** Which of the following pairs has heat of neutralisation
equal to 13.7 Kcals[DCE 2003]

(a)
$$HCl, NH_4OH$$
 (b) HNO_3, KOH

- (c) $NaOH, CH_3COOH$ (d) H_2SO_4, NH_4OH
- **167.** The enthalpies of combustion of carbon and carbon monodie are –393.5 and –283 *kJ mol*⁻¹ respectivley. The enthalpy of formation of carbon monoxide per mole is

[AIEEE 2004]

(a)	–676.5 kJ	(b)	676.5 kJ
(c)	110.5 <i>kJ</i>	(d)	-110.5 kJ

- **168.** The enthalpy of combustion of methane at $25^{\circ}C$ is 890 kJ. The heat liberated when 3.2 g of methane is burnt in air is [KCET 2004] (a) 445 kJ (b) 278 kJ
 - (c) $-890 \ kJ$ (d) $178 \ kJ$
- **169.** If (i) $C+O_2 \rightarrow CO_2$, (ii) $C+1/2O_2 \rightarrow CO$, (iii) $CO+1/2O_2 \rightarrow CO_2$, the heats of reaction are Q, -12, -10respectively. Then Q = [Orissa JEE 2004] (a) -2 (b) 2(c) -22 (d) -16
- **170.** How much energy is released when 6 mole of octane is burnt in air ? Given ΔH_f^o for $CO_2(g), H_2O(g)$ and $C_8H_{18}(l)$ respectively are -490, -240 and +160 kJ/mol

[AIIMS 2004]

(a)	– 6.2 <i>kJ</i>	(b) – 37.4 <i>kJ</i>
(c)	-35.5 kJ	(d) $-20.0 kJ$

171. Given : $2Fe + \frac{3}{2}O_2 \rightarrow Fe_2O_3$, $\Delta H = -193.4 kJ$;

$$Mg + \frac{1}{2}O_2 \rightarrow MgO, \Delta H = -140.2 \, kJ$$

What is the ΔH of the reaction

$Mg + Fe_2O_3 \rightarrow 3MgO + 2Fe$	[Orissa JEE 2005]
(a) $-1802 kJ$	(b) +1802 <i>kJ</i>

(c) ·	– 800 kJ		(0	d) +	228	kJ
m 1.		.1	(C	.1.	

172. The enthalpy change (ΔH) for the neutralisation of 1M HCl by caustic potash in dilute solution at 298 K is

[DPMT 2005]

(a) 68 <i>kJ</i>	(b) 65 <i>kJ</i>
(c) $57.3 kJ$	(d) 50 <i>kJ</i>

Bond energy

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	10		
1.	If the bond dissociation energies of XY , X_2 and Y_2 (all diatomic molecules) are in the ratio of $1:1:0.5$ and $\Delta_f H$	10.	If the bond energies of $H - H$, $Br - Br$ and HBr are 433, 192 and 364 $kJ mol^{-1}$ respectively, the ΔH^o for the
	for the formation of XY is $-200 kJ mole^{-1}$. The bond		reaction, $H_2(g) + Br_2(g) \rightarrow 2HBr(g)$ is [CBSE PMT 2004]
	dissociation energy of X_2 will be [AIEEE 2005]		(a) $+ 261 kJ$ (b) $- 103 kJ$
	(a) $100 \ kJ \ mol^{-1}$ (b) $800 \ kJ \ mol^{-1}$		(c) $-261 kJ$ (d) $+103 kJ$
	(c) $300 \ kJ \ mol^{-1}$ (d) $400 \ kJ \ mol^{-1}$		Free energy and Work function
2.	Energy required to dissociate 4 gm of gaseous hydrogen		The free energy shange for a reversible reaction of
	into free gaseous atoms is 208 kcal at $25^{\circ}C$. The bond energy of $H - H$ bond will be	1.	The free energy change for a reversible reaction at equilibrium is [NCERT 1984; Kurukshetra CEE 1998; AMU 1999]
	[CPMT 1989; MP PET 2000; J & K 2005]		(a) Large positive (b) Small negative
	(a) 104 kcal (b) 10.4 kcal		(c) Small positive (d) o
3.	(c) 1040 <i>kcal</i> (d) 104 <i>cal</i> The bond dissociation energy needed to form benzyl	2.	For a spontaneous change, free energy change ∆G is [MNR 1983; BHU 1981, 95; AMU 1999;
	radical from toluene is than the formation of the methyl radical from methane [IIT 1994]		DCE 2000, 01; BHU 2000; MP PMT 2003] (a) Positive
	(a) Less (b) Much		(b) Negative
	(c) Equal (d) None of the above		(c) Zero
4.	Which one of the following bonds has the highest average		(d) Can be positive or negative
	bond energy (<i>kcal/mol</i>) [CPMT 1994]	3.	A minus sign of the free energy change denotes that
	(a) $S = O$ (b) $C \equiv C$		(a) The reaction tends to proceed spontaneously
	(c) $C \equiv N$ (d) $N \equiv N$		(b) The reaction is non-spontaneous
5.	The bond dissociation energies of gaseous H_2, Cl_2 and		(c) The system is in equilibrium(d) The reaction is very much unlikely
	<i>HCl</i> are 104, 58 and 103 <i>kcal</i> respectively. The enthalpy	4.	The relation between ΔG and ΔH is
	of formation of <i>HCl</i> gas would be	4.	[MP PMT 1994, 95, 97; AFMC 1997;
	[MP PET 1997; MP PMT 1999, 2001]		Kurukshetra CEE 1998]
	(a) $-44 kcal$ (b) $44 kcal$		(a) $\Delta H = \Delta G - T \Delta S$ (b) $\Delta G = \Delta H - T \Delta S$
	(c) $-22 kcal$ (d) $22 kcal$		(c) $T \Delta S - \Delta G = \Delta H$ (d) $\Delta H = T \Delta G + \Delta S$
6.	The first ionization energy for <i>Li</i> is 5.4 <i>eV</i> and electron affinity of <i>Cl</i> is 3.61 <i>eV</i> . The ΔH (in <i>kJ/mol</i>) for the reaction $Li(g) + Cl(g) \rightarrow Li^+ + Cl^-$ is (if resulting ions do not combine	5.	At 300 <i>K</i> , the reactions which have following values of thermodynamic parameters occur spontaneously
	with each other) $(1 eV = 1.6 \times 10^{-19} J)$ [MP PMT 2000, 03]		[Roorkee 1999]
			(a) $\Delta G^{\circ} = -400 \ kJ \ mol^{-1}$
	(a) 70 (b) 100 (c) 170 (d) 270		(b) $\Delta H^{\circ} = 200 \ kJ \ mol^{-1}, \ \Delta S^{\circ} = -4 \ JK^{-1} mol^{-1}$
-	(c) 170 (d) 270 Given that $C(g) + 4H(g) \rightarrow CH_4(g); \Delta H = -166 kJ$		
/•	The bond energy $C - H$ will be [AMU 2002]		(c) $\Delta H^{o} = -200 \ kJ \ mol^{-1}, \ \Delta S^{o} = 4 \ JK^{-1} \ mol^{-1}$
	(a) $208 kJ/mole$ [AMO 2002]		(d) $\Delta H^{\circ} = 200 \ J \ mol^{-1}, \ \Delta S^{\circ} = 40 \ JK^{-1} \ mol^{-1}$
	(b) $-41.6 \ kJ/mole$	6.	The relation $\Delta G = \Delta H - T \Delta S$ was given by
	(c) $832 kJ/mole$		[MP PMT 2000; KCET 2002]
	(d) None of these		(a) Boltzmann (b) Faraday
8.	The $H-H$ bond energy is 430 kJ mol and $Cl-Cl$		(c) Gibbs–Helmholtz (d) Thomson
	bond energy is $240 \ kJ \ mol^{-1}$. ΔH for HCl is $-90 \ kJ$. The	7.	For precipitation reaction of Ag^+ ions with <i>NaCl</i> , which
	H - Cl bond energy is about [BVP 2003]	/•	of the following statements is correct [CPMT 1988]
	(a) $180 kJ mol^{-1}$ (b) $360 kJ mol^{-1}$		(a) ΔH for the reaction is zero
	(c) $213 kJ mol^{-1}$ (d) $425 kJ mol^{-1}$		(b) ΔG for the reaction is zero
_			(c) ΔG for the reaction is negative
9.	If enthalpies of methane and ethane are respectively 320 and 360 calories then the bond energy of $C - C$ bond is		(d) $[\Delta G] = [\Delta H]$
	[UPSEAT 2003]	8.	At constant pressure and temperature, the direction to the
	(a) 80 calories (b) 40 calories		result of any chemical reaction is where, there is less amount of
	(c) 60 calories (d) 120 calories		

> (c) Gibb's free energy (d) None of the above

(b) Enthalpy

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(a) Entropy

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9. The dependence of Gibbs free energy on pressure for an isothermal process of an ideal gas is given by

[MP PET 1996]

(a)
$$\Delta G_T = nRT \ln \frac{P_2}{P_1}$$
 (b) $\Delta G_T = nRT \ln \frac{V_2}{V_1}$
(c) $\Delta G_T = nRT \log \frac{P_1}{P_2}$ (d) $\Delta G_T = nRT \log \frac{V_2}{V_1}$

- **10.** For the change $H_2O(l) \rightarrow H_2O(g); P = 1 atm$, T = 373 K, the free energy change $\Delta G = 0$. This indicates that
 - (a) $H_2O(l)$ is in equilibrium with $H_2O(g)$
 - (b) Water boils spontaneously at 373 K
 - (c) Water does not boil spontaneously at 373 K
 - (d) Condensation of water vapour occurs spontaneously at 373 *K*
- **11.** What is the free energy change ΔG when 1.0 mole of water at $100^{\circ}C$ and 1 atm pressure is converted into steam at $100^{\circ}C$ and 1 atm pressure [MP PET/PMT 1998]

	(a)	540 cal	(b) –9800 <i>cal</i>
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- (c) $9800 \ cal$ (d) $0 \ cal$
- **12.** ΔG° for the reaction $X + Y \Rightarrow Z$ is -4.606 kcal. The value of equilibrium constant of the reaction at 227 ° C is $(R = 2.0 \text{ cal.mol}^{-1} \text{K}^{-1})$ [Roorkee 1999]
 - (a) 100 (b) 10

- **13.** The standard enthalpy of the decomposition of N_2O_4 to NO_2 is 58.04 kJ and standard entropy of this reaction is 176.7 J/K. The standard free energy change for this reaction at 25 ° C is [AIIMS 1999] (a) 539 kJ (b) -539 kJ (c) -5.39 kJ (d) 5.39 kJ
- 14. Spontaneity of a chemical reaction is decided by the negative change in

 [MP PET 2001]
 - (a) Internal energy(b) Enthalpy(c) Entropy(d) Free energy
- **15.** For a reaction at $25 \degree C$ enthalpy change and entropy changes are $-11.7 \times 10^3 J mol^{-1}$ and $-105 J mol^{-1}K^{-1}$ respectively. What is the Gibbs free energy **[BHU 2001]** (a) 15.05 kJ (b) 19.59 kJ

(c) $2.55 kJ$ (d) :	22.55 kJ
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- 16.Born-Haber cycle is used to determine[UPSEAT 2001](a)Crystal energy(b)Electron affinity
 - (c) Lattice energy (d) All of these
- 17. Gibbs free energy *G*, enthalpy *H* and entropy *S* are interrelated as in [MP PMT 2002]
 (a) G = H + TS (b) G = H TS
 - (c) G TS = H (d) G = S = H
- **18.** The essential condition for the feasibility of a reaction is that

[JIPMER 2002]

(a) The reaction should be exothermic

- (b) The entropy of products must be larger than that of reactants
- (c) The reaction is to be accompanied with free energy decrease
- (d) The reaction has to possess high activation energy
- **19.** The correct relationship between free energy change in a reaction and the corresponding equilibrium constant K_c is

(a)
$$\Delta G = RT \ln K_c$$
 (b) $-\Delta G = RT \ln K_c$

(c)
$$\Delta G^o = RT \ln K_c$$
 (d) $-\Delta G^o = RT \ln K_c$

20. In an irreversible process taking place at constant T and P and in which only pressure-volume work is being done, the change in Gibbs free energy (dG) and change in entropy (dS), satisfy the criteria **[AIEEE 2003]**

(a)
$$(dS)_{V,E} < 0, (dG)_{T,P} < 0$$

- (b) $(dS)_{VF} > 0, (dG)_{TP} < 0$
- (c) $(dS)_{V,E} = 0, (dG)_{T,P} = 0$
- (d) $(dS)_{V,E} = 0, (dG)_{T,P} > 0$
- **21.** The densities of graphite and diamond at 298 *K* are 2.25 and 3.31 *g cm*⁻³, respectively. If the standard free energy difference (ΔG^{o}) is equal to 1895 *J* mol^{-1} , the pressure at which graphite will be transformed diamond at 298 *K* is

[CBSE PMT 2003]

(a) $9.92 \times 10^5 Pa$	(b) $9.92 \times 10^8 Pa$
(c) $9.92 \times 10^7 Pa$	(d) $9.92 \times 10^6 Pa$

22. The free energy change for the following reactions are given below,

$$C_2H_2(g) + \frac{5}{2}O_2(g) \rightarrow 2CO_2(g) + H_2O(l); \Delta G^o = -1234 \ kJ$$

 $C(s) + O_2(g) \rightarrow CO_2(g) \Delta G^o = -394 \ kJ$

$$H_2(g) + \frac{1}{2}O_2(g) \rightarrow H_2O(l) \ \Delta G^o = -237 \, kJ$$

What is the standard free energy change for the reaction $H_2(g) + 2C(s) \rightarrow C_2H_2(g)$ [Kerala (Med.) 2002]

(a) -209 kJ (b) -2259 kJ

(c)
$$+ 2259 kJ$$
 (d) $209 kJ$

23. The equilibrium concentration of the species in the reaction $A + B \equiv C + D$ are 3, 5, 10 and 15 mole L^{-1} respectively at 300 K the ΔG for the reaction is **[Pb. PMT 2004]**

- **24.** Gibb's free enrgy (*G*) is defined as **[Pb. CET 2001]**
 - (a) $\Delta G = \Delta H T\Delta S$ (b) $\Delta G = \Delta H + \frac{T}{\Delta S}$ (c) $\Delta H = \Delta G - T\Delta S$ (d) $\Delta G = \Delta H + T.C_p$
- **25.** Standard enthalpy and standard entropy changes for the oxidation of ammonia at 298 K are $-382.64 \text{ kJ mol}^{-1}$



				(a) More than $10 = \frac{1}{10}$ (b) $10 = \frac{1}{10}$	
		espectively. Standard Gibbs	F	(c) More than 13.7 <i>kcal</i>(d) 108 kcalWhen a gas undergoes adiabatic expansion, it gets coordinates the set of the set	مامط
	energy change for the same		5.	due to [DCE 20	
		[CBSE PMT 2004]		(a) Loos of kinetic energy	
	(a) -439.3 kJ mol^{-1}	(b) $-523.2 \ kJ \ mol^{-1}$		(b) Fall in temperature	
	(c) $-221.1 \ kJ \ mol^{-1}$	(d) $-339.3 \ kJ \ mol^{-1}$		(c) Decrease in velocity	
26.	For spontaneity of a cell, wh	lich is correct		(d) Energy used in doing work	
		[Orissa JEE 2004]	6.	The heat content of a system is called	
	(a) $\Delta G = 0, \Delta E = 0$			(a) Internal energy (b) Entropy	
	(b) $\Delta G = -ve, \Delta E = 0$		-	(c) Free energy(d) EnthalpyTo calculate the amount of work done in joules during	rina
	(c) $\Delta G = +ve, \Delta E = +ve$		7•	reversible isothermal expansion of an ideal gas,	
	(d) $\Delta G = -ve, \Delta E = +ve$			volume must be expressed in [KCET (Med.) 20	
	, ,			(a) m^3 only (b) dm^3 only	
27.		ction having $\Delta H = 31400 \ ca;$.		(c) cm^3 only (d) Any one of them	
		00°C is [Orissa JEE 2005]	8.	In an isobaric process, the ratio of heat supplied to	
	(a) -9336 cal	(b) -7386 cal		system (dQ) and work done by the system (dW)	
	(c) -1936 cal	(d) + 9336 cal		diatomic gas is [AFMC 20	02]
28.		reaction at one atmospheric		(a) 1:1 (b) 7:2 (c) 7:5 (d) 5:7	
		and $0.066 \ kJk^{-1}$ respectively. the free energy change will be	9.	(c) 7:5 (d) 5:7 The enthalpy change for the reaction of 50.00 ml	l of
		berature the nature of reaction	9.	ethylene with 50.00 ml of H_2 at 1.5 atm pressure	
	would be	[Kerala CET 2005]		$\Delta H = -0.31 kJ$. The value of ΔE will be [DPMT 20	
	(a) 483 <i>K</i> , spontaneous			(a) $-0.3024 kJ$ (b) $0.3024 kJ$	
	(b) $443 K$, non-spontaneou	S		(c) $2.567 kJ$ (d) $-0.0076 kJ$	
	(c) 443 <i>K</i>, spontaneous(d) 463 <i>K</i>, non-spontaneou	5	10.	Enthalpy of solution of NaOH (solid) in water	is is
	(e) $463 K$, spontaneous			$-41.6 kJ mol^{-1}$. When <i>NaOH</i> is dissolved in water,	
	(c) 400 R, spontaneous			temperature of water [UPSEAT 20	04]
	Critica	l Thinking		(a) Increase (b) Decreases	
		l Thinking		(c) Does not change (d) Fluctuates indefinited	y
		Objective Ouestiens	11.	In which of the following entropy decreases ?	-
	(Objective Questions		[CPMT 1988, 94; MP PMT 20 (a) Crystallization of sucrose from solution	00]
1.	Adsorption of gases on	solid surface is generally		(b) Rusting of iron	
1.	exothermic because	[IIT JEE (Screening) 2004]		(c) Melting of ice	
	(a) Enthalpy is positive	(b) Entropy decreases		(d) Vaporization of camphor	
	(c) Entropy increases	(d) Free energy increase	12.	For conversion <i>C</i> (graphite) \rightarrow <i>C</i> (diamond) the ΔS is	
2.		is expanded isothermally and		[MP PMT 2001; MP PET 20	03]
	reversibly from 1 litre of 10 change (in kJ) for the proce	b litre at $300 K$. The enthalpy		(a) Zero (b) Positive	
	change (in x,) for the proce	[IIT JEE (Screening) 2004]		(c) Negative (d) Unknown	
	(a) 11.4 <i>kJ</i>	(b) -11.4 <i>kJ</i>	13.	For a reaction $\Delta H = 9.08 \ kJ \ mol^{-1}$ and	
	(c) $o kJ$	(d) 4.8 <i>kJ</i>		$\Delta S = 35.7 \ JK^{-1} mol^{-1}$	
3.		ong acid against strong base is		Which of the following statements is correct for	the
	constant and is equal to	ADMARADA DDMA		reaction	0
		980; AFMC 1989; DPMT 1991; 99; BHU 1999; MP PMT 1995]		[AMU (Engg.) 20	00]
	(a) 13.7 kcal	(b) 57 kJ		(a) Reversible and Isothermal	
	(c) $5.7 \times 10^4 J$	(d) All of the above		(b) Reversible and Exothermic	
	.,			(c) Spontaneous and Endothermic(d) Spontaneous and Exothermic	
4.	<i>gm</i> of <i>NaOH</i> reacts during	d when 36.5 <i>gm HCl</i> and 40 neutralization	14.	For a reaction to occur spontaneously [CBSE PMT 19	10 ≂ 1
	J	[NCERT 1984; CPMT 1993]	-4.	(a) $(\Delta H - T\Delta S)$ must be negative	-901
	(a) 76.5 <i>kcal</i>	(b) 13.7 <i>kcal</i>		(.) (

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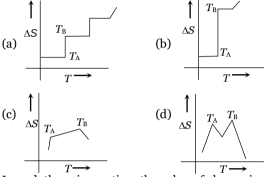
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(b) $(\Delta H + T\Delta S)$ must be negative

- (c) ΔH must be negative
- (d) ΔS must be negative

15. The total amount of energy in the universe is fixed, but

- [AMU (Engg.) 1999]
- (a) Disorder is increasing (b) Lightning is increasing
- (c) Matter is increasing (d) Gravitation is decreasing
- **16.** If for a given substance melting point is T_B and freezing point is T_A , then correct variation shown by graph between entropy change and temperature is **[DCE 2001]**



- 17. In endothermic reaction, the value of change in enthalpy (ΔH) is [Pb. CET 2001]
 - (a) Positive (b) Negative
 - (c) Zero (d) None of these
- 18. Which of the following would be expected to have the largest antropy per mole [MP PMT 2004]

(a) $SO_2Cl_2(s)$ (b) $SO_2Cl_2(g)$

(c)
$$SO_2Cl_2(l)$$
 (d) $SO_2(g)$

19. The enthalpies of formation of Al_2O_3 and Cr_2O_3 are -1596 kJ and -1134 kJ respectively. ΔH for the reaction $2Al + Cr_2O_3 \rightarrow 2Cr + Al_2O_3$ is **[KCET 2003]**

(a) -2730 kJ (b) -462 kJ

- (c) -1365 kJ (d) +2730 kJ
- **20.** Heat of reaction at constant volume is measured in the apparatus

(a) Bomb calorimeter ((b) Calorimeter
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- (c) Pyknometer (d) Pyrometer
- **21.** Which of the following gas has the highest heat of combustion

(a)]	Methane	(b)	Ethane
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- (c) Ethylene (d) Acetylene
- 22. The amount of heat measured for a reaction in a bomb calorimeter is [AIIMS 1991]
 - (a) ΔG (b) ΔH
 - (c) ΔE (d) $P\Delta V$
- **23.** For an endothermic reaction where ΔH represents the enthalpy of the reaction in *kJ/mole*, the minimum value for the energy of activation will be

[IIT JEE	1992; MP	PET 1993]
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- (a) Less than ΔH (b) Zero
- (c) More than ΔH (d) Equal to ΔH

- **24.** The heat of neutralization of a strong acid by a strong base is nearly equal to [MP PET 1993; BCECE 2005](a) + 57.32 J (b) - 57.32 kJ
 - (c) + 13.7 J (d) 13.7 kJ
- 25. A Beckmann thermometer is used to measure
 - [Kurukshetra CEE 2002]
 - (a) High temperature (b) Low temperature
 - (c) Normal temperature (d) All temperature
- **26.** The heat required to raise the temperature of a body by 1 K is called [AIEEE 2002]
 - (a) Specific heat (b) Thermal capacity
 - (c) Water equivalent (d) None of these
- 27. Mechanical work is specially important in system that contain [Kerala (Med.) 2002]
 - (a) Solid-liquid
- (b) Liquid-liquid(d) Amalgam
- (c) Solid-solid
- (e) Gases
- **28.** "The quantity of heat which must be supplied to decompose a compound into its element is equal to the heat evolved during the formation of that compound from the elements." This statement is known as
 - (a) Hess's law
 - (b) Joule's law
 - (c) Le-chatelier's principle
 - (d) Lavoiser and Laplace law
- **29.** Hess law deals with
 - (a) Changes in heat of reaction
 - (b) Rate of reaction
 - (c) Equilibrium constant
 - (d) Influence of pressure on volume of a gas
- **30.** For which one of the following reactions, ΔH is not equal to ΔE [IIT JEE 1995]
 - (a) $H_{2(g)} + I_{2(g)} \rightleftharpoons 2HI_{(g)}$
 - (b) $C_{(s)} + O_{2(g)} \rightleftharpoons CO_{2(g)}$
 - (c) $N_{2(g)} + 3H_{2(g)} \Rightarrow 2NH_{3(g)}$
 - (d) $HCl_{(aa)} + NaOH_{(aa)} \Rightarrow NaCl_{(aa)} + H_2O$
- **31.** The heat liberated when 1.89 *g* of benzoic acid is burnt in a bomb calorimeter at $25^{\circ}C$ increases the temperature of 18.94 *kg* of water by $0.632^{\circ}C$. If the specific heat of water at $25^{\circ}C$ is 0.998 *cal/g-deg*, the value of the heat combustion of benzoic acid is **[CPMT 1999; BHU 2000]**
 - (a) 771.1 *kcal* (b) 871.2 *kcal*
 - (c) 881.1 *kcal* (d) 981.1 *kcal*
- **32.** For a hypothetic reaction $A \rightarrow B$, the activation energies for forward and backward reactions are 19 *kJ/mole* and 9 *kJ/mole* respectively. The heat of reaction is

[CBSE PMT 2000]

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[BHU 1979]

(a) 28 kJ (b) 19 kJ(c) 10 kJ (d) 9 kJ

33.		tralization of <i>HCN</i> by <i>NaOH</i> is enthalpy of ionisation of <i>HCN</i> will	41.	The equilibrium constant of a reaction at 298 K 5×10^{-3} and at 1000 K is 2×10^{-5} . What is the size
	be	entitalpy of follisation of <i>Helv</i> with		5×10^{-3} and at 1000 K is 2×10^{-5} . What is the sign
		[BHU 1997]		$\Delta H \text{ for the reaction} \qquad [Pb. CET 199]$ (a) $\Delta H = 0$ (b) ΔH is negative
	(a) 4.519 <i>kJ</i>	(b) 45.10 <i>kJ</i>		(c) ΔH is positive (d) None of these
	(c) 451.9 <i>kJ</i>	(d) 45.19 <i>kJ</i>	42.	Which of the following has lowest fusion temperature
34∙	In thermodynamics, a p	rocess is called reversible when		[MP PET 200
		[AIIMS 2001]		(a) Naphthalene(b) Diamond(c) NaCl(d) Mn
	•	l system change into each other	43.	Consider the reactions
	(b) There is no surroundings	boundary between system and		$C(s) + 2H_2(g) \rightarrow CH_4(g), \Delta H = -x kcal$
	-	are always in equilibrium with the		$C(g) + 4H(g) \rightarrow CH_4(g), \Delta H = -x_1 kcal$
	system			$CH_4(g) \rightarrow CH_3(g) + H(g), \Delta H = +y kcal$
	(d) The system c spontaneously	hanges into the surroundings		The bond energy of $C - H$ bond is [JIPMER 199
85.		g unit represents largest amount of [AMU 2001]		(a) $y k cal mol^{-1}$ (b) $x_1 k cal mol^{-1}$
	(a) <i>Calorie</i>	(b) Joule		(c) $x/4 k cal mol^{-1}$ (d) $x_1/4 k cal mol^{-1}$
	(c) Erg	(d) Electron volt	44.	Given the bond energies $N \equiv N, H - H$ and $N - H$ bor
6.	Which of the following	g will have the highest bond energy		are 945,436 and $391 kJ mole^{-1}$ respectively, the enthal
		[MP PMT 1990]		of the following reaction $N_2(g) + 3H_2(g) \rightarrow 2NH_3(g)$ is
	(a) F_2	(b) Cl_2		[EAMCET 1992; JIPMER 199
	(c) Br_2	(d) I_2		(a) $-93 kJ$ (b) $102 kJ$
7.	$C_{(\text{graphite})} + O_2(g) \rightarrow CO$	$_{2}(g)$		(c) $90 kJ$ (d) $105 kJ$
	$\Delta H = -94.05 \ kcal \ mol$	-1	45.	The relation between ΔG and <i>E</i> for a cell is $\Delta G = -nF$ the cell reaction will be spontaneous if [MP PET 199]
	$C_{(\text{diamond})} + O_2(g) \rightarrow C_2$	$O_2(g); \Delta H = -94.50 \ kcal \ mol^{-1}$		(a) G is negative (b) G is positive
	therefore	[DPMT 2001]	46	(c) <i>E</i> is negative (d) <i>E</i> is positive Which of the following is not a correct statement?
		{nd)} ; $\Delta H{298 K}^{o} = -450 \ calmol^{-1}$	46.	[AMU (Engg.) 200
		{ite}); $\Delta H{298K}^{o} = +450 \ calmol^{-1}$		(a) When ΔG is negative, the process is spontaneous (b) When ΔG is zero, the process is in a state
	(c) Graphite is the sta	,		equilibrium
	(d) Diamond is harde	-		(c) When ΔG is positive, the process is non-spontaneo (d) None of these
8.		of two compounds x and y are -84	47.	If at 298 K the bond energies of $C - H, C - C, C = C$ a
	kJ and $-156 kJ$ restatements is correct	spectively. Which of the following [CBSE PMT 2001]		H-H bonds are respectively 414, 347, 615 and 435
	(a) x is more stable the formula $(x = 1)^{-1} = 1$	-		mol^{-1} , the value of enthalpy change for the reacting $H_2C = CH_2(g) + H_2(g) \rightarrow H_3C - CH_3(g)$ at 298 <i>K</i> will
	(b) x is less stable that	-		$H_2C = CH_2(g) + H_2(g) \rightarrow H_3C - CH_3(g)$ at 296 K with
	(c) Both x and y are u(d) x and y are endot			[AIEEE 200
39 .	For the process dry ice	-		(a) $+ 250 kJ$ (b) $- 250 kJ$ (c) $+ 125 kJ$ (d) $- 125 kJ$
	(a) ΔH is positive where ΔH		48.	1
	(b) Both ΔH and $\Delta \rho$			$\frac{1}{2}H_2 + \frac{1}{2}Cl_2 \to HCl \ (\Delta H_{298} = -22060 \ kcal \) \text{ means}$
	(c) Both ΔH and $\Delta \rho$	-		[CMC Vellore 19
	(d) ΔH is negative w			(a) The heat absorbed when one gram molecule of H is formed from its elements at 25 ° C is an of a keep
.	For melting of 3 mole	s of water at $0^{\circ} C$ the ΔG° is [MP PMT 2001]		 is formed from its elements at 25 ° C is 22.060 kca (b) The heat given out when one gram molecule of H is formed from its elements at 298 K is 22.060 kcal
	(a) Zero	(b) $+ ve$		(c) The heat absorbed when one atom of hydrogen rea
	(c) – <i>ve</i>	(d) Unpredictable		with one atom of chlorine to form one molecule HCl at $25^{\circ}C$ and one atmospheric pressure
				22.060 kcal



- (d) The heat absorbed when one gram equivalent of *HCl* is formed from its elements at 298 *K* is 22.060 *kcal*
- (e) The intrinsic heat of one molecule of *HCl* is 22.060 *kcal* more than the intrinsic heats of one atom of hydrogen and one atom of chlorine
- **49.** The $H_2O(g)$ molecule dissociates as
 - (i) $H_2O(g) \rightarrow H(g) + OH(g); \Delta H = 490 \ kJ$
 - (ii) $OH(g) \rightarrow H(g) + O(g); \Delta H = 424 \ kJ$
 - The average bond energy (in kJ) for water is (a) 490 (b) 424
 - (c) 914 (d) 914/2
- **50.** When $50 cm^3$ of $0.2 N H_2 SO_4$ is mixed with $50 cm^3$ of 1 N KOH, the heat liberated is
 [KCET 2004]

 (a) 11.46 kJ
 (b) 57.3 kJ

 (c) 573 kJ
 (d) 573 J
- **51.** Following reaction occurring in an automobile $2C_8H_{18}(g) + 25O_2(g) \rightarrow 16CO_2(g) + 18H_2O(g)$. The sign of $\Delta H, \Delta S$ and ΔG would be **[CBSE PMT 1994; KCET 1999]** (a) +, -, + (b) -, +, -
 - (c) -, +, + (d) +, +, -
- **52.** For the reaction

 $H_2(g) + \frac{1}{2}O_2(g) \rightarrow H_2O(l), \Delta H = -285.8 \text{ kJ mol}^{-1}$

 $\Delta S = -0.163 \ kJ \ mol^{-1} K^{-1}$. What is the value of free energy

- change at 27 ° C for the reaction [KCET 1999] (a) $-236.9 kJ mol^{-1}$ (b) $-281.4 kJ mol^{-1}$
- (c) $-334.7 kJ mol^{-1}$ (d) $+334.7 kJ mol^{-1}$

R Assertion & Reason For ANMS Aspirants

Read the assertion and reason carefully to mark the correct option out of the options given below :

- (a) If both assertion and reason are true and the reason is the correct explanation of the assertion.
- (b) If both assertion and reason are true but reason is not the correct explanation of the assertion.
- (c) If assertion is true but reason is false.
- (d) If the assertion and reason both are false.
- (e) If assertion is false but reason is true.

1.	Assertion	:	The enthalpy of formation of gaseous oxygen molecules at $298 K$ and under pressure of $1 atm$ is zero.
	Reason	:	The entropy of formation of gaseous oxygen molecules under the same condition is zero. [AIIMS 1996]
2.	Assertion	:	We feel cold on touching the ice.
	Reason	:	Ice is a solid form of water. [AIIMS 1999]
3.	Assertion	:	Entropy of ice is less than water.
	Reason	:	Ice have cage like structure. [AIIMS 2000]
4.	Assertion	:	The heat absored during the isothermal expansion of an ideal gas against vacuum is zero.

			na memochemistry 437
	Reason	:	The volume occupied by the molecules of an ideal gas is zero. [AIIMS 2002]
5.	Assertion	:	Absolute values of internal energy of substance can not be determined.
	Reason	:	It is impossible to determine exact values of constituent energies of the substances.
6.	Assertion	:	[AIIMS 2002] Mass and volume are extensive
	Reason	:	properties. Mass / volume is also an extensive parameter. [AIIMS 2002]
7.	Assertion	:	Molar entropy of vaporization of water is different from ethanol.
	Reason	:	Water is more polar than ethanol.
8.	Assertion	:	The increase in internal energy (ΔE) for
			the vaporiation of one mole of water at 1 atm and $373K$ is zero.
	Reason	:	For all isothermal processes $\Delta E = 0$.
			[AIIMS 2003]
9.	Assertion	:	ΔH and ΔE are almost the same for the reaction. $N_2(g) + O_2(g) \rightleftharpoons 2NO(g)$.
	Reason	:	All reactants and products are gases. [AIIMS 2003]
10.	Assertion	:	The enthalpies of neutralisation of strong acids and strong bases are always same.
	Reason	:	Neutralisation is heat of formation of water.
	A		[AIIMS 1996]
11.	Assertion	:	Zeroth law can also be termed as law of thermal equilibirum.
	Reason	:	Two objects in thermal equilibrium with the third one, are in thermal equilibrium with each other.
12.	Assertion	:	There is no reaction known for which ΔG is positive, yet it is spontaneous.
	Reason	:	For photochemical reactions ΔG is negative.
13.	Assertion	:	Heat of neutralisation of perchloric acid, $HClO_4$ with $NaOH$ is same as is that of HCl with $NaOH$.
	Deeger		
	Reason	:	Both HCl and $HClO_4$ are strong acids.
14.	Assertion	:	Heat absorbed in a reaction at constant temperature and constant volume is $-\Delta G$.
	Reason	:	ΔG should be negative for the reaction to be spontaneous.
15.	Assertion	:	T, P and V are state variables or state functions.
	Reason	:	Their values depends on the state of the system and how it is reached.
16.	Assertion	:	Internal energy is an extensive property.
	Reason	:	Internal energy depends upon the amount of the system.
17.	Assertion	:	For the combustion reactions, the value of ΔH is always negative.





	Reason	:	The combustions reactions are always endothermic.
18.	Assertion	:	For an isothermal reversible process $Q = -w$ i.e. work done by the system equals the heat absorbed by the system.
	Reason	:	Enthalpy change (ΔH) is zero for isothermal process.
19.	Assertion	:	Enthalpy of formation of graphite is zero but of diamond is not zero.
	Reason	:	Enthalpy of formation of most stable allotrope is taken as zero.
20.	Assertion	:	If a refrigerator's door is kept open room gets cooled.
	Reason	:	Material kept inside the refrigerator reamins cool.
21.	Assertion	:	Enthalpy and entropy of any elementary substance in the standard state are taken as zero.
	Reason	:	At zero degree absolute, the constituent particles become completely motionless.
22.	Assertion	:	A process is called adiabatic if the system does not exchange heat with the surroundings.
	Reason	:	It does not involve increase or decreae in temperature of the system.

41	d	42	a	43	С	44	a	45	c
46	С	47	b	48	С	49	d	50	b
51	а	52	d	53	b	54	C	55	a
56	b	57	d	58	d				

IInd & IIIrd Law of thermodynamics and Entropy

1	с	2	d	3	d	4	b	5	b
6	d	7	a	8	d	9	b	10	C
11	C	12	C	13	d	14	d	15	C
16	b	17	d	18	а	19	а	20	a
21	d	22	C	23	а	24	d	25	C
26	d	27	C	28	b	29	C	30	b
31	b	32	d	33	d	34	а	35	C
36	b	37	C	38	a	39	C	40	C
41	c	42	C	43	c	44	d	45	b
46	a	47	C	48	d	49	d	50	C
51	C	52	b	53	а	54	C	55	C
56	С	57	a	58	b	59	d	60	b
61	b	62	d	63	а	64	C	65	a
66	c	67	b	68	b				



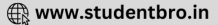
Basic concepts

1	b	2	d	3	b	4	c	5	с
6	d	7	d	8	c	9	c	10	c
11	d	12	d	13	а	14	C	15	b
16	b	17	d	18	а	19	b	20	b
21	d	22	b	23	С	24	C	25	d
26	c	27	c	28	c				

First law of thermodynamics and Hess law

1	d	2	b	3	a	4	b	5	c
6	a	7	C	8	a	9	b	10	d
11	b	12	c	13	c	14	c	15	b
16	b	17	C	18	b	19	c	20	а
21	С	22	C	23	a	24	а	25	b
26	a	27	a	28	a	29	bc	30	C
31	b	32	c	33	b	34	а	35	c
36	b	37	c	38	b	39	d	40	c





Heat of reaction										
1	b	2	b	3	d	4	d	5	d	
6	b	7	с	8	a	9	b	10	а	
11	bcd	12	a	13	d	14	b	15	с	
16	а	17	b	18	а	19	b	20	с	
21	с	22	a	23	d	24	c	25	с	
26	b	27	b	28	b	29	a	30	с	
31	а	32	а	33	d	34	b	35	b	
36	a	37	а	38	d	39	b	40	b	
41	с	42	с	43	b	44	b	45	b	
46	b	47	а	48	с	49	е	50	b	
51	а	52	b	53	с	54	c	55	b	
56	с	57	с	58	d	59	с	60	b	
61	d	62	с	63	b	64	a	65	b	
66	b	67	а	68	с	69	а	70	d	
71	а	72	b	73	d	74	с	75	b	
76	b	77	а	78	d	79	d	80	а	
81	b	82	b	83	a	84	b	85	b	
86	c	87	а	88	b	89	d	90	d	
91	a	92	с	93	b	94	а	95	d	
96	d	97	с	98	a	99	с	100	с	
101	b	102	с	103	d	104	d	105	с	
106	a	107	с	108	d	109	d	110	с	
111	a	112	b	113	a	114	a	115	а	
116	a	117	b	118	c	119	b	120	а	
121	а	122	a	123	d	124	b	125	b	
126	b	127	с	128	d	129	d	130	b	
131	d	132	b	133	d	134	a	135	а	
136	a	137	d	138	c	139	d	140	b	
141	c	142	d	143	c	144	b	145	с	
146	d	147	b	148	c	149	b	150	b	
151	d	152	a	153	b	154	b	155	b	
156	b	157	a	158	d	159	c	160	d	
161	а	162	с	163	c	164	c	165	с	
166	b	167	d	168	d	169	с	170	b	
171	a	172	с							

b 2 a 3 a 4 d 5 c

d

9

b

10

b

Free energy and Work function

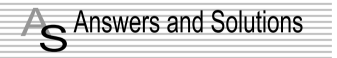
1	d	2	b	3	a	4	b	5	acd
6	C	7	C	8	С	9	a	10	a
11	d	12	а	13	d	14	d	15	b
16	d	17	b	18	c	19	d	20	b
21	b	22	d	23	b	24	а	25	d
26	d	27	a	28	d				

Critical Thinking Questions

1	b	2	c	3	d	4	b	5	b
6	d	7	d	8	b	9	а	10	a
11	а	12	C	13	C	14	а	15	a
16	а	17	а	18	d	19	b	20	a
21	b	22	С	23	C	24	b	25	b
26	b	27	е	28	d	29	а	30	с
31	а	32	С	33	d	34	ac	35	а
36	b	37	c	38	b	39	c	40	a
41	b	42	а	43	а	44	а	45	d
46	d	47	d	48	b	49	d	50	d
51	b	52	а						

Assertion & Reason

1	b	2	b	3	b	4	c	5	a
6	b	7	b	8	а	9	b	10	a
11	а	12	d	13	а	14	е	15	c
16	а	17	С	18	b	19	а	20	е
21	C	22	С						



Basic concepts

- (b) Internal energy of an ideal gas is a function of temperature only.
- 2. (d) In cyclic process, a system in a given state goes through a series of different processes, but in the end returns to its initial state.
- **4.** (c) $\Delta E = 0$ for isothermal reversible cycle.
- 5. (c) In isolated system neither exchange of matter nor exchange of energy is possible with surroundings.

8

1

6

7

b

с





- 9. (c) It is the definition of calorific value.
- (d) When a real gas is forced through a porous plug into a region of low pressure, it is found that due to expansion, the gas on the side of low pressure gets cooled.

The phenomenon of producing lowering of temperature when a gas is made to expand adiabatically from a region of high pressure into a region of low pressure is known as Joule-Thomson effect.

- **12.** (d) In isothermal reversible process ideal gas has constant volume and so $\Delta E = 0$ and $\Delta H = \Delta E = 0$.
- **13.** (a) The compressor has to run for longer time releasing more heat to the surroundings.
- 14. (c) This is based on Joule-Thomson effect.
- **15.** (b) Enthalpy is an extensive property.
- **16.** (b) dV = 0 for an isochoric process.
- 17. (d) Total energy of an isolated system is constant.
- **18.** (a) For isochoric process $\Delta V = 0$ so $q_v = \Delta E$ i.e.

heat given to a system under constant volume is used up in increasing ΔE .

- **19.** (b) The less energy of a system and more is its stability.
- 20. (b) The functions whose value depends only on the state of a system are known as state functions.
- **21.** (d) For adiabatic process q = 0.
- **22.** (b) The intensive property is mass/volume.
- 23. (c) Volume is not an intensive property.
- 24. (c) An isolated system neither shows exchange of heat nor matter with surroundings.
- **25.** (d) ΔQ is not a state function.
- **26.** (c) For adiabatic process $\Delta Q = 0$.
- 27. (c) Work is not a state function as during a process its value depends on the path followed. The value of enthalpy, internal energy and entropy depends on the state and not on the path followed to get that state, hence these are state functions.
- 28. (c) Surface tension is an intensive property which do not depend upon the quantity of matter present in the system.

First law of thermodynamics and Hess law

- (d) First low of thermodynamics is also known as Law of conservation of mass and energy.
- 2. (b) Formation of CO_2 from CO is an exothermic reaction; heat is evolved from the system, *i.e.*, energy is lowered. Thus, exothermic reactions occur spontaneously on account of decrease in enthapy of system. Thus, $\Delta E > \Delta H$.
- **4.** (b) $\Delta H = \Delta E + P \Delta V$.

- 5. (c) $\Delta n_g = 1 \frac{3}{2} = \frac{-1}{2}$, As Δn_g is negative, thus $\Delta H < \Delta E$.
- 6. (a) Bomb calorimeter is commonly used to find the heat of combustion of organic substances which consists of a sealed combustion chamber, called a bomb. If a process is run in a sealed container then no expansion or compression is allowed, so w = 0 and $\Delta U = q$. $\Delta U < 0, w = 0$
- 9. (b) $\Delta H = \Delta E + \Delta nRT$ Since $\Delta n = -2$ Than $\Delta H = \Delta E - 2RT$.
- **10.** (d) If $\Delta n = -ve$ than $\Delta H < \Delta E$.
- (c) Hess's law is an application of first law of thermodynamics.
- **13.** (c) At constant *P* or *T* $\Delta H = \Delta U + \Delta nRT \Rightarrow \Delta n = n_p - n_R = 2 - 4 = -2$ $\therefore \Delta H < \Delta U$.
- **16.** (b) It is a combustion reaction, $\Delta H = -ve$.
- 17. (c) During isothermal expansion of ideal gas, $\Delta T = 0$

$$\Delta H = \Delta E + P \Delta V = \Delta E + nR \Delta T = 0 + 0 = 0.$$

18. (b)
$$W = 2.303 \ nRT \log \frac{V_2}{V_1}$$

= 2.303 ×1 ×8.314 ×10⁷ × 298

$$= 298 \times 10^{7} \times 8.314 \times 2.303 \log 2$$
.

20. (a) The enthalpies of all elements in their standard state at 25° C or 298K are zero.

 $\log \frac{20}{10}$

- **21.** (c) $\Delta E_v = E_P E_R$.
- **22.** (c) $\Delta E = q + w$.
- **23.** (a) $\Delta E = 0$ for reversible isothermal process.
- **25.** (b) Hess law includes initial reactants and final products.
- **26.** (a) At constant *T* and *P* internal energy of ideal gas remains unaffected.
- **27.** (a) ΔE increases with temperature.
- **28.** (a) $\Delta H = \Delta E + W$ or $\Delta H = \Delta E + P\Delta V$.
- **29.** (bc) Heat of neutralisation of a strong acid and strong base is equal to $-13.7 \, kcal$.

31. (b) Joule-Thomson expansion is isoenthalpic.

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32. (c) Here $\Delta n = 0$ so, $\Delta E = \Delta H$.

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- **33.** (b) $q = \Delta E W$ if q = 0 for adiabatic process, than $\Delta E = W$.
- **34.** (a) For this reaction $\Delta n = 0$ than $\Delta E = \Delta H$.
- **35.** (c) As the system is closed and insulated no heat It we enter or leave the system, *i.e.* q = 0; $\therefore \Delta E = Q + W = W$. system.
- **37.** (c) $\Delta H \Delta E = \Delta nRT$; $\Delta n = -3$ so, $\Delta H - \Delta E = -3RT$.
- 38. (b) According to Hess's law. The heat of reaction depends upon initial and final conditions of reactants.
- **39.** (d) $\Delta H \Delta E = \Delta nRT$ also, $2NH_{3(g)} \rightarrow N_{2(g)} + 3H_{2(g)}$
 - $\Delta n = 2$.
- **40.** (c) $\Delta n = 0$ for this reaction so, $\Delta E = \Delta H$.
- **41.** (d) W = 0 is not true.

42. (a)
$$W = 2.303 \ nRT \log \frac{r_2}{P_1}$$

$$= 2.303 \times 1 \times 2 \times 300 \log \frac{10}{2} = 965.84$$

10

at constant temperature, $\Delta E = 0$.

 $\Delta E = q + w$; q = -w = -965.84 cal.

43. (c) q = 40 J w = -8 J (work done by the system) $\Delta E = q + w = 40 - 8 = 32 J$.

44. (a) We know that $\Delta E = Q + W = 600 + (-300) = 300 J$ W = 300, because the work done by the system.

45. (c) Given that

 $P_1 = 10 \ atm$, $P_2 = 1 \ atm$, $T = 300 \ K$, n = 1 $R = 8.314 \ J / K / mol$ Now, by using $W = 2.303 \ nRT \ \log_{10} \frac{P_2}{P_1}$

 $= 2.303 \times 1 \times 8.314 \times 300 \log_{10} \frac{1}{10}$

W = 5744.1 Joule

46. (c) We know that internal energy of a gas depends upon its pressure and temperature. Thus if a gas expands at constant temperature and pressure, then its internal energy remains same.

48. (c) Enthalpy (*H*) is defined as the sum of internal energy E + PV, H = E + PV.

49. (d)
$$(CH_3)_2 C = CH_2(g) + 6O_2(g) \rightarrow 4CO_2(g) + 4H_2O(l)$$

 $\Delta ng = 4 - 6 = -2$ (i.e., negative)
we know that $\Delta H = \Delta E + \Delta ngRT$
 $= \Delta E - (\Delta ng)RT$ (:: $\Delta ng = -ve$)
 $\therefore \Delta H < \Delta E$

(b) Given number of moles =1 Initial temperature $= 27^{\circ} C = 300 K$ Work done by the system = 3 KJ = 3000 KIt will be (-) because work is done by the system.

Heat capacity at constant volume (Cv) = 20 J/k

We know that work done

$$\begin{split} W &= -nC_V(T_2 - T_1) \; ; \quad 3000 \; = -1 \times 20 \; (T_2 - 300) \\ 3000 \; = -20 T_2 \; + \; 6000 \end{split}$$

$$20T_2 = 3000$$
; $T_2 = \frac{3000}{20} = 150 K$

- 51. (a) Internal energy of a system is a state function and extensive property and is independent of the path by which it is obtained.
- **52.** (d) $N_2O_4(g) \to 2NO_2(g)$

50.

For this reaction $\Delta ng = 2 - 1 = 1$ Δng is positive *i.e.*, there is an increase in the number of gaseous moles then $\Delta H > \Delta E$

53. (b)
$$2C + O_2 \rightarrow 2CO$$
; $\Delta H = -220 \text{ KJ}$

This reaction does not represent complete combustion of carbon, hence heat of combustion of carbon will not be equal to 110 kJ. The negative sign of ΔH indicates that this reaction is exothermic. Also, despite being spontaneous reaction, it requires initiation.

54. (c)
$$W = -P\Delta V = -1 \times 10^5 (1 \times 10^{-2} - 1 \times 10^{-3})$$

 $= -1 \times 10^{5} \times 9 \times 10^{-3} = -900 J$

55. (a) The exact value of internal energy is not known as it includes all type of energies of molecules constituting the given mass of matter such as translational, vibrational, rotational. The kinetic & potential energy of the nuclei & electron with in the individual molecules and the manner in which the molecule are linked together etc.

$$E = E_{translational} + E_{rotational} + E_{vibrational}$$

Thus, we can say that internal energy is partly potential and partly kinetic.

56. (b) $N_2(g) + O_2(g) \rightarrow 2NO(g); \Delta n_g = 2 - 2 = 0$ $\Delta H = \Delta E + \Delta n_g RT; \Delta H = \Delta E + 0; \Delta H = \Delta E$

57. (d)
$$W = -p\Delta V$$
; $W = -3 \times (6-4)$
 $W = -6 \times 101.32$ ($\therefore 1 Latm = 101.32 J$)
 $W = -608 J$

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58. (d)
$$A(s) + 3B(g) \rightarrow 4C(s) + D(l)$$

 $\Delta ng = 0 - 3 = -3$; $\Delta H = \Delta E + \Delta ngRT$ [:: $\Delta E = \Delta U$]
 $\therefore \Delta H = \Delta U + (-3)(RT)$
 $\Delta H = \Delta U - 3(RT)$; $\Delta H = \Delta U - 3RT$

IInd & IIIrd Law of thermodynamics and Entropy

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- **2.** (d) When $\Delta S = +ve$ the change is spontaneous.
- 3. (d) Heat is always flow from the higher to lower temperature.
- **4.** (b) Mixing of non-reacting gases increases randomness and so increase entropy .
- 5. (b) Entropy of the system increases as the process occur irreversibly and isothermally in an isolated system.
- **6.** (d) $\Delta S^o = 2S^o_{HCl} (S^o_{H_2} + S^o_{Cl_2})$

 $= 2 \times 186.7 - (130.6 + 223.0) = 19.8 JK^{-1}mol^{-1}$

9. (b) For adiabatic expansion q = 0 than according

to following relation $\Delta S = \frac{q}{T}, \Delta S = 0$.

- **10.** (c) It is the third law of thermodynamics.
- 11. (c) Entropy of gases is highest.
- **12.** (c) $\Delta G = \Delta H T \Delta S$

for spontaneous process ΔG should be negative in option (3) $\Delta H = -ve$ and $\Delta S = +ve$ than $\Delta G = (-ve) - T(+ve) = -ve$

- **15.** (c) For reverse reaction sign will be change.
- **16.** (b) Solid \longrightarrow Gas, ΔS is maximum.
- **17.** (d) $+ ve \Delta H$ and $ve \Delta S$ both oppose the reaction.
- **18.** (a) $\Delta S_{vap} = 186.5 / 373 = 0.5 \ JK^{-1} \ mol^{-1}$.
- **20.** (a) When $\Delta H = -ve$, $\Delta S = +ve$ and $\Delta G = -ve$ than reaction is spontaneous.
- **21.** (d) For endothermic process ΔS increases.
- **22.** (c) Calculation of change in entropy is done at constant temperature and pressure both.
- 23. (a) When the value of entropy is greater, then ability of work is maximum.
- 24. (d) At equilibrium, $\Delta G = 0$ Hence $0 = \Delta H - T\Delta S$ or $\Delta H = T\Delta S$. 25. (c) $\Delta S_{un} = \frac{\Delta H_{vap}}{\Delta S_{un}} = \frac{37.3 \text{ KJ mol}^{-1}}{37.3 \text{ KJ mol}^{-1}}$

(c)
$$\Delta S_{vap} = \frac{T}{T} = \frac{373 \ K}{373 \ K}$$

$$= 0.1 \, kJ \, mol^{-1} \, K^{-1} = 100 \, J \, mol^{-1} \, K^{-1}$$
.

- **26.** (d) Solid \rightarrow Liquid, ΔS increases.
- **27.** (c) $\Delta S = +ve$ than process is spontaneous.
- **28.** (b) $\Delta S = +ve$ than the system is more disordered.
- **29.** (c) Because solid \rightarrow solid, ΔS is same and ΔH is *ve*.

30. (b)
$$\Delta S_{vap} = \frac{\Delta H_{vap}}{T} = \frac{386}{298} = 1.2 kJ$$

- **31.** (b) Processes (a) and (c) take place with the increase of no. of moles of gaseous species and hence the disorder or entropy increases. (b) on increasing pressure, disorder or randomness decreases and so also the entropy (d) is endothermic process and ΔS is positive.
- **32.** (d) This is the statement of third law of thermodynamics.
- **33.** (d) For isothermal expansion of ideal gas, $\Delta E = 0$.

34. (a)
$$\Delta S_{vap} = \frac{(900 \times 18)}{373} = 43.4 \ JK^{-1} \ mol^{-1}$$

- **35.** (c) Entropy of the mixture increases due to increase in impurity.
- **37.** (c) If $\Delta H = +ve$ and $\Delta S = -ve$ than the reaction is non-spontaneous *i.e.* can not occur.
- **38.** (a) $T_2 = 150 + 273 = 423 K$
 - $T_1 = 25 + 273 = 298 K$ Q = 500 K $\frac{W}{Q} = \frac{T_2 T_1}{T_2} ; W = 500 \left(\frac{423 298}{423}\right) = 147.7 J.$
- 39. (c) According to III law of thermodynamics.
- **40.** (c) When $\Delta H = +ve$ and $\Delta S = -ve$ reaction is non-spontaneous.

41. (c)
$$T_m = \frac{\Delta H_{\text{fusion}}}{\Delta S_{\text{fusion}}} = \frac{9.2}{0.008} = 1150 \text{ K}$$
.

42. (c) Here: Change in Volume (V) = 500 - 300 = 200 cc = 0.2 litre, Pressure (P) = 0.6 atm and heat liberated (a) =

Work done (*W*) = $P\Delta V = (0.2 \times 0.6) = 0.12 \, litre - atm$

But 1 litre - atm = 101.3 J.

hence $W = 0.12 \times 101.3 = 12.156 J$. We also know that heat is liberated, therefore it would be negative. Thus change in $\Delta E = q + W = -10 + 12.16 = 2.16 J$.

43. (c) Formation of CO_2 is,

$$C_{(s)} + O_{2(g)} \rightarrow CO_{2(g)}$$

 $\Delta S^{o} = 213.5 - 5.690 - 205 = 2.81 \ JK^{-1}.$

45. (b) $H_2 O_{(g)} = H_2 O_{(l)}$ we know $\Delta G = \Delta H - T \Delta S$

at equilibrium
$$\Delta G = 0$$

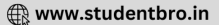
Therefore $\Delta H = T\Delta S$.

46. (a) We know that work done,
$$W = C_v(T_1 - T_2)$$

 $3 \times 1000 = 20(300 - T_2);$ $\therefore 3000 = 6000 - 20T_2$

$$T_2 = \frac{3000}{20} = 150 \ K$$
.

- **48.** (d) It does not violates the first law of thermodynamics but violates the II law of thermodynamics.
- **49.** (d) $C_v = \frac{3}{2}RT$; $C_p = \frac{5}{2}RT$ for monoatomic gas $C_v = \frac{5}{2}RT$; $C_p = \frac{7}{2}RT$ for diatomic gas Thus for mixture of 1 *mole* each, $C_v = \frac{\frac{3}{2}RT + \frac{5}{2}RT}{2}$ and $C_p = \frac{\frac{5}{2}RT + \frac{7}{2}RT}{2}$



Therefore,
$$C_p / C_v = \frac{3RT}{2RT} = 1.5$$
.

51. (c)
$$\Delta S = \frac{q_{rev}}{T}$$
 : unit of S is $JK^{-1}mol^{-1}$.

52. (b)
$$H_2 O_{(l)} \rightleftharpoons H_2 O_{(g)}, \Delta S = \frac{\Delta H_{vap}}{T},$$

 $\Delta H_{vap.} = 2.257 \ KJ / g$
or $\Delta H_{vap} = 2.257 \times 18 \ kJ / mol. = 40.7 \ KJ / mol$
hence, $\Delta S = \frac{40.7}{373} = 0.109 \ kJ / mol/K.$

- (a) Liquid \longrightarrow Vapour, entropy increases. 53.
- (c) *NaNO*₃ is a solid, which is converted to liquid 54. ions.
- (c) Heat capacity of water per gram = $\frac{75}{18}$ = 4.17 J 55.

$$Q = mst ; 1000 = 100 \times 4.17 \times t$$
$$t = \frac{1000}{100 \times 4.17} = 2.4 K.$$

56. (c) As the work is done on system, it will be As the work is done on ejected, positive i.e. W = +462 joule, E = -128 joule (heat is evolving) **68.** (b) $dS = \frac{dQ_{rev.}}{T}$; $T = \frac{30 \times 10^3}{75}$; T = 400 KFrom the Ist law of thermodynamics

 $\Delta E = q + w = (-128) + (+462) = +334$ Joules.

(a) Gases show highest entropy. 57.

58. (b)
$$\begin{array}{c} H^{+}{}_{(aq)} + OH^{-}{}_{(aq)} \to H_{2}O_{(l)} \\ S^{o}(298 \ K)K^{-}mol^{-1} & {}^{-10.7} & {}^{+70} \end{array}$$
$$\Delta S^{o}(298 \ K) = \Delta S_{P} - \Delta S_{R} = 70 - (-10.7 + 0) \\ 80.7 \ JK^{-1} \ mol^{-1} \end{array}$$

- heat of vaporisation (d) The entropy change =59. temperatur e
 - Here, heat of vaporisation $= 540 \, cal/gm$

$$= 540 \times 18 \ cal \ mol^{-1}$$

Temperature of water
$$= 100 + 273 = 373 K$$

$$\therefore \qquad \text{entropy} \qquad \text{change}$$

$$= \frac{540 \times 18}{373} = 26.06 \ cal \ mol^{-1} K^{-1}$$

- (b) Given that, $T_1 = 500 K$, $T_2 = 300 K$ 60. By using, $\eta = \frac{T_1 - T_2}{T_1} = \frac{500 - 300}{500} = \frac{200}{500} = 0.4$.
- (b) It is molar heat capacity. 61.

=

- (d) Entropy is the measure of randomness in the 62. molecules. Randomness is maximum in case of gases. Hence, entropy is maximum for water vapours.
- (a) Standard entropy of formation of $CO_2(g) =$ 63. standard entropy of $CO_2(g)$ – [Standard entropy of C(s) – standard entropy of $O_2(g)$] = 213.5 - [5.740 + 205] = 2.76 J/K.
- (c) Third law of thermodynamics help in 64. calculating entropy of different temperatures.

Thermodynamics and Thermochemistry 443

- (a) In case of gas randomness is maximum 65. therefore entropy is maximum in case of steam.
- 66. (c) $\Delta S_{system} + \Delta S_{surroundin\,gs} > O$ (for spontaneity)

ecause of
$$\Delta S = R \ln \frac{V_2}{V_1}$$

R

Here the volume of gas increase from V_1 to V_2 at constant temperature T.

The total increase in entropy of the system and its surrounding during the spontaneous process of expansion considered above is,

thus
$$R \ln \left(\frac{V_2}{V_1} \right)$$
 since $V_2 > V_1$ it is obvious that

the spontaneous (irreversible) isothermal expansion of a gas is accompanied by an increase in the entropy of the system and its surrounding considered together.

 $\Delta S_{system} + \Delta S_{surroundin\,gs} > 0$.

67. (b) $\Delta G = \Delta H - T \Delta S$ at constant temperature and pressure $\Delta G = 0$

$$0 = \Delta T - T\Delta S \text{ so } \Delta H = T\Delta S$$

Heat of reaction

1. (b)
$$C + O_2 \rightarrow CO_2 + 94.2$$
 Kcal.(i)

$$H_2 + \frac{1}{2}O_2 \to H_2O + 68.3$$
 Kcal.(ii)

On multiplication of eq. (ii) by 2 and than adding in eq. (i)

$$C + 2H_2 + 2O_2 \rightarrow CO_2 + 2H_2O + 230.8 \text{ Kcal}$$
 ...(iii)

On subtracting eq. (iii) by following eq.

 $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O + 210.8$ Kcal. we get,

$$C + 2H_2 \rightarrow CH_4 \quad \Delta H = 20 \quad Kcal.$$

2. (b)
$$\Delta S = 16 J mole^{-1} K^{-1}$$

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$$T_{b.p.} = \frac{\Delta H_{\text{vapour}}}{\Delta S_{\text{vapour}}} = \frac{6 \times 1000}{16} = 375 \, K$$

- (d) Heat of neutralisation between strong acid 4. and a strong base is about -13.7 Kcal.
- (b) Effect of temperature in heat of reaction is 6. given by Kirchoff's equation.
- (c) Heat of neutralisation between strong acid 7. and a strong base is about -13.7 Kcal.
- 8. (a) For exothermic reactions $H_p < H_R$. For endothermic reactions $H_p > H_R$.
- 9. (b) 78g of benzene on combustion produces heat = $-3264.6 \, kJ$

:. 39 g will produce =
$$\frac{-3264.6}{2} = -1632.3 \, kJ$$
.

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10. (a) eq. (i) + eq. (ii) gives the required result.

- **11.** (bcd) (b,c,d) are endothermic reactions because they proceeds by the absorption of heat.
- 12. (a) Change of liquid to vapour takes energy in the form of heat so it is endothermic reaction.
- **13.** (d) In exothermic reactions heat is evolute.
- **15.** (c) $\Delta H = -ve$ for exothermic reaction. $\Delta H = +ve$ for endothermic reaction

Enthalpy of fusion is + ve.

16. (a) Heat of neutralisation will be less than -57.33 kJ/mole because some amount of this energy will be required for the dissociation of weak base (M_gO)

18. (a)
$$H_2 + O_2 \to H_2O_2 \quad \Delta H_f^o = -188 \ kJ \ mole \qquad \dots$$
(i)

$$H_2 + \frac{1}{2}O_2 \rightarrow H_2O \quad \Delta H_f^o = -286 \ kJ \ / \ mole \qquad \dots (ii)$$

eq. (i) – eq. (ii)
$$\times$$
 2 gives the required result.

(b) Graphite
$$\longrightarrow$$
 diamond $\Delta H_t = (x - y)kJ mol^{-1}$.

(c) Heats of combustions are always exothermic except oxidation of N as,

$$N_2 + \frac{1}{2}O_2 \rightarrow N_2O \quad \Delta H = +ve$$

 $N_2 + O_2 \rightarrow 2NO \quad \Delta H = +ve$

- **22.** (a) For exothermic reactions $H_p < H_R$. For endothermic reactions $H_p > H_R$.
- 23. (d) Aim: $2C + H_{2(g)} \rightarrow C_2 H_{2(g)}$. eq. (ii) + eq. (iii) \rightarrow eq. (iv) - eq. (i) find the required result.
- **24.** (c) Enthalpy of formation of *HCl*.
- **25.** (c) Heat of neutralisation between strong acid and a strong base is about -13.7 Kcal.
- **26.** (b) For endothermic reaction, $\Delta H = +ve$.
- 27. (b) Heat of neutralisation is less than 56.1 *Kcal* when a strong base and a weak acid reacts.

28. (b) Aim:
$$CO + \frac{1}{2}O_2 \rightarrow CO_2$$

$$\Delta H = \Delta H_f^0 (CO_2) - \left[\Delta H_f^0 (CO) + \frac{1}{2}\Delta H_f^0 (O_2)\right]$$
$$= -94.0 - (-26.4) = -67.6 \ kcal.$$

29. (a)

19.

(i)
$$H_2 + \frac{1}{2}O_2 \rightarrow H_2O, \Delta H = -241 \, kJ$$

(ii) $C_6H_{10} + \frac{17}{2}O_2 \rightarrow 6CO_2 + 5H_2O, \Delta H = -3800 \, kJ$
(iii) $C_6H_{12} + 9O_2 \rightarrow 6CO_2 + 6H_2O, \Delta H = -3920 \, kJ$
 $C_6H_{10} + H_2 \rightarrow C_6H_{12}$
Eq. (i) + Eq. (ii) - Eq. (iii)
 $\Delta H = -241 - 3800 - (-3920)$
 $= -4041 + 3920 = -121 \, kJ$

- 30. (c) NH₄OH is a weak base. Heat of neutralisation < 13.7kcal.
- **31.** (a) CH_4 is the best fuel because its calorific value = $\frac{-212.8}{16} = -13.3 \, k \, cal/g$ is higher among the other gases.
- **32.** (a) Find ΔH for $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O_{(g)}$.
- **34.** (b) In neutralisation of a strong acid and base only H^+ and OH^- ions react.
- **35.** (b) When both acid and base are strong than heat of neutralisation is $57.1 kJ mol^{-1}$.

36. (a)
$$C_2H_6 + \frac{7}{2}O_2 \rightarrow 2CO_2 + 3H_2O$$
 substitute the values.

- **37.** (a) Decomposition is an endothermic process.
- **38.** (d) ΔH for this reaction is Heat of reaction.
- **40.** (b) One mole of H_2O is formed from its initial components.
- 41. (c) One mole of a substance is completely oxidized in air.
- **42.** (c) *S* (rhombic) $+O_2 \rightarrow SO_2$, $\Delta H = 70960$ cal. ...(i) *S* (monoclinic) $+O_2 \rightarrow SO_2 \Delta H = 71030$ cal ...(ii) Aim: S (rhombic) $\rightarrow S$ (monoclinic) *eq.* (i) -eq. (ii) gives the required result.
- **43.** (b) When $H_2O_{(l)}$ is convert to form $H_2O_{(g)}$, heat is absorbed hence $\Delta H_1 > \Delta H_2$.
- **44.** (b) Out of given substances, kerosene oil has maximum calorific value.

45. (b)
$$C_{(S)} + O_{2(g)} \to CO_{2(g)}$$
, $\Delta n_g = 1 - 1 = 0$
 $\therefore \Delta E = \Delta H = -94 \ kcal$
 $\Delta E = E_{CO_2} - (E_C + E_{O_2}) = E_{CO_2} - (0 + 0)$
or $E_{CO_2} = \Delta E = -94 \ kcal$.

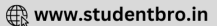
- **46.** (b) 0.2 *mole* will neutralize 0.2 mole of HNO_3 , heat evolved = $57 \times 0.2 = 11.4 kJ$.
- **47.** (a) Suppose heat evolved in I^{st} case is Q_1 and that

in the IInd case it is Q_2 . Then $Q_2 = \frac{1}{2}Q_1$. But $Q_2 = 1000 T$ and $Q_2 = 500 T$.

But
$$Q_1 = 1000 I_1$$
 and $Q_2 = 500 I_2$

:.500
$$T_2 = \frac{1}{2} \times 1000 T_1 i.e. T_2 = T_1$$

- **48.** (c) Enthalpy of a compound = Heat of reaction of that compound.
- **49.** (e) It is the definition of heat of neutralization.
- **50.** (b) $\Delta H = -ve$ for exothermic compound.
- (a) If acid or base or both are strong, heat of neutralization = 13.7 *kcal*.
- **52.** (b) Both NH_4OH and CH_3COOH are weak.
- **53.** (c) $57.1 \times 0.25 = 14.3 \ kJ \ mol^{-1}$.



54. (c) Heat of formation is for 1 mole.
Hence
$$\Delta H_f^0(HI) = -12.40/2 = -6.20 \, kcal.$$

55. (b) Chemical dissociations are reversible and
endothermic.
57. (c) Given,
(i) $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O$, $\Delta H = +20 \, kcal$
(ii) $C + O_2 \rightarrow CO_2$, $\Delta H = -40 \, kcal.$
(iii) $H_2 + \frac{1}{2}O_2 \rightarrow H_2O$, $\Delta H = -10 \, kcal.$
Aim: $C + 2H_2 \rightarrow CH_4$
(i) $+ 2 \times (iii) - (i)$ gives.
 $\Delta H = -40 + 2(-10) - (+20) = -80 \, kcal.$
58. (d) For exothermic reaction heat is evolved.
59. (c) $H_2O_{(f)} - \frac{electrolys is}{2} \rightarrow H_2(g) + \frac{1}{2}O_2(g).$
61. (d) All other are combustion phenomena.
62. (c) $\Delta H_{combusion} = -ve$, so exothermic process.
63. (b) Aim: $K_{(5)} + \frac{1}{2}O_2(g) + \frac{1}{2}H_2(g) \rightarrow KOH_{(5)}$
 $eq. (ii) + eq. (i) - eq. (iii) gives
 $\Delta H = -48 + (-68.39) - (-14) = -68.39 - 48 + 14.$
64. (a) By definition of heat of formation.
65. (c) $C_D \rightarrow C_G , \Delta H = -453.5 \, cal.$
i.e. energy of C_G is less and thus more stable.
69. (a) $\Delta H_{combusion}$ is always negative.
70. (d) $\Delta H_{reaction} = \Delta H_f^0 (N_2 O_4) - 2\Delta H_f^0 (NO_2)$
 $= 2 - 2(8) = -14 \, kcal.$
71. (a) $\Delta H - \Delta E = \Delta nRT = 3 \times 8.314 \times 298$
 $= -7432 \, J = -7.43 \, kJ.$
72. (b) Aim: $C_{(graphile)} \rightarrow C_{(diamond)}$; $eq.$ (i) – (ii) gives the result.
73. (d) In the formation of a compound, more the heat absorbed, less stable is the compound.
74. (c) $C_{(graphile)} + 2H_{2(g)} = CH_{4(g)}.$
75. (b) Heat of formation is the formation of one mole of the substance from its elements.
76. (b) $\Delta H_{(reaction)} = 2\Delta H_f^0 (M_g O) = -\Delta H_f^0 (SiO_2)$
 $= 2(-34.7) - (-48.4) = -21 \, kJ.$
77. (a) $eq.$ (i) $+ eq.$ (ii) $+ eq.$ (iii) gives $X = 131 - 282 - 242 = -393 \, kJ.$
78. (d) It is the definition of heat of transition.
79. (d) $C_{(graphile)} \rightarrow C_{(diamond)}, \Delta H = 1.9 \, kJ$
 $C_{(graphile)} + O_2 \rightarrow CO_2, \Delta H = -\Delta H_1$$

$$\begin{split} C_{\text{(graphite)}} + O_2 &\rightarrow CO_2, \Delta H = -\Delta H_1 \\ C_{\text{(diamond)}} + O_2 &\rightarrow CO_2, \Delta H = -\Delta H_2 \\ (-\Delta H_1) - (-\Delta H_2) = 1.9 \, kJ \text{ or } \Delta H_2 = \Delta H_1 + 1.9 \\ \text{For combustion of } 6g, \Delta H_2 &> \Delta H_1 \quad \text{by} \\ 1.9 / 2 = 0.95 \, kJ. \end{split}$$

- **80.** (a) $H_{\text{product}} \times H_{\text{reactant}}$ for exothermic reaction.
- **81.** (b) Due to high electron affinity of *Cl* the highest energy is evolved.

82. (b) $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O$ Molecular weight of $CH_4 = 12 + 4 = 16$ \therefore On the combustion of 2.0gm of methane= 25.0 kcal \therefore On the combustion of $16.0gm = \frac{25 \times 16}{2} = 200$ kcal.

- **83.** (a) $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O$.
- **84.** (b) $C_2H_4 + 3O_2 \rightarrow 2CO_2 + 2H_2O$

$$\Delta H_{\text{reaction}} = [2 \times \Delta H_f^o(CO_2) + 2 \times \Delta H_f^o(H_2O)]$$
$$-[\Delta H_f^o(C_2H_4) + 3 \times \Delta H_f^o(O_2)]$$

$$= [2(-394) + 2(-286)] - [52 + 0] = -1412 kJ.$$

85. (b) $C + O_2 \rightarrow CO_2 + 394 \, kJ$. $2C + 2O_2 \rightarrow 2CO_2 + 788 \, kJ$

$$2CO + O_2 \rightarrow 2CO_2 + 569 \, kJ$$
(ii)

....(i)

eq. (i) + eq. (iii) = $-109.5 \ kJ \ mol^{-1}$.

- **86.** (c) $\Delta H_f = \frac{44}{2} kcal = 22 kcal$.
- 87. (a) $CaO_{(s)} + H_2O_{(l)} \rightarrow Ca(OH)_2$ is exothermic.
- **88.** (b) $\Delta H = -ve$ in exothermic reaction.
- **89.** (d) According to the definition of heat of formation.
- **90.** (d) Multiplying *eq*. (ii) by (iii) and *eq*. (iii) by (vi), and then add

$$6C + 3H_2 + \frac{15}{2}O_2 \rightarrow 6CO_2 + 6H_2O; \Delta H = 3218.7$$

Subtract *eq*. (i) from the above equation and find the required result.

- **91.** (a) *HF* is more stable than *HCl*.
- **92.** (c) $\Delta H(H_P H_R) = q_p$.
- **93.** (b) $\Delta H = +ve$ for endothermic reaction.

94. (a)
$$\Delta H = \frac{13.95 \times 44}{2.2016} = 278.7 \ kcal$$
.

- **95.** (d) eq. (i) + eq. (ii) gives the required result.
- 96. (d) Standard molar heat enthalpy (H°) of a compound is equal to its standard heat of formation from most stable states of initial components.
- **97.** (c) In the complete combustion of butanol $\Delta H > \Delta E$.
- **98.** (a) X Y and find the required result.

99. (c)
$$S + O_2 \to SO_2$$
 $\Delta H = -298.2 \ kJ$ (i)

$$SO_2 + \frac{1}{2}O_2 \to SO_3 \quad \Delta H = -98.2 \ kJ \qquad \dots (ii)$$

eq. (i) – (ii) and find the required result.

100. (c) $C + 2S \rightarrow CS_2 \quad \Delta H = ?$

$$C + O_2 \rightarrow CO_2 \quad \Delta H = -393.3 \ kJ \qquad \dots (i)$$

$$S + O_2 \rightarrow SO_2 \quad \Delta H = -293.72 \ kJ \qquad \dots (ii)$$

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$$CS_2 + 3O_2 \rightarrow CO_2 + 2SO_2 \quad \Delta H = -1108.76 \, kJ \dots$$
 (iii)
eq. (ii) × 2 + eq. (i) - (iv) eq. (iii) - eq. (iv)
and find required result.

- **101.** (b) Fermentation is exothermic reaction.
- **102.** (c) $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O$.
- **103.** (d) Use $\Delta H = \Delta E + \Delta n RT$

$$\Delta H = 19 + 2 \times 2 \times 10^{-3} \times 300 = 20.2 \, kcal; \, \Delta n = 2.$$

- **104.** (d) $-20.6 8.8 \text{ KJ mol}^{-1} = -29.4 \text{ kJ}$.
- **106.** (a) Subtract equation (ii) from (i).
- **107.** (c) $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O, \Delta H = -890.4 \, kJ$ (i)

$$C + O_2 \rightarrow CO_2, \quad \Delta H = -395.5 \, kJ \qquad \dots (ii)$$

$$C + O_2 \rightarrow CO_2, \quad \Delta H = -395.5 \, kJ \qquad \dots \dots (11)$$

$$H_2 + \frac{1}{2}O_2 \to H_2O, \ \Delta H = -285.8 \, kJ$$
(iii)

From (1), (11), (11),

$$\Delta H_{comb.}(CH_4) = \Delta H_f(CO_2) + 2\Delta H_f(H_2O) - \Delta H_f(CH_4) - 2\Delta H_f(O_2) = -890.4 = -395.5 + 2(-285.5) - \Delta H_f(CH_4) - 2 \times 0$$

$$\Delta H_f(CH_4) = -76.7 \, kJ \, mol^{-1}$$

- 108. (d) As methanoic acid is weak acid, heat of neutralization < x.</p>
- **109.** (d) Neutralization of a strong acid by a strong base is always same.
- 110. (c) $S + O_2 \rightarrow SO_2$, $\Delta H_f = -4.6 \, kJ$

 \because 0.5 gm. of sulphur on burning produce 1gm of SO $_2$

 \therefore 32 gm of sulphur on burning produce 64 gm.. of SO $_2$

$$\therefore \Delta H = (-4.6 \, kJ) \times 64 = -294.4 \, kJ$$

111. (a) $3O_2 = 2O_3$ – energy is given out.

112. (b)
$$\Delta H \ per \ 1.6g = \frac{72 \times 1.6}{180} = 0.64 \ kcal$$
.
113. (a) $C + Q_2 \rightarrow CQ_2 \quad \Delta H = -394 \ kJ$

13. (a)
$$C + O_2 \to CO_2$$
, $\Delta H = -394 \, kJ$ (i)
 $2H_2 + O_2 \to 2H_2O$, $\Delta H = -568 \, kJ$ (ii)

$$CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O, \ \Delta H = -892 \ kJ \quad \dots (iii)$$

- (i) + (ii) (iii) and find the required result.
- 115. (a) No doubt heat evolved during neutralisation of $250 \text{ } cm^3$ of each acid and base is five time the heat evolved during neutralisation of $50 \text{ } cm^3$ of each acid and base but the quantity if solution taking heat is also five time thus same temperature rise is noticed.

116. (a)
$$H_2 + \frac{1}{2}O_2 \rightarrow H_2O_{(l)}; \ \Delta H = -285.8 \ KJ$$

 $H_2O_{(l)} \rightarrow H_2O_{(g)}; \ \Delta H = 44 \ KJ$
 $\therefore H_2 + \frac{1}{2}O_2 \rightarrow H_2O_{(g)}; \ \Delta H^o = -241.8 \ KJ$

117. (b) Given: Weight of iron burnt = 4g; Heat liberated = 29.28 *KJ* and atomic weight of iron (Fe) = 56. We know that in ferric oxide $(Fe_2O_3), 2 \text{ moles}$ of iron or $2 \times 56 = 112 \text{ gram}$ of iron are burnt. We also know that when 4grams of iron are burnt, then heat liberated = 29.28 *kJ*, therefore when 112 grams of the iron are burnt, then heat liberated $= \frac{29.28 \times 112}{4} = -819.8 \text{ kJ}$ (Minus sign due to liberation of heat).

118. (c)
$$H^+ + OH^- \rightarrow H_2O$$
, $\Delta H_{\text{neutralization}} = 13.7 \, kcal$.

119. (b)
$$\Delta H / mole$$
 of $FeS = \frac{3.77 \times 56}{2.1} = 100.5$.

- **120.** (a) Heat of formation $=\frac{194}{2} = 97 \, kJ$.
- **121.** (a) $\Delta H = \Delta H_{\text{ioniz}} + \Delta H_{\text{neu}} = -50.6 = \Delta H_{\text{ioniz}} + (-55.9)$ $\Delta H_{\text{ioniz}} = +5.3 \, kJ / mol$.
- **122.** (a) Strong acid (HNO_3) and strong base (LiOH).

123. (d)
$$\Delta H = \Delta E + \Delta nRT$$
 or $\Delta E = \Delta H - \Delta nRT$
 $\therefore \Delta E = +7.3 - \frac{1}{2} \times 0.002 \times 298 = 7.3 - 0.298 = 7$ kcal.

124. (b) Heat evolved during burning of 39g

$$C_6H_6 = \frac{781.0 \times 39}{78} = 390.5 \, kcal \, mol^{-1}$$

125. (b) By (i) + (ii) ;
$$Na + \frac{1}{2}Cl_2 \rightarrow NaCl, \Delta H = -196 \ kcal$$
.

126. (b)
$$C + O_2 \to CO_{2(g)}, \Delta H_f^o = -393.5 \, kJ \, mol^{-1}$$
(i)

$$C + \frac{1}{2}O_2 \to CO_{(g)}, \ \Delta H_f^o = -110.5 \, kJ \, mol^{-1}$$
(ii)

$$H_{2} + \frac{1}{2}O_{2} \rightarrow H_{2}O_{(g)}, \ \Delta H_{f}^{o} = -241.8 \ kJmol^{-1} \quad \dots (iii)$$

By (ii) + (iii) - (i)
$$CO_{2(g)} + H_{2(g)} \rightarrow CO_{(g)} + H_{2}O_{(g)}, \ \Delta H = +41.2.$$

127. (c) More +ve is ΔH_s more is heat of solution.

128. (d) CS_2 is formed from its initial components carbon and sulphur so, ΔH is heat of formation of CS_2 .

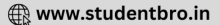
129. (d)
$$\Delta H = 18500 = \Delta E + \Delta nRT$$

or $18500 = \Delta E + (-1) \times 2 \times 298 = 19096 \ cal$.

- **130.** (b) Heat evolved during combustion of 0.39*g* $C_6H_6 = \frac{3250 \times 0.39}{78} = 16.25 \, kJ.$
- **131.** (d) By (i) (ii): $C + \frac{1}{2}O_2 \rightarrow CO; \Delta H = -229.2.$
- **132.** (b) Strong acid (*HCl*) and strong base (*NaOH*) shows $\Delta H_{\text{neutralization}} = -57.3 \text{ kJ}$.
- 133. (d) By (i) (ii) and find required result.

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134. (a)
$$CO_{(g)} + \frac{1}{2}O_{2(g)} \rightarrow CO_{(g)}, \Delta n = 1 - 1 + \frac{1}{2} = -\frac{1}{2}$$

 $\Delta H = \Delta E + \Delta nRT$
 $\Delta H = -283.3 - \frac{1}{2} \times \frac{8.314}{1000} \times 290 = -284.5kJ$.
135. (a) $H_2 + \frac{1}{2}O_2 \rightarrow H_2O$ $\Delta H = -188 \ kJ/mole$ (i)
 $H_2 + O_2 \rightarrow H_2O_2; \ \Delta H = -286 \ kJ/mole$ (ii)
By 2 × (i) and (ii)
 $2H_2 + O_2 \rightarrow 2H_2O; \ \Delta H = -376 \ kJ/mole$ (iii)
 $2H_2 + 2O_2 \rightarrow 2H_2O_2 \ \Delta H = -572 \ kJ/mole$ (iv)
By (iii) - (iv)
 $2H_2O_2 \rightarrow 2H_2O + O_2 \ \Delta H = +196 \ kJ$.
136. (a) ΔH for $C_2H_6 = -341.1K \ cal$
its calorific value $= \frac{-341.1}{30} = -11.37 \ kcal/g$.
 ΔH for $C_2H_2 = -310.0 \ kcal$
its Calorific value $= \frac{-310.0}{26} = -11.92$,
hence C_2H_2 is a better fuel.

138. (c) For the decomposition of 9gm of water heat required = 142.5 kJ

we know $H_2O = 2 + 16 = 18$

Therefore heat required for decomposition of 18gm water $=\frac{18}{9} \times 142.5 = 285$ *KJ*

Than, enthalpy of formation of water is reverse of heat required = -285 kJ.

139. (d)
$$C_6 H_{6(g)} + \frac{15}{2} O_{2(g)} \rightarrow 6CO_{2(g)} + 3H_2O_{(g)}$$

$$\Delta n = 6 + 3 - 1 - \frac{15}{2} = +\frac{1}{2}.$$

15

- **140.** (b) By $2 \times (i) + (ii) (iii)$ ΔH of methane = 20.0 kcal.
- 141. (c) Heat of neutralization of strong acid and weak base is less than 13.7 kcalmol⁻¹.
- **142.** (d) $\Delta E = 0$ for a cyclic process.
- **143.** (c) $22.0 \times 2 = 44 \ kcal$.
- 144. (b) 0.3 mole OH^- , neutralize 0.3 mole of HNO_3 Evolved heat, = $57.1 \times 0.3 = 17.13kJ$
- 146. (d) Compounds with high heat of formation are less stable because energy rich state leads to instability.

147. (b)
$$CH_4$$
 required = $\frac{445.15 \times 16}{890.3} = 8 gm$.

148. (c)
$$C_{(s)} + 2H_{2(g)} \rightarrow CH_{4(g)}$$
(i)

$$C_{(s)} + O_{2(g)} \to CO_{2(g)} \Delta H = -94 \ kcal \ mol^{-1}$$
(ii)

$$\begin{split} H_{2(g)} + \frac{1}{2} O_{2(g)} &\to H_2 O_{(l)} \quad \Delta H = -68 \ kcal \ mol^{-1} \dots \ \text{(iii)} \\ CH_4 + 3/_2 O_2 &\to CO_2 + 2H_2 O \end{split}$$

$$\Delta H = -213 \ k cal \ mol^{-1} \qquad \dots (iv)$$

to obtain equation (i) operate-(ii) + 2 \times (iii) – (iv).

- **149.** (b) $\Delta H = +ve$ for endothermic reactions.
- **150.** (b) It pertains to neutralization of strong acid weak base.
- **151.** (d) Strong base (KOH) and strong acid (HCl)
- **153.** (b) This reaction absorbed heat, so it is endothermic reaction.
- **154.** (b) $C + O_2 \rightarrow CO_2$, $\Delta H = -94.3$

This is also heat of formation of CO₂

$$C + \frac{1}{2}O_2 \rightarrow CO, \ \Delta H = -26.0$$
.

155. (b) $X = \frac{1}{2}Y$.

157. (a) Molecular weight of $NH_4NO_3 = 80$

Heat evolved $= 1.23 \times 6.12$

 \therefore Molar heat capacity = $1.23 \times 6.12 \times C$.

- **158.** (d) Both X_e and F_2 are gaseous elements at $25^{\circ}C$ and in their standard states and form $XeF_{4(g)}$ hence $\Delta H_{(f)}^{\circ} = \Delta H_{react}^{\circ}$.
- **159.** (c) According to Hess low, enthalpy change for a reaction does not depend on the nature of inter mediate reaction steps.
- **160.** (d) By (ii) (i), $C_{gr} \rightarrow C_{dia}$, $\Delta H = +1.9$.
- **162.** (c) Heat of neutralization of strong acid and strong base is equal to the $-57.32 \text{ KJ mol}^{-1}$

163. (c)
$$C + O_2 \rightarrow CO_2 \Delta H = -393.5 \text{ KJ} / \text{mol}$$

:: 44 gm of CO_2 form by which heat released

= -393.5 kJ

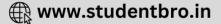
 \therefore 1gm of CO_2 form by which heat released 393.5

 $\therefore~35.2\,gm$ (given) of CO_2 form by which heat released

$$=-\frac{393.5}{44} \times 35.2 = -315 \ kJ$$

- **164.** (c) $C_{(graphite)} + 2H_2(g) \rightarrow CH_4(g)$
- **165.** (c) Heat of formation of a substance is the heat exchanged when one mole of that substance is formed by its constituent elements.

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: Heat evolved when 1 mole $(44g)CO_2$ is formed = 393 kJ

 $\therefore \qquad \text{Heat evolved when } 0.156 \, Kg(156 \, g) \text{ is}$ formed = $\frac{393 \times 156}{44}$

- $\therefore \Delta H$ for the process = 1572 kJ = -1572.0kJ
- **166.** (b) It is heat of neutralisation (13.7 *Kcal*) for strong acid and strong base.
- **167.** (d) $C(s) + O_2(g) \to CO_2(g)$

 $\Delta H = -393.5 \text{ KJ mol}^{-1}$ (i) $CO(g) + 1/2O_2(g) \rightarrow CO_2(g)$

$$\Delta H = -283 \text{ KJ mol}^{-1} \qquad \dots \dots (ii)$$

On substracting equation (ii) from equation

We get

(i)

$$C(s) + O_2(g) \rightarrow CO(g); \Delta H = -110.5 \text{ KJ mol}^{-1}$$

The enthalpy of formation of carbon monooxide per mole = $-110.5 KJ mol^{-1}$

168. (d) 1 mole (i.e.,) 16 gm of methane on combustion liberate 890 kJ

:
$$3.2 \, gm$$
 will liberate $=\frac{890 \times 3.2}{16} = 178 \, KJ$

- **169.** (c) $C + O_2 \rightarrow CO_2$; $\Delta H = q$
 - $C+1/2O_2 \rightarrow CO$; $\Delta H = -12$ (i)

$$CO + 1/2O_2 \rightarrow CO_2$$
; $\Delta H = -10$ (ii)

adding equation (i) and (ii) we can get $\Delta H = -12 + (-10) = -22$

170. (b)
$$C + O_2 \to CO_2$$
; $\Delta H_F = -490 \text{ KJ/mol}$ - (I)

$$H_2 + 1/2O_2 \rightarrow H_2O; \Delta H_F = -240 \text{ KJ/mol} - (II)$$

$$8C + 18H \rightarrow C_8H_{18}; \Delta H_F = +160 \text{ KJ/mol}$$
 - (III)

applying

(I) ×8 + (II) ×9 + (III)

$$C_8 H_{18} + \frac{25}{2} O_2 \rightarrow 8CO_2 + 9H_2O$$

 $\Delta H^o = -3920 - 2160 - 160 = 6240 \ KJ / mol$
 $\Delta H^o = \text{ for 6 moles of octane } = 6240 \times 6$
 $= 37440 \ KJ / mol = -37.4 \ KJ$

171. (d)
$$2Fe + \frac{3}{2}O_2 \rightarrow Fe_2O_3; \Delta H = -193.4 \, kJ$$
(i)

$$Mg + \frac{1}{2}O_2 \rightarrow MgO; \Delta H = -140.2 \, kJ$$
(ii)

On multiplying eq. (ii) by 3

$$3Mg + \frac{3}{2}O_2 \rightarrow 3MgO; \Delta H = -420.6 \, kJ \qquad \dots \dots (iiii)$$

Resulting equation can be obtained by subtracting eq. (i) from (iii)

$$3Mg + \frac{3}{2}O_2 \rightarrow 3MgO; \Delta H = -420.6 \, kJ$$
$$2Fe + \frac{3}{2}O_2 \rightarrow Fe_2O_3; \Delta H = -193.4 \, kJ$$
Subtraction :

$$3Mg + Fe_2O_3 \rightarrow 2Fe + 3MgO; \Delta H = -227.2 kJ$$

172. (c) HCl and KOH both are strong

Bond energy

1. **(b)** $XY \longrightarrow X_{(g)} + Y_{(g)}$; $\Delta H = +a \ kJ \ mole$ (i) $X_2 \longrightarrow 2X$; $\Delta H = +a \ kJ \ mole$

$$Y_2 \longrightarrow 2Y$$
; $\Delta H = +0.5 a \ kJ \ / \ mole$(iii)

$$\frac{1}{2} \times (ii) + \frac{1}{2} \times (iii) - (i), \text{ gives}$$
$$\frac{1}{2} X_2 + \frac{1}{2} Y_2 \longrightarrow XY \text{ ;}$$
$$\Delta H = \left(+ \frac{a}{2} + \frac{0.5}{2} a - a \right) kJ / mole$$
$$+ \frac{a}{2} + \frac{0.5a}{2} - a = -200$$
$$a = 800$$

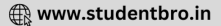
- 2. (a) $4gH_2 = 2$ moles. Bond energy for 1 mole of $H_2 = 208 / 2 = 104$ kcal.
- 4. (d) By definition of Hess's law.

5. (c) Aim:
$$\frac{1}{2}H_2 + \frac{1}{2}Cl_2 \rightarrow HCl$$

 $\Delta H = \sum B.E.(\text{Products}) - \sum B.E.(\text{Reactants})$
 $= B.E.(HCl) - \left[\frac{1}{2}B.E.(H_2) + \frac{1}{2}B.E.(Cl_2)\right]$
 $= -103 - \left[\frac{1}{2}(-104) + \frac{1}{2}(-58)\right]$
 $= -103 - (-52 - 29) = -22 \text{ kcal}$.

6. (c) First ionization potential of
$$Li = 5.4eV$$

Electron affinity of $Cl = 3.61eV$
We have $\Delta H = I.P. - E.A.$
 $= 5.4 - 3.61 = 1.80eV = 1.80 \times 1.6 \times 10^{-22} kJ$



$$\Delta H = 2.86 \times 10^{-22} kJ,$$

for 1mol Avogadro's number = 6.02×10^{23} .
 $\Delta H = 2.86 \times 10^{-22} \times 6.02 \times 10^{23} = 170 kJ/mole.$
7. (b) $\frac{-166}{4} = -41.5 kJ/mole$.
8. (d) $\frac{1}{2}H_2 + \frac{1}{2}Cl_2 \rightarrow HCl, \ \Delta H = -90 KJ$
 $\therefore \ \Delta H = \frac{1}{2}E_{H-H} + \frac{1}{2}E_{Cl-Cl}$
or $-90 = \frac{1}{2} \times 430 + \frac{1}{2} \times 240 - E_{HCl}$
 $\therefore E_{H-Cl} = 425 kJ mol^{-1}.$
9. (b) $CH_4 \rightarrow C + 4H, \ \Delta H = 320$
 $E_{C-H} = 90 cal$
 $C_2H_6 \rightarrow 2C + 6H, \ \Delta H = 360$
 $\therefore 360 = E_{C-C} + 6E_{C-H}$
 $\therefore E_{C-C} = 360 - 320 = 40 cal.$
10. (b) $H - H + Br - Br \rightarrow 2H - Br$
 $433 + 192$ 2×364
 625 728
Energy absorbed = Energy released
Net energy released = 728 - 625 = 103 kJ
i.e., $= \Delta H = -103 KJ$

Free energy and Work function

- **1.** (d) ΔG at equilibrium = 0.
- **2.** (b) For spontaneous change $\Delta G = -ve$.
- **3.** (a) When $\Delta G = -ve$ than the reaction is spontaneous in nature.
- **5.** (acd)When $\Delta H = +ve$ and $\Delta S = -ve$ than the reaction is non-spontaneous.
- 7. (c) Because the reaction is spontaneous .
- **8.** (c) $\Delta G = -ve$.

9. (a)
$$\Delta G_T = nRT \ln \frac{P_2}{P_1}$$
.

- 10. (a) $\Delta G = 0$ for equilibrium.
- **11.** (d) At equilibrium $\Delta G = 0$.
- **12.** (a) $\Delta G^o = -2.303 RT \log K$

 $-4.606 = -2.303 \times 0.002 \times 500 \log K$

 $\log K = 2, K = 100$.

14. (d) Spontaneous change shows $\Delta G = -ve$.

15. (b)
$$\Delta G = \Delta H - T \Delta S$$
, $T = 25 + 273 = 298 K$

$$=-11.7 \times 10^{3} - 298 \times (-105) = 19590 J = 19.59 kJ$$

18. (c) If $\Delta G = -ve$ reaction is spontaneous.

- **20.** (b) $(dS)_{V,E} > 0, (dG)_{T,P} < 0$.
- **21.** (b) Only high pressure favours the conversion.
- **22.** (d) By 2× (ii) (i) (iii)

$$H_{2(g)} + 2C_{(s)} \rightarrow C_2 H_{2(g)}, \ \Delta G^{\circ} = 209 \ kJ.$$

- 23. (b) $\Delta G = -2.303 \text{ RT} \log K'$, Here R = 2 cal, T = 300 K $K' = \frac{10 \times 15}{3 \times 5} = 10$; $\Delta G = -2.303 \times 2 \times 300 \times \log_{10} 10$ $= -2.303 \times 2 \times 300 \times 1 = -1381.8 \text{ cal}$
- **24.** (a) $\Delta G = \Delta H T \Delta S$ (Gibb's free energy equation)

25. (d)
$$\Delta G = \Delta H - T \Delta S$$

 $\Delta G = -382.64 - (-145.6) \times 10^{-3} \times 298$
 $= -339.3 \text{ KJ mol}^{-1}$

26. (d) $\Delta G = \Delta H - T\Delta S$; $\Delta G = \Delta E + P\Delta V - T\Delta S$ For spontaneity $(\Delta G = -ve)$ $\therefore \Delta G = -ve$, $\Delta E = +ve$

27. (a)
$$\Delta G = \Delta H - T\Delta S = 31400 - 1273 \times 32$$

= 31400 - 40736 = -9336 cal

28. (d)
$$\Delta G = \Delta H - T\Delta S$$

 $0 = +30.558 - T \times 0.066$
or $T = \frac{30.558}{0.066} = 463 K$

2

If $(dG)_{T,P} = 0$ sign '=' mean. If is reversible process

Critical Thinking Questions

- (b) Due to randomness of particles is reduced since entropy decreases.
- **2.** (c) $\Delta H = nCp \Delta T$

The process is isothermal therefore $\Delta G = 0$; $\therefore \Delta H = 0$

- **3.** (d) 13.7 $kcal = 57 kJ = 5.7 \times 10^4 J$.
- (b) When strong acid and strong base neutralize each other than the value of heat generated is about 13.7 *kcal*.
- **5.** (b) Due to fall in temperature.
- **6.** (d) $q_p = \Delta H$.
- 7. (d) $w = 2.303 RT \log \frac{V_2}{V_1}$. As it involves the ratio $\frac{V_2}{V_1}$, volume can be used in any units. (R should be in joules).

8. (b)
$$-\frac{dQ}{dW} = \frac{dQ}{dQ - dE}$$
; $dE = dW + dQ$; $-dW = dQ - dE$
$$= \frac{nc_p dT}{nc_p dT - nc_y dT} = \frac{c_p}{(c_p - c_y)}$$

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 $= \frac{7R}{2R} \{ \text{for diatomic of gas } c_p = \frac{7R}{2} \}$ (a) $C_2H_2(g) + H_2(g) \rightarrow C_2H_4(g)$

$$\Delta ng = 1 - 2 = -1$$
; $\Delta H = -0.31 \, KJ \, mol^{-1}$

P = 1.5 atm, $\Delta V = -50 mL = -0.050 L$

 $\Delta H = \Delta E + P \Delta V$

9.

 $-0.31 = \Delta E - 0.0076$; $\Delta E = -0.3024 \ KJ$

- (a) Since process is exothermic then heat is evolved due to this temperature of water increases.
- 11. (a) Because randomness is decreases.
- 12. (c) Conversion of graphite into diamond is an endothermic reaction. So, heat of diamond is higher than that of graphite. But ΔS would be negative for the conversion of graphite into diamond.
- **13.** (c) ΔH and ΔS both are +ve for spontaneous change, and $\Delta H = +ve$ for endothermic reaction.
- **14.** (a) $\Delta G = \Delta H T \Delta S$ is negative for spontaneity.
- 15. (a) Entropy (a measure of disorder) of universe is increasing toward maximum. This is II law of thermodynamics.
- **16.** (a) For a pure substance T_A and T_B represent the same temperature. Hence A is a correct choice.
- 17. (a) For endothermic reaction enthalpy of products (H_P) > enthalpy of reactant (H_R)

Hence, change in enthalpy,

 $\Delta H = H_P - H_R = \text{Positive}$

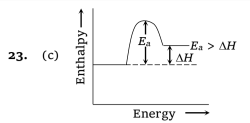
- 18. (d) Suppose the no. of gm, 1gm for each case so the max. no of mole expected for SO₂(g) which has to expected maximum entropy.
- **19.** (b) $2Al + \frac{3}{2}O_2 \rightarrow Al_2O_3$, $\Delta H = -1596 \ kJ$ (i)

$$2Cr + \frac{3}{2}O_2 \to Cr_2O_3, \ \Delta H = -1134 \ kJ \qquad \dots (ii)$$

By (i) - (ii)

 $2Al + Cr_2O_3 \rightarrow 2Cr + Al_2O_3, \Delta H = -462 kJ.$

- **20.** (a) Bomb calorimeter is used to measure heat of reaction.
- **21.** (b) Larger the number of *C* and *H* atoms, greater is the heat of combustion.
- **22.** (c) The bomb calorimeter is used to measure ΔE .



- **24.** (b) $\Delta H_{\text{Neutralization}} = -57.1 kJ = -13.7 kcal$.
- **25.** (b) Backmann thermometer is used to measure low temperature.
- 26. (b) Heat required to rise the temperature of a body by 1K called thermal capacity of the body.
- **27.** (e) Mechanical work is important only in gases as they undergo appreciable change in volume.
- 28. (d) It is Lavoisier and Laplace law. This is another law of thermochemistry which was put forward before Hess's law.
- 29. (a) It is also known as "constant heat summation" law. It state that the total amount of heat evolved or absorbed in a reaction is same whether the reaction takes place in one step or in a number of steps.

30. (c) In this reaction
$$\Delta n = 2 - 4 = -2$$
 so $\Delta H \neq \Delta E$.

31. (a)
$$Q = \frac{18.94 \times 0.632 \times 0.998 \times 122}{1.89} = 771.1 Kcall$$

- **32.** (c) $\Delta H = E_a$ for forward reaction $-E_a$ for backward reaction $= 19 9 = 10 \, kJ$.
- **33.** (d) $Q = 18.94 \times 0.632 \times 0.998 \times 1000$

$$-57.4 + x = -12.13$$

 $x = 45.2$

34. (c) Thermodynamics is a reversible process in which surroundings are always equilibrium with system.

35. (a)
$$1 \, cal = 4.18 \, J = 4.18 \times 10^7 \, erg = \frac{4.18}{1.602} \times 10^{19} \, eV$$
.

- **36.** (b) Order of bond energy : $Cl_2 > Br_2 > F_2 > I_2$.
- **37.** (c) Heat energy is also involved when one allotropic form of an element is converted in to another. graphite is the stabler allotrope because the heat of transformation of $C_{(diamond)} \rightarrow C_{(graphile)}$.

(i)
$$C_{(dia)} + O_{2(g)} = CO_{2(g)}\Delta H = -94.5kcal$$

(ii)
$$C_{(graphite)} + O_{2(g)} = CO_{2(g)}\Delta H = -94.0 k cal$$

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$$\Delta H_{transformation} = -94.5 - (-94.0)$$

 $= -0.5k \, cal$.

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- **38.** (b) The compound *Y* possess less energy than the *X* and thus *Y* is more stable than *X*.
- **39.** (c) Both ΔH and Δp are positive.
- **40.** (a) For equilibrium $\Delta G^o = 0$.
- **41.** (b) Equilibrium constant is decreases with temperature and ΔH is also decreases so it is *-ve*.
- **42.** (a) The fusion temperature of napthalene is minimum, because it is non-polar covalent compound and has less fusion temperature.
- **43.** (a) The bond energy of C H bond is y kcal mol^{-1} .

44. (a)
$$N \equiv N + 3H - H \longrightarrow 2N - H$$

Energy absorbed $U = 2(3 \times 391) = 2346$
Energy released

Net. energy released = 2346 - 2253 = 93 kJ

77

i.e.
$$\Delta H = -93 \ kJ$$
.

- **45.** (d) If E = +ve than the cell reaction will be spontaneous.
- **46.** (d) All are the correct statements.
- **47.** (d) $CH_2 = CH_{2(g)} + H_{2(g)} \rightarrow H_3C CH_{3(g)}$ $414 \times 4 = 1656$ $414 \times 6 = 2484$ $615 \times 1 = 615$ $347 \times 1 = 347$ $435 \times 1 = \frac{435}{2706}$ 2831 $\Delta H = 2706 - 2831 = -125 \, kJ$
- **48.** (b) It means that the heat absorbed when one gram molecule of HCl is formed from its elements at 298 K is
 - $= 22.060 \, k \, cal$.
- **49.** (d) *eq*. (i) + *eq*. (ii) find the required result and divide by 2.
- **50.** (d) For complete neutralization of strong acid and strong base energy released is 57.32 *KJ*/*mol*

No. of mole of
$$H_2SO_4 = \frac{0.2 \times 50}{1000} = 10^{-2}$$

No. of mole of
$$KOH = \frac{1}{1000} \times 50 = 5 \times 10^{-2}$$

So =
$$57.32 \times 10^{-2} = 0.5732 \text{ KJ} = 573.2 \text{ Joule}$$
.

51. (b) For Combustion reaction, ΔH is negative, $\Delta n = (16 + 18) - (25 + 2) = +7$, so ΔS is +ve, reaction is spontaneous, hence ΔG is -ve. **52.** (a) $\Delta G = \Delta H - T \Delta S$, T = 27 + 273 = 300 K

 $\Delta G = (-285.8) - (300)(-0.163) = -236.9 \, kJ \, mol^{-1}$

Assertion & Reason

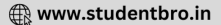
- 1. (b) Assertion and reason show, reaction at equilibrium state in which $\Delta G = 0$, $\Delta S = 0$, $\Delta H = 0$.
- 2. (b) It is correct that on touching the ice we feel cold because ice absorb heat from our hand.
- 3. (b) Entropy of ice is less than water because water molecules in solid state lose kinetic energy and hence their tendency of movement minimise. Hence entropy decrease in solid state. The reason that ice have cage like structure is also correct but reason is not or correct explanation for assertion.
- (c) During isothermal expansion of an ideal gas against vacuum is zero because expansion is isothermal. The reason, that volume occupied by the molecules of an ideal gas is zero, is false.
- 5. (a) it is fact that absolute values of internal energy of substances can not be determined. It is also true that to determine exact values of constituent energies of the substance is impossible.
- 6. (b) Mass and volume are extensive properties. mass/volume is also an extensive parameter. Here, both assertion and reason are true.
- 7. (b) The molar entropy of vaporization of water it differ from ethanol due to hydrogen bonding according to VSEPR theory water molecule having two lone pair of electron by which it angular and show some polarity which is higher than that of ethanol so both assertion and reason are correct but reason is not explaination assertion.
- 8. (a) The assertion that the increase in internal energy for the vaporisation of one mole of water at 1 *atm* and 373 *K* is zero is true and this is because that for all isothermal process internal energy is zero.
- **9.** (b) $N_2(g) + O_2(g) \Rightarrow 2NO(g)$

We know that $\Delta H = \Delta E + \Delta nRT$

 $\Delta n = 2 - 2 = 0$ hence, $\Delta nRT = 0$

Therefore, $\Delta H = \Delta E$

10. (a) The enthalpies of neutralisation of strong acids and strong bases are same they are 13.7



Kcal. The reason is that it is heat of formation of water from H^+ and OH^- ions. $H^+ + OH^- \rightarrow H_2O, \Delta H = 13.7 \ Kcal$. Thus, both assertion and reason are true.

- 11. (a) Zeroth (law of temperature) can also be summarized as two objects at different temperature in thermal contact with each other tend to move towards the same temperature.
- **12.** (d) Photo chemical reactions have $+ve \Delta G$.

If a system undergoes a change in which internal energy of the system remains constant (*i.e.*, $\Delta E = 0$), then -W = Q.

This means that work done by the system equals the heat absorbed by the system.

- **20.** (e) If a refrigerator's door is kept open room gets heated as heat energy is released to the surroundings.
- 21. (c) Enthalpy is zero but entropy is not zero.Vibrational motion exists even at absolute zero.
- 22. (c) It may involve increase or decrease in temperature of the system. Systems in which such process occur, are thermally insulated from the surroundings.

- **14.** (e) Heat absorbed in a reaction at constant temperature and constant volume (Qv) = E.
- (c) Values of state functions depend only on the state of the system and not on how it is reached.
- 16. (a) The properties whose magnitude depends upon the quantity of matter present in the system are called extensive properties.
- 17. (c) Combustion reactions are always accompanied by the evolution of heat therefore, for such reactions the value of ΔH is always negative.
- 18. (b) In an isothermal process change in internal energy (ΔE) is zero (as it is a fucntion of temperature). ∴ According to first law of thermodynamics

 $\therefore q + w = \Delta E$, hence Q = -W (if $\Delta E = 0$)

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