

### Course Objectives

The course enables the students to

1. Explain the nature of the electrochemical terms and concepts
2. Understand the nature of electrochemical reactions
3. Understand the phase equilibrium.
4. Understand different electrochemical cells and EMF measurements.
5. Understand the fundamentals of surface chemistry

### Course outcome

The course enabled the students to

1. Understand the nature of the electrochemical terms and concepts
2. Explain the nature of electrochemical reactions
3. Understand the phase equilibrium.
4. Differentiate different electrochemical cells and able to do EMF measurements.
5. Understand the fundamentals of surface chemistry

### UNIT I

**Phase Equilibria:** Concept of phases, components and degrees of freedom, derivation of Gibbs Phase Rule for nonreactive and reactive systems; Clausius-Clapeyron equation and its applications to solid-liquid, liquid-vapour and solid-vapour equilibria, phase diagram for one component systems ( $H_2O$  and S), with applications. Phase diagrams for systems of solid-liquid equilibria involving eutectic, congruent and incongruent melting points.

### UNIT II

**Three component systems:** triangular plots, water-chloroform-acetic acid system. Binary solutions: Gibbs-Duhem-Margules equation, its derivation and applications to fractional distillation of binary miscible liquids (ideal and non ideal), azeotropes, lever rule, partial miscibility of liquids, CST, miscible pairs, steam distillation. Nernst distribution law: its derivation and applications.

### UNIT III

**Electrochemical Cells:** Rules of oxidation/reduction of ions based on half-cell potentials, applications of electrolysis in metallurgy and industry. Chemical cells, reversible and irreversible cells with examples. Electromotive force of a cell and its measurement, Nernst equation; Standard electrode (reduction) potential and its application to different kinds of half-cells.

## UNIT IV

### Application of EMF measurements

Application of EMF measurements in determining (i) free energy, enthalpy and entropy of a cell reaction, (ii) equilibrium constants, and (iii) pH values, using hydrogen, quinone-hydroquinone, glass and  $\text{SbO/Sb}_2\text{O}_3$  electrodes. Concentration cells with and without transference, liquid junction potential; determination of activity coefficients and transference numbers. Qualitative discussion of potentiometric titrations (acid-base, redox, precipitation).

## UNIT V

**Surface chemistry:** Physical adsorption, chemisorption, adsorption isotherms (Langmuir and Freundlich). nature of adsorbed state. Qualitative discussion of BET.

### Suggested Readings:

#### Text Books:

1. Peter Atkins & Julio De Paula. (2010). *Physical Chemistry*. 9th Ed. Oxford University Press.
2. Castellan, G. W. (2004). *Physical Chemistry*. 4th Ed. Narosa
3. McQuarrie, D. A. & Simon, J. D. (2004). *Molecular Thermodynamics*. New Delhi : Viva Books Pvt. Ltd. • Engel, T. & Reid, P. (2012). *Physical Chemistry*. 3rd Ed. Prentice-Hall

#### Reference Books

1. Assael, M. J., Goodwin, A. R. H., Stamatoudis, M., Wakeham, W. A. & Will, S. (2011). *Commonly Asked Questions in Thermodynamics*. NY : CRC Press.
2. Zundhal, S.S. (2011). *Chemistry concepts and applications*. Cengage India • Ball, D. W. (2012). *Physical Chemistry*. Cengage India.
3. Mortimer, R. G. (2009). *Physical Chemistry*. 3rd Ed. Elsevier: NOIDA, UP.
4. Levine, I. N. (2011). *Physical Chemistry*. 6th Ed. Tata McGraw-Hill.
5. Metz, C. R. (2009). *Physical Chemistry*. 2nd Ed. Tata McGraw-Hill.

## UNIT I

### Phase Equilibria

**Phase Equilibria:** Concept of phases, components and degrees of freedom, derivation of Gibbs phase Rule for nonreactive and reactive systems: Clausius-Claypeyron equation and its applications to solid-liquid, liquid-vapour and solid-vapour equilibria, Phase diagram for one component systems ( $H_2O$  and S), with applications. Phase diagrams for systems of solid-liquid equilibria involving eutectic, congruent and incongruent melting points.

### PHASE EQUILIBRIA

Gibbs' phase rule was proposed by Josiah Willard Gibbs in his landmark paper titled On the Equilibrium of Heterogeneous Substances, published from 1875 to 1878. The rule applies to non-reactive multi-component heterogeneous systems in thermodynamic equilibrium and is given by the equality

$$F = C - P + 1$$

where F is the number of degrees of freedom, C is the number of components and P is the number of phases in thermodynamic equilibrium with each other.

The number of degrees of freedom is the number of independent intensive variables, i.e. the largest number of thermodynamic parameters such as temperature or pressure that can be varied simultaneously and arbitrarily without affecting one another. An example of one-component system is a system involving one pure chemical, while two-component systems, such as mixtures of water and ethanol, have two chemically independent components, and so on. Typical phases are solids, liquids and gases.

**Phase:** It is defined as any homogeneous, physically distinct and mechanically separable portion of a system, which is separated from other such parts of the system by definite boundary surfaces in a system.

**Component-** It is defined as the smallest number of independently variable constituents taking part in the state of equilibrium by means of which the composition of each phase can be expressed directly or in the form of chemical equation.

**Degree of freedom-** It is defined as the minimum number of the independently variable factors such as the temperature, pressure and composition of the phases which must be arbitrarily specified in order to represent perfectly the condition of a system.

A phase is a form of matter that is homogeneous in chemical composition and physical state. Typical phases are solid, liquid and gas. Two immiscible liquids (or liquid mixtures with different compositions) separated by a distinct boundary are counted as two different phases, as are two immiscible solids.

The number of components (C) is the number of chemically independent constituents of the system, i.e. the minimum number of independent species necessary to define the composition of all phases of the system.<sup>[2]</sup> For examples see component (thermodynamics).

The number of degrees of freedom (F) in this context is the number of intensive variables which are independent of each other.

The basis for the rule (Atkins and de Paula,<sup>[2]</sup> justification 6.1) is that equilibrium between phases places a constraint on the intensive variables. More rigorously, since the phases are in thermodynamic equilibrium with each other, the chemical potentials of the phases must be equal. The number of equality relationships determines the number of degrees of freedom. For example, if the chemical potentials of a liquid and of its vapour depend on temperature (T) and pressure (p), the equality of chemical potentials will mean that each of those variables will be

dependent on the other. Mathematically, the equation  $\mu_{\text{liq}}(T, p) = \mu_{\text{vap}}(T, p)$ , where  $\mu$  = chemical potential, defines temperature as a function of pressure or vice versa. (Caution: do not confuse  $p$  = pressure with  $P$  = number of phases.)

To be more specific, the composition of each phase is determined by  $C - 1$  intensive variables (such as mole fractions) in each phase. The total number of variables is  $(C - 1)P + 2$ , where the extra two are temperature  $T$  and pressure  $p$ . The number of constraints is  $C(P - 1)$ , since the chemical potential of each component must be equal in all phases. Subtract the number of constraints from the number of variables to obtain the number of degrees of freedom as

$$F = (C - 1)P + 2 - C(P - 1) = C - P + 2.$$

The rule is valid provided the equilibrium between phases is not influenced by gravitational, electrical or magnetic forces, or by surface area, and only by temperature, pressure, and concentration.

## CLAUSIUS CLAPEYRON EQUATION

The vaporization curves of most liquids have similar shape. The vapour pressure steadily increase as the temperature increases. A good approach is to find a mathematical model for the pressure increase as a function of temperature. Experiments showed that the pressure  $P$ , enthalpy of vaporization,  $\Delta H_{\text{vap}}$ , and temperature  $T$  are related,

$$P = A \exp (- \Delta H_{\text{vap}} / R T)$$

where  $R (= 8.3145 \text{ J mol}^{-1} \text{ K}^{-1})$  and  $A$  are the gas constant and unknown constant. This is known as the Clausius- Clapeyron equation. If  $P_1$  and  $P_2$  are the pressures at two temperatures  $T_1$  and  $T_2$ , the equation has the form:

$$\ln \frac{P_1}{P_2} = \frac{\Delta H_{\text{vap}}}{R} \left( \frac{1}{T_2} - \frac{1}{T_1} \right)$$

The Clausius-Clapeyron equation allows us to estimate the vapor pressure at another temperature, if the vapor pressure is known at some temperature, and if the enthalpy of vaporization is known.

## Phase diagram for water

Water is a unique substance in many ways. One of these special properties is the fact that solid water (ice) is less dense than liquid water just above the freezing point. The phase diagram for water is shown in the Figure below .

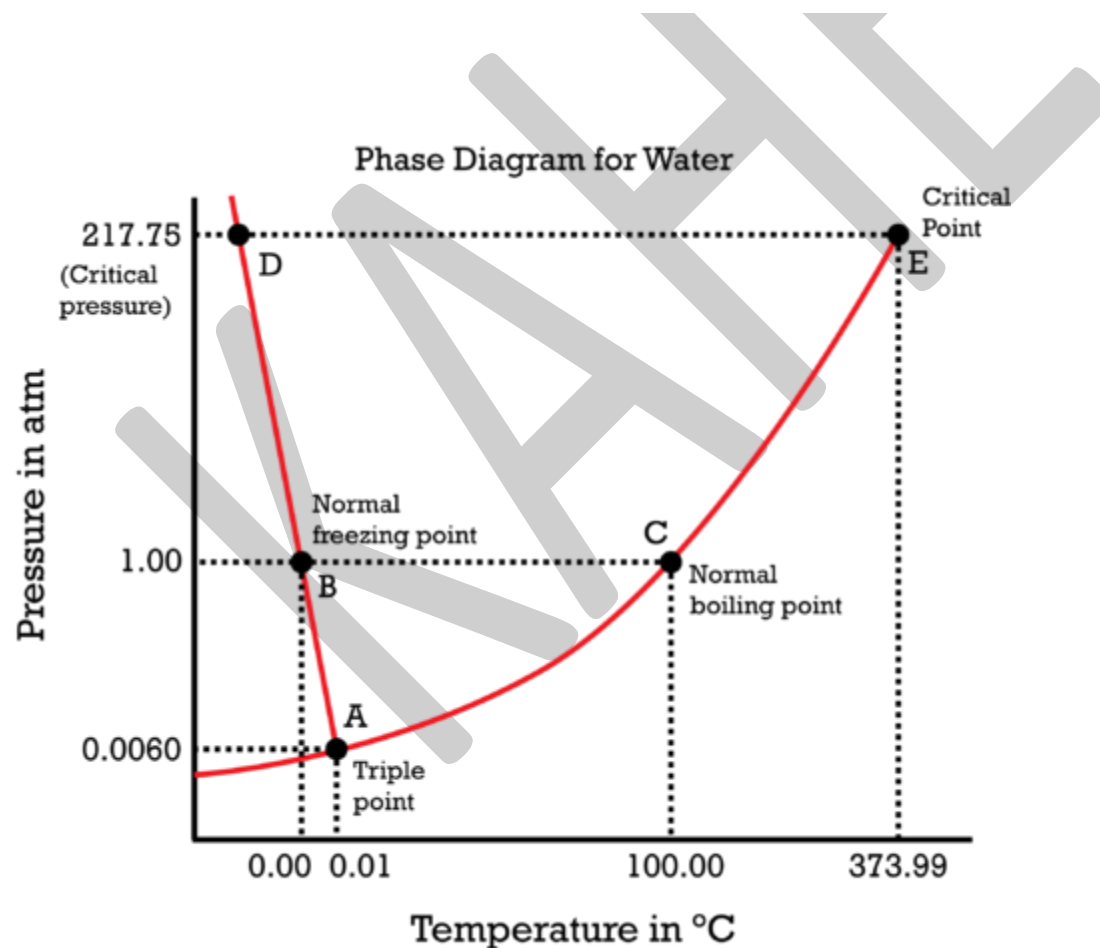


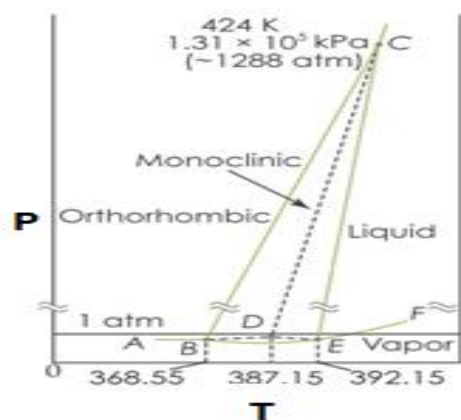
Figure 1

Notice one key difference between the general phase diagram and the phase diagram for water. In water's diagram, the slope of the line between the solid and liquid states is negative rather than positive. The reason is that water is an unusual substance in that its solid state is less dense than the liquid state. Ice floats in liquid water. Therefore, a pressure change has the opposite effect on those two phases. If ice is relatively near its melting point, it can be changed into liquid water by the application of pressure. The water molecules are actually closer together in the liquid phase than they are in the solid phase.

The point E in the above phase diagram labeled the critical point. At 373.99°C, particles of water in the gas phase are moving very, very rapidly. At any temperature higher than that, the gas phase cannot be made to liquefy, no matter how much pressure is applied to the gas. The critical pressure ( $P_c$ ) is the pressure that must be applied to the gas at the critical temperature in order to turn it into a liquid. For water, the critical pressure is very high, 217.75 atm. The critical point is the intersection point of the critical temperature and the critical pressure.

## PHASE DIAGRAM OF SULPHUR

Phase Diagram of Sulphur In the following example, we will use the phase diagram of sulphur in the form of overhead projector (OHPI) transparencies to illustrate such overlapping. Fig.6.1 shows the phase diagram of sulphur pressure. Except that a different presentation of the lines (eg. the dotted lines, the continuous lines and a solid dark line) is used to show the three fundamental parts of this phase diagram.



The curves AB, BC, CD, BE, CE, EG divide the diagram into four areas. Curve AB, the Vapour Pressure curve of SR. It shows the vapour pressure of solid rhombic sulphur (SR) at different temperatures. Along this curve the two phases SR and sulphur vapour (SV) are in equilibrium. The system SR/SV has one degree of freedom,  $F = C - P + 2 = 1 - 2 + 2 = 1$  i.e., it is monovariant.

Curve BC, the Vapour Pressure curve of SM. It shows variation of the vapour pressure of monoclinic sulphur (SM) with temperature. SM and SV coexist in equilibrium along this curve. The system SM/SV is monovariant. Curve CD, the Vapour Pressure curve of SL. It depicts the variation of the vapour pressure of liquid sulphur (SL) with temperature. SL and SV are in equilibrium along CD. The two phase system SL/SV is monovariant. One atmosphere line meets this curve at a temperature (444.6°C) which is the boiling point of sulphur. Curve BE, the Transition curve. It shows the effect of pressure on the transition temperature for SR and SM. As two solid phases are in equilibrium along the curve, the system SR/SM is monovariant. The transformation of SR and SM is accompanied by increase of volume (density of SR = 2.04; SM = 1.9) and absorption of heat i.e.,  $SR + Q \text{ (heat energy)} \rightarrow SM$ . Thus the increase of pressure will shift the equilibrium to the left (Le Chatelier's Principle) and the transition temperature will, therefore, be raised. This is why the line BE slopes away from the pressure axis showing thereby that the transition temperature is raised with increase of pressure. Curve CE, the

Fusion curve of SM. It represents the effect of pressure on the melting point of SM. The two phases in equilibrium along this curve are SM and SL. The system SM/SL is monovariant. As the melting or fusion of SM is accompanied by a slight increase of volume, the melting point will rise by increase of pressure (Le Chatelier's principle). Thus the curve CE slopes slightly away from the pressure axis. The curve ends at E because SM ceases to exist beyond this point. Curve EG, the Fusion curve for SR. Here the two phases in equilibrium are SR and SL. The number of phases being two, the system SR/SL is monovariant.

The Triple points B, C, E Triple point B. This is the meeting point of the three curves AB, BC and BE. Three phases, solid SR, solid SM and SV are in equilibrium at the point B. There being three phases and one component, the system SR/SM/SL is nonvariant.  $F = C - P + 2 = 1 - 3 + 2 = 0$  At B, SR is changed to SM and the process is reversible. Thus the temperature corresponding to B is the transition temperature (95.6°C). Triple point C. The curves BC, CD, CE meet at this point. The three phases in equilibrium are SM, SL and SV. There being three phases and one component, the system SM/SL/SV is nonvariant. The temperature corresponding to C as indicated on the phase diagram is 120°C. This is the melting point of SM. Triple point E. The two lines CE and BE, having different inclinations away from the pressure axis, meet at E where a third line EG also joins. The three phases SR, SM and SL are in equilibrium and the system at the point E is nonvariant. This point gives the conditions of existence of the system SR/SM/SL at 155°C and 1290 atmospheres pressure. (3) The Areas The phase diagram of the sulphur system has four areas or regions. These are labelled as rhombic sulphur, monoclinic sulphur, liquid sulphur and vapour. These represent single phase systems which have two degrees of freedom,  $F = C - P + 2 = 1 - 1 + 2 = 2$  That is, each of the systems SR, SM, SL, and SV are bivariant.

Metastable Equilibria The change of SR to SM takes place very slowly. If enough time for the change is not allowed and SR is heated rapidly, it is possible to pass well above the transition point without getting SM. In that case, there being three phases (SR, SL, SV) only and one component, the phase diagram, like that of water system, will consist of three curves, one triple

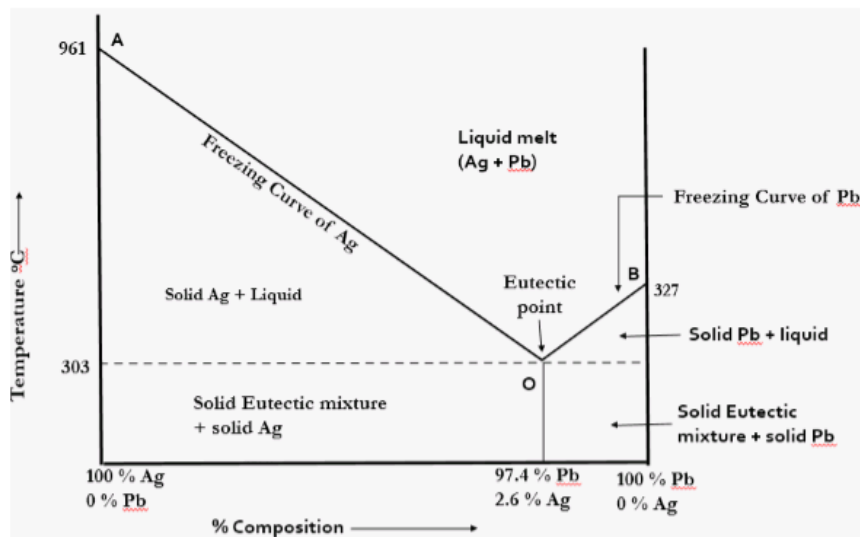
point and three areas. The dashed curve BF, the Vapour Pressure curve of metastable SR. This is a continuation of the vapour pressure curve AB of stable SR. The metastable phases SR and SV are in equilibrium along this curve. It is a monovariant system. The dashed curve CF, the Vapour Pressure curve of supercooled SL. On supercooling liquid sulphur, the dashed curve CF is obtained. It is, in fact, the back prolongation of DC. The curve CF represents the metastable equilibrium between supercooled SL and SV. Thus it may be designated as the vapour pressure curve of supercooled SL. It meets the dashed curve BF at F. The dashed curve FE, the Fusion curve of metastable SR. The two metastable phases SR and SL are in equilibrium along this curve and the system is monovariant. This shows that the melting point of metastable SR is increased with pressure. Beyond E, this curve depicts the conditions for the stable equilibrium SR/SL as the metastable SR disappears. The metastable Triple point F. At this point, three metastable phases SR, SL and SV are in equilibrium. The system is a metastable triple point with no degree of freedom. The corresponding temperature is the melting point of metastable SR (114°C)

Phase diagrams for systems of solid liquid equilibria involving eutectic points.

## Lead-Silver system

**TWO-COMPONENT SYSTEMS** When a single phase is present in a two-component system, the degree of freedom is three,  $F = 2 - 1 + 2 = 3$  This means that three variables must be specified in order to describe the condition of the phase.

## Phase diagram of Ag-Pb system



In two component systems there are four possible phases solid Ag, solid Pb, solution of Ag, + Pb and vapour. Since the pressure has no effect on equilibrium so the system can be represented by temperature concentration diagram at constant atmospheric pressure. As pressure is neglected the phase rule is called condensed phase rule.

- 1) Curve AO.** It is a freezing point curve of Ag. Ag Co exists as solid and liquid. Melting point of Ag falls gradually on adding Pb till the lowest point is reached. The solution gets saturated with respect to lead.
- 2) Curve BO.** It is a freezing point curve of Pb. At this curve the melting point gradually falls on the addition of Ag till lowest point it reach.
- 3) Point O.** It is eutectic point. Here 3 phases co-exists and point O represents a fixed composition and system is in variant.

Below the temperature line of eutectic temperature, we have two regions.

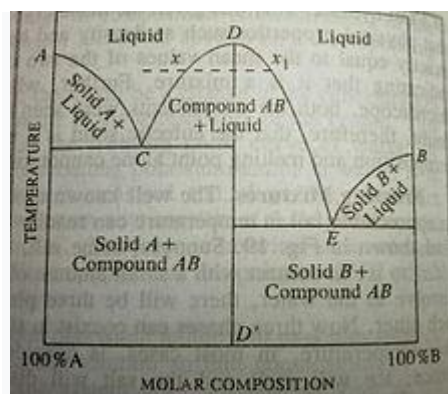
a) The region marked eutectic plus solid Ag in which crystalline silver and solid eutectic are stable.

b) The region marked eutectic plus solid Pb in which crystalline lead and solid eutectic are stable.

**4) Area AOB.** It represents solution of Pb Ag. On lowering temperature the lead begins to separate out till the point O is reached.

## Phase diagrams for systems of congruent melting point

Congruent melting occurs during melting of a compound when the composition of the liquid that forms is the same as the composition of the solid. It can be contrasted with incongruent melting. This generally happens in two- component systems. To take a general case, let A and B be the two components and AB a stable solid compound formed by their chemical combination. If we draw a phase diagram for the system, we notice that there are three solid phases, namely A, B and compound AB. Accordingly, there will be three fusion or freezing point curves AC, BE and CDE for the three solid phases. In the phase diagram, we can notice that the top point D of the phase diagram is the congruent melting point of the compound AB because the solid and liquid phases now have the same composition. Evidently, at this temperature, the two-component system has become a one-component system because both solid and liquid phases contains only the compound AB.



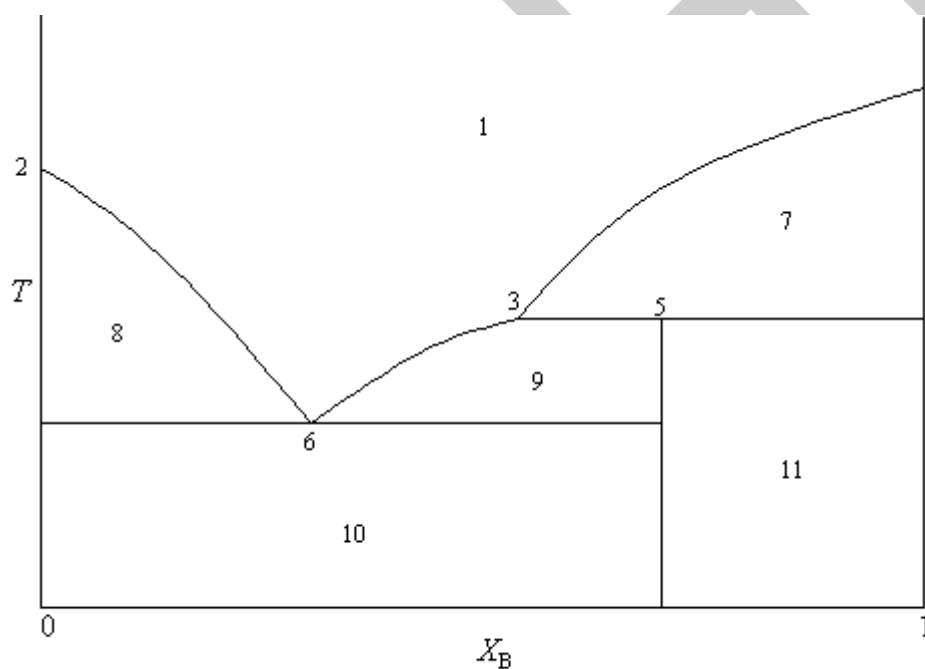
## Phase Diagram for the Formation of a compound with congruent Melting Point

Congruent melting point represents a definite temperature just like the melting points of pure components. In the phase diagram, the congruent melting point D of compound AB lies above the melting points of pure components A and B. But it is not necessarily true. There are different types of systems known in which the congruent melting point is observed to be less than melting points of pure components.

This happens for inter-metallic compounds. For example, MgSi

## Phase diagram for system of incongruent melting point

Incongruent Melting Point (melting with decomposition)



This phase diagram shows an incongruent melting point. The vertical line at point 5 represents formation of a compound. It looks like the composition of the compound is  $X_B = 2/3$ , from which we conclude using the previous arguments,

$$X_B = \frac{2}{3} = \frac{n_B}{n_A + n_B} = \frac{2}{1 + 2},$$

so that the compound is  $AB_2$ . Point 5 is the melting point of  $AB_2$ , but notice that melting  $AB_2$  does not give liquid of the same composition. Rather, melting of  $AB_2$  gives liquid with the composition at point 3 and pure B(s). So the compound,  $AB_2$ , melts and decomposes at the same time. An analysis of the points and regions is:

## POSSIBLE QUESTIONS

### Part B

1. Define Phase.
2. Define Component.
3. Define Degree of freedom.
4. State reduced phase rule.
5. What is meant by triple point?
6. What is meant by eutectic point?
7. What is meant by congruent melting point?
8. What is meant by incongruent melting point?

### Part C

9. What is phase rule. Define the terms phase, component and degrees of freedom with suitable examples.
10. Explain the phase diagram of water system.
11. Derive Clausius-clapeyron equation and apply to solid-liquid, liquid-vapour and solid-vapour equilibria.
12. Explain simple eutectic system by taking lead-silver as an example
13. Explain incongruent melting point by taking sodium-potassium system.
14. Explain congruent melting point by taking Zn-Mg alloy system.

**CLASS: II- B.SC CHEMISTRY**

**COURSE NAME: PHYSICAL CHEMISTRY III**

**COURSE CODE:18CHU301**

**UNIT-I**

**BATCH: 2018-2021**



| S.No | Question                                                                                                                    | Option 1      | Option 2      | Option 3            | Option 4       | Answer        |
|------|-----------------------------------------------------------------------------------------------------------------------------|---------------|---------------|---------------------|----------------|---------------|
| 1    | J.W.Gibbs enunciated the phase rule in                                                                                      | 1876          | 1875          | 1874                | 1873           | 1876          |
| 2    | A gas mixture constitutes                                                                                                   | single phase  | two phase     | three phase         | four phase     | single phase  |
| 3    | CCl <sub>4</sub> and water forms                                                                                            | three phase   | two phase     | four phase          | single phase   | two phase     |
| 4    | Benzene and alcohol constitutes                                                                                             | single phase  | two phase     | three phase         | four phase     | single phase  |
| 5    | Calcium component is an example for                                                                                         | one component | two component | three component     | four component | two component |
| 6    | Sulphur exist in                                                                                                            | three phase   | one phase     | two phase           | four phase     | four phase    |
| 7    | Sugar and water is and example for                                                                                          | two component | one component | three component     | four component | two component |
| 8    | In a chemically reactive system, the number of components is given by                                                       | $C=N-m-N-R$   | $C=n-M-N-r$   | $C=N-m-n-R$         | $C=R-N-n-m$    | $C=N-m-n-R$   |
| 9    | If $F=0$ , the system is called                                                                                             | bivariant     | univariant    | invariant           | trivariant     | invariant     |
| 10   | If $F=1$ , the system is called                                                                                             | bivariant     | univariant    | invariant           | trivariant     | univariant    |
| 11   | If $F=2$ , the system is called                                                                                             | bivariant     | univariant    | invariant           | trivariant     | bivariant     |
| 12   | The greater the number of components in a system, the ----- is the number of degrees of freedom for a given number of phase | greater       | lower         | greater and lower   | constant       | greater       |
| 13   | The greater the number of phases in a system, the ----- is the number of degrees of freedom for a given number of phase     | greater       | smaller       | greater and smaller | constant       | smaller       |
| 14   | For a given number of components, the number of phases is ----- when the                                                    | maximum       | minimum       | constant            | maximum and    | maximum       |

|    |                                                                                                                                  |                           |                         |                           |                     |                         |
|----|----------------------------------------------------------------------------------------------------------------------------------|---------------------------|-------------------------|---------------------------|---------------------|-------------------------|
|    | number of degrees of freedom is zero                                                                                             |                           |                         |                           | minimum             |                         |
| 15 | All the phases must be at the ----- temperature otherwise there will be flow of heat from one phase to another                   | greater                   | same                    | lower                     | same or lower       | same                    |
| 16 | All the phases must be at the ----- pressure otherwise the volume of one phase will increase at the expense of another           | greater                   | same                    | lower                     | same or lower       | same                    |
| 17 | Gibbs phase rule equation for one component system is                                                                            | $F=C-P+2$                 | $F=C-P+1$               | $F=C-P$                   | $F=P-C+2$           | $F=C-P+2$               |
| 18 | Gibbs phase rule equation for two component system is                                                                            | $F=C-P+2$                 | $F=C-P+1$               | $F=C-P$                   | $F=P-C+2$           | $F=C-P+1$               |
| 19 | Water is an example for                                                                                                          | three component           | two component           | one component             | four component      | one component           |
| 20 | Sulphur is an example for                                                                                                        | four component            | two component           | three component           | one component       | one component           |
| 21 | The temperature at which all the three phases will be equilibrium is known as                                                    | eutectic point            | congruent melting point | incongruent melting point | triple point        | triple point            |
| 22 | The temperature at which two solid and one liquid phase are in equilibrium is known as                                           | incongruent melting point | congruent melting point | triple point              | eutectic point      | eutectic point          |
| 23 | Carbon di-oxide is an example for                                                                                                | three component           | two component           | one component             | four component      | one component           |
| 24 | Reduced phase rule is otherwise known as                                                                                         | condensed phase rule      | phase rule equation     | Gibbs phase rule equation | expanded phase rule | condensed phase rule    |
| 25 | Lead-Silver is an example for                                                                                                    | three component           | two component           | one component             | four component      | two component           |
| 26 | A compound which melts sharply at a constant temperature into a liquid of the same composition as the solid is said to possess a | incongruent melting point | congruent melting point | d)triple point            | eutectic point      | congruent melting point |

|    |                                                                                                                                                                                                                                                          |                           |                         |                           |                             |                           |
|----|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------|-------------------------|---------------------------|-----------------------------|---------------------------|
| 27 | Ferric chloride -water is an example for                                                                                                                                                                                                                 | congruent melting point   | triple point            | incongruent melting point | eutectic point              | congruent melting point   |
| 28 | The compounds formed by the combination of two components, instead of melting congruently, decompose when heated giving a new solid phase and a solution with a composition different from that of the solid phase. Such a compound is said to possess a | incongruent melting point | congruent melting point | triple point              | eutectic point              | incongruent melting point |
| 29 | Transition temperature is otherwise known as                                                                                                                                                                                                             | peritectic temperature    | melting temperature     | sublimation temperature   | vapour pressure temperature | peritectic temperature    |
| 30 | Phase rule becomes -----for a three component system                                                                                                                                                                                                     | $F=5-P$                   | $F=4-P$                 | $F=3-P$                   | $F=2-P$                     | $F=5-P$                   |
| 31 | Isothermal critical point is otherwise known as                                                                                                                                                                                                          | plait point               | melting point           | sublimation point         | plait and melting point     | plait point               |
| 32 | Acetic acid-chloroform and water is an example for                                                                                                                                                                                                       | three component           | two component           | one component             | four component              | three component           |
| 33 | An equation of fundamental importance which finds extensive application in one-component, two phase systems was derived independently by                                                                                                                 | Clausius                  | Einstein                | Gibbs                     | Henry                       | Clausius                  |
| 34 | By supplying heat infinitesimally slowly to the system, it is possible to change any desired amount of the substance from the-----at the same temperature and pressure                                                                                   | liquid to solid           | solid to vapour         | liquid to vapour          | solid to liquid             | liquid to vapour          |
| 35 | G is known as                                                                                                                                                                                                                                            | enthalpy                  | entropy                 | free energy               | work                        | free energy               |
| 36 | Clapeyron equation gives change in pressure $dP$ which accompany the change in-----                                                                                                                                                                      | temperature               | volume                  | composition               | temperature and composition | temperature               |

|    |                                                                                                       |                           |                            |                      |                        |                            |
|----|-------------------------------------------------------------------------------------------------------|---------------------------|----------------------------|----------------------|------------------------|----------------------------|
| 37 | $\Delta H_v$ is known as                                                                              | molar heat of sublimation | molar heat of vaporisation | molar heat of fusion | molar heat of enthalpy | molar heat of vaporisation |
| 38 | $\Delta H_f$ is known as                                                                              | molar heat of sublimation | molar heat of vaporisation | molar heat of fusion | molar heat of enthalpy | molar heat of fusion       |
| 39 | $\Delta S$ is known as                                                                                | change in free energy     | change in entropy          | change in enthalpy   | change in energy       | change in entropy          |
| 40 | Lewis introduce the concept of                                                                        | fugacity                  | volume                     | pressure             | temperature            | fugacity                   |
| 41 | Calculate F in water $\leftrightarrow$ vapour equilibria                                              | 1                         | 2                          | 3                    | 4                      | 1                          |
| 42 | Calculate F in solid $\leftrightarrow$ liquid equilibria                                              | 2                         | 1                          | 3                    | 4                      | 1                          |
| 43 | Calculate F in solid $\leftrightarrow$ vapour equilibria                                              | 3                         | 5                          | 4                    | 1                      | 1                          |
| 44 | calculate F in ice $\leftrightarrow$ liquid $\leftrightarrow$ vapour equilibria                       | 0                         | 1                          | 2                    | 4                      | 0                          |
| 45 | Pressure-temperature axis is drawn for-----<br>-----                                                  | two component             | one component              | three component      | one and two component  | one component              |
| 46 | Temperature-composition axis is drawn for                                                             | two component             | one component              | three component      | one and two component  | two component              |
| 47 | The simplest three component systems are those in which a liquid system breaks down into ----- phases | one                       | two                        | three                | four                   | two                        |
| 48 | Application of the phase rule to a systme corresponding to a point in the two phase region gives $F=$ | 3                         | 2                          | 1                    | 0                      | 3                          |
| 49 | Tie lines are drawn in                                                                                | one component             | two component              | three component      | four component         | three component            |
| 50 | A system of two salts and water furnishes an example of a                                             | two component             | three component            | one component        | four component         | three component            |

|    |                                                                                                             |                           |                         |                          |                             |                 |
|----|-------------------------------------------------------------------------------------------------------------|---------------------------|-------------------------|--------------------------|-----------------------------|-----------------|
| 51 | Bismuth-cadmium is an example for                                                                           | incongruent melting point | congruent melting point | eutectic system          | triple point                | eutectic system |
| 52 | Potassium iodide-water is an example for                                                                    | eutectic system           | one component           | three component          | four component              | eutectic system |
| 53 | The well known observation that the addition of salt to ice produces an appreciable fall in -----           | temperature               | pressure                | pressure and temperature | composition                 | temperature     |
| 54 | On adding increasing amounts of ferric chloride in ferric chloride -water system there will be fall in----- | pressure                  | temperature             | composition              | temperature and composition | temperature     |
| 55 | $S_1 \leftrightarrow S_2 + \text{-----}$                                                                    | solution                  | $S_3$                   | $S_2$                    | $S_1 + S_2$                 | solution        |
| 56 | Number of phases in water $\leftrightarrow$ solid is                                                        | 1                         | 2                       | 3                        | 4                           | 2               |
| 57 | Number of phases in water $\leftrightarrow$ vapour is                                                       | 1                         | 2                       | 3                        | 4                           | 2               |
| 58 | Number of phases in solid $\leftrightarrow$ vapour is                                                       | 2                         | 3                       | 1                        | 0                           | 1               |
| 59 | Number of component in NaCl and water is                                                                    | 1                         | 2                       | 3                        | 4                           | 2               |
| 60 | Number of component in alcohol and water is                                                                 | 2                         | 1                       | 3                        | 4                           | 2               |

## UNIT II

### Three Component System

Three Component Systems: Triangular plots, water-chloroform-acetic acid system. Binary solutions: Gibbs-Duhem-Margules equation, its derivation and applications to fractional distillation of binary miscible liquids (ideal and non ideal), azeotropes, lever rule, partial miscibility of liquids, CST, miscible pairs, steam distillation, Nernst distribution law: its derivation and applications

#### Three component systems

The system acetic acid - chloroform - water is one of the classical examples of partial miscibility in a ternary system, first studied by Wright, Thompson, and Leon in 1891 (1). Thanks to the subsequent workers Brancker, Hunter, and Nash (2), the equilibrium relations at 25° are very well known, as far as the compositions of equilibrium liquid layers are concerned. Nothing, however, is known about: (a) the solid-liquid equilibria at low temperatures; (b) equilibrium liquid compositions at temperatures other than 25°; (c) the critical phenomena, L1-V (L1 = chloroform layer) and L2-V (L2 = aqueous layer). So far no study has been made of critical phenomena in any system exhibiting partial miscibility in the liquid state and the results arrived at in this paper have general applicability to all systems of this kind. A previous paper (3) has dealt with the thermodynamics of this system, as exemplified by the vapor pressures and densities

Because of the chemical natures of the substances named in the title, formation of compounds or of solid solutions is inherently improbable and therefore the study of the freezing point curves was expected to yield little of interest. The study was, however, carried out for the binary systems chloroform - acetic acid and chloroform - water, and for the ternary system, using a double-junction copper-constantan thermocouple, a Brown Elektronik recorder, and liquid nitrogen as a coolant. The system acetic acid - water was previously studied by various workers (4) and the eutectic found to lie at 58.1% acetic acid and -28.5 to -27.0°. The eutectic of the

chloroform - acetic acid system was found to lie at 91.8% CHCl<sub>3</sub> by weight, and -67.5° (Fig. 1). In the system chloroform-water, the eutectic lies at 0.1% H<sub>2</sub>O and -64.0°. The ternary eutectic was found to lie at -70° and 90.7% chloroform, 8.3% acetic acid, and 1.0% water. The eutectic trough leading from the acetic acid -water eutectic to the acetic acid -chloroform eutectic was determined experimentally.

## Gibbs-Duhem –Margules equation

The Gibbs free energy can be defined in two different ways once by subtracting off combinations of entropy S, enthalpy H and temperature T and other as a sum of chemical potentials and amounts of species. The fact that they are equal gives a new relation known as “Gibbs-Duhem Relation.” The Gibbs-Duhem relation helps us to calculate relationships between quantities as a system which remains in equilibrium. One example is the Clausius-Clapeyron equation which states that two phases at equilibrium with each other having equal amount of a given substance must have exactly the same free energy i.e. it relates equilibrium changes in pressure to changes in temperature as a function of material parameters.

Deriving the Gibbs-Duhem equation from thermodynamics state equations is very easy. The Gibbs free energy G in equilibrium can be expressed in terms of thermodynamics as:

$$dG = \mu_1 dn_1 + n_1 d\mu_1 + \mu_2 dn_2 + n_2 d\mu_2 + \dots + \mu_j dn_j + n_j d\mu_j$$

$$= (\mu_1 dn_1 + \mu_2 dn_2 + \dots + \mu_j dn_j) + (n_1 d\mu_1 + n_2 d\mu_2 + \dots + n_j d\mu_j)$$

At constant temperature and pressure, the above equation can be written as:

$$n_1 d\mu_1 + n_2 d\mu_2 + \dots + n_j d\mu_j = 0$$

$$\sum n_i d\mu_i = 0 \dots \dots \dots (1)$$

Because at constant temperature and pressure,  $(\mu_1 dn_1 + \mu_2 dn_2 + \dots + \mu_j dn_j) = dG$

The equation (1) is known as the Gibbs-Duhem equation.

## Applications of Gibbs-Duhem equation:

(i) Gibbs-duhem equation is helpful in calculating partial molar quantity of a binary mixture by measuring the composition of the mixture which depends on the total molar quantity.

(ii) Gibbs-duhem equation is helpful in calculating the partial vapor pressures by calculating the total vapor pressure. All these calculations require a curve-fitting procedure. Using tabulated experimental data the accuracy of the calculated quantities was found to be comparable to the accuracy of the original experimental data.

## Lever rule

The **lever rule** is a tool used to determine weight percentages of each phase of a binary equilibrium phase diagram. It is used to determine the percent weight of liquid and solid phases for a given binary composition and temperature that is between the liquidus and solidus line.

In an alloy with two phases,  $\alpha$  and  $\beta$ , which themselves contain two elements, A and B, the lever rule states that the weight percentage of the  $\alpha$  phase is

$$X_{\alpha} = \frac{c-b}{a-b}$$

where

- a is the weight percentage of element B in the  $\alpha$  phase
- b is the weight percentage of element B in the  $\beta$  phase
- c is the weight percentage of element B in the entire alloy

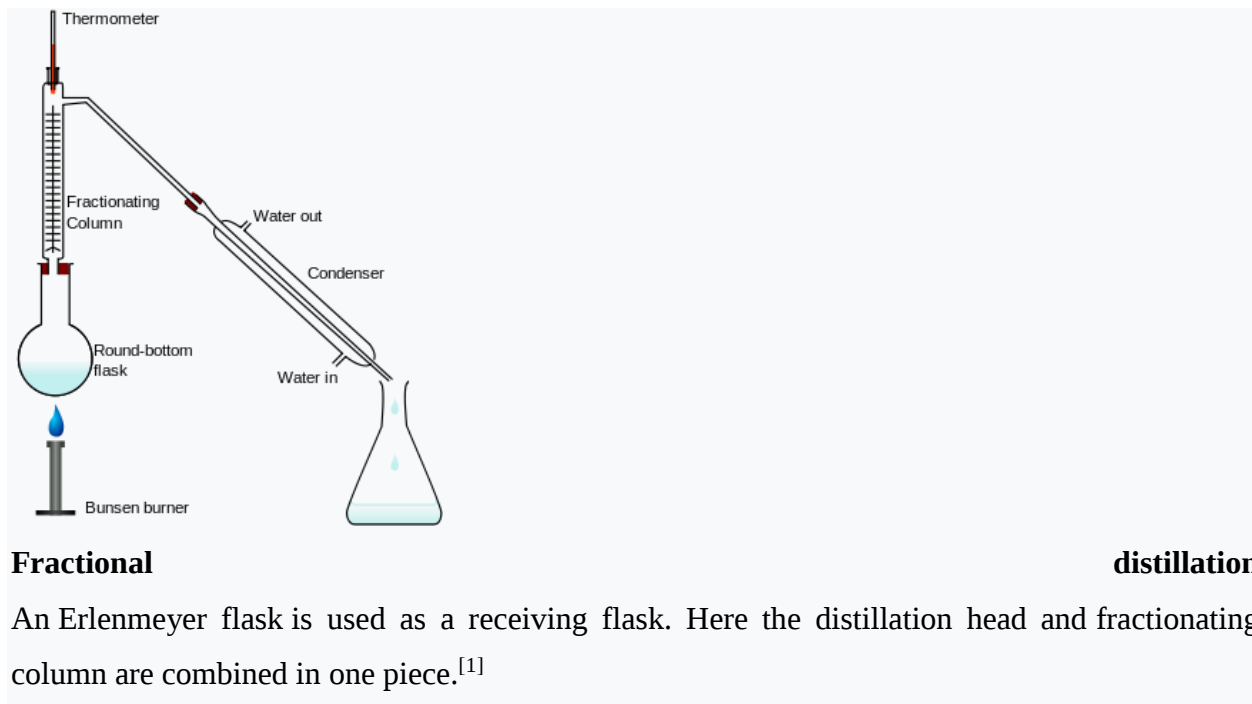
all at some fixed temperature.

## Fractional distillation of binary miscible liquids

**Fractional distillation** is the separation of a mixture into its component parts, or fractions. Chemical compounds are separated by heating them to a temperature at which one or more fractions of the compound will vaporize. It uses distillation to fractionate. Generally the component parts have boiling points that differ by less than 25 °C from each other under a pressure of one atmosphere. If the difference in boiling points is greater than 25 °C, a simple distillation is typically used.

Fractional distillation in a laboratory makes use of common laboratory glassware and apparatuses, typically including a Bunsen burner, a round-bottomed flask and a condenser, as well as the single-purpose fractionating column.

### Apparatus



- heat source, such as a hot plate with a bath, and ideally with a magnetic stirrer.
- distilling flask, typically a round-bottom flask
- receiving flask, often also a round-bottom flask
- fractionating column
- distillation head
- thermometer and adapter if needed
- condenser, such as a Liebig condenser or Allihn condenser
- vacuum adapter (not used in image to the right)
- boiling chips, also known as anti-bumping granules
- Standard laboratory glassware with ground glass joints, e.g. quickfit apparatus.

As an example consider the distillation of a mixture of water and ethanol. Ethanol boils at 78.4 °C while water boils at 100 °C. So, by heating the mixture, the most volatile component (ethanol) will concentrate to a greater degree in the vapor leaving the liquid. Some mixtures form azeotropes, where the mixture boils at a lower temperature than either component. In this example, a mixture of 96% ethanol and 4% water boils at 78.2 °C; the mixture is more volatile than pure ethanol. For this reason, ethanol cannot be completely purified by direct fractional distillation of ethanol-water mixtures.

The apparatus is assembled as in the diagram. (The diagram represents a batch apparatus as opposed to a continuous apparatus.) The mixture is put into the round bottomed flask along with a few anti-bumping granules (or a Teflon coated magnetic stirrer bar if using magnetic stirring), and the fractionating column is fitted into the top. The fractional distillation column is set up with the heat source at the bottom on the still pot. As the distance from the stillpot increases, a temperature gradient is formed in the column; it is coolest at the top and hottest at the bottom. As the mixed vapor ascends the temperature gradient, some of the vapor condenses and revaporizes along the temperature gradient. Each time the vapor condenses and vaporizes, the composition of the more volatile component in the vapor increases. This distills the vapor along the length of the column, and eventually the vapor is composed solely of the more volatile component (or an azeotrope). The vapor condenses on the glass platforms, known as trays, inside the column, and runs back down into the liquid below, refluxing distillate. The efficiency in terms of the amount of heating and time required to get fractionation can be improved by insulating the outside of the column in an insulator such as wool, aluminium foil or preferably a vacuum jacket. The hottest tray is at the bottom and the coolest is at the top. At steady state conditions, the vapor and liquid on each tray are at *equilibrium*. The most volatile component of the mixture exits as a gas at the top of the column. The vapor at the top of the column then passes into the condenser, which cools it down until it liquefies. The separation is more pure with the addition of more trays (to a practical limitation of heat, flow, etc.) Initially, the condensate will be close to the azeotropic composition, but when much of the ethanol has been drawn off, the condensate becomes

gradually richer in water. The process continues until all the ethanol boils out of the mixture. This point can be recognized by the sharp rise in temperature shown on the thermometer.

The above explanation reflects the theoretical way fractionation works. Normal laboratory fractionation columns will be simple glass tubes (often vacuum-jacketed, and sometimes internally silvered) filled with a packing, often small glass helices of 4 to 7 mm diameter. Such a column can be calibrated by the distillation of a known mixture system to quantify the column in terms of number of theoretical trays. To improve fractionation the apparatus is set up to return condensate to the column by the use of some sort of reflux splitter (reflux wire, gage, Magnetic swinging bucket, etc.) - a typical careful fractionation would employ a reflux ratio of around 4:1 (4 parts returned condensate to 1 part condensate take off).

In laboratory distillation, several types of condensers are commonly found. The Liebig condenser is simply a straight tube within a water jacket, and is the simplest (and relatively least expensive) form of condenser. The Graham condenser is a spiral tube within a water jacket, and the Allihn condenser has a series of large and small constrictions on the inside tube, each increasing the surface area upon which the vapor constituents may condense.

Alternate set-ups may use a multi-outlet distillation receiver flask (referred to as a "cow" or "pig") to connect three or four receiving flasks to the condenser. By turning the cow or pig, the distillates can be channeled into any chosen receiver. Because the receiver does not have to be removed and replaced during the distillation process, this type of apparatus is useful when distilling under an inert atmosphere for air-sensitive chemicals or at reduced pressure. A Perkin triangle is an alternative apparatus often used in these situations because it allows isolation of the receiver from the rest of the system, but does require removing and reattaching a single receiver for each fraction.

Vacuum distillation systems operate at reduced pressure, thereby lowering the boiling points of the materials. Anti-bumping granules, however, become ineffective at reduced pressures.

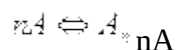
## **Nernst distribution law**

At constant temperature, a solute distributes itself between two immiscible solvents only in a particular ratio. This statement is a Nernst Distribution Law – i.e. the law that determines the relative distribution of a component that is soluble in two liquids, these liquids being immiscible or miscible to a limited extent. This law is one of the laws applying to ideal dilute solutions. It was discovered by W. Nernst in 1890. The Nernst distribution law states that, at equilibrium, the ratio of the concentrations of a third component in two liquid phases is constant. The law may be expressed in the form

$$c_1/c_2 = k$$

where  $c_1$  and  $c_2$  are the molar equilibrium concentrations of the third component in the first and second phase, respectively; the constant  $k$  is the distribution coefficient, which is temperature dependent.

The Nernst distribution law permits us to determine the most favorable conditions for the extraction of substances from solutions. If the dissolved compound in one of the solvents can associate:



than the ratio  $c_1/c_2$  is **not** stable at constant temperature

## Partial miscibility of liquids

Only pairs of liquids that are completely miscible have been considered so far. Many pairs of liquids, however, are only partially miscible in one another, the degree of miscibility often depending strongly on temperature. In most cases, rising temperature produces enhanced solubility, but this is not always so. For example, at 50° C the solubility (weight percent) of *n*-butyl alcohol in water is 6.5 percent, whereas that of water in *n*-butyl alcohol is 22.4 percent. At 127° C, the upper consolute temperature, complete miscibility is

attained: above  $127^{\circ}\text{C}$  the two liquids mix in all proportions, but below  $127^{\circ}\text{C}$  they show a miscibility gap. Thus, if *n*-butyl alcohol is added to water at  $50^{\circ}\text{C}$ , there is only one liquid phase until 6.5 weight percent of the mixture is alcohol; when more alcohol is added, a second liquid phase appears the composition of which is 22.4 weight percent water. When sufficient alcohol is present to make the overall composition 77.6 weight percent alcohol, the first phase disappears, and only one liquid phase remains. A qualitatively different example is the system water-triethylamine, which has a lower consolute temperature at  $17^{\circ}\text{C}$ . Below  $17^{\circ}\text{C}$  the two liquids are completely miscible, but at higher temperatures they are only partially miscible. Finally, it is possible, although rare, for a binary system to exhibit both upper and lower consolute temperatures. Above  $128^{\circ}\text{C}$  and below  $49^{\circ}\text{C}$  butyl glycol and water are completely miscible, but between these temperatures they do not mix in all proportions.

### Colligative properties

Colligative properties depend only on the concentration of the solute, not on the identity of the solute molecules. The concept of an ideal solution, as expressed by Raoult's law, was already well-known during the last quarter of the 19th century, and it provided the early physical chemists with a powerful technique for measuring molecular weights. (Reliable measurements of molecular weights, in turn, provided important evidence for the modern atomic and molecular theory of matter.)

### Rise in boiling point

It was observed that, whenever one component in a binary solution is present in large excess, the partial pressure of that component is correctly predicted by Raoult's law, even though the solution may exhibit departures from ideal behaviour in other respects. When Raoult's law is applied to the solvent of a very dilute solution containing a nonvolatile solute, it is possible to calculate the mole fraction of the solute from an experimental determination of the rise in boiling point that results when the solute is dissolved in the solvent. Since the separate weights of solute and solvent are readily measured, the procedure provides a simple experimental method for the determination of molecular weight. If a weighed amount of a nonvolatile substance,  $w_2$ , is dissolved in a weighed amount of a solvent,  $w_1$ , at constant pressure, the increase in the boiling temperature,  $\Delta T_{b1}$ , the gas constant,  $R$  (derived from the gas laws), the heat of vaporization of

the pure solvent per unit weight,  $l_1^{\text{vap}}$ , and the boiling temperature of pure solvent,  $T_{b1}$ , are related in a simple product of ratios equal to the molecular weight of the solute,  $M_2$ .

## Azeotropes

An **azeotrope** or a **constant boiling mixture** is a mixture of two or more liquids whose proportions cannot be altered by simple distillation.<sup>[1]</sup> This happens because when an azeotrope is boiled, the vapour has the same proportions of constituents as the unboiled mixture.

Because their composition is unchanged by distillation, azeotropes are also called (especially in older texts) **constant boiling mixtures**. The word azeotrope is derived from the Greek words (boil) and  $\tau\rho\acute{o}\pi o\varsigma$  (turning) combined with the prefix  $\alpha$ - (no) to give the overall meaning, "no change on boiling". The term "azeotrope" was coined in 1911 by English chemist John Wade (1864–1912) and Richard William Merriman.

Many azeotropic mixtures of pairs of compounds are known,<sup>[4]</sup> and many azeotropes of three or more compounds are also known.<sup>[5]</sup> In such a case it is not possible to separate the components by fractional distillation. There are two types of azeotropes: minimum boiling azeotrope and maximum boiling azeotrope. A solution that shows greater positive deviation from Raoult's law forms a minimum boiling azeotrope at a specific composition. For example, an ethanol-water mixture (obtained by fermentation of sugars)

## Steam distillation

**Steam distillation** is a special type of distillation (a separation process) for temperature sensitive materials like natural aromatic compounds. It once was a popular laboratory method for purification of organic compounds, but has become obsolete by vacuum distillation. Steam distillation remains important in certain industrial sectors.<sup>[1]</sup>

Many organic compounds tend to decompose at high sustained temperatures. Separation by distillation at the normal (1 atmosphere) boiling points is not an option, so water or steam is introduced into the distillation apparatus. The water vapor carries small amounts of the vaporized compounds to the condensation flask, where the condensed liquid phase separates, allowing for easy collection. This process effectively allows for distillation at lower temperatures, reducing

the deterioration of the desired products. If the substances to be distilled are very sensitive to heat, steam distillation may be applied under reduced pressure, thereby reducing the operating temperature further.

After distillation the vapors are condensed. Usually the immediate product is a two-phase system of water and the organic distillate, allowing for separation of the components by decantation, partitioning or other suitable methods.

## Applications



A boiling water distiller. Boiling tank on top and holding tank on the bottom.

Steam distillation is employed in the isolation of essential oils, for use in perfumes, for example. In this method, steam is passed through the plant material containing the desired oils. Eucalyptus oil and orange oil are obtained by this method on an industrial scale. Steam distillation is also sometimes used to separate intermediate or final products during the synthesis of complex organic compounds.

Steam distillation is also widely used in petroleum refineries and petrochemical plants where it is commonly referred to as "steam stripping".

Steam distillation also is an important means of separating fatty acids from mixtures and for treating crude products such as tall oils to extract and separate fatty acids, soaps and other commercially valuable organic compounds.

## Miscible pairs

**Miscibility** is the property of substances to mix in all proportions (that is, to fully dissolve in each other at any concentration), forming a homogeneous solution. The term is most often applied to liquids, but applies also to solids and gases. Water and ethanol, for example, are miscible because they mix in all proportions.

By contrast, substances are said to be immiscible if a significant proportion does not form a solution. Otherwise, the substances are considered miscible. For example, butanone is significantly soluble in water, but these two solvents are not miscible because they are not soluble in all proportions.

## Determination

Miscibility of two materials is often determined optically. When the two miscible liquids are combined, the resulting liquid is clear. If the mixture is cloudy the two materials are immiscible. Care must be taken with this determination. If the indices of refraction of the two materials are similar, an immiscible mixture may be clear and give an incorrect determination that the two liquids are miscible.

## POSSIBLE QUESTIONS

### Part A(Each carry one mark)

1. Isothermal critical point is otherwise known as  
a)eutectic point      b)triple point      **c)plait point**      d)melting point
2. In Gibbs-Duhem-Margules equation, if the vapour behaves as an ideal gas,the fugacity can be replaced by  
a)temperature      b)pressure      c)concentration      **d)vapour pressure**
3. For a three component system, the phase rule becomes  
a) $F=4-P$       **b) $F=5-P$**       c) $F=3-P$       d) $F=2-P$
4. Minimum boiling point is said to be  
**a)high volatile**      b)least volatile      c)no volatile      d)volatility increases and then decreases
5. complex ions are studied in

- a) **nernst distribution law**      b) gibbs distribution law      c) einstein distribution law  
d) albert distribution law
6. Existence of similar molecular species in the two phases in-----with other  
a) **contact**      b) no contact      c) contact and no contact      d) bonded
7. Application of the phase rule to a system corresponding to a point in the two phase region gives  
a)  $F=2$       **b)  $F=3$**       c)  $F=1$       d)  $F=0$
8. In Gibbs-Duhem-Margules equation, if the vapour behaves as an ideal gas, the fugacity can be replaced by  
a) temperature      b) pressure      c) concentration      **d) vapour pressure**
9. Tie lines are represented in  
a) two component system      b) one component system      **c) three component system**  
d) four component system
10. In type I of fractional distillation  
**a) possible to isolate both the pure constituents from each other**  
b) not possible to isolate both the pure constituents from each other  
c) possible as well as not possible to isolate both the pure constituents from each other  
d) possible to isolate one constituent
11. Ethanol in type II fractional distillation is obtained as  
a) residue      **b) distillate**      c) residue and distillate      d) vapour
12. Minimum boiling point is said to be  
**a) high volatile**      b) least volatile      c) no volatile      d) volatility increases and then decreases
13. An ----- solution is defined as the one in which the activity of each component is equal to its mole fraction under all conditions of temperature, pressure and concentration  
**a) ideal**      b) real      c) ideal and real      d) non-ideal
14. In type III of fractional distillation

- a)possible to isolate both the pure constituents from each other  
**b) not possible to isolate both the pure constituents from each other**  
c)possible as well as not possible to isolate both the pure constituents from each other  
d)possible to isolate one constituent
15. Water-ethanol system is an example for  
a)type I                      **b)type II**                      c)type III                      d)type IV
16. Solutions of different composition coexisting with one another are termed as  
**a)conjugate solutions**                      b)non-conjugate solutions                      c)mixed solutions                      d)non-mixed solutions
17. Application of the phase rule to a system corresponding to a point in the two phase region gives  
a) $F=2$                       **b) $F=3$**                       c) $F=1$                       d) $F=0$
18. In Gibbs-Duhem-Margules equation, if the vapour behaves as an ideal gas, the fugacity can be replaced by  
a)temperature                      b)pressure                      c)concentration                      **d)vapour pressure**
19. In type II constant boiling mixture obtained as  
a)residue                      **b)distillate**                      c)residue and distillate                      d)vapour
20. Critical solution temperature is otherwise known as  
**a)consolute temperature**                      b)solubility temperature                      c)miscibility temperature  
d)melting temperature

## Part B(Each carry two marks)

- 21.What is known as CST?
- 22.What is known as UCST?
- 23.What is known as LCST?
- 24.State Nernst distribution law.
- 25.Mention few examples for three component system.

26.Mention the applications of Nernst distribution law.

27.What are miscible pairs.

**Part C(Each carry 8 marks)**

28.Derive Gibbs- Duhem-Margules equation and mention its applications.

29.Derive Nernst distribution law and mention its important applications

30.Derive Gibbs-Duhem Margules equation.

31.Explain fractional distillation in detail

32.write notes on

(i)Fractional distillation

(ii)Steam distillation

33.What is triangular plot. Apply phase rule to water-chloroform-acetic acid system.

34.Analyse type I, type II and type III fractional distillation

KAHE

## UNIT-II

### Physical chemistry

| S.No | Question                                                                                                                                                                      | Option 1             | Option 2             | Option 3               | Option 4              | Answer                 |
|------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------|----------------------|------------------------|-----------------------|------------------------|
| 1    | For a three component system, the phase rule becomes                                                                                                                          | $F=4-P$              | $F=5-P$              | $F=3-P$                | $F=2-P$               | $F=5-P$                |
| 2    | The simplest three component systems are those in which a liquid system breaks down into                                                                                      | two phases           | three phases         | four phases            | five phases           | two phases             |
| 3    | Isothermal critical point is otherwise known as                                                                                                                               | eutectic point       | triple point         | plait point            | melting point         | plait point            |
| 4    | Application of the phase rule to a system corresponding to a point in the two phase region gives                                                                              | $F=2$                | $F=3$                | $F=1$                  | $F=0$                 | $F=3$                  |
| 5    | Tie lines are represented in                                                                                                                                                  | two component system | one component system | three component system | four component system | three component system |
| 6    | A system of two salts and water furnishes an example of a ----- system involving solids and liquids                                                                           | three component      | two component        | one component          | four component        | three component        |
| 7    | An ----- solution is defined as the one in which the activity of each component is equal to its mole fraction under all conditions of temperature, pressure and concentration | ideal                | real                 | ideal and real         | non-ideal             | ideal                  |

|    |                                                                                                                             |                                                                |                                                                    |                                                                                        |                                     |                                                                |
|----|-----------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------|--------------------------------------------------------------------|----------------------------------------------------------------------------------------|-------------------------------------|----------------------------------------------------------------|
| 8  | A binary solution of components A and B in equilibrium with their vapours, at constant temperature and pressure is given by | Gibbs equation                                                 | Margules equation                                                  | Gibbs-Duhem-Margules equation                                                          | Duhem equation                      | Gibbs-Duhem-Margules equation                                  |
| 9  | $n_A$ and $n_B$ are the numbers of moles of components A and B                                                              | Gibbs equation                                                 | Margules equation                                                  | Gibbs-Duhem-Margules equation                                                          | Duhem equation                      | Gibbs-Duhem-Margules equation                                  |
| 10 | In Gibbs-Duhem-Margules equation, if the vapour behaves as an ideal gas, the fugacity can be replaced by                    | temperature                                                    | pressure                                                           | concentration                                                                          | vapour pressure                     | vapour pressure                                                |
| 11 | The process of separating mixtures by repeated distillation and condensation is known as                                    | steam distillation                                             | fractional distillation                                            | distillation                                                                           | condensation                        | fractional distillation                                        |
| 12 | In type I of fractional distillation                                                                                        | possible to isolate both the pure constituents from each other | not possible to isolate both the pure constituents from each other | possible as well as not possible to isolate both the pure constituents from each other | possible to isolate one constituent | possible to isolate both the pure constituents from each other |
| 13 | In type II of fractional distillation                                                                                       | possible to isolate both the pure constituents from each other | not possible to isolate both the pure constituents from each other | possible as well as not possible to isolate both the pure constituents                 | possible to isolate one constituent | possible to isolate one constituent                            |

|    |                                                  |                                                                |                                                                    |                                                                                        |                                     |                                     |
|----|--------------------------------------------------|----------------------------------------------------------------|--------------------------------------------------------------------|----------------------------------------------------------------------------------------|-------------------------------------|-------------------------------------|
|    |                                                  |                                                                | other                                                              | from each other                                                                        |                                     |                                     |
| 14 | In type III of fractional distillation           | possible to isolate both the pure constituents from each other | not possible to isolate both the pure constituents from each other | possible as well as not possible to isolate both the pure constituents from each other | possible to isolate one constituent | possible to isolate one constituent |
| 15 | In type II constant boiling mixture obtained as  | residue                                                        | distillate                                                         | residue and distillate                                                                 | vapour                              | distillate                          |
| 16 | In type II Pure A or Pure B obtained as          | residue                                                        | distillate                                                         | residue and distillate                                                                 | vapour                              | residue                             |
| 17 | In type III constant boiling mixture obtained as | residue                                                        | distillate                                                         | residue and distillate                                                                 | vapour                              | residue                             |
| 18 | In type III pure A or pure B obtained as         | residue                                                        | distillate                                                         | residue and distillate                                                                 | vapour                              | distillate                          |
| 19 | The constant boiling point in type II is         | minimum                                                        | maximum                                                            | increases and then decreases                                                           | decreases and then increases        | minimum                             |
| 20 | The constant boiling point in type III is        | minimum                                                        | maximum                                                            | increases and then decreases                                                           | decreases and then increases        | maximum                             |
| 21 | Pure water boils at                              | 100 degree celsius                                             | 50 degree celsius                                                  | 120 degree celsius                                                                     | 110 degree celsius                  | 100 degree celsius                  |

|    |                                                                                                                                                                   |                      |                    |                        |                   |                      |
|----|-------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------|--------------------|------------------------|-------------------|----------------------|
| 22 | pure hydrogen chloride boils at                                                                                                                                   | 65 degree celsius    | 100 degree celsius | -85 degree celsius     | 85 degree celsius | -85 degree celsius   |
| 23 | In water-ethanol system, the constant boiling mixture corresponding to the point has a composition of                                                             | 100% ethanol         | 95.6% ethanol      | 93% ethanol            | 94% ethanol       | 95.6% ethanol        |
| 24 | Ethanol in type II fractional distillation is obtained as                                                                                                         | residue              | distillate         | residue and distillate | vapour            | distillate           |
| 25 | pure water is obtained as                                                                                                                                         | residue              | distillate         | residue and distillate | vapour            | residue              |
| 26 | Ethanol boils at----- under a pressure of 1 atmosphere                                                                                                            | 78.13 degree celsius | 80 degree celsius  | 100 degree celsius     | 20 degree celsius | 78.13 degree celsius |
| 27 | Water-ethanol system is an example for                                                                                                                            | type I               | type II            | type III               | type IV           | type II              |
| 28 | Mixture of acetone and chloroform is an example for                                                                                                               | type I               | type II            | type III               | type IV           | type III             |
| 29 | Mixture of water and hydrogen chloride are example for                                                                                                            | type I               | type II            | type III               | type IV           | type III             |
| 30 | Mixture of water and nitric acid are example for                                                                                                                  | type I               | type II            | type III               | type IV           | type III             |
| 31 | Mixture like a pure chemical compound boils at a constant temperature and distils over completely at the same temperature without change in composition is called | azeotropic mixture   | liquid mixture     | solid mixture          | gaseous mixture   | azeotropic mixture   |
| 32 | azeotropic mixture is otherwise known as                                                                                                                          | constant boiling     | liquid mixture     | gaseous mixture        | solid mixture     | constant boiling     |

|    |                                                                                                                                    |                         |                     |                                   |                                         |                      |
|----|------------------------------------------------------------------------------------------------------------------------------------|-------------------------|---------------------|-----------------------------------|-----------------------------------------|----------------------|
|    |                                                                                                                                    | mixture                 |                     |                                   |                                         | mixture              |
| 33 | maximum boiling point is said to be                                                                                                | high volatile           | least volatile      | no volatile                       | volatility increases and then decreases | least volatile       |
| 34 | minimum boiling point is said to be                                                                                                | high volatile           | least volatile      | no volatile                       | volatility increases and then decreases | high volatile        |
| 35 | ----- are used in fractional distillation                                                                                          | fractionating column    | distillation column | condensation column               | fractionating and distillation column   | fractionating column |
| 36 | composition of liquid and vapour phases can be determined with the help of                                                         | phase rule              | lever's rule        | gibbs rule                        | charles rule                            | lever's rule         |
| 37 | The masses of the liquids in the distillate will be in the ratio of their vapour pressures and molar masses                        | fractional distillation | steam distillation  | fractional and steam distillation | column distillation                     | steam distillation   |
| 38 | Any mixture of two immiscible liquids will boil at a temperature -----than that at which any pure constituent of the mixture boils | lower                   | higher              | lower and higher                  | constant                                | lower                |
| 39 | The boiling point of aniline is                                                                                                    | 180 degree celsius      | 100 degree celsius  | 120 degree celsius                | 85 degree celsius                       | 180 degree celsius   |
| 40 | A compound immiscible or nearly so in water is                                                                                     | benzene                 | aniline             | phenol                            | amide                                   | aniline              |

|    |                                                                                                                         |                                     |                                                                   |                               |                      |                                                                   |
|----|-------------------------------------------------------------------------------------------------------------------------|-------------------------------------|-------------------------------------------------------------------|-------------------------------|----------------------|-------------------------------------------------------------------|
| 41 | Partial miscibility increases on increasing the temperature                                                             | phenol-water                        | water-nicotine                                                    | benzene-water                 | methanol-water       | phenol-water                                                      |
| 42 | partial miscibility increases on lowering the temperature                                                               | phenol-water                        | (C <sub>2</sub> H <sub>5</sub> ) <sub>2</sub> NH-H <sub>2</sub> O | water-picotine                | ether-water system   | (C <sub>2</sub> H <sub>5</sub> ) <sub>2</sub> NH-H <sub>2</sub> O |
| 43 | partial miscibility increases on both raising as well as lowering the temperature                                       | water-nicotine                      | phenol-water                                                      | water-benzene                 | ether-water system   | water-nicotine                                                    |
| 44 | complete miscibility temperature cannot be obtained                                                                     | water-benzene                       | phenol-water                                                      | ether-water                   | water-nicotine       | ether-water                                                       |
| 45 | The temperature above which a pair of partially miscible liquids become miscible in all proportions is called           | lower critical solution temperature | higher critical solution temperature                              | critical solution temperature | critical temperature | critical solution temperature                                     |
| 46 | critical solution temperature is otherwise known as                                                                     | consolute temperature               | solubility temperature                                            | miscibility temperature       | melting temperature  | consolute temperature                                             |
| 47 | Liquid pairs attain complete miscibility above a certain temperature in which case they are said to have the            | LCST                                | UCST                                                              | CST                           | ST                   | UCST                                                              |
| 48 | Liquid pairs show complete miscibility below a certain temperature when they are said to have                           | LCST                                | UCST                                                              | CST                           | ST                   | LCST                                                              |
| 49 | The composition points of the conjugate phases are joined by                                                            | tie lines                           | conjugate lines                                                   | upper lines                   | lower lines          | tie lines                                                         |
| 50 | The two phases having dissimilar composition in equilibrium with each other at a given temperature constitute a pair of | conjugate phase                     | non-conjugate phase                                               | tie phase                     | gibbs phase          | conjugate phase                                                   |

|    |                                                                                          |                                                             |                                                           |                                                                                   |                                                     |                                                             |
|----|------------------------------------------------------------------------------------------|-------------------------------------------------------------|-----------------------------------------------------------|-----------------------------------------------------------------------------------|-----------------------------------------------------|-------------------------------------------------------------|
| 51 | Solutions of different composition coexisting with one another are termed as             | conjugate solutions                                         | non-conjugate solutions                                   | mixed solutions                                                                   | non-mixed solutions                                 | conjugate solutions                                         |
| 52 | aniline-hexane is an example for                                                         | partial miscibility increases on increasing the temperature | partial miscibility increases on lowering the temperature | partial miscibility increases on both raising as well as lowering the temperature | complete miscibility temperature cannot be obtained | partial miscibility increases on increasing the temperature |
| 53 | triethylamine-water system is an example for                                             | partial miscibility increases on increasing the temperature | partial miscibility increases on lowering the temperature | partial miscibility increases on both raising as well as lowering the temperature | complete miscibility temperature cannot be obtained | partial miscibility increases on lowering the temperature   |
| 54 | partition coefficient is otherwise known as                                              | distribution coefficient                                    | diffusion coefficient                                     | viscosity coefficient                                                             | freezing coefficient                                | distribution coefficient                                    |
| 55 | The Nernst distribution law is valid only for -----of single molecules in the two phases | concentrations                                              | pressure                                                  | volume                                                                            | temperature                                         | concentrations                                              |
| 56 | benzene-water                                                                            | mutually insoluble                                          | mutually soluble                                          | insoluble                                                                         | completely soluble                                  | mutually insoluble                                          |

|    |                                                                            |                         |                        |                           |                         |                         |
|----|----------------------------------------------------------------------------|-------------------------|------------------------|---------------------------|-------------------------|-------------------------|
| 57 | The mutual solubility is not altered by the presence of the solute in      | aniline-water           | benzene-water          | amide-water               | chloroform-water        | benzene-water           |
| 58 | complex ions are studied in                                                | nernst distribution law | gibbs distribution law | einstein distribution law | albert distribution law | nernst distribution law |
| 59 | One of the important validity in Nernst distribution law is                | constant temperature    | constant pressure      | constant volume           | constant composition    | constant temperature    |
| 60 | Existence of similar molecular species in the two phases in-----with other | contact                 | no contact             | contact and no contact    | bonded                  | contact                 |

### UNIT III

#### Electrochemical Cells

Electrochemical Cells: Rules of oxidation/reduction of ions based on half-cell potentials, applications of electrolysis in metallurgy and industry. Chemical cells, reversible and irreversible cells with examples. Electromotive force of a cell and its measurement. Nernst equation; Standard electrode(reduction)potential and its application to different kinds of half cells.

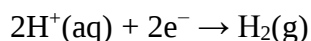
#### ELECTROCHEMICAL CELLS

Rules of oxidation/reduction of ions based on half-cell potentials

A **half-cell** is a structure that contains a conductive electrode and a surrounding conductive electrolyte separated by a naturally occurring Helmholtz double layer. Chemical reactions within this layer momentarily pump electric charges between the electrode and the electrolyte, resulting in a potential difference between the electrode and the electrolyte. The typical anode reaction involves a metal atom in the electrode dissolved and transported as a positive ion across the double layer, causing the electrolyte to acquire a net positive charge while the electrode acquires a net negative charge. The growing potential difference creates an intense electric field within the double layer, and the potential rises in value until the field halts the net charge-pumping reactions. This self-limiting action occurs almost instantly in an isolated half-cell; in applications two dissimilar half-cells are appropriately connected to constitute a Galvanic cell.

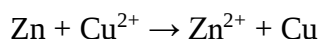
A standard half-cell, used in electrochemistry, consists of a metal electrode in a 1 molar (1 mol/L) aqueous solution of the metal's salt, at 298 kelvin (25 °C).<sup>[1]</sup> The electrochemical series, which consists of standard electrode potentials and is closely related to the reactivity series, was generated by measuring the difference in potential between the metal half-cell in a circuit with a standard hydrogen half-cell, connected by a salt bridge.

The standard hydrogen half-cell:

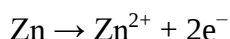


The half-cells of a Daniell cell:

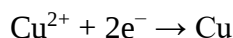
Original equation



Half-cell (anode) of Zn



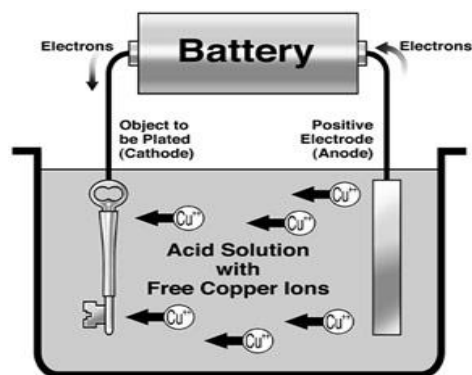
Half-cell (cathode) of Cu



Applications of electrolysis in metallurgy and industry

**Electroplating** is used to coat one metal with another metal by using electrolysis. Electroplating is usually done to improve the appearance of the metal or prevent the corrosion of the metal.

Chromium plating is used to prevent iron from rusting.



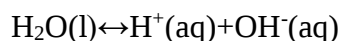
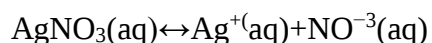
The cathode is the object to be plated, while anode is the desired metal to coat the object. The electrolyte solution must contain ions of the same metal for plating. During electrolysis, the anode will dissolve into the solution. The ions produced will migrate to the cathode where they are discharged and deposited as a layer on the cathode.

It is important to ensure that the cathode is electrically conductive. (If not, the electrolysis does not work.)

Example: Silver Plating

**Electrodes:** Silver anode, Cathode is object to be plated with silver

**Electrolyte:** Solution of soluble silver salt. (E.g. Silver nitrate,  $\text{AgNO}_3(\text{aq})$ )



**Ions present in solution:**  $\text{Ag}^+$ ,  $\text{H}^+$ ,  $\text{NO}_3^-$ ,  $\text{OH}^-$

Reaction at CATHODE:

- $\text{Ag}^+$  and  $\text{H}^+$  is attracted to the cathode.
- $\text{Ag}^+$  is preferentially discharged.
- Silver metal is deposited on the cathode.
- $\text{Ag}^+(\text{aq}) + \text{e}^- \rightarrow \text{Ag}(\text{s})$

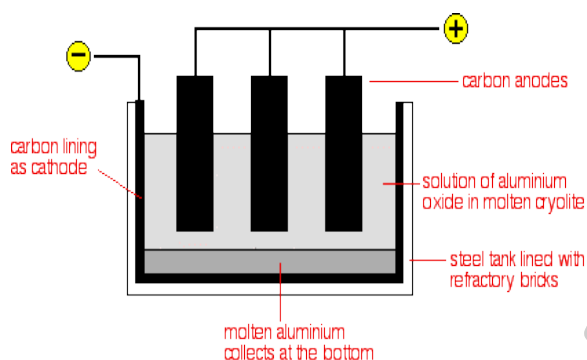
Reaction at ANODE:

- $\text{NO}_3^-$  and  $\text{OH}^-$  is attracted to the anode.
- Neither  $\text{NO}_3^-$  nor  $\text{OH}^-$  is discharged.
- Each silver atom loses one electron to form one  $\text{Ag}^+$  ion.
- The silver anode dissolves into the solution.
- $\text{Ag}(\text{s}) \rightarrow \text{Ag}^+(\text{aq}) + \text{e}^-$

## Extraction Of Reactive Metals

Reactive metals are the metals that occupy the top positions in the electrochemical series. Metals that are higher than zinc in the electrochemical series are extracted using electrolysis. These very

reactive metals cannot be extracted by other metals such as reduction with carbon. The reactive metals are obtained by electrolyzing a molten ionic compound of metal.



Example: Extraction Of Aluminium

Aluminium is extracted from aluminium oxide ( $\text{Al}_2\text{O}_3$ ) or known as bauxite.

The electrolytic cell is an iron tank lined with carbon, which acts as the cathode. The anodes are blocks of carbon dipped into the electrolyte. The electrolyte is a solution of molten aluminium oxide in molten cryolite. Cryolite acts as a solvent to dissolve aluminium oxide and as an impurity to lower the melting point of aluminium oxide. The electrolytic cell is maintained at around  $900^\circ\text{C}$

**Electrodes:** Carbon

**Ions present in electrolyte:**  $\text{Al}^{+3}$ ,  $\text{O}^{2-}$

**Reaction at cathode:**  $\text{Al}^{+3}(\text{l}) + 3\text{e}^- \rightarrow \text{Al}(\text{l})$

**Reaction at anode:**  $2\text{O}^{2-}(\text{l}) \rightarrow \text{O}_2(\text{g}) + 4\text{e}^-$

Aluminium ions are discharged at the cathode, forming a pool of molten aluminium at the bottom of the tank.

At high temperature, oxygen reacts with the carbon anode to form carbon dioxide gas. Hence, the anodes are slowly burnt away as carbon dioxide gas and needs to be replaced frequently.

**Chemical cells**

An **electrochemical cell** is a device capable of either generating electrical energy from chemical reactions or facilitating chemical reactions through the introduction of electrical energy. A common example of an electrochemical cell is a standard 1.5 - volt cell meant for consumer use. This type of device is known as a single galvanic cell. A battery consists of one or more cells, connected in either parallel or series pattern.

## Equilibrium reaction

Each half-cell has a characteristic voltage. Various choices of substances for each half-cell give different potential differences. Each reaction is undergoing an equilibrium reaction between different oxidation states of the ions: When equilibrium is reached, the cell cannot provide further voltage. In the half-cell that is undergoing oxidation, the closer the equilibrium lies to the ion/atom with the more positive oxidation state the more potential this reaction will provide. Likewise, in the reduction reaction, the closer the equilibrium lies to the ion/atom with the more negative oxidation state the higher the potential.

## Cell potential

The cell potential can be predicted through the use of electrode potentials (the voltages of each half-cell). These half-cell potentials are defined relative to the assignment of 0 volts to the standard hydrogen electrode (SHE). (See table of standard electrode potentials). The difference in voltage between electrode potentials gives a prediction for the potential measured. When calculating the difference in voltage, one must first rewrite the half-cell reaction equations to obtain a balanced oxidation-reduction equation.

1. Reverse the reduction reaction with the smallest potential ( to create an oxidation reaction/ overall positive cell potential)
2. Half-reactions must be multiplied by integers to achieve electron balance.

Note that the cell potential does not change when the reaction is multiplied by a constant.

Cell potentials have a possible range of roughly zero to 6 volts. Cells using water-based electrolytes are usually limited to cell potentials less than about 2.5 volts, because the very powerful oxidizing and reducing agents that would be required to produce a higher cell potential tend to react with the water. Higher cell potentials are possible with cells using other solvents instead of water. For instance, lithium cells with a voltage of 3 volts are commonly available.

The cell potential depends on the concentration of the reactants, as well as their type. As the cell is discharged, the concentration of the reactants decreases and the cell potential also decreases

Reversible and irreversible cells with examples

A cell is said to be reversible if the following two conditions are fulfilled

- (i) The chemical reaction of the cell stops when an exactly equal external emf is applied.
- (ii) The chemical reaction of the cell is reversed and the current flows in opposite direction when the external emf is slightly higher than that of the cell. Any other cell, which does not obey the above two conditions, is termed as irreversible. Daniell cell is reversible but  $Zn | H_2SO_4 | Ag$  cell is irreversible in nature

(5) **Types of electrochemical cells:** Two main types of electrochemical cells have been reported, these are,

(i) **Chemical cells:** The cells in which electrical energy is produced from the energy change accompanying a chemical reaction or a physical process are known as chemical cells. Chemical cells are of two types,

(a) **Chemical cells without transference:** In this type of chemical cells, the liquid junction potential is neglected or the transference number is not taken into consideration. In these cells, one electrode is reversible to cations while the other is reversible to the anions of the electrolyte.

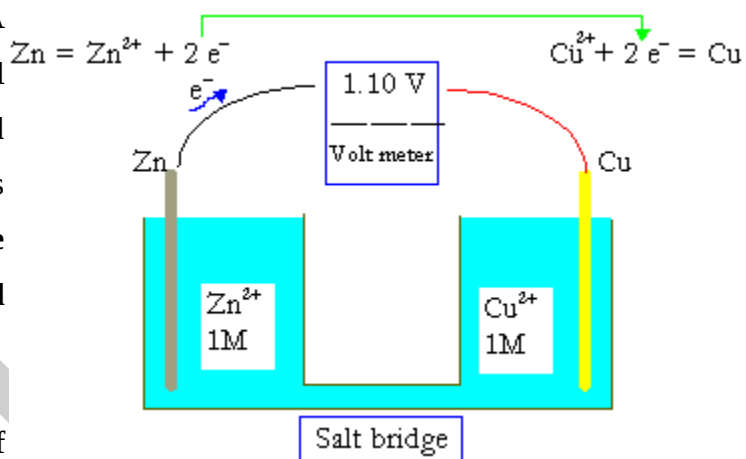
Electromotive force of a cell and its measurement

The **electromotive force (EMF)** is the maximum potential difference between two electrodes of a galvanic or voltaic cell. This quantity is related to the tendency for an element, a compound or

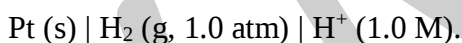
an ion to acquire (i.e. gain) or release (loss) electrons. For example, the maximum potential between Zn and Cu of a well known cell



has been measured to be 1.100 V. A concentration of 1 M in an ideal solution is defined as the standard condition, and 1.100 V is thus the **standard electromotive force,  $E^\circ$** , or **standard cell potential** for the Zn-Cu galvanic cell.

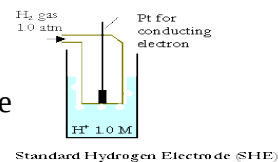


The standard cell potential,  $E^\circ$ , of the a galvanic cell can be evaluated from the standard reduction potentials of the two half cells  $E^\circ$ . The reduction potentials are measured against the standard hydrogen electrode (SHE):



**Its reduction potential or oxidation potential is defined to be exactly zero.**

The reduction potentials of all other half-cells measured in volts against the SHE are the difference in electrical potential energy per coulomb of charge.



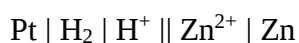
Note that the unit for energy  $J = \text{Coulomb volt}$ , and the Gibbs free energy  $G$  is the product of charge  $q$  and potential difference  $E$ :

$$G \text{ in } J = q E \text{ in } C V$$

for electric energy calculations.

## Evaluating Standard Cell Potential $E^\circ$ of Galvanic Cells

A galvanic cell consists of two half-cells. The convention in writing such a cell is to put the (reduction) cathode on the right-hand side, and the (oxidation) anode on the left-hand side. For example, the cell

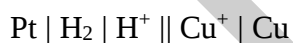


consists of the oxidation and reduction reactions:

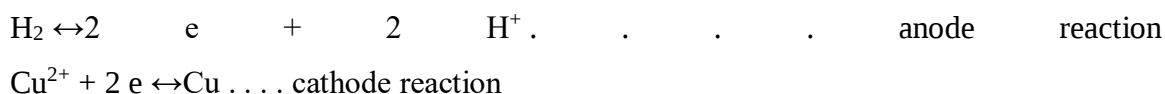


If the concentrations of  $\text{H}^+$  and  $\text{Zn}^{2+}$  ions are 1.0 M and the pressure of  $\text{H}_2$  is 1.0 atm, the voltage difference between the two electrodes would be -0.763 V (the Zn electrode being the negative electrode). The conditions specified above are called the **standard conditions** and the EMF so obtained is **the standard reduction potential**.

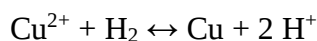
Note that the above cell is in reverse order compared to that given in many textbooks, but this arrangement gives the standard reduction potentials directly, because the Zn half cell is a reduction half-cell. The negative voltage indicates that the reverse chemical reaction is spontaneous. This corresponds to the fact that Zn metal reacts with an acid to produce  $\text{H}_2$  gas. As another example, the cell



consists of an oxidation and a reduction reaction:



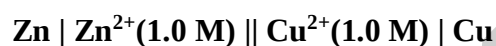
and the standard cell potential is 0.337 V. The positive potential indicates a spontaneous reaction,



but the potential is so small that the reaction is too slow to be observed.

## Example 1

**What is the potential for the cell**



## Solution

From a table of standard reduction potentials we have the following values



Add (1) and (2) to yield



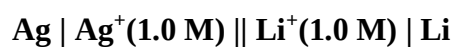
Note that  $E^*$  is the oxidation standard potential, and  $E^\circ$  is the reduction standard potential,  $E^* = -E^\circ$ . The standard cell potential is represented by  $\Delta E^\circ$ .

## Discussion

The positive potential confirms your observation that zinc metal reacts with cupric ions in solution to produce copper metal.

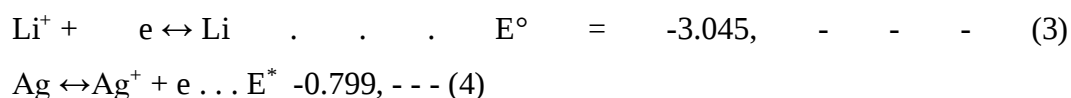
## Example 2

**What is the potential for the cell**



## Solution

From the table of standard reduction potentials, you find



According to the convention of the cell, the reduction reaction is on the right. The cell on your left-hand side is an oxidation process. Thus, you add (4) and (3) to obtain



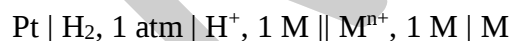
## Discussion

The negative potential indicates that the reverse reaction should be spontaneous.

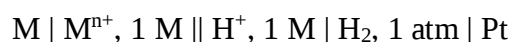
Some calculators use a lithium battery. The atomic weight of Li is 6.94, much lighter than Zn (65.4).

## Summary

- The **electromotive force (EMF)** is the maximum potential difference between two electrodes of a galvanic or voltaic cell.
- The standard reduction potential of  $\text{M}^{n+}$ , 1 M / M couple is the standard cell potential of the galvanic cell:



- The standard oxidation potential of  $\text{M} \mid \text{M}^{n+}$ , 1 M couple is the standard cell potential of the galvanic cell:



- If the cell potential is negative, the reaction is reversed. In this case, the electrode of the galvanic cell should be written in a reversed order.

## Nernst equation

In electrochemistry, the **Nernst equation** is an equation that relates the reduction potential of an electrochemical reaction (half-cell or full cell reaction) to the standard electrode potential, temperature, and activities (often approximated by concentrations) of the chemical species undergoing reduction and oxidation. It is the most important equation in the field of electrochemistry. It is named after the German physical chemist who first formulated it, Walther Nernst.

## Standard electrode potential and its application to different kinds of half-cells

In electrochemistry, the **standard electrode potential**, abbreviated  $E^\circ$  or  $E^\ominus$  (with a superscript plimsoll character, pronounced "standard" or "nought"), is the measure of individual potential of a reversible electrode at standard state, which is with solutes at an effective concentration of  $1 \text{ mol dm}^{-3}$ , and gases at a pressure of 1 atm. The reduction potential is an intensive property. The values are most often tabulated at 25 °C. The basis for an electrochemical cell such as the galvanic cell is always a redox reaction which can be broken down into two half-reactions: oxidation at anode (loss of electron) and reduction at cathode (gain of electron). Electricity is generated due to electric potential difference between two electrodes. This potential difference is created as a result of the difference between individual potentials of the two metal electrodes with respect to the electrolyte. (Reversible electrode is an electrode that owes its potential to changes of a reversible nature, in contrast to electrodes used in electroplating which are destroyed during their use.)

Although the overall potential of a cell can be measured, there is no simple way to accurately measure the electrode/electrolyte potentials in isolation. The electric potential also varies with temperature, concentration and pressure. Since the oxidation potential of a half-reaction is the

negative of the reduction potential in a redox reaction, it is sufficient to calculate either one of the potentials. Therefore, standard electrode potential is commonly written as standard reduction potential.

## Calculation of standard electrode potentials

The electrode potential cannot be obtained empirically. The galvanic cell potential results from a pair of electrodes. Thus, only one empirical value is available in a pair of electrodes and it is not possible to determine the value for each electrode in the pair using the empirically obtained galvanic cell potential. A reference electrode, standard hydrogen electrode (SHE), for which the potential is defined or agreed upon by convention, needed to be established. In this case SHE is set to 0.00 V and any electrode, for which the electrode potential is not yet known, can be paired with SHE—to form a galvanic cell—and the galvanic cell potential gives the unknown electrode's potential. Using this process, any electrode with an unknown potential can be paired with either the SHE or another electrode for which the potential has already been derived and that unknown value can be established.

Since the electrode potentials are conventionally defined as reduction potentials, the sign of the potential for the metal electrode being oxidized must be reversed when calculating the overall cell potential. Note that the electrode potentials are independent of the number of electrons transferred—they are expressed in volts, which measure energy per electron transferred—and so the two electrode potentials can be simply combined to give the overall cell potential even if different numbers of electrons are involved in the two electrode reactions.

For practical measurements, the electrode in question is connected to the positive terminal of the electrometer, while SHE is connected to the negative terminal.

## Standard reduction potential

The larger the value of the standard reduction potentials, the easier it is for the element to be reduced (accept electrons); in other words, they are better oxidizing agents. For example,  $F_2$  has 2.87 V and  $Li^+$  has  $-3.05$  V.  $F$  reduces easily and is therefore a good oxidizing agent. In contrast,

$\text{Li}_{(s)}$  would rather undergo oxidation (hence a good reducing agent). Thus  $\text{Zn}^{2+}$  whose standard reduction potential is  $-0.76 \text{ V}$  can be oxidized by any other electrode whose standard reduction potential is greater than  $-0.76 \text{ V}$  (e.g.  $\text{H}^+(0 \text{ V})$ ,  $\text{Cu}^{2+}(0.16 \text{ V})$ ,  $\text{F}_2(2.87 \text{ V})$ ) and can be reduced by any electrode with standard reduction potential less than  $-0.76 \text{ V}$  (e.g.  $\text{H}_2(-2.23 \text{ V})$ ,  $\text{Na}^+(-2.71 \text{ V})$ ,  $\text{Li}^+(-3.05 \text{ V})$ ).

In a galvanic cell, where a spontaneous redox reaction drives the cell to produce an electric potential, Gibbs free energy  $\Delta G^\circ$  must be negative, in accordance with the following equation:

$$\Delta G^\circ_{\text{cell}} = -nFE^\circ_{\text{cell}}$$

where  $n$  is number of moles of electrons per mole of products and  $F$  is the Faraday constant,  $\sim 96485 \text{ C/mol}$ . As such, the following rules apply:

If  $E^\circ_{\text{cell}} > 0$ , then the process is spontaneous (galvanic cell)

If  $E^\circ_{\text{cell}} < 0$ , then the process is nonspontaneous (electrolytic cell)

Thus in order to have a spontaneous reaction ( $\Delta G^\circ < 0$ ),  $E^\circ_{\text{cell}}$  must be positive, where:

$$E^\circ_{\text{cell}} = E^\circ_{\text{cathode}} - E^\circ_{\text{anode}}$$

where  $E^\circ_{\text{anode}}$  is the standard potential at the anode and  $E^\circ_{\text{cathode}}$  is the standard potential at the cathode as given in the table of standard electrode potential.

## EMF and its measurement

**EMF measurements** are measurements of ambient (surrounding) electromagnetic fields that are performed using particular sensors or probes, such as EMF meters. These probes can be generally considered as *antennas* although with different characteristics. In fact probes should not perturb the electromagnetic field and must prevent coupling and reflection as much as possible in order to obtain precise results. There are two main types of EMF measurements:

- *broadband measurements* performed using a broadband probe, that is a device which senses any signal across a wide range of frequencies and is usually made with three independent diode detectors;



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4. What is the potential of a half cell consisting of zinc electrode in 0.01M  $\text{ZnSO}_4$  solution  $25^\circ\text{C}$ .

$E^\circ = 0.763 \text{ V}$

- a) **0.0591 V**                      b) 0.6521 V    c) 0.7532 V                      d) 0.8221

V

5. The salt bridge is filled with a solution of

- a) potassium chromate    b) sodium chloride    **c) potassium chloride**    d) zinc chloride

6. An example for metal-metal ion electrodes is

- a) daniel cell**                      b) hydrogen electrode    c) chlorine electrode                      d) calomel electrode

7. Which one is Metal-insoluble metal salt electrode

- a) calomel electrode**                      b) standard hydrogen electrode    c) silver-silver chloride electrode    d) Gas electrode

8. The reduction potential of the electrode is 1.5v then its oxidation potential is

- a) 0 V                      b) 1 V    **c) negative 1.5**                      d) 2

9. The Nernst equation is

- a)  $E = 2.303 \text{ RT/nF} \log K$     **b)  $E = 2.303/\text{nF} \text{ RT} \log K$**     c)  $E = 2.303 \text{ RT/nF} - E^\circ$   
d)  $E = E^\circ - 2.303 \text{ RT/nF} \log K$

10. The EMF generated by an electrochemical cell is given by the symbol

- a) E**                      b)  $E^\circ$                       c) V                      d)  $V^\circ$

11. calomel is a

- a)potassium chloride      b)sodium chloride      **c)mercurous chloride**  
d)barium chloride

12. The device in which the free energy of a physical or chemical process is converted into electrical energy is called

- a)daniel cell      **b)galvanic cell**      c)laclanche cell      d)voltaic cell

13. The reference electrode employed is usually the -----in glass electrode

- a)hydrogen      **b)calomel**      c)quinine      d)hydroquinone

14. The EMF is measured in

- a)volts**      b)coulomb      c)faraday      d)joules

15. The electrochemical or electrolytic processes are carried out in a device known as

- a)battery      **b)galvanometer**      c)potentiometer      d)electrolyte

16. The electrode in which oxidation occurs is

- a)anode**      b)cathode      c)Anode and Cathode      d)Electrolyte

17. The electrode in which reduction occurs is

- a)anode      **b)cathode**      c)Anode and Cathode      d)Electrolyte

18. The value of standard electrode potential arranged in the decreasing order is called

- a)chemical series      b)potential series      **c)electrochemical series**  
d)electricity series

19. The relationship between free energy change and emf of a cell is

- a) $\Delta G = -nFE$**       b) $\Delta H = -nFE$       c) $\Delta E = nFG$       d) $\Delta F = nEG$

20. An example for gas electrode is

- a)hydrogen electrode                      b)chlorine electrode                      c)oxygen electrode  
d)hydrogen, Chlorine and oxygen electrode

**Part B (Each carry two marks)**

- 21.What are electrochemical cells?  
22.What are reversible and irreversible cells.  
23.Define EMF.  
24.What is known as standard electrode potential?  
25.What is known as single electrode potential?  
26.What are half cells?  
27.Write down Nernst equation for oxidation potential.  
28.Write down Nernst equation for reduction potential.

**Part C (Each carry 8 marks)**

- 29.Derive Nernst equation for EMF of a cell.  
30. Discuss the following  
    (i)Reversible and irreversible cells  
    (ii)Standard electrode potential  
31.Explain electromotive force of a cell and its measurement.  
32.write notes on  
    (i)reversible and irreversible cells  
    (ii)rules of oxidation/reduction of ions based on half-cell potentials

33.What is known as standard electrode potential and derive an expression for Nernst equation for an oxidation potential.

34.What is known as electrochemical cell and write down the rules of oxidation/reduction of ions based on half-cell potentials.

KAHE

### UNIT-III

#### Physical chemistry

| S.No | Question                                                                                                            | Option 1                 | Option 2           | Option 3              | Option 4                                | Answer                                  |
|------|---------------------------------------------------------------------------------------------------------------------|--------------------------|--------------------|-----------------------|-----------------------------------------|-----------------------------------------|
| 1    | The device in which the free energy of a physical or chemical process is converted into electrical energy is called | daniel cell              | galvanic cell      | laclanche cell        | voltaic cell                            | galvanic cell                           |
| 2    | The electrode in which oxidation occurs is                                                                          | anode                    | cathode            | Anode and Cathode     | Electrolyte                             | anode                                   |
| 3    | The salt bridge is filled with a solution of                                                                        | potassium chromate       | sodium chloride    | potassium chloride    | zinc chloride                           | potassium chloride                      |
| 4    | If the electricity produced by the cell is equal to the EMF, the cell is                                            | reversible               | irreversible       | Sometimes reversible  | Sometimes irreversible                  | irreversible                            |
| 5    | In quinhydrone electrode, platinum wire is placed                                                                   | hydroquinone and quinone | only hydroquinone  | only quinone          | Water                                   | only hydroquinone                       |
| 6    | An example for metal-metal ion electrodes is                                                                        | daniel cell              | hydrogen electrode | chlorine electrode    | calomel electrode                       | daniel cell                             |
| 7    | An example for gas electrode is                                                                                     | hydrogen electrode       | chlorine electrode | oxygen electrode      | hydrogen, Chlorine and oxygen electrode | hydrogen, Chlorine and oxygen electrode |
| 8    | calomel is a                                                                                                        | potassium chloride       | sodium chloride    | mercurous chloride    | barium chloride                         | mercurous chloride                      |
| 9    | The wire used in the calomel electrode is made of                                                                   | platinum                 | copper             | titanium              | iron                                    | platinum                                |
| 10   | An example for oxidation-reduction electrode is                                                                     | calomel electrode        | chlorine electrode | quinhydrone electrode | hydrogen electrode                      | quinhydrone electrode                   |
| 11   | In quinhydrone electrode, the platinum wire is placed in a solution containing                                      | hydroquinone and quinone | only hydroquinone  | only quinone          | Water                                   | only hydroquinone                       |

|    |                                                                                                              |                     |                             |                                  |                         |                        |
|----|--------------------------------------------------------------------------------------------------------------|---------------------|-----------------------------|----------------------------------|-------------------------|------------------------|
| 12 | The tendency of an electrode to lose or gain electrons when contact with its own ions in solution, is called | electrode potential | reduction potential         | oxidation potential              | Concentration potential | electrode potential    |
| 13 | The electrode in which reduction occurs is                                                                   | anode               | cathode                     | Anode and Cathode                | Electrolyte             | cathode                |
| 14 | The value of standard electrode potential arranged in the decreasing order is called                         | chemical series     | potential series            | electrochemical series           | electricity series      | electrochemical series |
| 15 | Any two suitable half cells can be combined to form a                                                        | daniel cell         | electrochemical cell        | galvanic cell                    | leclanche cell          | galvanic cell          |
| 16 | If the electricity produced by the cell is greater than the applied EMF, then the cell is                    | reversible          | irreversible                | Sometimes reversible             | Sometimes irreversible  | reversible             |
| 17 | In the calomel electrode, the wire used is made of                                                           | platinum            | copper                      | titanium                         | iron                    | platinum               |
| 18 | The salt bridge is made of                                                                                   | potassium chromate  | sodium chloride             | potassium chloride               | zinc chloride           | potassium chloride     |
| 19 | An example for metal-insoluble metal salt electrode is                                                       | calomel electrode   | standard hydrogen electrode | silver-silver chloride electrode | Gas electrode           | calomel electrode      |
| 20 | The voltaic cell Zn-Cu, the standard EMF is                                                                  | 1.20 v              | 1.15 v                      | 1.25 v                           | 1.10 v                  | 1.10 v                 |
| 21 | Which one is Metal-insoluble metal salt electrode                                                            | calomel electrode   | standard hydrogen electrode | silver-silver chloride electrode | Gas electrode           | calomel electrode      |
| 22 | The EMF is measured in                                                                                       | volts               | coulomb                     | faraday                          | joules                  | volts                  |
| 23 | The electrical energy is measured in                                                                         | volts               | joules                      | coulomb                          | meter                   | joules                 |
| 24 | The EMF generated by an electrochemical cell is given by the symbol                                          | E                   | $E^\circ$                   | V                                | $V^\circ$               | E                      |

|    |                                                                                            |                                                |                               |                                                |                                         |                                                |
|----|--------------------------------------------------------------------------------------------|------------------------------------------------|-------------------------------|------------------------------------------------|-----------------------------------------|------------------------------------------------|
| 25 | The EMF is measured by                                                                     | voltmeter                                      | galvanometer                  | potentiometer                                  | ammeter                                 | potentiometer                                  |
| 26 | The EMF of the unknown half cell is calculated from                                        | $E^\circ = E_R - E_L$                          | $E^\circ = E_L - E_R$         | $E^\circ = E_L + E_R$                          | $E^\circ = E_L + E_s$                   | $E^\circ = E_R - E_L$                          |
| 27 | The potential of a single electrode in a half cell is called the                           | electromotive force                            | single electrode potential    | standard reduction potential                   | cell potential                          | single electrode potential                     |
| 28 | If the standard emf $E^\circ$ is positive, the reaction is                                 | feasible                                       | not feasible                  | reversible                                     | Irreversible                            | feasible                                       |
| 29 | The Nernst equation is                                                                     | $E = E^\circ - \frac{2.303}{nF} \log K$        | $E = \frac{2.303}{nF} \log K$ | $E = E^\circ - \frac{2.303}{nF} \log K$        | $E = E^\circ - \frac{2.303}{nF} \log K$ | $E = \frac{2.303}{nF} \log K$                  |
| 30 | The Nernst equation for the oxidation half cell is                                         | $E = E^\circ - \frac{2.303}{n} \log [Zn^{2+}]$ | $E = \frac{2.303}{nF} \log K$ | $E = E^\circ - \frac{0.0591}{n} \log [Mn^{+}]$ | $E = \frac{2.303}{nF} \log K$           | $E = E^\circ - \frac{2.303}{n} \log [Zn^{2+}]$ |
| 31 | Which compound is used in the salt bridge                                                  | potassium chromate                             | sodium chloride               | potassium chloride                             | zinc chloride                           | potassium chloride                             |
| 32 | The relationship between free energy change and emf of a cell is                           | $\Delta G = -nFE$                              | $\Delta H = -nFE$             | $\Delta E = nFG$                               | $\Delta F = nEG$                        | $\Delta G = -nFE$                              |
| 33 | The metals near the bottom of the electrochemical series is                                | strong reducing agents                         | strong oxidising agent        | weak reducing agent                            | weak oxidising agents                   | strong reducing agents                         |
| 34 | The feasibility of a redox reaction can be predicted with the help of                      | electronegativity                              | electrochemical series        | electron affinity                              | equivalent conductance                  | electrochemical series                         |
| 35 | The emf of a cell with 1M solution of reactants and products in solution at 25°C is called | half cell potential                            | standard emf                  | single electrode potential                     | redox potential                         | standard emf                                   |
| 36 | The relationship between equilibrium constant and the standard emf of a cell is            | $E^\circ = \frac{0.0591}{n} \log k$            | $0.0591 E^\circ = \log k$     | $n E^\circ = \frac{0.0591}{n} \log k$          | $n E^\circ = \frac{0.0591}{n} \log k$   | $0.0591 E^\circ = \log k$                      |
| 37 | Which one of the following is not present in the calomel electrode                         | mercurous chloride                             | mercury                       | KCl                                            | ZnCl                                    | ZnCl                                           |

|    |                                                                                     |                             |                             |                                  |                              |                             |
|----|-------------------------------------------------------------------------------------|-----------------------------|-----------------------------|----------------------------------|------------------------------|-----------------------------|
| 38 | The electrochemical or electrolytic processes are carried out in a device known as  | battery                     | galvanometer                | potentiometer                    | electrolyte                  | galvanometer                |
| 39 | When the electrodes are connected externally, then the circuit is said to be        | open                        | closed                      | ideal                            | constant                     | open                        |
| 40 | What is SHE                                                                         | standard hydrogen electrode | standard helium electrode   | simple half electrode            | standard reference electrode | standard hydrogen electrode |
| 41 | The overall reaction taking place in the daniel cell is                             | oxidation                   | reduction                   | redox                            | forward                      | redox                       |
| 42 | If the reduction potential of the electrode is 1.5v then its oxidation potential is | 0 v                         | 1 v                         | negative 1.5                     | 2                            | negative 1.5                |
| 43 | Daniel cell is called as                                                            | voltaic cell                | half cell                   | anode cell                       | cathode half cell            | half cell                   |
| 44 | Metal-insoluble metal salt electrode is                                             | calomel electrode           | standard hydrogen electrode | silver-silver chloride electrode | Gas electrode                | calomel electrode           |
| 45 | The standard EMF of Zn-Cu voltaic cell is                                           | 1.20 v                      | 1.15 v                      | 1.25 v                           | 1.10 v                       | 1.10 v                      |
| 46 | The standard emf can be determined at standard temperature is                       | 32°C                        | 20°C                        | 25°C                             | 27°C                         | 25°C                        |
| 47 | The standard electrode potential at 25°C is zero for                                | Hydrogen electrode          | Gas electrode               | Calomel electrode                | Metal electrode              | Hydrogen electrode          |
| 48 | The standard hydrogen electrode can act as                                          | anode                       | cathode                     | Anode and Cathode                | Electrolyte                  | Anode and Cathode           |
| 49 | The emf of the standard hydrogen electrode is arbitrarily assigned the value of     | 0 v                         | 0.1 v                       | 0.001 v                          | 1 v                          | 0 v                         |
| 50 | The reduction potential of the electrode is 1.5v then its oxidation potential is    | 0 v                         | 1 v                         | negative 1.5                     | 2                            | negative 1.5                |

|    |                                                                                                                                                          |                      |                             |                                  |                      |                   |
|----|----------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------|-----------------------------|----------------------------------|----------------------|-------------------|
| 51 | The standard temperature at which the standard emf can be determined is                                                                                  | 32°C                 | 20°C                        | 25°C                             | 27°C                 | 25°C              |
| 52 | For Zn-Cu voltaic cell, the standard EMF is                                                                                                              | 1.20 v               | 1.15 v                      | 1.25 v                           | 1.10 v               | 1.10 v            |
| 53 | Which one the following is example for Metal-insoluble metal salt electrode                                                                              | calomel electrode    | standard hydrogen electrode | silver-silver chloride electrode | Gas electrode        | calomel electrode |
| 54 | The unit of electrical energy is                                                                                                                         | volts                | joules                      | coulomb                          | meter                | joules            |
| 55 | In IUPAC conventions, the double vertical line represents                                                                                                | two half cell        | cathode half cell           | salt bridge                      | anode half cell      | salt bridge       |
| 56 | Platinum is a                                                                                                                                            | positive electrode   | negative electrode          | Positive and negative electrode  | Inert electrode      | Inert electrode   |
| 57 | If the emf acts in the opposite direction through the cell circuit it is denoted as a                                                                    | positive             | negative                    | zero                             | cannot be determined | zero              |
| 58 | What is the potential of a half cell consisting of zinc electrode in 0.01M ZnSO <sub>4</sub> solution 25°C. E° = 0.763 V                                 | 0.0591 V             | 0.6521 V                    | 0.7532 V                         | 0.8221 V             | 0.0591 V          |
| 59 | What is R in Nernst equation                                                                                                                             | rate of the reaction | redox reaction              | gas constant                     | reduction of gas     | gas constant      |
| 60 | What is the free energy change for the reaction $\text{Sn}^{4+} + 2\text{e}^- \rightarrow \text{Sn}^{2+}$ . If its standard reduction potential is +0.15 | 25.59 kJ             | 29.52 kJ                    | 28.95 kJ                         | data inadequate      | 25.59 kJ          |

## UNIT IV

### Application of EMF Measurements

**APPLICATION OF EMF MEASUREMENTS:** Application of EMF measurements in determining (i) free energy, enthalpy and entropy of a cell reaction, (ii) equilibrium constants, and (iii)  $pH$  values, using hydrogen, quinone-hydroquinone glass and  $SbO/Sb_2O_3$  electrodes. Concentration cells with and without transference, liquid junction potential; determination of activity coefficients and transference numbers. Qualitative discussion of potentiometric titrations (acid-base, redox, precipitation)

### APPLICATION OF EMF MEASUREMENTS

Electrode at which oxidation takes place is anode and the electrode at which reduction takes place is cathode. When a metal is in contact with its own ion solution it develops a potential with respect to the electrolyte. The potential difference developed at the anode - electrolyte interface is called oxidation potential and the potential difference developed at the cathode - electrolyte interface is called reduction potential. The potential difference between the anode and cathode is called the EMF of the cell. The potential difference measured at standard conditions (1 atm pressure, 273K) is called standard electrode potential. Standard electrode potential gives the tendency of the electrode to get oxidized or reduced. If the electrolytes are different the two compartments are joined by a salt bridge, which is a tube containing a concentrated electrolyte solution in agar jelly that completes the electrical circuit and enables the cell to function.

#### Electrochemical Series:

A series in which metals are arranged in the decreasing order of reduction potential.

| Electrodes                           | $E^0$ in volt |
|--------------------------------------|---------------|
| $F_2(g) + 2e^- \rightarrow 2F^-(aq)$ | +2.87         |

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|                                                                                                                    |       |
|--------------------------------------------------------------------------------------------------------------------|-------|
| $\text{H}_2\text{O}_2(\text{aq}) + 2\text{H}^+(\text{aq}) + 2\text{e}^- \rightarrow 2\text{H}_2\text{O}(\text{l})$ | +1.77 |
| $\text{Au}^+(\text{aq}) + \text{e}^- \rightarrow \text{Au}(\text{s})$                                              | +1.68 |
| $\text{Cl}_2(\text{g}) + 2\text{e}^- \rightarrow 2\text{Cl}^-(\text{aq})$                                          | +1.36 |
| $\text{O}_2(\text{g}) + 4\text{H}^+(\text{aq}) + 4\text{e}^- \rightarrow 2\text{H}_2\text{O}(\text{l})$            | +1.23 |
| $\text{Br}_2(\text{l}) + 2\text{e}^- \rightarrow 2\text{Br}^-(\text{aq})$                                          | +1.09 |
| $\text{Ag}^+(\text{aq}) + \text{e}^- \rightarrow \text{Ag}(\text{s})$                                              | +0.80 |
| $\text{Fe}^{3+}(\text{aq}) + \text{e}^- \rightarrow \text{Fe}^{2+}(\text{aq})$                                     | +0.77 |
| $\text{I}_2(\text{s}) + 2\text{e}^- \rightarrow 2\text{I}^-(\text{aq})$                                            | +0.54 |
| $\text{O}_2(\text{g}) + 2\text{H}_2\text{O}(\text{l}) + 4\text{e}^- \rightarrow 4\text{OH}^-(\text{aq})$           | +0.40 |
| $\text{Cu}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Cu}(\text{s})$                                           | +0.34 |
| $\text{S}(\text{s}) + 2\text{H}^+(\text{aq}) + 2\text{e}^- \rightarrow \text{H}_2\text{S}(\text{g})$               | +0.14 |
| $2\text{H}^+(\text{aq}) + 2\text{e}^- \rightarrow \text{H}_2(\text{g})$                                            | 0.00  |
| $\text{Pb}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Pb}(\text{s})$                                           | -0.13 |
| $\text{Sn}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Sn}(\text{s})$                                           | -0.14 |
| $\text{Ni}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Ni}(\text{s})$                                           | -0.23 |
| $\text{Co}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Co}(\text{s})$                                           | -0.28 |
| $\text{Fe}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Fe}(\text{s})$                                           | -0.44 |
| $\text{Zn}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Zn}(\text{s})$                                           | -0.76 |
| $2\text{H}_2\text{O}(\text{l}) + 2\text{e}^- \rightarrow \text{H}_2(\text{g}) + 2\text{OH}^-(\text{aq})$           | -0.83 |
| $\text{Mn}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Mn}(\text{s})$                                           | -1.03 |
| $\text{Al}_3^+(\text{aq}) + 3\text{e}^- \rightarrow \text{Al}(\text{s})$                                           | -1.67 |
| $\text{Mg}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Mg}(\text{s})$                                           | -2.34 |
| $\text{Na}^+(\text{aq}) + \text{e}^- \rightarrow \text{Na}(\text{s})$                                              | -2.71 |
| $\text{Ca}_2^+(\text{aq}) + 2\text{e}^- \rightarrow \text{Ca}(\text{s})$                                           | -2.87 |
| $\text{K}^+(\text{aq}) + \text{e}^- \rightarrow \text{K}(\text{s})$                                                | -2.93 |
| $\text{Li}^+(\text{aq}) + \text{e}^- \rightarrow \text{Li}(\text{s})$                                              | -3.02 |

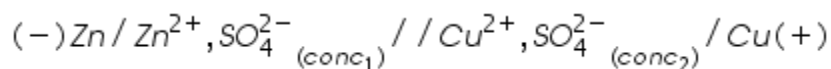
We can construct innumerable number of galvanic cells by taking combinations of different half cells. Each half cell consists of a metallic rod dipped in to an electrolyte. The metal with higher reduction potential act as cathode and the other will act as anode.

Standard EMF of the cell:

$$E_{cell}^0 = E_{cathode}^0 - E_{anode}^0$$

A galvanic cell is represented by putting a vertical line between metal and electrolyte solution and putting a double vertical line between the two electrolytes connected by a salt bridge.

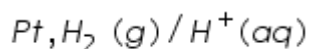
Eg: The symbolic representation of Daniel cell is given below,



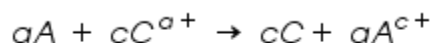
First, the reduced form of the metal to be oxidized at the anode (Zn) is written. This is separated from its oxidized form by a vertical line, which represents the limit between the phases (oxidation changes). The double vertical lines represent the saline bridge on the cell. Finally, the oxidized form of the metal to be reduced at the cathode, is written, separated from its reduced form by the vertical line. The electrolyte concentration is given as it is an important variable in determining the cell potential.

### Standard Hydrogen Electrode (S.H.E.):

The potential of Standard hydrogen electrode used as the reference electrode has been arbitrarily taken as zero. The electrode consist of a glass jacket consisting of dry hydrogen gas bubbled at one atmosphere. There is a platinum wire sealed in the glass jacket. The entire system is immersed in 1M HCl solution. Standard hydrogen electrode can be represented as,



Electrode potential at any concentration can be calculated using Nernst equation. For the reaction,



Nernst Equation,

$$E_{cell} = E_{cell}^0 - \frac{2.303RT}{nF} \log \frac{[C^{a+}]^c}{[A^{c+}]^a}$$

$$E_{cell}^0 = E_{cathode}^0 - E_{anode}^0$$

$$n=c \quad \text{if} \quad c=a$$

$$n=c \times a \quad \text{if} \quad c \neq a$$

Where;

n=number of electrons.

$E^0$ = electrode potential of cell at standard conditions.

T=temperature.

R=universal gas constant.

F=Faraday constant.

When a cell reaction takes place electrical energy is produced which results in decrease in the free energy of the system.

Electrical work = Decrease in free energy

In an electro chemical cell,

Electric work done = Quantity of electric charge produced x E.M.F of the cell

For one mole of electrons quantity of electric charge is 1F (96500 coulomb)

Therefore, for n moles it is nF.

$$\text{Electric work done} = nFE_{\text{cell}}$$

$$-\Delta G = nFE_{\text{cell}}$$

For a standard cell,

$$-\Delta G^0 = nFE_{\text{cell}}^0$$

By van 't Hoff relation,

$$\Delta G^0 = -RT \ln K$$

$$E_{\text{cell}}^0 = \frac{RT}{nF} \ln K$$

$$\ln K = \frac{nFE_{\text{cell}}^0}{RT}$$

K=equilibrium constant

Spontaneity or Feasibility of Reaction:

| $\Delta G$ | K  | $E_{\text{cell}}^0$ | Reaction    |
|------------|----|---------------------|-------------|
| Negative   | >1 | Positive            | Spontaneous |

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| $\Delta G$ | $K$ | $E_{cell}^0$ | Reaction          |
|------------|-----|--------------|-------------------|
| Zero       | =1  | Zero         | Equilibrium       |
| Positive   | <1  | Negative     | Non - spontaneous |

Application of Emf measurements in determining

Measurement of entropy and enthalpy of a cell reaction

From Gibbs-Helmholtz equation we have

$$\Delta G = \Delta H + T \left[ \frac{\partial(\Delta G)}{\partial T} \right]_P$$

$\Delta G = -nEF$  and differentiating  $\Delta G$  with respect to temperature at constant pressure yields

$$\left[ \frac{d(\Delta G)}{dT} \right]_P = -nF \left( \frac{\partial F}{\partial T} \right)_P$$

The quantity  $\left( \frac{\partial E}{\partial T} \right)_P$  is the temperature coefficient of the cell. Substituting the value of  $\left[ \frac{\partial(\Delta G)}{\partial T} \right]_P$  in equation, we get

$$\Delta G = \Delta H - nFT \left( \frac{\partial E}{\partial T} \right)_P$$

Also  $dG = dH - TdS$

$$\Delta S = nF \left( \frac{\partial E}{\partial T} \right)_P$$

Equation can be used for calculating the entropy changes for the cell reaction in terms of the temperature coefficient of the cell emf. Equation can also be written as

$$-nFE = \Delta H - nFT \left( \frac{\partial E}{\partial T} \right)_P$$

$$\Delta H = nF \left[ T \left( \frac{\partial E}{\partial T} \right)_P - E \right]$$

or

From equation, the enthalpy change for cell reaction can be determined from the measurement of cell emf and the temperature coefficient of the emf.

### Determination of equilibrium constant

To calculate the equilibrium constant for an electrochemical cell we need to know:

- the standard state potential for a cell
- the half-reactions involved

The Nernst equation is used in calculating the equilibrium constant.

$$E^\circ_{\text{cell}} = \frac{RT}{nF} \ln Q$$

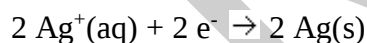
At equilibrium  $Q = K$ . Substituting in  $K$  for  $Q$ , and the values for  $R$ ,  $T$ , and  $F$ , we get:

$$E^\circ_{\text{cell}} = \frac{0.0257}{n} \ln K = \frac{0.0592}{n} \log K$$

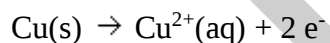
**Example:** Find the value of the equilibrium constant at 25°C for the cell reaction for the following electrochemical cell:



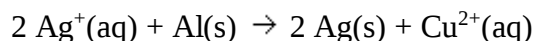
- Write the equations for the cell half-reactions, calculate the standard cell potential, and determine the number of electrons transferred.



$$E^\circ_{\text{reduction}} = + 0.799 \text{ V}$$



$$E^\circ_{\text{oxidation}} = - 0.518 \text{ V}$$



$$E^\circ_{\text{cell}} = + 0.281 \text{ V}$$

$n = 2$  moles of electrons

- Substitute into the above equation and solve for  $K$
- Substitute into the above equation and solve for  $K$ .

$$E^{\circ}_{\text{cell}} = \frac{0.0592}{n} \log K$$

$$0.281 = \frac{0.0592}{2} \log K$$

$$\log K = 9.49$$

$$K = 10^{9.49} = 3.1 \times 10^9$$

Note: values for the equilibrium constant for electrochemical cell reactions are sometimes very large.

### Determining the Standard State Free Energy Change from $E^{\circ}_{\text{cell}}$

To determine the standard state free energy change for a cell reaction

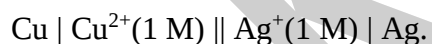
- determine the  $E^{\circ}_{\text{cell}}$
- determine the number of moles of electrons transferred in the reaction.
- solve for  $dG^{\circ}$  using the equation

$$dG^{\circ} = -nFE^{\circ}_{\text{cell}}$$

|                |          |           |          |         |        |                       |
|----------------|----------|-----------|----------|---------|--------|-----------------------|
| $DG^{\circ} =$ | standard | state     | free     | energy  | change | (joules)              |
| $n$            | =        | number    | of       | moles   | of     | electrons transferred |
| $F$            | =        | Faraday's | constant | (96,485 | C/mol  | $e^{-}$ )             |

$$E^{\circ}_{\text{cell}} = \text{standard state cell potential (volts or joules/C)}$$

**Example:** Find the value of the equilibrium constant at 25°C for the cell reaction for the following electrochemical cell:



(The solution for the determination of the  $E^{\circ}_{\text{cell}}$  and the number of moles of electrons,  $n$ , are shown in the example in the previous section. Click [HERE](#) to see the solution.)

- Determine the  $E^{\circ}_{\text{cell}}$ .  
 $E^{\circ}_{\text{cell}} = + 0.281 \text{ volts}$
- Determine the number of moles of electrons transferred.  
 $n = 2 \text{ moles of } e^{-}$
- Substitute into the equation and solve.

$$DG^{\circ} = - (2 \text{ mol } e^{-})(96,485 \text{ C/mol } e^{-})(0.281 \text{ J/C})$$

$$DG^{\circ} = - 54,200 \text{ J or } - 54.2 \text{ kJ}$$

## Determining the Non-Standard State Free Energy Change

To determine the non-standard state free energy change:

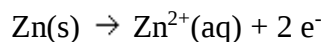
- calculate the standard cell potential,  $E^\circ_{\text{cell}}$
- determine the number of moles of electrons transferred,  $n$
- calculate the reaction quotient,  $Q$
- calculate the non-standard cell potential,  $E_{\text{cell}}$ , using the Nernst equation
- Calculate the non-standard free energy change using the equation:

$$\Delta G = -nFE_{\text{cell}}$$

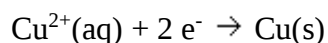
**Example:** Calculate the free energy change for the following electrochemical cell.



- Calculate  $E^\circ_{\text{cell}}$ .



$$E^\circ_{\text{oxidation}} = + 0.762 \text{ volts}$$



$$E^\circ_{\text{reduction}} = + 0.339 \text{ volts}$$



$$E^\circ_{\text{cell}} = + 1.101 \text{ volts}$$

- Determine "n".

$$n = 2 \text{ moles of electrons}$$

- Calculate  $Q$

$$Q = \frac{[\text{Zn}^{2+}]}{[\text{Cu}^{2+}]} = \frac{1.50}{0.25} = 6.0$$

- Calculate  $E_{\text{cell}}$

$$E_{\text{cell}} = 1.101 \text{ volts} - \frac{0.0257}{2} \ln 6 = 1.078 \text{ volts}$$

- Calculate  $\Delta G$ .

$$\Delta G = -nFE_{\text{cell}} = - (2 \text{ mole e}^-)(96,485 \text{ C/mole e}^-)(1.078 \text{ volts})$$

$$\Delta G = - 208, 000 \text{ joules or } -208 \text{ kJ}$$

## Application of EMF measurements in determining pH values

Hydrogen ions play a central role in the lives of cells. For example, changes in hydrogen ion concentration are intimately tied to the charge of side chains in proteins. This charge state, in turn, affects the activity of enzymes as well as their folding and even localization. Further, the

famed ATP synthases that churn out the ATPs that power many cellular processes are driven by gradients in hydrogen ions across membranes.

The abundance of these ions and, as a result, the charge state of many compounds is encapsulated in the pH defined as

$$\text{pH} = -\log_{10}([\text{H}^+]/1)$$

where  $[\text{H}^+]$  denotes the concentration or more formally the activity of the charged hydrogen ions ( $\text{H}^+$ , or more accurately the sum of hydronium,  $\text{H}_3\text{O}^+$ , as well as the functionally important but often overlooked Zundel,  $\text{H}_5\text{O}_2^+$ , and Eigen,  $\text{H}_7\text{O}_3^+$ , cations). We are careful to divide the hydrogen ion concentration by a so-called “standard state” concentration, the agreed upon value is 1M, in order to ensure that when taking the log we have a unitless quantity.

The integer 7 is often etched in our memory from school as the pH of water, but there is nothing special about the integral value of 7. Water has a neutral pH of about 7, with the exact value varying with temperature, ionic strength and pressure. What is the pH inside the cell? Just like with other parameters describing the “state” of molecules and cells, the answer depends on physiological conditions and which compartment within the cell we are considering (i.e. which organelle). Despite these provisos, crude generalizations about the pH can be a useful guide to our thinking.

**Concentration cells with and without transference:** A concentration cell is an electrolytic cell that is comprised of two half-cells with the same electrodes, but differing in concentrations. A concentration cell acts to dilute the more concentrated solution and concentrate the more dilute solution, creating a voltage as the cell reaches an equilibrium. This is achieved by transferring the electrons from the cell with the lower concentration to the cell with the higher concentration. The standard electrode potential, commonly written as  $E^\circ_{\text{cell}}$ , of a concentration cell is equal to zero because the electrodes are identical. But, because the ion concentrations are different, there is a potential difference between the two half-cells. One can find this potential difference via the Nernst Equation,

$$E_{\text{cell}} = E^\circ_{\text{cell}} - 0.0592n \log Q$$

at 25°C. The E stands for the voltage that can be measured using a voltmeter (make sure if the voltmeter measures it in millivolts that you convert the number before using it in the equation). Note that the Nernst Equation indicates that cell potential is dependent on concentration, which

results directly from the dependence of free energy on concentration. Remember that to find  $Q$  you use this equation:



$$Q = \frac{(C)^c (D)^d}{(A)^a (B)^b} \quad Q = \frac{(C)^c (D)^d}{(A)^a (B)^b}$$

When  $Q=1$ , meaning that the concentrations for the products and reactants are the same, then taking the log of this equals zero. When this occurs, the  $E_{\text{cell}}$  is equal to the  $E^{\circ}_{\text{cell}}$ .

Another way to use the  $E^{\circ}_{\text{cell}}$ , or to find it, is using the equation below.

$$E^{\circ}_{\text{cell}} = E^{\circ}_{\text{cathode}} - E^{\circ}_{\text{anode}}$$

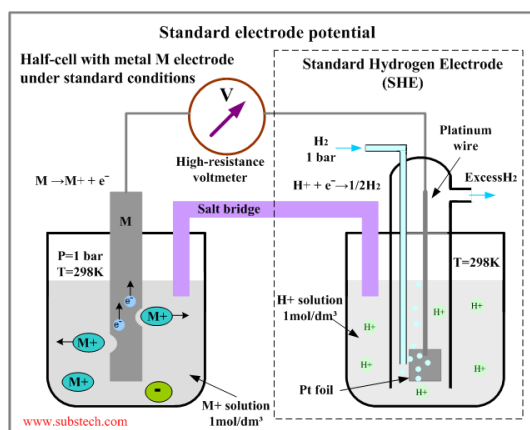


Fig.1 An example of a concentration cell

### Connected Information

These concepts are useful for understanding the electron transfer and what occurs in half-cells.

### Use of a Salt Bridge

The two compartments of a cell must be separated so they do not mix, but cannot be completely separated with no way for ions to be transferred. A wire cannot be used to connect the two compartments because it would react with the ions that flow from one side to another. Because of this, a salt bridge is an important part of a concentration cell. It solves the major problem of electrons beginning to pile up too much in the right beaker. This buildup is due to electrons moving from the left side, or left beaker, to the right side, or right beaker. The salt bridge itself can be in a few different forms, such as a salt solution in a U-tube or a porous barrier (direct contact). It evens the charge by moving ions to the left side, or left beaker. In the written expression which shows what is occurring in specific reactions, the salt bridge is represented by the double lines. An example of this would be:



The double lines between the  $\text{Zn}^{2+}(\text{1M})$  and the  $\text{Cu}^{2+}(\text{1M})$  signify the salt bridge. The single lines, however, do not represent bridges; they represent the different phase changes, for example, from solid zinc to liquid zinc solution. If there is a comma where you would expect to see a single line, this is not incorrect. It simply means that no phase changes occurred.

### Electrode Use

In this type of reaction, there are two electrodes which are involved. These are known as the anode and the cathode, or the left and right side, respectively. The anode is the side which is losing electrons (oxidation) while the cathode is the side which is gaining electrons (reduction).

### Use of a Voltmeter

A voltmeter (not to be confused with a different kind of voltmeter which also measures a type of energy) is used to measure the cell potential that is passed between the two sides. It is typically located in between the two cells. This cell potential (also known as an electromotive force) occurs due to the flow of electrons. The value it shows can be negative or positive depending on the direction in which the electrons are flowing. If the potential is positive then the transfer of electrons is spontaneous, but the reverse reaction is NONspontaneous. Conversely, if the value of the potential is negative, the transfer of electrons is nonspontaneous and the reverse reaction. The voltmeter measures this potential in volts or millivolts.

### Electrons

The tendency of electrons to flow from one chemical to another is known as electrochemistry. This is what occurs in a concentration cell. The electrons flow from the left side (or left beaker) to the right side (or right beaker). Because the left side is losing electrons and the right is gaining them, the left side is called the oxidation side and the right side is the reduction side. Although you could switch the two to be on the opposite sides, this is the general way in which the set up is done. The oxidation side is called the anode and the reduction side is the cathode. It is the flow of the electrons that cause one side to be oxidized and the other to be reduced.

### Corrosion

Corrosion can occur on a concentration cell when the metal being used is in contact with different concentrations, causing parts of the metal to have different electric potential than the

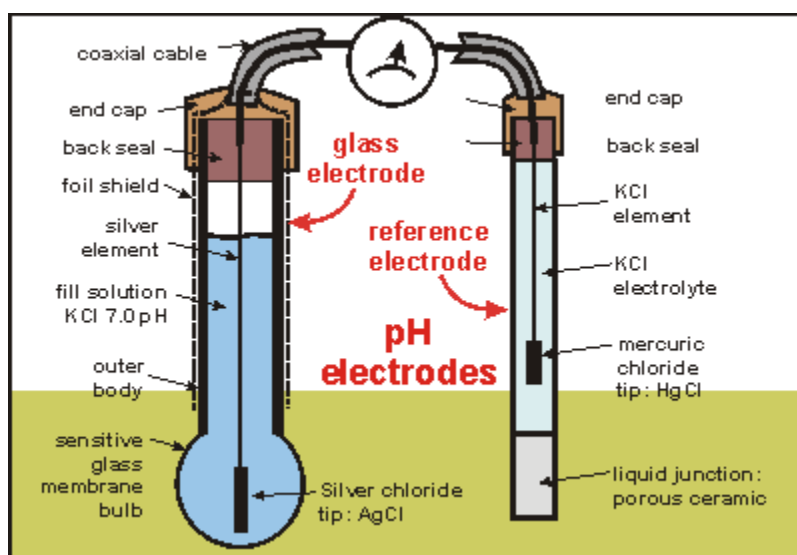
other parts. One element that is often linked to this corrosion is oxygen. In the areas in which there is a low oxygen concentration corrosion occurs.

This can be somewhat prevented through sealing off the cell and keeping it clean, but even this cannot prevent any corrosion from occurring at some point.

Corrosion is most frequently a problem when the cell is in contact with soil. Because of the variations that occur within soil, which is much greater than the variations that occur within a fluid, contact with soil often causes corrosion for the cell.

### Uses of Concentration Cells

A pH meter is a specific type of concentration cell that uses the basic setup of a concentration cell to determine the pH, or the acidity/basicity, of a specific solution. It is comprised of two electrodes and a voltmeter. One of the electrodes, the glass one has two components: a metal (commonly silver chloride) wire and a separate semi-porous glass part filled with a potassium chloride solution with a pH of 7 surrounding the AgCl. The other electrode is called the reference electrode, which contains a potassium chloride solution surrounding a potassium chloride wire. The purpose of this second electrode is to act as a comparison for the solution being tested. When the glass electrode comes into contact with a solution of different pH, an electric potential is created due to the reaction of the hydrogen ions with the metal ions. This potential is then measured by the voltmeter, which is connected to the electrode. The higher the voltage, the more hydrogen ions the solution contains, which means the solution is more acidic.



### Liquid junction potential

**Liquid junction potential** occurs when two electrolytic solutions of different concentrations are in contact with each other. The more concentrated solution will have a tendency to diffuse into the comparatively less concentrated one. The rate of diffusion of each ion will be roughly proportional to its speed in an electric field. If the anions diffuse more rapidly than the cations, they will diffuse ahead into the dilute solution, leaving the latter negatively charged and the concentrated solution positively charged. This will result in an electrical double layer of positive and negative charges at the junction of the two solutions. Thus at the point of junction, a potential difference will develop because of the ionic transfer. This potential is called liquid junction potential or diffusion potential which is non-equilibrium potential. The magnitude of the potential depends on the relative speeds of the ions' movement.

## Calculation

The liquid junction potential cannot be measured directly but calculated. The Electromotive force (EMF) of a concentration cell with transference includes the liquid junction potential.

where  $a_1$  and  $a_2$  are activities of HCl in the two solutions,  $R$  is the Universal Gas Constant,  $T$  is the temperature and  $F$  is Faraday's Constant.

where  $a_2$  and  $a_1$  are activities of HCl solutions of right and left hand electrodes respectively and  $t_M$  be transport number of  $Cl^-$

$$\text{Liquid Junction potential} = E_{wt} - E_{nt} = (t_M - 1) \frac{RT}{F} \cdot \ln (a_2/a_1)$$

## Elimination of liquid junction potential

Go

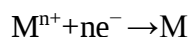
The liquid junction potential interferes with the exact measurement of the electromotive force of a chemical cell, so its effect should be minimized as much as possible for accurate measurement. The most common method of eliminating the liquid junction potential is to place a salt bridge consisting of a saturated solution of potassium chloride(KCl) and ammonium nitrate( $NH_4NO_3$ ) with lithium acetate( $CH_3COOLi$ ) between the two solutions constituting the junction. When such a bridge is used, the ions in the bridge are present in large excess at the junction and they carry almost the whole of the current across the boundary. The efficiency of KCl/ $NH_4NO_3$  is connected with the fact that in these salts, the transport numbers cation.

## Potentiometric titrations

**Potentiometric titration** is a technique similar to direct titration of a redox reaction. It is a useful means of characterizing an acid. No indicator is used; instead the potential is measured

across the analyte, typically an electrolyte solution. To do this, two electrodes are used, an indicator electrode (the glass electrode and metal ion indicator electrode) and a reference electrode. Reference electrodes generally used are hydrogen electrodes, calomel electrodes, and silver chloride electrodes. The indicator electrode forms an electrochemical half cell with the interested ions in the test solution. The reference electrode forms the other half cell,

The overall electric potential is calculated as  $E_{\text{cell}} = E_{\text{ind}} - E_{\text{ref}} + E_{\text{sol}}$ .  $E_{\text{sol}}$  is the potential drop over the test solution between the two electrodes.  $E_{\text{cell}}$  is recorded at intervals as the titrant is added. A graph of potential against volume added can be drawn and the end point of the reaction is halfway between the jump in voltage.  $E_{\text{cell}}$  depends on the concentration of the interested ions with which the indicator electrode is in contact. For example, the electrode reaction may be



As the concentration of  $M^{n+}$  changes, the  $E_{\text{cell}}$  changes correspondingly. Thus the potentiometric titration involve measurement of  $E_{\text{cell}}$  with the addition of titrant. types of potentiometric titration: acid-base titration (total alkalinity and total acidity), redox titration (HI/HY and cerate), precipitation titration (halides), and complexometric titration (free EDTA)

The first potentiometric titration was carried out in 1893 by Robert Behrend at Ostwald's Institute in Leipzig. He titrated mercurous solution with potassium chloride, potassium bromide, and potassium iodide. He used a mercury electrode along with a mercury/mercurous nitrate reference electrode. He found that in a cell composed of mercurous nitrate and mercurous nitrate/mercury, the initial voltage is 0. If potassium chloride is added to mercurous nitrate on one side, mercury (I) chloride is precipitated. This decreased the osmotic pressure of mercury (I) ions on the side and creates a potential difference. This potential difference increases slowly as additional potassium chloride is added, but then increases more rapidly. He found the greatest potential difference is achieved once all of the mercurous nitrate has been precipitated. This was used to discern end points of titrations.

Wilhelm Bottger then developed the tool of potentiometric titration while working at Ostwald's Institute. He used potentiometric titration to observe the differences in titration between strong and weak acids, as well as the behavior of polybasic acids. He introduced the idea of using potentiometric titrations for acids and bases that could not be titrated in conjunction with a colorimetric indicator

Potentiometric titrations were first used for redox titrations by Crotono. He titrated halide ions using potassium permanganate using a shiny platinum electrode and a calomel electrode. He says that if an oxidizing agent is added to a reducing solution then the equilibrium between the reducing substance and reaction product will shift towards the reaction product. The changes the potential very slowly until the amount of reducing substance becomes very small. A large change in potential will occur then once a small addition of the titrating solution is added, as the final amounts of reducing agent are removed and the potential corresponds solely to the oxidizing agent. This large increase in potential difference signifies the endpoint of the reaction.

## Redox titration

A **redox titration** is a type of titration based on a redox reaction between the analyte and titrant. Redox titration may involve the use of a redox indicator and/or a potentiometer. A common example of a redox titration is treating a solution of iodine with a reducing agent to produce iodide using a starch indicator to help detect the endpoint. Iodine ( $I_2$ ) can be reduced to iodide ( $I^-$ ) by e.g. thiosulfate ( $S_2O_3^{2-}$ ), and when all iodine is spent the blue colour disappears. This is called an iodometric titration.

Most often of all, the reduction of iodine to iodide is the last step in a series of reactions where the initial reactions are used to convert an unknown amount of the solute (the substance being analyzed) to an equivalent amount of iodine, which may then be titrated. Sometimes other halogens (or haloalkanes) than iodine are used in the intermediate reactions because they are available in better measurable standard solutions and/or react more readily with the solute. The extra steps in iodometric titration may be worth while because the equivalence point, where the blue turns a bit colourless, is more distinct than some other analytical or may be by volumetric methods.

To evaluate a redox titration we need to know the shape of its titration curve. In an acid-base titration or a complexation titration, the titration curve shows how the concentration of  $H_3O^+$  (as pH) or  $M^{n+}$  (as pM) changes as we add titrant. For a redox titration it is convenient to monitor the titration reaction's potential instead of the concentration of one species.

The Nernst equation relates a solution's potential to the concentrations of reactants and products participating in the redox reaction. Consider, for example, a titration in which a titrand in a reduced state,  $A_{red}$ , reacts with a titrant in an oxidized state,  $B_{ox}$ .



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Titration involving precipitation at end of process is called as precipitation titration. Most of metallic halides are titrated by precipitation method.

It is also called as argentimetric titration. There are three methods used for determining end point in precipitation titration.

Use for determination of halides and pseudo-halides

To determine solubility constant of compounds

To determine electrode potential

For determining chloride, cyanides and thiosulphites

**Part A(Each carry one mark)**

1.If the -----of a particular cell is known the equilibrium constant of the cell reaction can be calculated

- a) standard EMF      b) entropy      c) free energy  
d) enthalpy

2. Quinhydrone electrode is preferred to the -----electrode

3. Electrode-concentration cell are evidently independent of
- a) **concentration of electrolyte**    b) concentration of ions    c) concentration in electrode  
d) electrolyte concentration cell
4. Which of the following shows a metal being oxidized

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5. Emf is measured in

a)volts

b)amperes

c)coulombs

d)ohm

6. A solution of -----molar HCl which furnishes a constant hydrogen ion concentration

a)0.01

b)0.1

c)0.2

d)0.001

7. Glass electrodes can be used in

a)strong oxidising solutions b)weak oxidising solutions c)strong reducing solutions

d)weak reducing solutions

8. In given reaction  $2\text{Cr(s)} + 3\text{Cu}^{2+}(\text{aq}) \rightarrow 3\text{Cr}^{3+}(\text{aq}) + 3\text{Cr(s)}$  which reaction occurs of the cathode in an electrochemical cell

a)reduction of  $\text{Cu}^{2+}(\text{aq})$

b)reduction of  $\text{Cu(s)}$

c)oxidation of  $\text{Cu}^{3+}(\text{aq})$

d)oxidation of  $\text{Cr(s)}$

9. The reference electrode employed is usually the -----in glass electrode

a)hydrogen

b)calomel

c)quinine

d)hydroquinone

10. Two like electrodes with different concentration are immersed in an electrolyte are known as

a)electrode-concentration cell

b)electrolyte concentration cell c)cell concentration

d)cell constant

11. Transfer of ions from one electrolytic solution to the other does not take place at

a)concentration cell with transfer

b)concentration cell without transfer

c)emf is

partially negative

d)concentration cell without transfer and concentration cell with transfer

12. Electrical energy forced in electrochemical cell is

a)Spontaneous

b)Non-Spontaneous

c)Exothermic

d)Endothermic

13. . If the -----of a particular cell is known the equilibrium constant of the cell reaction can be calculated

a)standard EMF

b)entropy c)free energy

d)enthalpy

14. The electrical energy produced by a galvanic cell is given by the ----- of its electromotive force and the quantity of electricity which passes

a)sum

b)product

c)sum and product

d)divided

15. Electricity is measured in

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a)volts                      b)amperes                      c)coulombs                      d)ohm

16 The potential of a hydrogen electrode in contact with a solution of  $H^+$  ions involving the reaction is given by

a)Nernst equation      b)Gibbs equation      c)Duhem equation      d)Clausius equation

17. Glass electrodes can be used in

a)strong oxidising solutions      b)weak oxidising solutions      c)strong reducing solutions  
d)weak reducing solutions

18. Why not pH cannot be measured through potentiometer or voltmeter, but can be done through electronic voltmeter

a)resistance of the glass membrane is very high and the current is small  
b) resistance of the glass membrane is low and the current is large  
c)different salt bridges are used  
d) glass electrode compares pH, while electrode measures P H

19. Liquid junction potential depends on

a)transference number of anion and cation      b)removal of anion and cation  
c)transfer of only anion                      d)transfer of only cation

20. When does L.J.P becomes negative

a) $t_- = t_+$                       b) $t_- \neq t_+$                       c) $t_- > t_+$                       d) $t_- < t_+$

## Part A(Each carry two marks)

21.What is known as liquid junction potential?

22.What is the principle behind potentiometric titration?

23.Mention the application of EMF measurements in determining free energy.

24. Mention the application of EMF measurements in determining enthalpy.

25. Mention the application of EMF measurements in determining entropy.
26. Mention the application of EMF measurements in determining equilibrium constant.
27. Define transport number.
28. What is known as activity coefficient?

**Part B(Each carry 8 marks)**

29. Explain potentiometric precipitation reaction.
30. Explain the following
  - (i) application of EMF measurements in determining enthalpy
  - (ii) application of EMF measurements in determining free energy of a cell
31. Explain concentration cells with and without transference.
32. Describe potentiometric titrations of a redox reaction
33. Explain concentration cells with and without transference.
34. Explain the following
  - (i) application of EMF measurements in determining enthalpy
  - (ii) application of EMF measurements in determining free energy of a cell
35. How to determine pH using the following electrodes
  - (i) Hydrogen electrode
  - (ii) Quinhydrone electrode
  - (iii) Glass electrode
36. Explain potentiometric redox titration.
37. Write short note on
  - (i) Liquid junctional potential
  - (ii) EMF measurements in determining equilibrium constant
38. Explain potentiometric acid-base titration..

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KARPAGAM

## UNIT-IV

### Physical chemistry

| S. No | Question                                                                                                                                           | Option 1          | Option 2         | Option 3                     | Option 4                    | Answer               |
|-------|----------------------------------------------------------------------------------------------------------------------------------------------------|-------------------|------------------|------------------------------|-----------------------------|----------------------|
| 1     | The electrical energy produced by a galvanic cell is given by the ----- of its electromotive force and the quantity of electricity which passes    | sum               | product          | sum and product              | divided                     | product              |
| 2     | Electricity is measured in                                                                                                                         | volts             | amperes          | coulombs                     | ohm                         | coulombs             |
| 3     | Electrical energy of a reversible cell originated from the ----- in the enthalpy of the cell reaction                                              | decrease          | increase         | decreases and then increases | increases and the decreases | decrease             |
| 4     | The standard free energy change of a cell reaction is given by                                                                                     | $\Delta G = -nFE$ | $\Delta F = nFE$ | $\Delta G = nFE$             | $\Delta G = FE$             | $\Delta G = -nFE$    |
| 5     | Knowing the EMF of the cell and the concentrations of reactants and products of the cell reaction, we can calculate the ----- of the cell reaction | free energy       | entropy          | enthalpy                     | equilibrium constant        | equilibrium constant |
| 6     | If the ----- of a particular cell is known the equilibrium constant of the cell reaction can be calculated                                         | standard EMF      | entropy          | free energy                  | enthalpy                    | standard EMF         |
| 7     | The potential of a hydrogen electrode in contact with a solution of $H^+$ ions involving the reaction is given by                                  | Nernst equation   | Gibbs equation   | Duhem equation               | Clausius equation           | Nernst equation      |
| 8     | Instead of taking quinone and hydroquinone, a small amount of ----- is taken                                                                       | quinhydrone       | glass            | hydroquinone                 | quinone                     | quinhydrone          |

|    |                                                                                                                       |                            |                                                        |                                                       |                                                                                      |                                                                                      |
|----|-----------------------------------------------------------------------------------------------------------------------|----------------------------|--------------------------------------------------------|-------------------------------------------------------|--------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------|
| 9  | Quinhydrone electrode is preferred to the -----electrode                                                              | hydrogen                   | calomel                                                | glass                                                 | platinum                                                                             | hydrogen                                                                             |
| 10 | The quinhydrone electrode is combined with a saturated-----electrode to form a cell                                   | hydrogen                   | calomel                                                | glass                                                 | platinum                                                                             | calomel                                                                              |
| 11 | The quinhydrone electrode cannot be used for solutions p H more than                                                  | 7                          | 9                                                      | 8                                                     | 5                                                                                    | 8                                                                                    |
| 12 | Hydroquinone ionises appreciably as an acid and also gets -----partly by atmospheric oxygen                           | oxidised                   | reduced                                                | oxidised and then reduced                             | reduced and then oxidised                                                            | oxidised                                                                             |
| 13 | The glass electrode is made of a special glass of relatively -----melting point and then -----electrical conductivity | high and low               | low and high                                           | high and high                                         | low and low                                                                          | low and high                                                                         |
| 14 | A solution of -----molar HCl which furnishes a constant hydrogen ion concentration                                    | 0.01                       | 0.1                                                    | 0.2                                                   | 0.001                                                                                | 0.1                                                                                  |
| 15 | The reference electrode employed is usually the -----in glass electrode                                               | hydrogen                   | calomel                                                | quinone                                               | hydroquinone                                                                         | calomel                                                                              |
| 16 | The EMF can be determined by means of a -----                                                                         | potentiometer              | conductivity meter                                     | d)p H meter                                           | voltmeter                                                                            | potentiometer                                                                        |
| 17 | Glass electrodes can be used in                                                                                       | strong oxidising solutions | weak oxidising solutions                               | strong reducing solutions                             | weak reducing solutions                                                              | strong oxidising solutions                                                           |
| 18 | calomel electrode consists of mercury,solid mercurous chloride and a solution of -----                                | KCl                        | NaCl                                                   | NaOH                                                  | KOH                                                                                  | KCl                                                                                  |
| 19 | Standard cell potential is measured                                                                                   | at a temperature of 25°C   | when ion concentration of aqueous reactants are 1.00 M | under the condition of 1.00 atm for gaseous reactants | at a temperature of 25°C, when ion concentration of aqueous reactants are 1.00 M and | at a temperature of 25°C, when ion concentration of aqueous reactants are 1.00 M and |

|    |                                                                                                                  |                                                                        |                                                                  |                                                               |                                                               |                                                                      |
|----|------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------|------------------------------------------------------------------|---------------------------------------------------------------|---------------------------------------------------------------|----------------------------------------------------------------------|
|    |                                                                                                                  |                                                                        |                                                                  |                                                               | under the condition of 1.00 atm for gaseous reactants         | under the condition of 1.00 atm for gaseous reactants                |
| 20 | The standard electrode commonly used are                                                                         | hydrogen electrode                                                     | quinhydrone electrode                                            | glass electrode                                               | nitrogen electrode                                            | hydrogen electrode                                                   |
| 21 | Which of the following is a different between glass electrode and hydrogen electrode                             | glass electrode measures $pH$ while hydrogen electrode compares $pH$   | different salt bridges are used                                  | glass electrode compares $pH$ , while electrode measures $pH$ | different electrolyte are used                                | glass electrode measures $pH$ while hydrogen electrode compares $pH$ |
| 22 | Why not $pH$ cannot be measured through potentiometer or voltmeter, but can be done through electronic voltmeter | resistance of the glass membrane is very high and the current is small | resistance of the glass membrane is low and the current is large | different salt bridges are used                               | glass electrode compares $pH$ , while electrode measures $pH$ | resistance of the glass membrane is low and the current is large     |
| 23 | $pH$ of the cell can be found out                                                                                | given the $E_{cell}$                                                   | given the $E^\circ G$                                            | given the $E^\circ$ value                                     | given the $E^\circ G$ and given the                           | given the $E^\circ G$ and given the                                  |

|    |                                                                                              |                                  |                                     |                            |                                                                          |                                     |
|----|----------------------------------------------------------------------------------------------|----------------------------------|-------------------------------------|----------------------------|--------------------------------------------------------------------------|-------------------------------------|
|    |                                                                                              | value                            |                                     |                            | Ecell value                                                              | Ecell value                         |
| 24 | How does electrical energy is obtained from the cell                                         | due to ions                      | chemical reaction inside the cell   | due to water               | chemical reaction outside the cell                                       | chemical reaction inside the cell   |
| 25 | Transfer of matter with respect to concentration is known as                                 | cell equilibrium                 | cell constant                       | cell concentration         | Electrolyte concentration                                                | cell concentration                  |
| 26 | Two like electrodes with different concentration are immersed in an electrolyte are known as | electrode-concentration cell     | electrolyte concentration cell      | cell concentration         | cell constant                                                            | electrolyte concentration cell      |
| 27 | Electrode-concentration cell are evidently independent of                                    | concentration of electrolyte     | concentration of ions               | concentration in electrode | electrolyte concentration cell                                           | concentration of electrolyte        |
| 28 | The whole process will be spontaneous only when the                                          | emf is negative                  | emf is partially negative           | emf is partially positive  | emf is positive                                                          | emf is negative                     |
| 29 | Solution with same electrolyte with different concentration is found in                      | electrolyte concentration cell   | electrode-concentration cell        | concentration of ions      | electrode-concentration cell and electrolyte concentration cell          | electrode-concentration cell        |
| 30 | Transfer of ions from one electrolytic solution to the other does not take place at          | concentration cell with transfer | concentration cell without transfer | emf is partially negative  | concentration cell without transfer and concentration cell with transfer | concentration cell without transfer |

|    |                                                                                            |                                         |                                         |                                         |                                                                          |                                         |
|----|--------------------------------------------------------------------------------------------|-----------------------------------------|-----------------------------------------|-----------------------------------------|--------------------------------------------------------------------------|-----------------------------------------|
| 31 | Transfer of ions takes place directly at                                                   | concentration cell without transfer     | concentration cell with transfer        | emf is partially negative               | concentration cell without transfer and concentration cell with transfer | concentration cell with transfer        |
| 32 | Liquid junction potential depends on                                                       | transference number of anion and cation | removal of anion and cation             | transfer of only anion                  | transfer of only cation                                                  | transference number of anion and cation |
| 33 | When does L.J.P becomes null/zero                                                          | $t_+ = t_-$                             | $t_+ \neq t_-$                          | $t_+ > t_-$                             | $t_+ < t_-$                                                              | $t_+ = t_-$                             |
| 34 | When does L.J.P becomes negative                                                           | $t_+ = t_-$                             | $t_+ \neq t_-$                          | $t_+ > t_-$                             | $t_+ < t_-$                                                              | $t_+ > t_-$                             |
| 35 | When does L.J.P becomes positive                                                           | $t_+ = t_-$                             | $t_+ \neq t_-$                          | $t_+ > t_-$                             | $t_+ < t_-$                                                              | $t_+ < t_-$                             |
| 36 | Which among the following electrolytes have the same transfer number of anions and cations | potassium chloride and ammonium nitrate | potassium nitrate and ammonium chloride | potassium sulphate and ammonium nitrate | potassium nitrate and ammonium sulphate                                  | potassium sulphate and ammonium nitrate |
| 37 | Reversibility of electrode with respect to cation will determine                           | transfer number of anion                | transfer number of cation               | emf is partially negative               | concentration cell with transfer                                         | transfer number of anion                |
| 38 | Reversibility of electrode with respect to anion will determine                            | transfer of anion                       | transfer of cation                      | emf is partially negative               | concentration cell with transfer                                         | transfer of cation                      |
| 39 | Potential of any electrode depends on                                                      | concentration of ions                   | concentration of anion                  | concentration of cation                 | transfer of cation                                                       | concentration of ions                   |

|    |                                                                                                                             |                                          |                      |                                          |                                                                  |                                                                  |
|----|-----------------------------------------------------------------------------------------------------------------------------|------------------------------------------|----------------------|------------------------------------------|------------------------------------------------------------------|------------------------------------------------------------------|
| 40 | Why an indicator is not required for a potentiometric titration that is carried out with a coloured solution                | no indicator required                    | self indicator       | no indicator required and self indicator | External indicator                                               | no indicator required and self indicator                         |
| 41 | In a potentiometric titration, the EMF slowly changes when                                                                  | the end point changes                    | before the end point | when the end point approaches            | emf is partially negative                                        | when the end point approaches                                    |
| 42 | Which among the following electrode is used as oxidation- reduction electrode                                               | copper                                   | silver-silver        | platinum                                 | tin                                                              | silver-silver                                                    |
| 43 | The Liquid junction potential becomes positive                                                                              | $t = t^+$                                | $t \neq t^+$         | $t > t^+$                                | $t < t^+$                                                        | $t < t^+$                                                        |
| 44 | Which among the following will come under potentiometric titration                                                          | acid-base titration                      | redox titration      | precipitation titration                  | acid-base titration, redox titration and precipitation titration | acid-base titration, redox titration and precipitation titration |
| 45 | Which among the following is taken as indicator in the precipitation titration of silver nitrate against potassium chloride | potassium electrode                      | silver electrode     | hydrogen electrode                       | silver-silver electrode                                          | silver electrode                                                 |
| 46 | Organic compounds which can exist in oxidised form as well as reduced form are said to be                                   | amphoteric indicator                     | redox indicator      | acid indicator                           | base indicator                                                   | redox indicator                                                  |
| 47 | Which is that one main feature of redox indicators                                                                          | different concentration                  | different colour     | different cell                           | same colour                                                      | different colour                                                 |
| 48 | When does a redox indicator can be successfully used                                                                        | measured potential to be about 0.03 volt | more than 0.03 volt  | 0 potential                              | 1                                                                | more than 0.03 volt                                              |
| 49 | Which among the following is a redox indicator for titration of ferrous ions against dichromate                             | diphenylamine                            | KCl                  | NaCl                                     | CaCl <sub>2</sub>                                                | diphenylamine                                                    |

|    |                                                                                                                                                                                       |                                                                          |                                                        |                                                                              |                                                                                                                                     |                                                                                                                                     |
|----|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------|--------------------------------------------------------|------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------|
|    | ions                                                                                                                                                                                  |                                                                          |                                                        |                                                                              |                                                                                                                                     |                                                                                                                                     |
| 50 | Determination of transport number, valency of a ion, coefficient of electrolyte are purely based upon                                                                                 | cell constant                                                            | equilibrium constant                                   | EMF                                                                          | Concentration                                                                                                                       | cell constant                                                                                                                       |
| 51 | In given reaction $2\text{Cr(s)} + 3\text{Cu}^{2+}(\text{aq}) \rightarrow 3\text{Cr}^{3+}(\text{aq}) + 3\text{Cr(s)}$ which reaction occurs of the cathode in an electrochemical cell | reduction of $\text{Cu}^{2+}(\text{aq})$                                 | reduction of $\text{Cu(s)}$                            | oxidation of $\text{Cu}^{3+}(\text{aq})$                                     | oxidation of $\text{Cr(s)}$                                                                                                         | oxidation of $\text{Cu}^{3+}(\text{aq})$                                                                                            |
| 52 | The site of oxidation in an electrochemical cell is                                                                                                                                   | the anode                                                                | cathode                                                | electrodes                                                                   | salt bridge                                                                                                                         | the anode                                                                                                                           |
| 53 | Which statement below is not true for the reaction $\text{Fe}^{3+} + \text{e}^{-} \rightarrow \text{Fe}^{2+}$                                                                         | $\text{Fe}^{3+}$ is being reduced                                        | the oxidation state of Fe has charged                  | $\text{Fe}^{3+}$ could be referred to as an oxidizing agent in this reaction | both $\text{Fe}^{2+}$ and $\text{Fe}^{2+}$ are called anions                                                                        | both $\text{Fe}^{2+}$ and $\text{Fe}^{2+}$ are called anions                                                                        |
| 54 | In any electrochemical cell, the cathode is always                                                                                                                                    | non metal                                                                | attached to battery                                    | the electrode at which some species gain electrons                           | the electrode at which some species lose electrons                                                                                  | non metal                                                                                                                           |
| 55 | Electrical energy forced in electrochemical cell is                                                                                                                                   | Spontaneous                                                              | Non-Spontaneous                                        | Exothermic                                                                   | Endothermic                                                                                                                         | Non-Spontaneous                                                                                                                     |
| 56 | Which of the following shows a metal being oxidized                                                                                                                                   | $2\text{Na} + 2\text{H}_2\text{O} \rightarrow 2\text{NaOH} + \text{H}_2$ | $\text{Cu} \rightarrow \text{Cu}^{2+} + 2\text{e}^{-}$ | $\text{Cu}^{2+} + 2\text{e}^{-} \rightarrow \text{Cu}$                       | $\text{Cu} \rightarrow \text{Cu}^{2+} + 2\text{e}^{-}$ and $2\text{Na} + 2\text{H}_2\text{O} \rightarrow 2\text{NaOH} + \text{H}_2$ | $\text{Cu} \rightarrow \text{Cu}^{2+} + 2\text{e}^{-}$ and $2\text{Na} + 2\text{H}_2\text{O} \rightarrow 2\text{NaOH} + \text{H}_2$ |
| 57 | A voltaic cell has an $E^{\circ}$ value of -1.00 v the reaction is                                                                                                                    | spontaneous                                                              | has positive $\Delta G^{\circ}$                        | has negative $\Delta G^{\circ}$                                              | 0                                                                                                                                   | has positive $\Delta G^{\circ}$                                                                                                     |
| 58 | Which of the following can we use to measure Ph?                                                                                                                                      | a glass electrode                                                        | a concentration cell                                   | a hydrogen electrode                                                         | a glass electrode, a concentration                                                                                                  | a glass electrode, a concentration                                                                                                  |

|    |                                                                                                                                                                  |                                                                        |                                                                  |                                 |                                                          |                                                                  |
|----|------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------|------------------------------------------------------------------|---------------------------------|----------------------------------------------------------|------------------------------------------------------------------|
|    |                                                                                                                                                                  |                                                                        |                                                                  |                                 | cell and a hydrogen electrode                            | cell and a hydrogen electrode                                    |
| 59 | What is $\Delta G^\circ$ at 298k for the reaction $\text{Hg(l)} + 2\text{Fe}^{3+}(\text{aq}) \rightarrow \text{Hg}^{2+}(\text{aq}) + 2\text{Fe}^{2+}(\text{aq})$ | positive 314 Kelectrode                                                | negative 16 KJ                                                   | negative 314 Kelectrode         | positive 16 KJ                                           | positive 16 KJ                                                   |
| 60 | Why not pH cannot be measured through potentiometer or voltmeter, but can be done through electronic voltmeter                                                   | resistance of the glass membrane is very high and the current is small | resistance of the glass membrane is low and the current is large | different salt bridges are used | glass electrode compares pH, while electrode measures Ph | resistance of the glass membrane is low and the current is large |

## UNIT V

### Surface Chemistry

**Surface Chemistry:**Physical adsorption, chemisorptions adsorption isotherms(Langmuir and Freundlich), nature of adsorbed state. Qualitative discussion of BET

#### Surface chemistry

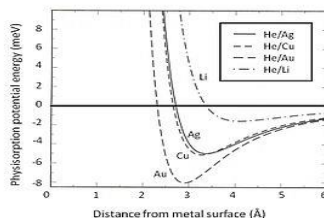
##### Physical adsorption

The fundamental interacting force of physisorption is caused by van der Waals force. Even though the interaction energy is very weak ( $\sim 10\text{--}100$  meV), physisorption plays an important role in nature. For instance, the van der Waals attraction between surfaces and foot-hairs of geckos provides the remarkable ability to climb up vertical walls. Van der Waals forces originate from the interactions between induced, permanent or transient electric dipoles.

In comparison with chemisorption, in which the electronic structure of bonding atoms or molecules is changed and covalent or ionic bonds form, physisorption, generally speaking, can only be observed in the environment of low temperature (thermal energy at room temperature  $\sim 26$  meV) and the absence of the relatively strong chemisorptions. In practice, the categorisation of a particular adsorption as physisorption or chemisorption depends principally on the binding energy of the adsorbate to the substrate.

To give a simple illustration of physisorption, we can first consider an adsorbed hydrogen atom in front of a perfect conductor, as shown in Fig. 1. A nucleus with positive charge is located at  $\mathbf{R} = (0, 0, Z)$ , and the position coordinate of its electron,  $\mathbf{r} = (x, y, z)$  is given with respect to the nucleus. The adsorption process can be viewed as the interaction between this hydrogen atom and its image charges of both the nucleus and electron in the conductor. As a result, the total electrostatic energy is the sum of attraction and repulsion terms:

##### Physisorption potential



Even though the van der Waals interaction is attractive, as the adsorbed atom moves closer to the surface the wavefunction of electron starts to overlap with that of the surface atoms. Further the energy of the system will increase due to the orthogonality of wavefunctions of the approaching atom and surface atoms.

This Pauli exclusion and repulsion are particularly strong for atoms with closed valence shells that dominate the surface interaction. As a result, the minimum energy of physisorption must be found by the balance between the long-range van der Waals attraction and short-range Pauli repulsion. For instance, by separating the total interaction of physisorption into two contributions- a short-range term depicted by Hartree–Fock theory and a long-range van der Waals attraction, the equilibrium position of physisorption for rare gases adsorbed on jellium substrate can be determined.<sup>[5]</sup> Fig. 2 shows the physisorption potential energy of He adsorbed on Ag, Cu, and Au substrates which are described by the jellium model with different densities of smear-out background positive charges. It can be found that the weak van der Waals interaction leads to shallow attractive energy wells (<10 meV). One of the experimental methods for exploring physisorption potential energy is the scattering process, for instance, inert gas atoms scattered from metal surfaces. Certain specific features of the interaction potential between scattered atoms and surface can be extracted by analyzing the experimentally determined angular distribution and cross sections of the scattered particles.

## Comparison with chemisorption

- Physisorption is a general phenomenon and occurs in any solid/fluid or solid/gas system. Chemisorption is characterized by chemical specificity.

- In physisorption, perturbation of the electronic states of adsorbent and adsorbate is minimal. For chemisorption, changes in the electronic states may be detectable by suitable physical means.
- Typical binding energy of physisorption is about 10–100 meV. Chemisorption usually forms bonding with energy of 1–10 eV.
- The elementary step in physisorption from a gas phase does not involve an activation energy. Chemisorption often involves an activation energy.
- For physisorption, under appropriate conditions, gas phase molecules can form multilayer adsorption. In chemisorption, molecules are adsorbed on the surface by valence bonds and only form monolayer adsorption.

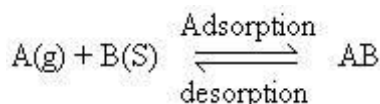
## Langmuir adsorption isotherm

In 1916, Irving Langmuir proposed another Adsorption Isotherm which explained the variation of Adsorption with pressure. Based on his theory, he derived Langmuir Equation which depicted a relationship between the number of active sites of the surface undergoing adsorption and pressure.

### Assumptions of Langmuir Isotherm

Langmuir proposed his theory by making following assumptions.

1. Fixed number of vacant or adsorption sites are available on the surface of solid.
2. All the vacant sites are of equal size and shape on the surface of adsorbent.
3. Each site can hold maximum of one gaseous molecule and a constant amount of heat energy is released during this process.
4. Dynamic equilibrium exists between adsorbed gaseous molecules and the free gaseous molecules.



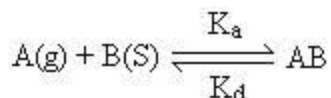
Where A (g) is unadsorbed gaseous molecule, B(s) is unoccupied metal surface and AB is Adsorbed gaseous molecule.

5. Adsorption is monolayer or unilayer.

## Derivations of the Langmuir Adsorption Equation

### Calculation of Equilibrium Constant

Langmuir proposed that dynamic equilibrium exists between adsorbed gaseous molecules and the free gaseous molecules. Using the equilibrium equation, equilibrium constant can be calculated.



Where  $K_a$  represents equilibrium constant for forward reaction and  $K_d$  represents equilibrium constant for backward direction.

According to Kinetic theory,

Rate of forward reaction =  $K_a [A] [B]$

Rate of backward reaction =  $K_d [AB]$

At equilibrium, Rate of forward reaction is equal to Rate of backward reaction

$$K_a [A] [B] = K_d [AB]$$

$$\text{Or, } \frac{K_a}{K_d} = \frac{[AB]}{[A][B]}$$

$$K = \frac{K_a}{K_d} = \frac{[AB]}{[A][B]}$$

The above equation represents the equilibrium constant for distribution of adsorbate between the surface and the gas phase.

### ***Derivation***

Langmuir Equation which depicts a relationship between the number of active sites of the surface undergoing adsorption (i.e. extent of adsorption) and pressure.

To derive Langmuir Equation and new parameter '  $\theta$  ' is introduced. Let  $\theta$  the number of sites of the surface which are covered with gaseous molecules. Therefore, the fraction of surface which are unoccupied by gaseous molecules will be  $(1 - \theta)$ .

Now, Rate of forward direction depends upon two factors: Number of sites available on the surface of adsorbent,  $(1 - \theta)$  and Pressure,  $P$ . Therefore rate of forward reaction is directly proportional to both mentioned factors.

$$\text{Rate of forward reaction} \propto P (1 - \theta)$$

$$\text{Rate of adsorption} \propto P (1 - \theta)$$

$$\text{Or, Rate of adsorption} = K_a P (1 - \theta)$$

Similarly, Rate of backward reaction or Rate of Desorption depends upon number of sites occupied by the gaseous molecules on the surface of adsorbent.

$$\text{Rate of desorption} \propto \theta$$

$$\text{Or, Rate of desorption} = K_d \theta$$

At equilibrium, rate of adsorption is equal to rate of desorption.

$$K_a P (1 - \theta) = K_d \theta$$

We can solve the above equation to write it in terms of  $\theta$ .

$$K_a P - K_a P \theta = K_d \theta$$

$$K_a P = K_a P \theta + K_d \theta$$

$$K_a P = (K_d + K_a P) \theta$$

$$\theta = \frac{K_a P}{K_d + K_a P}$$

Divide numerator and denominator on RHS by  $K_d$ , we get

$$\theta = \frac{\frac{K_a P}{K_d}}{\frac{K_d}{K_d} + \frac{K_a P}{K_d}}$$

Now put

$$K = \frac{K_a}{K_d}$$

in above equation we get

$$\theta = \frac{KP}{1 + KP}$$

Langmuir Adsorption Equation

This is known as Langmuir Adsorption Equation.

### ***Alternate form of Langmuir Adsorption Equation***

Langmuir adsorption equation can be written in an alternate form in terms of volume of gas adsorbed. Let  $V$  be volume of gas adsorbed under given sets of conditions of temperature and pressure and  $V_{\text{mono}}$  be the adsorbed volume of gas at high pressure conditions so as to cover the surface with a unilayer of gaseous molecules.

$$\theta = \frac{V}{V_{mono}}$$

Substituting the value of  $\theta$  in Langmuir equation

$$\frac{V}{V_{mono}} = \frac{KP}{1 + KP}$$

$$\text{Or } V_{mono} = 1 + \frac{1}{KP}$$

Or in terms of pressure P we get,

$$\frac{P}{V} = \frac{P}{V_{mono}} + \frac{1}{KV_{mono}}$$

Langmuir Adsorption Equation in alternate form

Thus, if we plot a graph between  $P/V$  Vs  $P$ , we will obtain a straight line with

$$\text{slope} = \frac{1}{V_{mono}} \quad \text{and} \quad \text{Intercept} = \frac{1}{KV_{mono}}$$

## Limitations of Langmuir Adsorption Equation

1. The adsorbed gas has to behave ideally in the vapor phase. This condition can be fulfilled at low pressure conditions only. Thus Langmuir Equation is valid under low pressure only.
2. Langmuir Equation assumes that adsorption is monolayer. But, monolayer formation is possible only under low pressure condition. Under high pressure condition the assumption breaks down as gas molecules attract more and more molecules towards each other. *BET theory* proposed by Brunauer, Emmett and Teller explained more realistic multilayer adsorption process.
3. Another assumption was that all the sites on the solid surface are equal in size and shape and have equal affinity for adsorbate molecules i.e. the surface of solid is homogeneous. But we all know that in real solid surfaces are heterogeneous.

4. Langmuir Equation assumed that molecules do not interact with each other. This is impossible as weak force of attraction exists even between molecules of same type.
5. The adsorbed molecules has to be localized i.e. decrease in randomness is zero ( $\Delta S = 0$ ). This is not possible because on adsorption liquefaction of gases taking place, which results into decrease in randomness but the value is not zero.

From above facts we can conclude that, **Langmuir equation is valid under low pressure conditions.**

## Freundlich Adsorption Equation: A Special Case of Langmuir Equation

We consider Langmuir Equation

$$\theta = \frac{KP}{1 + KP}$$

At low pressure value of  $KP \ll 1$ . Therefore,

$$\theta = KP \quad \text{Or} \quad \theta \propto P \quad \dots (1)$$

The above equation shows linear variation between extent of adsorption of gas and pressure.

At high pressure value of  $KP \gg 1$

$$\therefore \theta = \frac{KP}{KP} = 1 \quad \dots (2)$$

The extent of adsorption,  $\theta$  is independent of pressure at high pressure conditions. The reaction at this stage becomes zero order

Combining the results of equation (4) and (5), we can conclude that

$$\theta = Kp^{0-1}$$

$$\text{or } \theta = Kp^{1/n} \quad \dots (3)$$

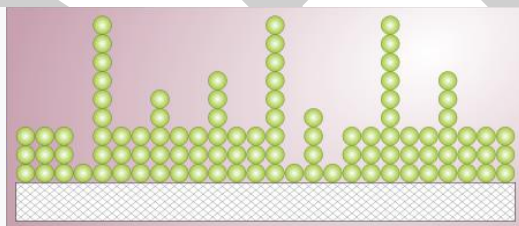
Equation (3) is in agreement with Freundlich adsorption equation.

We can say that Freundlich adsorption equation is a special case of Langmuir equation.

## Qualitative discussion of BET

**Brunauer–Emmett–Teller (BET) theory** aims to explain the physical adsorption of gas molecules on a solid surface and serves as the basis for an important analysis technique for the measurement of the specific surface area of materials. In 1938, Stephen Brunauer, Paul Hugh Emmett, and Edward Teller published the first article about the BET theory in the Journal of the American Chemical Society.<sup>[1]</sup> The BET theory applies to systems of multi layer adsorption, and usually utilizes probing gases that do not chemically react with material surfaces as adsorbates to quantify specific surface area. Nitrogen is the most commonly employed gaseous adsorbate used for surface probing by BET methods. For this reason, standard BET analysis is most often conducted at the boiling temperature of N<sub>2</sub> (77 K). Further probing adsorbates are also utilized, albeit with lower frequency, allowing the measurement of surface area at different temperatures and measurement scales. These have included argon, carbon dioxide, and water. Specific surface area is a scale-dependent property, with no single true value of specific surface area definable, and thus quantities of specific surface area determined through BET theory may depend on the adsorbate molecule utilized and its adsorption cross section.

## Concept



BET model of multilayer adsorption, that is, a random distribution of sites covered by one, two, three, etc., adsorbate molecules.

The concept of the theory is an extension of the Langmuir theory, which is a theory for monolayer molecular adsorption, to multilayer adsorption with the following hypotheses:

1. gas molecules physically adsorb on a solid in layers infinitely;
2. there is no interaction between each adsorption layer; and
3. the Langmuir theory can be applied to each layer.

## POSSIBLE QUESTIONS

### Part A(Each question carry one mark)

1. The substance adsorbed or attached is called the-----

a)adsorbent

**b)adsorbate** c)sorbate

d)desorption

2. Ps called as

- a)pressure      **b)saturation pressure**      c)temperature      d)volume

3. In physical adsorption, there a regular decrease in extent of adsorption as-----increases

- a)pressure      b)concentration      c)volume      **d)temperature**

4. By application of the BET theory it is possible to determine the inner surface of hardened -----paste

- a)composite      **b)cement**      c)matrix      d)concrete

5. . charcoal and silica gel are excellent-----

- a)adsorbate      **b)adsorbent**      c)sorbent      d)absorbate

6. Freundlich adsorption isotherm fails if concentration of adsorbate is very-----

- a)high**      b)low      c)medium      d)average

7. According to Langmuir if a bond is weak, a-----adsorption takes place

- a)chemical      **b)physical**      c)oxidation      d)reduction

8. The surface area of catalysts is an important factor in----- activity

- a)activity coefficients      b)viscosity coefficients      c)diffusion coefficients  
**d)catalytic**

9. charcoal and silica gel are excellent-----

- a)adsorbate      **b)adsorbe**      c)sorbent      d)absorbate

10. Freundlich adsorption isotherm fails if concentration of adsorbate is very-----

- a)high**      b)low      c)medium      d)average

11. Activated charcoal is used for removing-----

- a)colouring matter**      b)dust      c)sediments      d)coagulant

12. In order to increase the rate of adsorption,-----is very necessary

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a)catalyst b)reactant c)**activation** d)sorption

13. The adsorption is accompanied by a -----in residual surface forces

a)**decrease** b)increase c)increase and then decreases d)decrease and then increases

14.  $x/m$  is known as

a)nature of adsorption b)**extent of adsorption** c)rate of adsorption  
d)magnitude of adsorption

15. According to Langmuir if a bond is weak, a-----adsorption takes place

a)chemical b)**physical** c)oxidation d)reduction

16. The surface area of catalysts is an important factor in----- activity

a)activity coefficients b)viscosity coefficients c)diffusion coefficients  
d)**catalytic**

17. The adsorption is accompanied by a -----in residual surface forces

a)**decrease** b)increase c)increase and then decreases d)decrease and then increases

18.  $x/m$  is known as

a)nature of adsorption b)**extent of adsorption** c)rate of adsorption  
d)magnitude of adsorption

19. BET finds application in

a)**catalysis** b)activity of coefficients c)diffusion coefficients d)viscosity coefficient

20. The surface area of catalysts is an important factor in----- activity

a)activity coefficients b)viscosity coefficients c)diffusion coefficients d)**catalytic**

**Part B(Each question carry 2 marks)**

- 21.What is known as physical adsorption?
- 22.What is known as chemical adsorption?
- 23.Define adsorption.
- 24.Define absorption.
- 25.Define adsorbate.
- 26.Define adsorbent.
- 27.Write down an equation for BET.
- 28.What is the condition for Freundlich adsorption isotherm at high pressure.
- 29.Write down the condition for Langmuir adsorption isotherm at low pressure.

**Part C (Each question carry 8 marks)**

30. Differentiate physical adsorption with chemical adsorption.
- 31.Explain the postulates of Langmuir adsorption isotherm and derive the same .
- 32.What is adsorption and absorption.Derive Freundlich adsorption isotherm and mention its conditions.
33. Discuss BET in detail.
- 34.What is adsorption and absorption.Derive Freundlich adsorption isotherm and mention its conditions.
- 35.What is adsorption and absorption.Derive Freundlich adsorption isotherm and mention its conditions.
- 36.What is adsorption and absorption. Explain BET

**CLASS: II- B.SC CHEMISTRY**  
**COURSE CODE:18CHU301**

**COURSE NAME: PHYSICAL CHEMISTRY III**  
**UNIT-V**

**BATCH: 2018-2021**

KAHE

## UNIT-V

### Physical chemistry

| S.No | Question                                                                                                                       | Option 1            | Option 2            | Option 3                    | Option 4                    | Answer              |
|------|--------------------------------------------------------------------------------------------------------------------------------|---------------------|---------------------|-----------------------------|-----------------------------|---------------------|
| 1    | The phenomenon of concentration or assimilation of gas at the surface of a solid with which it is in close proximity is called | adsorption          | absorption          | sorption                    | desorption                  | adsorption          |
| 2    | The adsorption is accompanied by a -----<br>-----in residual surface forces                                                    | decrease            | increase            | increase and then decreases | decrease and then increases | decrease            |
| 3    | The adsorption of a gas on a solid is sometimes called                                                                         | inclusion           | occlusion           | sorption                    | desorption                  | occlusion           |
| 4    | The material providing the surface upon which adsorption occurs is known as                                                    | adsorbent           | adsorbate           | sorbate                     | desorption                  | adsorbent           |
| 5    | The substance adsorbed or attached is called                                                                                   | adsorbent           | adsorbate           | sorbate                     | desorption                  | adsorbate           |
| 6    | The removal of adsorbed substance from the surface is called                                                                   | inclusion           | occlusion           | sorption                    | desorption                  | desorption          |
| 7    | The amount of heat evolved when 1 mole of any gas is adsorbed on solid adsorbent surface is called-----of adsorption           | entropy             | enthalpy            | free energy                 | work function               | enthalpy            |
| 8    | When the concentration of the adsorbate is more on the surface of the adsorbent than the bulk, it is called                    | positive adsorption | negative adsorption | inclusion                   | occlusion                   | positive adsorption |
| 9    | When the concentration of the adsorbate is less on the surface of the adsorbent than the bulk, it is called                    | positive adsorption | negative adsorption | inclusion                   | occlusion                   | negative adsorption |
| 10   | Adsorption is a                                                                                                                | surface phenomena   | bulk phenomena      | layer phenomena             | negative phenomena          | surface phenomena   |

|    |                                                                                                               |                      |                      |                    |                          |                      |
|----|---------------------------------------------------------------------------------------------------------------|----------------------|----------------------|--------------------|--------------------------|----------------------|
| 11 | Adsorption is a                                                                                               | slow process         | fast process         | negative process   | positive process         | fast process         |
| 12 | Absorption is a-----process compared to adsorption                                                            | slow                 | fast                 | medium             | average                  | slow                 |
| 13 | Adsorption involves the-----into the interior of the matter                                                   | diffusion            | penetration          | occlusion          | inclusion                | diffusion            |
| 14 | In-----, a gas gets adsorbed on the solid only if it forms chemical bonds                                     | adsorption           | desorption           | chemisorption      | absorption               | chemisorption        |
| 15 | Greater the surface area of the adsorbent, -----is its adsorption capacity                                    | greater              | lower                | average            | medium                   | greater              |
| 16 | charcoal and silica gel are excellent                                                                         | adsorbate            | adsorbent            | sorbent            | absorbate                | adsorbent            |
| 17 | charcoal is -----porous and hence possess-----surface areas                                                   | high and large       | low and small        | high and small     | low and large            | high and large       |
| 18 | The extent of adsorption depends upon the                                                                     | temperature          | pressure             | volume             | concentration            | pressure             |
| 19 | Adsorption isotherm is a graph plotted between magnitude of adsorption and -----at constant temperature       | temperature          | pressure             | volume             | concentration            | pressure             |
| 20 | $P_s$ is called as                                                                                            | pressure             | saturation pressure  | temperature        | volume                   | saturation pressure  |
| 21 | $P_s$ becomes ----- to the rate of desorption and further increase of pressure does not alter the equilibrium | equal                | greater              | lower              | greater than or equal to | equal                |
| 22 | $x/m$ is known as                                                                                             | nature of adsorption | extent of adsorption | rate of adsorption | magnitude of adsorption  | extent of adsorption |
| 23 | At low pressure in Freundlich isotherm the graph is almost a                                                  | parallel             | perpendicular        | straight line      | slope                    | straight line        |
| 24 | At high pressure in Freundlich isotherm the graph is almost-----to X-axis                                     | parallel             | perpendicular        | straight line      | slope                    | parallel             |
| 25 | At intermediate pressure in Freundlich isotherm $x/m$ depends on                                              | 0 to 1               | 1 to 2               | 0 to 2             | 0 to 3                   | 0 to 1               |

|    |                                                                                                           |                   |               |                              |                             |                   |
|----|-----------------------------------------------------------------------------------------------------------|-------------------|---------------|------------------------------|-----------------------------|-------------------|
| 26 | Freundlich adsorption isotherm fails if pressure is                                                       | high              | low           | medium                       | average                     | high              |
| 27 | Freundlich adsorption isotherm fails if concentration of adsorbate is very                                | high              | low           | medium                       | average                     | high              |
| 28 | Adsorption isobar is a graph plotted between magnitude of adsorption and ----- at constant pressure       | temperature       | volume        | concentration                | composition                 | temperature       |
| 29 | The amount absorbed x/m should -----with an increase in temperature                                       | decrease          | increase      | remain constant              | decrease and then increases | decrease          |
| 30 | chemisorption requires                                                                                    | activation energy | entropy       | enthalpy                     | free energy                 | activation energy |
| 31 | adsorption-----can be used to distinguish between physical and chemical adsorption                        | isobar            | isotherm      | isochore                     | isotherm and isochore       | isobar            |
| 32 | In physical adsorption, there a regular decrease in extent of adsorption as-----increases                 | pressure          | concentration | volume                       | temperature                 | temperature       |
| 33 | In chemisorption, there is initial increase and then decrease in extent of adsorption as ----- -increases | pressure          | concentration | volume                       | temperature                 | temperature       |
| 34 | In order to increase the rate of adsorption,----- is very necessary                                       | catalyst          | reactant      | activation                   | sorption                    | activation        |
| 35 | mechanical rubbing can be done to increase                                                                | speed             | activation    | reaction                     | speed of catalyst           | activation        |
| 36 | Adsorption from solutions decreases with rise of temperature and decrease in-----of solution              | increase          | decrease      | decreases and then increases | increases and the decreases | decrease          |
| 37 | Activated charcoal is used for removing                                                                   | colouring matter  | dust          | sediments                    | coagulant                   | colouring matter  |
| 38 | Activated charcoal is used in                                                                             | sedimentation     | coagulation   | colouration                  | gas masks                   | gas masks         |
| 39 | Adsorption process is used in production of vacuum by using activated charcoal in -----                   | condenser         | distillation  | Dewar's                      | air condenser               | Dewar's           |

|    |                                                                                                                                                                  |                 |                     |                              |                                |                     |
|----|------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------|---------------------|------------------------------|--------------------------------|---------------------|
|    | flask                                                                                                                                                            |                 |                     |                              |                                |                     |
| 40 | Lake test for aluminium is based upon -----<br>-of litmus colour                                                                                                 | absorption      | adsorption          | sorption                     | desorption                     | adsorption          |
| 41 | According to Langmuir, valencies at the surface of adsorbent atoms are                                                                                           | fully satisfied | not fully satisfied | partially satisfied          | satisfied                      | not fully satisfied |
| 42 | According to Langmuir if a bond is weak, a-----<br>-----adsorption takes place                                                                                   | chemical        | physical            | oxidation                    | reduction                      | physical            |
| 43 | According to Langmuir if a bond is strong, a-----<br>-----adsorption takes place                                                                                 | chemical        | physical            | oxidation                    | reduction                      | chemical            |
| 44 | The residual valency force on the surface of adsorbent is effective only up to a small distance and hence, the adsorbed gas layer is only-----<br>molecule thick | one             | two                 | zero                         | three                          | one                 |
| 45 | The phenomenon of adsorption consists of two opposing processes namely condensation and -----<br>-----                                                           | distillation    | evaporatio<br>n     | circulation                  | floatation                     | evaporatio<br>n     |
| 46 | When the adsorption starts the whole adsorbent surface is bare and consequently the initial rate of absorption is-----                                           | lowest          | highest             | constant                     | medium                         | highest             |
| 47 | When the adsorption starts the whole adsorbent surface is bare and consequently the rate of condensation-----                                                    | increases       | decreases           | decreases and then increases | increases and the decreases    | decreases           |
| 48 | The rate of evaporation of the condensed molecules gradually                                                                                                     | increases       | decreases           | decreases and then increases | increases and the decreases    | increases           |
| 49 | A dynamic equilibrium is set up, when the rate of condensation becomes equal to the rate of -----<br>-----                                                       | distillation    | evaporatio<br>n     | circulation                  | floatation                     | evaporatio<br>n     |
| 50 | P/w against P, we should get a                                                                                                                                   | curve           | straight line       | deviated line                | curve as well as deviated line | straight line       |

|    |                                                                                                                        |                       |                           |                         |                       |                   |
|----|------------------------------------------------------------------------------------------------------------------------|-----------------------|---------------------------|-------------------------|-----------------------|-------------------|
| 51 | Brunauer–Emmett–Teller (BET) theory aims to explain the physical adsorption of gas molecules on a                      | solid                 | liquid                    | gaseous                 | semi liquid           | solid             |
| 52 | The concept of BET is extension of                                                                                     | Freundlich isotherm   | Langmuir isotherm         | Emmett isotherm         | Teller isotherm       | Langmuir isotherm |
| 53 | The BET method is widely used in                                                                                       | surface               | adsorption                | material                | polymer               | surface           |
| 54 | According to BET,adsorptions occur only on well-defined sites of the sample -----                                      | bulk                  | surface                   | surface as well as bulk | layered               | surface           |
| 55 | According to BET,the uppermost molecule layer is in equilibrium with which of the following?                           | liquid                | gas                       | solid                   | gas as well as liquid | gas               |
| 56 | A kinetically-limited process is                                                                                       | adsorption            | desorption                | sorption                | adsorbent             | desorption        |
| 57 | At the saturation pressure, the molecule layer number tends to                                                         | infinity              | low                       | high                    | low and high          | infinity          |
| 58 | <a href="#">By application of the BET theory it is possible to determine the inner surface of hardened ----- paste</a> | composite             | cement                    | matrix                  | concreate             | cement            |
| 59 | BET finds application in                                                                                               | catalysis             | activity of coefficient s | diffusion coefficient s | viscosity coefficient | catalysis         |
| 60 | The surface area of catalysts is an important factor in which of the activity?                                         | activity coefficients | viscosity coefficient s   | diffusion coefficient s | catalytic             | catalytic         |

Reg. No.:-----

[18CHU301]

**KARPAGAM ACADEMY OF HIGHER EDUCATION**  
(Deemed to be University) (Established  
Under Section 3 of UGC Act 1956)  
**COIMBATORE-21**

**B.Sc. Degree Internal Examination, July 2019**  
(For the candidates admitted from 2018& onwards)

**II B.Sc CHEMISTRY**  
**INTERNAL EXAM I**  
**PHYSICAL CHEMISTRY III**  
(Phase Equilibria and Chemical Kinetics)

**Time: 2 hours**

**Maximum: 50marks**

**PART- A (20 x 1= 20 Marks)**

**Answer ALL the Questions**

1. Calcium component is an example for  
a) one component    b) two component    c) three component    d) four component
2. Sulphur exist in  
a) three phase    b) one phase    c) two phase    d) four phase
3. Water is an example for  
a) three component    b) two component    c) one component    d) four component
4. J.W.Gibbs enunciated the phase rule in  
a) 1876    b) 1875    c) 1874    d) 1873
5. Ferric chloride -water is an example for  
a) congruent melting point    b) triple point    c) incongruent melting point    d) eutectic point
6. Sugar and water mixture is an  
a) two component    b) one component    c) three component    d) four component
7. If  $F=1$ , the system is called  
a) bivariant    b) univariant    c) invariant    d) trivariant
8. Reduced phase rule is otherwise known as  
a) condensed phase rule    b) phase rule equation  
c) Gibbs phase rule equation    d) expanded phase rule
9.  $\Delta H_v$  is known as  
a) molar heat of sublimation    b) molar heat of vaporization  
b) c) molar heat of fusion    d) molar heat of enthalpy
10. Temperature-composition axis is drawn for  
a) two component    b) one component    c) three component    d) one and two component

11. Application of the phase rule to a system corresponding to a point in the two phase region gives  $F =$   
 a) 3                      b) 2                      c) 1                      d) 0
12. Number of phases in water  $\leftrightarrow$  vapour is  
 a) 1                      b) 2                      c) 3                      d) 4
13. Potassium iodide-water is an example for  
 a) Eutectic system    b) one component    c) three component    d) four component
14. Transition temperature is otherwise known as  
 a) peritectic temperature                      b) melting temperature  
 c) sublimation temperature                      d) vapour pressure temperature
15. The process of separating mixtures by repeated distillation and condensation is known as  
 a) steam distillation    b) fractional distillation    c) distillation    d) condensation
16. Pure water boils at  
 a) 100 degree Celsius    b) 50 degree Celsius    c) 120 degree Celsius    d) 110 degree Celsius
17. Isothermal critical point is otherwise known as  
 a) eutectic point    b) triple point    c) plait point    d) melting point
18. For a three component system, the phase rule becomes  
 a)  $F = 4 - P$     b)  $F = 5 - P$     c)  $F = 3 - P$     d)  $F = 2 - P$
19. The simplest three component systems are those in which a liquid system breaks down into  
 a) two phases    b) three phases    c) four phases    d) five phases
20. An equation of fundamental importance which finds extensive application in one-component, two-phase systems, was derived by  
 a) J. Willard Gibbs    b) Albert Einstein    c) Clausius-Clapeyron    d) Dmitri Mendeleev

### **PART- B (3 x 2 = 6 Marks)**

#### **Answer ALL the Questions**

21. What is meant by triple point of water? Why is it different from the normal melting point of ice?
22. Define congruent melting point?
23. What the number of components, phases and degrees of freedom in the following equilibrium system?  $\text{CaCO}_3 (\text{s}) \leftrightarrow \text{CaO} (\text{s}) + \text{CO}_2 (\text{g})$

**PART- C(3 x 8= 24Marks)**

**Answer ALL the Questions**

24.a. State the phase rule. Discuss the application of phase rule for the water system.

**(OR)**

b. draw the phase diagram of sulphur system and calculate the degrees of freedom.

25.a. Derive Gibbs-Duhem-Margules equation.

**(OR)**

b. Derive Clausius-clapeyron equation and apply to solid-liquid, liquid-vapour and solid-vapour equilibria

26.a.State reduced phase rule. Justify reduced phase rule for Lead- Silver system with a neat diagram.

**(OR)**

b. Explain incongruent melting point by taking sodium-potassium system.

Reg.No.....

[18CHU302]

**KARPAGAM ACADEMY OF HIGHER EDUCATION**  
**COIMBATORE-641 021**

(For the candidates admitted from 2018 & Onwards)

**B.Sc., DEGREE EXAMINATION**

**THIRD SEMESTER**

**DEPARTMENT OF CHEMISTRY**

*Internal-II*

**PHYSICAL CHEMISTRY III**

**(Phase Equilibria and Chemical Kinetics)**

**Date: 28.8.2019 (AN)**

**Time: 2 hours**

**Marks: 50 marks**

**PART-A (20 x 1 = 20 marks)**

Answer All the Questions

1. Composition of liquid and vapour phases can be determined with the help of  
a) phase rule      b) lever's rule      c) gibbs rule      d) charles rule
2. The boiling point of aniline is  
a) 180 degree celsius      b) 100 degree celsius      c) 120 degree celsius      d) 85 degree celsius
3. Critical solution temperature is otherwise known as  
a) consolute temperature      b) solubility temperature      c) miscibility temperature  
d) melting temperature
4. The composition points of the conjugate phases are joined by  
a) tie lines      b) conjugate lines      c) upper lines      d) lower lines
5. Aniline-hexane is an example for  
a) partial miscibility increases on increasing the temperature  
b) partial miscibility increases on lowering the temperature  
c) partial miscibility increases on both raising as well as lowering the temperature  
d) complete miscibility temperature cannot be obtained
6. Complex ions are studied in  
a) Nernst distribution law      b) Gibbs distribution law  
c) Einstein distribution law      d) Albert distribution law

7. One of the important validity in Nernst distribution law is  
a) constant temperature    b) constant pressure    c) constant volume    d) constant composition
8. Partition coefficient is otherwise known as  
a) distribution coefficient    b) diffusion coefficient  
c) viscosity coefficient    d) freezing coefficient
9. The electrode in which oxidation occurs is  
a) anode    b) cathode    c) Anode and Cathode    d) Electrolyte
10. Calomel is a  
a) potassium chloride    b) sodium chloride    c) mercurous chloride    d) barium chloride
11. An example for oxidation-reduction electrode is  
a) calomel electrode    b) chlorine electrode  
c) quinhydrone electrode    d) hydrogen electrode
12. In the calomel electrode, the wire used is made of  
a) platinum    b) copper    c) titanium    d) iron
13. The EMF is measured in  
a) volts    b) coulomb    c) faraday    d) joules
14. When the electrodes are connected externally, then the circuit is said to be  
a) open    b) closed    c) ideal    d) constant
15. Daniel cell is called as  
a) voltaic cell    b) half cell    c) anode cell    d) cathode half cell
16. The standard hydrogen electrode can act as  
a) anode    b) cathode    c) Anode and Cathode    d) Electrolyte
17. Platinum is a  
a) positive electrode    b) negative electrode  
c) Positive and negative electrode    d) Inert electrode
18. The unit of electrical energy is  
a) volts    b) joules    c) coulomb    d) meter
19. The overall reaction taking place in the daniel cell is  
a) oxidation    b) reduction    c) redox    d) forward

20. The salt bridge is filled with a solution of

- a) potassium chromate      b) sodium chloride      c) zinc chloride      d) potassium chloride

**PART- B      (3 x 2= 6 Marks)**

**Answer ALL the Questions**

21. How do we use distribution law in desilverisation of lead?

22. Define electrode potential?

23. Differentiate between electrochemical cell and electrolytic cell

**PART- C      (3 x 8= 24Marks)**

**Answer ALL the Questions**

24.a. Define Nernst distribution law. Explain the application and limitations of Nernst distribution law.

**(OR)**

b. How to purify the organic liquids by Steam distillation

25. a. Write a note on reversible and irreversible cells

**(OR)**

b. What is known as electrochemical cell and write down the rules of oxidation/reduction of ions based on half-cell potentials.

26. a. A zinc electrode is placed in 0.1 M solution of zinc sulphate at 25 °C. If the degree of dissociation of salt at this concentration is found to be 0.95, calculate the electrode potential of the electrode at 25 °C.  $E^0 (\text{Zn}^{+2}, \text{Zn}^2) = -0.76$  volt.

**(OR)**

b. How to measure the electrode potential of Zinc and copper electrode using hydrogen electrode as a reference electrode.

Reg.No.....

[18CHU302]

**KARPAGAM ACADEMY OF HIGHER EDUCATION**

**COIMBATORE-641 021**

(For the candidates admitted from 2018 & Onwards)

**B.Sc., DEGREE EXAMINATION**

**THIRD SEMESTER**

**DEPARTMENT OF CHEMISTRY**

*Internal-III*

**PHYSICAL CHEMISTRY III**

**(Phase Equilibria and Chemical Kinetics)**

**Date:**

**Time: 2 hours**

**Marks: 50 marks**

**PART-A (20 x 1 = 20 marks)**

Answer All the Questions

1.The standard free energy change of a cell reaction is given by

- a)  $\Delta G = -nFE$       b)  $\Delta F = nFE$       c)  $\Delta G = nFE$       d)  $\Delta G = FE$

2.Glass electrodes can be used in

- a) weak oxidising solutions      b) strong reducing solutions  
c) weak reducing solutions      d) strong oxidising solutions

3.calomel electrode consists of mercury,solid mercurous chloride and a solution of -----

- a) KCl      b) NaCl      c) NaOH      d) KOH

4.The quinhydrone electrode is combined with a saturated-----electrode to form a cell

- a) hydrogen      b) calomel      c) glass      d) platinum

5.The standard electrode commonly used are

- a) quinhydrone electrode      b) glass electrode  
c) nitrogen electrode      d) hydrogen electrode

6.Liquid junction potential depends on

- a) removal of anion and cation      b) transfer of only anion  
c) transfer of only cation      d) transference number of anion and cation

7.Electrical energy forced in electrochemical cell is

- a) Spontaneous      b) Non-Spontaneous      c) Exothermic      d) Endothermic

8.Electrode-concentration cell are evidently independent of

- a) concentration of electrolyte      b) concentration of ions  
c) concentration in electrode      d) electrolyte concentration cell

9. The quinhydrone electrode cannot be used for solutions pH more than

- a) 7                      b) 9                      c) 8                      d) 5

10. Quinhydrone electrode is preferred to the -----electrode

- a) hydrogen                      b) calomel                      c) glass                      d) platinum

11. The substance adsorbed or attached is called

- a) adsorbent                      b) adsorbate                      c) sorbate                      d) desorption

12. At low pressure in Freundlich isotherm the graph is almost a

- a) parallel                      b) perpendicular                      c) straight line                      d) slope

13. Chemisorption requires

- a) activation energy                      b) entropy                      c) enthalpy                      d) free energy

14. Activated charcoal is used in

- a) sedimentation                      b) coagulation                      c) colouration                      d) gas masks

15. The BET method is widely used in

- a) surface                      b) adsorption                      c) material                      d) polymer

16. The removal of adsorbed substance from the surface is called

- a) inclusion                      b) occlusion                      c) sorption                      d) desorption

17. At the saturation pressure, the molecule layer number tends to

- a) Infinity                      b) low                      c) high                      d) low and high

18. Mechanical rubbing can be done to increase

- a) speed                      b) activation                      c) reaction                      d) speed of catalyst

19. Electricity is measured in

- a) volts                      b) amperes                      c) coulombs                      d) ohm

20. Organic compounds which can exist in oxidised form as well as reduced form are said to be

- a) amphoteric indicator                      b) redox indicator                      c) acid indicator                      d) base indicator

### **PART- B (3 x 2= 6 Marks)**

**Answer ALL the Questions**

21. What is known electrolytic cell?

22. Differentiate physical adsorption and chemical adsorption?

23. What is said to be liquid junction potential?

**PART- C (3 x 8= 24 Marks)**

**Answer ALL the Questions**

24. a. How to determine pH using the following electrodes

- (i) Hydrogen electrode
- (ii) Quinhydrone electrode

**(OR)**

b. Explain concentration cells with and without transference.

25. a. Explain potentiometric redox titration.

**(OR)**

b. Explain the following

- (i) application of EMF measurements in determining enthalpy
- (ii) application of EMF measurements in determining free energy of a cell

26. a. What is adsorption and absorption. Explain BET

**(OR)**

b. Derive Freundlich adsorption isotherm and mention its conditions.