



### First/Second Semester B.E./B.Tech. Degree Examination, Dec.2023/Jan.2024 **Chemistry for EEE Stream**

Time: 3 hrs.

Max. Marks: 100

Note: 1. Answer any FIVE full questions, choosing ONE full question from each module. 2. VTU Formula Hand Book is permitted.<br>3. M : Marks, L: Bloom's level, C; Course outcomes.





### **BCHEE102/202**





### Solution

### **Answer 1a.**

**Conductors** are the materials or substances which allow electricity to flow through them. They conduct electricity because they allow electrons to flow easily inside them from atom to atom. Also, conductors allow the transmission of heat or light from one source to another.

**Insulators** are the materials or substances which resist or don't allow the current to flow through them. In general, they are solid in nature. Also, insulators are finding use in a variety of systems. As they do not allow the flow of heat. The property which makes insulators different from conductors is its resistivity.



#### **Band Theory of conduction:**

Conductors: According to band theory, a conductor is essentially a substance with its conduction bands and valence bands overlapping, permitting electrons to move freely between the valence band and the conduction band. In conductors, conduction band is only partially filled. This means there are spaces for electrons to move into conduction band and hence thee materials acts as conductors.

Insulators: An insulator has a large gap between the valence band and the conduction band. The valence band is full and no electrons can move up to the conduction band, hence these material can't conduct.

### **Answer 1b. Czochralski process**

**(CZ):** The Czochralski process, is a method of crystal growth used to obtain single crystals of semiconductors (e.g. silicon, germanium and gallium arsenide), metals (e.g. palladium, platinum, silver, gold), salts and synthetic gemstones. In this process high-purity polycrystalline silicon is placed in the quartz crucible and melted using a RF coil in an atmosphere of argon. The temperature is maintained at the melting point of the silicon (around 1,412 °C). Dopant impurity atoms such as boron or phosphorus can be added to the molten silicon in precise amounts to dope the silicon, thus changing it into p-type or n-type silicon, with different electronic properties. A rotating puller rod with a





seed crystal at the bottom is lowered such that the seed crystal just touches the surface of molten silicon. A slight temperature drop initiates the crystallization of silicon on the seed crystal. The puller rod is pulled out at the rate of 1.5-5 cm/hour and simultaneously rotated at a speed of 100 rpm. As the rod is pulled away from the surface, silicon solidifies and a single crystal of silicon having the same crystal structure as that of the seed crystal is obtained. By precisely controlling the temperature gradients, rate of pulling and speed of rotation, it is possible to extract a large, single-crystal, cylindrical ingot from the melt.

### **Answer 1c. Graphene Oxide**

- $\triangleright$  Graphene oxide (GO) is a layered carbon structure with oxygen-containing functional groups (=O, -OH, -O-, -COOH) attached to both sides of the layer as well as the edges of the plane.
- $\triangleright$  As with any 2D carbon material, GO can also have either single layer or multilayer structure.
- $\triangleright$  A structure with one layer is graphene oxide; two layers of graphene oxide are referred to as a two-layered GO, GO with five to ten layers is called multi layered GO, and material with eleven or more layers is called graphite oxide.
- $\triangleright$  In contrary to graphene, GO is hydrophilic, and it is hence relatively simple to prepare a water- or organic solvent-based suspensions.
- $\triangleright$  Highly oxidized forms of GO are electric insulators with a bandgap of approximately 2.2 eV.
- $\triangleright$  Simplistically, GO is a monolayer sheet of graphite containing hydroxyl, carboxyl, and epoxy oxygen groups on its basal plane and edges, resulting in a mixture of sp2 and sp3 hybridized carbon atoms.

#### **Properties of Graphene Oxide:**

- $\triangleright$  The properties of graphene can be changed by the functionalization of graphene oxide. The chemically-altered graphene's could possibly be used in several applications.
- $\triangleright$  Graphene Oxide has a high surface area, and so it can be fit for use as electrode material for batteries, capacitors and solar cells.
- $\triangleright$  Graphene Oxide is cheaper and easier to manufacture than graphene, and so may enter mass production and use sooner.
- $\triangleright$  GO can easily be mixed with different polymers and other materials, and enhance properties of composite materials like tensile strength, elasticity, conductivity and more.

#### **Synthesis of Graphene Oxide:**

- $\triangleright$  There are several ways to prepare graphite oxide/graphene oxide. The most common way is to use an oxidizing agent in an acidic environment.
- $\triangleright$  In this procedure, phosphoric acid is mixed with sulphuric acid in the ratio 1:9 and potassium permanganate and graphite added in the ratio 6:1 in an ice bath.
- $\triangleright$  The mixture is then heated at 50<sub>o</sub>C and stirred for 12 h
- $\triangleright$  After cooling down, the mixture is poured onto ice
- $\triangleright$  Finally, 30% H2O2 is added in order to remove the excess of potassium permanganate.
- $\triangleright$  Phosphoric acid works as a dispersive and etching agent, as well as a stabilizer of the oxidation process, which makes the synthesis of GO safe.
- $\triangleright$  This route produces a higher yield of GO with a higher level of oxidation and a more regular structure.

### **Applications of Graphene Oxide:**



- $\triangleright$  Air pollution caused by the industrial release of harmful gases such as CO<sub>2</sub>, CO, NO<sub>2</sub>, and NH3.
- $\triangleright$  GO can be employed in catalysis for converting polluting gases during industrial processing.
- $\triangleright$  The approach of GO application in this area can be divided into two paths: pollutant adsorption and conversion.
- $\triangleright$  The functional groups of few-layered GO composites exhibit unique adsorption behaviour towards different gases like acetone, formaldehyde,  $H_2S$ ,  $SO_2$ , and NO<sub>x</sub> can be adsorbed by GO-based composites.
- $\triangleright$  Small-molecule drug delivery seems to be another promising medical application of GO. Small molecules of drugs can be attached to a GO surface using pH-sensitive linkers.

**Answer 2a.** Polymers which can conduct electricity are called conducting polymers. Mechanism of conduction in polyacetylene:

- $\triangleright$  (i) By Oxidative doping (p-doping): In this process,  $\pi$ -back bone of polymer is partially oxidized using a suitable oxidizing agent such as I2 in CCl4. The removal of an electron from the polymer π-back bone leads to the formation of delocalized radical ion called polaron. A second oxidation of chain containing polaron produces bipolaron which on radical recombination yields two charge carries on each chain. The positive charge sites on the polymer chain are compensated by anion I3- formed by the oxidizing agent during doping. The delocalized positive charges on the polymer chain are mobile, not the dopant anions. Thus these delocalized positive charges are current carriers for conduction. These charges must move from chain to chain as well as along the chain for bulk conduction .
- $\triangleright$  (v) On doping polyacetylene using I2 in CCl<sub>4</sub>, the conductivity increases from 10<sup>-5</sup> S cm-1 to 10<sup>3</sup>-10<sup>5</sup> S cm<sup>-1</sup>.





 $\triangleright$  If polyacetylene is heavily doped, polarons form pairs called solitons. In polyacetylenes the solitons are delocalized over 12 carbon atoms. Due to the formation of soliton, a new localized electronic state appears in the middle of the energy gap. When the doping is high, several charged solitons form soliton band. This band can later merge with edges of valence and conduction bands thus exhibiting conductivity.



Answer 2b. Electroless plating is the controlled deposition of a continuous film of a metal from its salt solution on a catalytically active surface of the substrate by a suitable reducing agent, without the use of electrical energy.

#### **Electroless-plating of Copper on PCB**

Activation of surface: The base of a printed circuit board is a plastic material such as epoxy or phenolic polymer or a glass fiber reinforced polymer composite. It is activated by treatment with acidified SnCl<sub>2</sub> and then with acidified PdCl2 leads to deposition of Pd.

#### $SnCl<sub>2</sub> + PdCl<sub>2</sub> \rightarrow Pd + SnCl<sub>4</sub>$

In the manufacture of double sided PCB, the board is clad on either side with thin electroformed copper foils. Then both sides of the copper clad board are printed with etchresistant circuit patterns. Rest of the unprotected copper foil is etched (formation of tracks) away by using an acid. This leaves only the circuit patterns on both sides of the board. Electrical connection between the two sides of PCB is made by drilling a hole through the board. The hole is then activated & electroless plated with copper, as it can't be electroplated. The composition of the electroless plating bath & the procedure given below.







#### Following reactions takes place during the process:

#### **Reactions:**

 $\overline{?}$  2HCHO + 4OH  $\longrightarrow$  2HCOO + 2H<sub>2</sub>O + H<sub>2</sub> + 2e<sup>-</sup> Oxidation of reducing reagent Reduction of metal ion over object surface :  $Cu^{2+} + 2e^-$  —  $\longrightarrow Cu$ 

Overall reaction

:  $Cu^{2+} + 2HCHO + 4OH$   $\longrightarrow Cu + 2HCOO + 2H_2O + H_2\uparrow$ 

### **Answer 2c. Solution:**

Number average molecular weight =  $N_1M_1 + N_2M_2 + N_3M_3$  $N_1 + N_2 + N_3$  $= (20 \times 15000) + (45 \times 25000) + (35 \times 27000)$  $20+45+35$ **=** 23700 g/mol Weight average molecular weight =  $N_1M_1^2 + N_2M_2^2 + N_3M_3^2$  $N_1M_1 + N_2M_2 + N_3M_3$  $= (20 \times 15000 \times 15000) + (45 \times 25000 \times 25000) + (35 \times 27000 \times 27000)$  $(20 \times 15000) + (45 \times 25000) + (35 \times 27000)$ **=** 24531.6 g/mol

**Answer 3a.** Battery is a device consisting of two or more galvanic cells arranged in series or parallel or both, that can generate electrical energy.

Batteries are classified under the following categories:

● **Primary battery:** In primary batteries, chemical energy stored in the battery is converted to electrical energy spontaneously as long as active materials are present. This battery cannot be recharged, because cell reaction is irreversible. *Example:* Zn-MnO<sub>2</sub> battery, Li-MnO<sub>2</sub> battery.

● **Secondary battery:** This battery can be recharged by passing electric current, because cell reactions are reversible. The redox reaction is reversed during recharing. Electrical energy is stored in the form of chemical energy in these batteries and used when needed. *Example:* Lead acid battery, Ni-MH battery, Li-ion battery. Primary cells act only as galvanic cell, whereas, a secondary cell can act both as galvanic cell and electrolytic cell. During discharging it acts as galvanic cell converting chemical energy to electrical energy and during charging process it acts as electrolytic cell converting electrical energy to chemical energy

● **Reserve battery:** In this battery, one of the key component is stored separately, and is incorporated into battery when required. When long storage is required, reserve batteries are often used, since the active component of the cell is separated until needed, thus reducing selfdischarge.

*Example:* Mg-AgCl battery. They are activated by adding sea water. These batteries have high reliability and long shelf life, hence they find applications in missiles and submarines. Another example is zinc-air batteries where the cell is sealed until use, the seal is removed to admit air and activate the cell when needed.



### **Answer 3b. Photovoltaic Cells:**

Photovoltaic cells or solar cells are semiconductor device that converts sunlight into direct current (DC) electricity. As long as light is shining on the solar cell, it generates electrical power. When light stops, electricity stops.

#### **Construction & Working of PhotoVoltaic Cells Construction:**

- PhotoVoltaic Cells consist of a semiconductor diode (p-n junction) made of a silicon.
- Silicon wafer or very thin silicon slices are made by silicon blocks and they are doped by p-type and n-type dopents to make p-n junction.
- It has two electrical



contact, on one of its sides, a mettalic grid is used and on the other side a layer of noble metal (such as Ag) is used.

• The metal grid permits the light to fall on the diode between the grid lines.

The part between the metallic grid is coated with antireflective compound. eg  $TiO<sub>2</sub>$ 

**Working:** 

 Electromagnetic radiation consists of particle called photon (hv). They carry a certain amount of energy given by the Plank quantum equation.

#### $E = hc/\lambda$

Where, h = Planck's constant, c = velocity of light,  $\lambda$  = wavelength of the radiation

- The electromagnetic radiation (sunlight) falls normal to the plane of the solar cell, the photons which possess energy sufficient to overcome the barrier potential are absorbed, electrons are ejected and electron-hole pairs are formed.
- The electrons move towards the n-region (as it is positively charged). The electrons are driven into the external circuit and used for various applications or appliances.

 $\bullet$ 

**Answer 3c. Lithium polymer (LiPo) Batteries** The science behind LiPo batteries is the same as in other Li-ion batteries: chemical energy is converted to electrical energy when electrons travel from the battery's anode to its cathode, creating an electrical current. The cathode contains a lithium metal oxide (such as lithium-cobalt oxide  $(LiCoO<sub>2</sub>)$ ), which provides lithium ions, whereas the anode contains a lithium carbon (such as graphite).

The anode and cathode are separated by an electrolyte that interacts with the anode to generate electrons, which creates a charge gradient in the cell. As the anode becomes negatively charged, the electrons travel along a conducting wire to the cathode. The whole system thus undergoes an electrochemical redox reaction (reduction/oxidation): the anode loses electrons and becomes oxidized while the cathode gains electrons and is reduced.





### **Figure: LiPo Battery**

**Anode:** Lithium intercalated Carbon or graphite

**Cathode:** lithium transition metal oxides (such as lithium-cobalt oxide (LiCoO2)),

**Electrolyte & Separator:** High conductivity semisolid (gel) polymers electrolyte polymer

electrolytes and include poly(methyl methacrylate) (PMMA), poly(acrylonitrile) (PAN)

### **Working:**



**Applications:** These batteries provide higher specific energy than other lithium battery types and are used in applications where weight is a critical feature, such as mobile devices, radiocontrolled aircraft and some electric vehicles.

**Answer 4a. Flow battery:** A flow battery, or redox flow battery, is a type of electrochemical cell where chemical energy is provided by two chemical components dissolved in liquids that are pumped through the system on separate sides of a membrane. Example: **Vanadium redox flow battery**

**Vanadium redox flow battery**: The vanadium redox battery (VRB), also known as the vanadium flow battery (VFB) or vanadium redox flow battery (VRFB), is a type of rechargeable flow battery. It employs vanadium ions as charge carriers.

**Construction:** A vanadium redox battery consists of an assembly of power cells in which two electrolytes are separated by a proton exchange membrane.

**Electrodes:** The electrodes in a VRB cell are carbon based. The most common types are carbon felt, carbon paper, carbon cloth, and graphite felt. Recently, carbon nanotube-based electrodes have attracted interest from the scientific community.

**Electrolytes:** Both electrolytes are vanadium-based. The electrolyte in the positive half-cells contains  $VO_2^+$  and  $VO^{2+}$  ions, while the electrolyte in the negative half-cells consists of  $V^{3+}$  and  $V^{2+}$  ions. The electrolytes can be prepared by several processes, including electrolytically



dissolving vanadium pentoxide  $(V_2O_5)$  in sulfuric acid  $(H_2SO_4)$ . The solution remains strongly acidic in use.

**Membrane:** The most common membrane material is perfluorinated sulfonic acid (PFSA) (Nafion). polytetrafluoroethylene (Teflon).



Figure. A diagram of a vanadium redox flow battery



Figure. Vanadium redox flow battery: (a) charge reaction and (b) discharge reaction.

#### **Working:**

Discharge

Negative electrode:

Charge

 $\leftrightarrow$  v

Discharge

 $3^{+}$  + e−

Positive electrode:  $VO_2^+ + e^- + 2H^+$ 

 $V^{2+}$ 

 $\leftrightarrow \quad \text{VO}^{2+} + \text{H}_2\text{O}$ 

Charge

Discharge

Overall reaction: 
$$
VO_2^+ + V^{2+} + 2H^+
$$
  
  $VO^{2+} + V^{3+} + H_2O$  (E<sup>0</sup>=1.26 V)  
 Charge



**Application:** Vanadium batteries are typically used for grid energy storage, i.e., attached to

power plants/electrical grids.

**Answer 4b. Definition:** A fuel cell is a device that converts the chemical energy of a fuel (hydrogen, natural gas, methanol, gasoline, etc.) and an oxidant (air or oxygen) into electricity. **Construction, working and applications of methanol–oxygen fuel cell.**

It is good example for liquid fuel cell. They use either acidic or alkaline medium. The preferred electrolyte is the acidic. Methanol is an efficient active organic fuel at low temperature.

**Construction:** Methanol – oxygen fuel cell consist of

- 1. Anode It is a porous Nickel (Ni) electrode impregnated with Pt/Pd catalyst.
- 2. Cathode It is a porous Nickel (Ni) electrode coated with silver (Ag) catalyst.
- 3. Electrolyte Aqueous sulphuric acid  $(H_2SO_4)$ , 3.7 M.
- 4. Active components: (a) Fuel Methanol mixed with sulphuric acid supplied at anode. (b) Oxidant – Pure oxygen is supplied at cathode.



**Figure:** Methanol – oxygen fuel cell

#### **Working:**

Cell reaction; Anode Reaction:  $CH_3OH + H_2O \longrightarrow CO_2 + 6H^+ + 6e$ Cathode Reaction:  $3/2$  O<sub>2</sub> + 6 H<sup>+</sup> + 6e-  $\longrightarrow$  3 H<sub>2</sub>O Overall Cell Reaction:  $CH_3OH + 3/2 O_2 \longrightarrow CO_2 + 2 H_2O$ 

 $H<sub>2</sub>O$  and  $CO<sub>2</sub>$  are formed as by-products but they do not harm the cell functioning because they are removed as and when they are formed.

Cell Potential : 1.2 V

#### **Applications:**

1. used in automobiles, military applications.

- 2. Power backup and portable instruments.
- 3. In large scale power production.



**Answer 4c.** Battery is a device which convert chemical energy into electrical energy on demand through redox titrations. OR

Battery is a device consisting of two or more galvanic cells arranged in series or parallel or both, that can generate electrical energy.

#### **Construction:**

**Anode:** Non-graphitic anodes, which consist largely of various carbonaceous materials (such as carbon black, pitch-based carbon-fibers, hard carbons etc)

**Cathode:** layer and tunnel type transition metal oxides, transition metal sulfides and fluorides etc

**Electrolyte:** The most common electrolyte formulations for SIBs are NaClO<sub>4</sub> or NaPF<sub>6</sub> salts in carbonate ester solvents (particularly propylene carbonate).



#### **Figure. Na-ion Battery (SIB)**

#### **Working principles of SIBs**

When the battery is being charged, Na atoms in the cathode release electrons to the external circuit and become ions which migrate through the electrolyte toward the anode, where they combine with electrons from the external circuit while reacting with the layered anode material. This process is reversed during discharge.

At anode: 
$$
Na_{x}C_{6} \longrightarrow
$$
 
$$
Na_{x}C_{6}
$$
 
$$
Na_{x} + xe^{-} + C_{6}
$$

At cathode: 
$$
NaMO_2 + xNa^+ + xe^-
$$

$$
\overrightarrow{\text{Discharge}} \quad \text{Na}_{1+x} \text{ MO}_2 + \text{C}_6
$$
\nChange



Overall cell reaction:  $Na<sub>x</sub>C<sub>6</sub> + Na MO<sub>2</sub>$   $\longrightarrow$   $Na<sub>1+x</sub>MO<sub>2</sub> + C<sub>6</sub>$ 

 Discharge Charge

**Applications:** Sodium-ion batteries can be used for a broad range of battery applications, including renewable energy storage for homes and businesses, grid storage, and backup power for data and telecom companies.

**Answer 5a. Definition**: Corrosion is defined as the **destruction of metal or alloys from its** 

**surface by the surrounding environment through chemical or electrochemical changes**.

Eg. 1. Rusting of Fe

**Electrochemical theory of corrosion:** Most of the corrosion problems are best explained on the basis of electrochemical theory.



**(i)** According to electrochemical theory, corrosion of metals takes place due to the formation of minute galvanic cells over the surface of metal. Thus anodic and cathodic regions are formed on the same metal surface or when two metals are in contact with each other in the presence of a conducting medium.

**(ii)** At the anodic region oxidation reaction takes place and the **metal gets converted into its ions by liberating electrons**. Consequently, **metal undergoes corrosion at the anodic region**.

 $M \longrightarrow M^+ + ne^-$ 

**Ex:** Fe  $\longrightarrow$  Fe  $^{2+}$  + 2e

**(iii)** The electrons flow from the anodic to cathodic area and at the cathodic region, reduction takes place. Since metal cannot be reduced further**, metal atoms at the cathodic region are unaffected by the cathodic reaction**. Some constitutions of the corrosion medium take part in the cathodic reaction. There are three possible ways in which the reduction can take place.

(i) If the solution is aerated and almost neutral, oxygen is reduced in presence of water to OH-



ions according to the equation.

 $\frac{1}{2}O_2 + H_2O + 2e^-$  2OH

(ii) If the solution is deaerated and almost neutral the cathodic reaction involves formation of hydroxyl ions with evolution of hydrogen gas.

$$
2H_2O + 2e^- \longrightarrow H_2 + 2OH^-
$$

(iii) If the solution is deaerated and acidic, the H<sup>+</sup> ions are reduced to H<sub>2</sub>

 $2H^{+} + 2e^{-}$   $\longrightarrow H_{2}$ 

(iv) The electrons liberated at the anodic region migrates to the cathodic region constituting corrosion current, the metal ions liberated at the anode and some anions liberated at the cathode diffuse towards each other through the conducting medium-range and form a corrosion products.

Corrosion of iron produced Fe<sup>2+</sup> ions and OH<sup>-</sup> ions at the anode and cathode sites respectively. These ions diffuse towards each other and produce insoluble Fe (OH)2.

 $Fe^{2+}$  + 2 OH  $^{-}$  $\longrightarrow$  Fe (OH)<sub>2</sub> (Ferrous hydroxide)

In an oxidizing environment, it is oxidized to ferric oxide and the rust is hydrated ferric oxide.

2Fe (OH) $2 + \frac{1}{2}$  O<sub>2</sub> + (n-2) H<sub>2</sub>O  $\longrightarrow$  [Fe<sub>2</sub>O<sub>3</sub>.nH<sub>2</sub>O] [Rust]

Corrosion of metal continues as long as both anodic and cathodic reactions take place simultaneously

**Answer 5b.** Corrosion Penetration Rate (CPR) is defined in three ways: (1) the speed at which any metal in a specific environment deteriorates due to a chemical reaction in the metal when it is exposed to a corrosive environment, (2) the amount of corrosion lost per year in thickness, (3) the speed at which corrosion spreads to the inner portions of a material.

The CPR is calculated as follows:  $CPR = (k x W) / (D x A x T)$ where  $k = a$  constant depends on unit used. when K = 534 the mpy will be used. when K = 87.6, mm/yr will be used. W = total weight lost  $(m_0 - m)$  m = weight after corroded  $m_0$  = weight before corroded  $T =$  time taken for the loss of metal  $A =$  the surface area of the exposed metal D = the metal density in  $g/cm^3$ 

Corrosion penetrating rate in mpy  $CPR = KW/DAT$ 



Weight loss, W=  $485 \times 10^3$  mg Density,  $D = 7.9g/cm^3$ ; Time,  $T = 24 \times 365 \times 1$  hours Area A =  $100 \text{ in}^2$ CPR =  $534 \times 485 \times 10^3$  $7.9\times100\times1\times24\times365$ **CPR = 37.42 mpy**

Corrosion penetrating rate mm/y  $CPR = KW/DAT$ Weight loss,  $W = 485 \times 10^3$  mg Density,  $D = 7.9g/cm3$ ; Time,  $T = 24 \times 365 \times 1$ Area A =  $100 \text{ in}^2$  =  $100 \times 6.45 \text{ cm}^2$  $CPR = 87.6 \times 485 \times 10^3$  7.9×100×6.45×24×365×1 **CPR = 0.9518 mm/y**

**Answer 5c.** Extraction of copper and gold from e-waste: The electronic waste also contains fair percentage of precious metals like Cu, Ag, Au, Pd, Rh etc. These metals can be recovered from E-waste at cheaper cost than from the usual ores. The recycling of metals such as gold and copper from discarded e-waste is an important aspect to develop the environmental friendly manufacturing processes. Techniques such as Pyrometallurgy, Hydrometallurgy and Biohydrometallurgy are used to recover precious metals like copper, silver and gold.

The extraction of copper and gold from e-waste using hydrometallurgical route comprises the liberation of the metallic fractions from downsized PCBs, a two-stage acid leaching process to provide a bulk separation of copper and gold from the other metals present, and subsequent purification of the copper and gold-containing solutions by solvent extraction using highly selective phenolic oxime and amide extractants, respectively.

Following steps are involved in the process.

1. Pre-procesing: After collection of e-waste, it is segregated, delaminated, cleaned with water and acetone and treated with N,N-dimethylacetamind (DMA) to remove the orgamic components (epoxy resins) present in the e-waste. This pretreated e-waste is shredded into small pieces and powdered, further it is subjected to recovery of precious metals. The metals were dissolved in acid and non-metallic component such as plastic and ceramic is separated by filtration.

2. Leaching: The liberated metallic fraction collected from the pre-processing step was leached in two stages to provide a bulk separation of copper and gold from the other metals present.

(i) First stage leaching process involves the selective dissolution of copper over gold. The samples were dissolved in dilute nitric acid, which almost exclusively leached the copper.

(ii) The filtered residue was then subjected to a second stage leaching process using sulfuric acid and halide salts (NaCl, NaBr) to selectively dissolve, silver and tin.

### 3. Solvent extraction:

(i) Recovery of Cu: Distilled water and ammonia solution was added to the leach solutions obtained after stage-one leaching process. To selectively recover the copper, the phenolic oxime dissolved in kerosene was used as the extractant. The Cu metal loaded organic phase was separated from the acidic aqueous leach liquor and washed with either sulfuric acid or nitric acid.



(ii) Recovery of Au: In the second-stage solvent extraction, an organic amide diluted in toluene was used as the extractant. The gold-loaded organic phase was then separated from the aqueous phase and stripped of the metal using either water or sodium hydroxide.

### **Answer 6a.** Differential metal corrosion (Galvanic corrosion)

- Differentian metal corrosion arises when two dissimilar metals are in contact with each other. Under this condition a potential difference is set up resulting in a galvanic current.
- A metal placed high in the electrochemical series is said to be anodic to the one placed below it.
- Eg: Fe is anodic to Cu but cathodic to Zn, Mg.
- Thus, Fe corrodes when it is in contact with Cu and Ni; But Zn and Mg gets corroded when they are in contact with Fe.
- The rate of differential metal corrosion depends primarily on the amount of the current passing from anode to cathode i.e; on the difference in electrochemical potential, higher the difference faster is the rate of corrosion.
- Consider a galvanic cell, in this cell, iron and copper in contact with an aerated solution of NaCl. A current flows between the two electrodes spontaneously and electrons flow from Fe to Cu, indicating that Fe is the anode and Cu is the cathode. The reaction that take place as follows-

$$
\text{Fe} \longrightarrow \text{Fe}^{2+} + 2\text{e}^{\cdot} \quad \text{(anode)}
$$
\n
$$
\frac{1}{2}O_2 + 2H_2O + 2\text{e}^{\cdot} \longrightarrow 2OH^{\cdot}
$$
\n
$$
\text{Fe}^{2+} + 2OH^{\cdot} \longrightarrow \text{Fe} (OH)_{2}
$$

(Ferrous hydroxide)

2 Fe (OH)  $_2$  + (n-2) H<sub>2</sub>O + O<sub>2</sub>  $\longrightarrow$  [Fe<sub>2</sub>O<sub>3</sub>.nH<sub>2</sub>O] [Rust]



### Fig. Differential metal corossion

*Differential aeration corrosion (Concentration cell corrosion):* Differential aeration corrosion occurs when the metal surface is exposed to two different concentration of air. The part of the metal exposed to lower oxygen concentration acts as anodic region and other acts as cathodic



region. The difference in the oxygen concentration produces a potential difference which causes the corrosion. At anode: Fe  $\rightarrow$  Fe<sup>2+</sup> + 2e<sup>-</sup> (lower  $O_2$  concentration) At cathode:  $\frac{1}{2}O_2 + H_2O + 2e^- \rightarrow 2OH^-$ (higher  $O_2$  concentration) Rust formation:  $Fe^{2+} + 2OH^{-} \rightarrow Fe(OH)_2$  (Ferrous hydroxide) Further oxidation leads to rust formation.  $2Fe(OH)<sub>2</sub> + \frac{1}{2}O<sub>2</sub> + H<sub>2</sub>O \rightarrow [Fe<sub>2</sub>O<sub>3</sub>.3H<sub>2</sub>O]$  .....[Rust] Cathodic region (higher oxygen concentration) Anodic region (Lower oxygen concentration -----------<br>**---------**-Pure iron partially immersed in an aerated solution of NaCl.

### **Answer 6b. Sacrificial anodic method:**

In this method, the base metal structure is converted into a cathode by connected it to a more active [higher up in galvanic series or less noble] metal. This active metal actually acts as an auxiliary anode. Mostly Zn, Mg, Al are the common auxiliary anodes used in this method. These metals being more active, acts as anode and undergo destruction and protect the metal structure. Since the anode material sacrifice to protect the metal structure, the method is known as sacrificial anode method. After continuous use exhausted sacrificial anodes are replaced by new ones as and when required. Eg: A Mg/ Zn block connected to buried oil storage tank, Mg/Zn base areas fixed to the side of ocean going ships, Mg/ Zn blocks area connected to the buried pipe lines.





Fig: Cathodic Protection by sacrificial anode

Advantage:

- (i)The method is simple.
- (ii) low installation cost.
- (iii) minimum maintenance cost.
- (iv) Doesn't require power supply.

Disadvantages: Involves recurring expenditure for Replacement of consumed anodes.

**Answer 6c. Definition:** E-waste or electronic waste means discarded electrical or electronic devices or components.

**Effects of e-waste on environment and human health:** E-waste is a serious issue for our environment because it releases harmful toxic chemicals from the metals due to chemical reactions and these toxic chemicals harm our environment, therefore they must be handled with care when no longer wanted or needed.

Electronic scrap components, such as CPUs, contain potentially harmful materials such as lead, cadmium, beryllium, or brominated flame retardants. Improper disposal of e-waste is highly dangerous to the global environment involving significant risk to the health of workers and their communities

Effects on Air quality: Contamination in the air occurs when e-waste is informally disposed of by dismantling, shredding, or melting the materials, releasing dust particles or toxins, such as dioxins, into the environment that cause air pollution and damage respiratory health. Chronic diseases and cancers are at a higher risk to occur when burning e-waste because it also releases fine particles, which can travel thousands of miles, creating numerous negative health risks to humans and animals. The air pollution caused by e-waste impacts some animal species more than others, which may be endangering these species and the biodiversity of certain regions that are chronically polluted. Over time, air pollution can hurt water quality, soil, and plant species, creating irreversible damage to ecosystems.

Effects on Soil: When the improper disposal of e-waste in regular landfills or in places where it is dumped illegally, both heavy metals and flame retardants can seep directly from the e-waste into the [soil,](https://www.clearias.com/causes-soil-degradation-methods-soil-conservation/) causing contamination of underlying groundwater or contamination of crops that may be planted nearby or in the area in the future. When the soil is contaminated by heavy metals, the crops become vulnerable to absorbing these toxins, which can cause many illnesses and doesn't allow the farmland to be as productive as possible.

Effects on Water: After soil contamination, heavy metals from e-waste, such as mercury, lithium, lead, and barium, then leak through the earth even further to reach groundwater. When these heavy metals reach groundwater, they eventually make their way into ponds, streams, rivers, and lakes. Through these pathways, acidification and toxicity are created in the water, which is unsafe for animals, plants, and communities even if they are miles away from a recycling site. Clean drinking water becomes problematic to find.

Effects on Humans: Electronic waste contains toxic components that are dangerous to human health, such as mercury, lead, cadmium, polybrominated flame retardants, barium, and lithium. The negative health effects of these toxins on humans include brain, heart, liver, kidney, and skeletal system damage. It can also considerably affect the nervous and reproductive systems of the human body, leading to disease and birth defects.

**Answer 7a.** Nanotechnology can be defined as the manipulation of atoms and molecules (one billionth) scale (1-100 nm) to produce devices, structures, or systems with at least one novel or superior property. Materials having at least one dimension in the nanoscale are called **nanomaterials**.



**Size-dependent properties** 1). Surface area: In chemical reactions, this surface-to-volume ratio plays an important role. There is an enormous change in the properties of materials due to the increased surface

area-to-volume ratio. The nanomaterials have a relatively larger surface area when compared to the same volume of the material produced in a larger form. So we know that material has high surface energy if it is small in size and vice versa. Therefore, nanoparticles have a large surface area to volume ratio and they possess large surface energy. Due to high surface energy materials



are more reactive and also nanoparticles show enhanced stability and a broader scope of applications. In some cases, materials that are inert in their larger form are reactive when produced in their nanoscale form. This affects their strength or electrical properties. The ratio of surface area to volume of a material is given by area/volume =  $4\pi r^2/ 4/3 \pi r^2 = 3/r$ 

2) Catalytic properties: The factors such as very small size, very high surface-to-volume ratio, and the increasing number of atoms on the surface are the most important reasons for the emergence of catalytic properties in nanomaterials. Basically, when particles become very small (nanoscale), due to the very high curvature they find, they have many atoms on their surface, which are very weakly bonded to the lattice atoms of the lattice. Therefore, these particles have very high surface energy and are highly active, and it is said that surface atoms are in a state of physical instability and are chemically active, and are prone to perform many chemical

reactions. It can be said that the main and determining reason for the emergence of catalytic properties in nanomaterials is their very high surface-to-volume ratio. The higher this ratio, the higher the catalytic properties in nanomaterials due to the increase in surface energy. In principle, the reason for these changes is due to changes in the electronic structure of materials, which can be justified by quantum mechanics.The effect of gold nanoparticle size on catalytic activity in the propene epoxidation reaction has been investigated, which shows that by reducing the size of gold nanoparticles, the yield of the product increases.



**Answer 7b.** Perovskite Materials is a material that has the same crystal structure as the mineral calcium titanium oxide (Perovskite).

Properties of Perovskite Materials:

- $\triangleright$  ABX3 Type Crystal Structure; where A = cation (such as methyl ammonium ion), B = metal cation, and  $X =$  halide ion.
- $\triangleright$  Optical Properties: As one of the unique properties of metal-organic halide perovskites, the optical properties of photo-generated charge carriers have been researched. The specific excitonic absorption peaks of the metal-organic halide perovskite could be transited to various absorption spectra, and it changed significantly in visible light through the adjustment of metal atoms and halogens
- $\triangleright$  Electroneutrality; the perovskite formula must have a neutral balanced charge therefore the product of the addition of the charges of A and B ions should be equivalent to the whole charge of the oxygen ions.
- $\triangleright$  Dielectric properties: There are some properties inherent to dielectric materials like ferroelectricity, piezoelectricity, electrostriction, and pyroelectricity. One of the important characteristics of perovskites is ferroelectric behavior.



- $\triangleright$  Electrical conductivity: Some perovskites exhibited great electronic conductivity
- $\triangleright$  Superconductivity: One of the obvious properties of perovskites is superconductivity.
- $\triangleright$  Catalytic Activity: Perovskites exhibited high catalytic activity.
- $\triangleright$  Piezoelectric property: Piezoelectric Effect is the ability of certain materials to generate an electric charge in response to applied mechanical stress. Various applications such as capacitors, piezoelectric devices, and ferroelectric devices have been designed by using traditional inorganic perovskite materials.

#### **Applications in Optoelectronic devices**

1. Solar cells: A perovskite solar cell is a type of solar cell, which includes a perovskite structured compound, most commonly a hybrid organic-inorganic lead or tin halide-based

material, as the light-harvesting active layer.

Perovskite materials are usually cheap to produce and relatively simple to manufacture. They possess intrinsic properties like broad absorption spectrum, fast charge separation, long transport distance of electrons and holes, long carrier separation lifetime, which make them very promising materials for solid-state solar cells.



2. Photodetectors: Lead halide perovskites have also been used to fabricate high-performance photodetectors.

3. Apart from these perovskite materials can be used in Light-emitting devices, Lasers, Water splitting applications such as Oxygen reduction and oxygen evolution reactions, Hydrogen evolution reactions and solid oxide fuel cells.

4. Nanoperovskites are recently utilized in electrochemical sensing of alcohols, gases, amino acids, acetone, glucose,  $H_2O_2$ , and neurotransmitters exhibiting good selectivity, sensitivity, unique long-term stability, excellent reproducibility, and anti-interference ability.

**Answer 7c.** Co-precipitation method: The commonly used solution method for the synthesis of multi-component oxide ceramics is the co-precipitation method, which produces a "mixed" precipitate comprising two or more insoluble species that are simultaneously removed from the solution. The precursors used in this method are mostly inorganic salts (nitrate, chloride, sulfate, etc.) that are dissolved in water or any other suitable medium to form a homogeneous solution with clusters of ions. The solution is then subjected to pH adjustment or evaporation to force those salts to precipitate as hydroxides, hydrous oxides, or oxalates. The crystal growth and their aggregation are influenced by the concentration of salt, temperature, the actual pH, and the rate of pH change. After precipitation, the solid mass is collected, washed, and gradually dried by heating to the boiling point of the medium. The washing and drying procedures applied for coprecipitated hydroxides affect the degree of agglomeration in the final powder and must be considered when nanosized powders are the intended product. Generally, a calcination step is necessary to transform the hydroxide into crystalline oxides. In most of the binary, ternary and quaternary systems, a crystallization step is necessary, which is generally achieved by calcination or, more elegantly, by a hydrothermal procedure in high-pressure autoclaves.

Example: Precursor solution: 0.1 M Fe ion solution, prepared by dissolving ferrous and ferric chlorides in a 1:2 FeCl<sub>2</sub>·4H<sub>2</sub>O: FeCl<sub>3</sub>·6H<sub>2</sub>O molar ratio in deionised (DI) water (pHprec = 1.8).



Base solution: DI water mixed with 2 M NaOH in a 10:4 volumetric ratio (0.57 M NaOH). Neutralisation solution:  $0.316$  M citric acid solution, pHneutr = 1.8. All chemicals, the provider and lot numbers are listed in table S1. 5 IONPs were co-precipitated by mixing the precursor and base solution of the (standard) concentrations stated above (if not mentioned otherwise). Mixing was performed at a reaction temperature of 60 °C for all syntheses. The sequence of neutralisation solution addition and the added quantity varied with the experiment

**Answer 8a.** The sol-gel process, involves the evolution of inorganic networks through the formation of a colloidal suspension (sol) and gelation of the sol to form a network in a continuous liquid phase (gel). The precursors for synthesizing these colloids consist usually of a metal or metalloid element surrounded by various reactive ligands. The starting material is processed to form a dispersible oxide and forms a sol in contact with water or dilute acid. Removal of the liquid from the sol yields the gel, and the sol/gel transition controls the particle size and shape. Calcination of the gel produces the oxide.



Step1: Formation of different stable solutions of the alkoxide or solvated metal precursor (the sol).

**M**-OR + H2O -----------------► M-OH + HOR

Step 2: Gelation resulting from the formation of an oxide or alcohol bridged network (the gel) by polycondensation or poly esterification reaction. This results in a dramatic increase in the viscosity of the solution.

M-OH + M-OR ----------------- ► M-O-M + HOR

 $M-OH + M-OH$ ----------------  $M-O-M + HOR$ 

Step 3: Aging of the gel(synthesis), during which the polycondensation reactions continue until the gel transforms into a solid mass. This is accompanied by the contraction of the gel network and the expulsion of solvent from gel pores.

Step 4: Drying of the gel, when water and other volatile liquids are removed from the gel network. If isolated by thermal evaporation, the resulting is termed a xerosal. If the solvent (such as water) is extracted under supercritical or near supercritical conditions, the product is an aerogel.

Step 5: Dehydration, during which surface-bound M-OH groups are removed. This is normally achieved by calcination of the monolith at temperatures up to 800 0 C. Step 6: Densification and decomposition of the gels at high temperature  $(T>800^{\circ}C)$ . The pores of the gel network collapse and the remaining organic species are volatilized.



The typical steps that are involved in sol gel processing are shown in fig. By different processes, one can get either nano film coating or nanopowder or dense ceramic with nanograins.

**Example: Sol-Gel synthesis process ZnO NPs** First of all, in a 100ml beaker 30 ml of water was added with 35 ml of triethanolamine(TEA) and drop wise ethanol was added with continuous stirring to get a homogeneous solution. After addition of 100 drops ethanol that was about 3 ml and continuous stirring results a homogeneous solution. Keeping the stoichiometry in mind a 2.0 gm batch of zinc oxide was prepared. Firstly, 30ml of water was mixed with 20 ml of TEA with constant stirring and drop wise addition of ethanol. The obtained homogeneous solution was kept at rest for 3.0 hours. For 2.0 gm batch of zinc oxide 5.49gm of zinc acetate dihydrate was mixed with 50ml water and 0.5M of solution was prepared which was subjected to continuous stirring to get a homogeneous solution. After that the two solutions were mixed together in 500ml beaker and drop wise ammonium hydroxide was added with continuous heating and stirring for 20minutesvia hot plate. Nearly 10ml of distil water was added during stirring. Then the solution was left for half an hour which results in the formation of white bulky solution. The obtained solution was then washed 10-12 times with distil water and filtered in a filter paper. The residue obtained waskept for drying in an oven at a temperature of about 95°C for 6 hours. The yellowish white powder obtained was then subjected to calcinations at a temperature of600°C for 5 hours

**Answer 8b.** Quantum dot light-emitting diode (QLED) use quantum dots for emission and attracted much attention for the next generation of display

Properties:

- $\triangleright$  high color saturation
- $\triangleright$  tunable color emission
- $\triangleright$  high stability.
- $\triangleright$  QLED display has advantages in flexible and robust application, which makes wearable and stretchable display possible in the future.
- $\triangleright$  energy efficient, thin-film display and solid-state lighting applications.
- $\triangleright$  facile scale-up capability
- $\triangleright$  Wide-viewing angle

Applications of QLED

- $\triangleright$  quantum dot light-emitting diodes (QLED) are cost-effective electroluminescence devices ideal for large-area display and lighting applications.
- $\triangleright$  For photomedical applications: facilitate widespread clinical applications of photodynamic therapy (PDT) or photobiomodulation (PBM).
- $\triangleright$  Consumer electronics: Used in smart phones, tablets, laptops, and televisions.
- $\triangleright$  Lighting: Used in lighting applications
- Wearable devices:Used in smart watches and fitness trackers.
- $\triangleright$  Automotive displays: Used in automotive displays
- $\triangleright$  Medical devices: OLEDs are used in medical devices

**Answer 8c.** Nanophotonics is a field that studies the behavior of light on the nanoscale, typically structures that are smaller than the wavelength of light. Some properties of nanophotonics include:

- 1. Enhanced light-matter interactions: Nanophotonic structures can enhance the interaction between light and matter, allowing for more efficient absorption, emission, and manipulation of light.
- 2. Sub-wavelength confinement: Nanophotonic structures can confine light to subwavelength dimensions, leading to improved spatial resolution and sensitivity in imaging and sensing applications.



- 3. Tunability: The properties of nanophotonic structures can be tuned by varying their size, shape, and composition, allowing for the design of structures with specific optical properties.
- 4. Nonlinear effects: Nanophotonic structures can exhibit nonlinear optical effects, such as harmonic generation, which can be useful in applications such as frequency conversion.
- 5. Plasmonics: Nanophotonics often involve the study of plasmonic effects, which arise from the interaction between light and electrons in metallic nanostructures. Plasmonics can be used for applications such as sensing, imaging, and data storage.
- 6. Quantum effects: Nanophotonics can be used to study and exploit quantum effects, such as single-photon emission, quantum entanglement, and quantum computing.

Applications of nanophotonics: Optoelectronics: Nanophotonics is used in the design of optoelectronic devices such as light-emitting diodes (LEDs), solar cells, and photodetectors. These devices can be made more efficient and compact using nanophotonic structures.

Imaging and sensing: Nanophotonics is used in imaging and sensing applications, including super-resolution microscopy, single-molecule detection, and biosensing. Nanophotonic structures can enhance the sensitivity and spatial resolution of these techniques.

Data storage: Nanophotonics is used in data storage applications, including high-density data storage and non-volatile memory. Nanophotonic structures can be used to manipulate light to encode and read data.

Quantum technologies: Nanophotonics is used in the development of quantum technologies, including quantum computing, quantum cryptography, and quantum sensing. Nanophotonic structures can be used to create and manipulate quantum states of light and matter.

Energy: Nanophotonics is used in energy-related applications, including light harvesting for solar energy conversion and light management in solid-state lighting. Nanophotonic structures can enhance the absorption and emission of light in these applications.

Biomedical applications: Nanophotonics is used in biomedical applications, including biosensing, drug delivery, and imaging. Nanophotonic structures can be used to target specific cells or tissues and enhance the sensitivity and specificity of biomedical techniques.

Nanosensors are sensors that make use of the unique properties of nanomaterials and nanoparticles to detect and measure materials and components on the nanoscale. The signals can be biomedical, optical, electronic, electrical, physical or mechanical.

### **Properties of Nanosensors:**

- Available in small sizes
- Requires less power to operate
- Less weight
- Works as data storage systems
- Great sensitivity, accuracy, scalability, efficiency, precision, and specificity
- Easy to execute.
- Provides a high-volume ratio.

### Response time is low.

### **Nanosensor Applications**

They are used:

• To detect various chemicals in gases for pollution monitoring.

Nanosensors have potential applications in the food sector, in food processing monitoring, food quality assessment, food packaging, food storage, shelf-life monitoring, and viability, as indicators of food safety and microbial contamination.

• For medical diagnostic purposes either as bloodborne sensors or in lab-on-a-chip type devices. The nanosensors provide an understanding of a person's health status through



noninvasive detection of clinically relevant biomarkers in several biofluids such as tears, saliva, and sweat without sampling, complex manipulation, and treatment steps.

- To monitor physical parameters such as temperature, displacement and flow
- To monitor plant signaling and metabolism to understand plant biology
- To study neurotransmitters in the brain for understanding neurophysiology

**Answer 9a. Reference Electrodes :** Reference electrodes are the electrodes whose potentials are known and they are used for the determination of potentials of other electrodes.

Calomel electrode: It is a metal – metal salt ion electrodes and also a secondary reference electrode.

Construction: It consists of mercury, mercurous chloride and a solution of KCl. It is made up of glass tube, Hg is placed at the bottom of the glass tube. It is covered with a paste of calomel  $(HgCl<sub>2</sub>)$ and KCl solution. A solution of KCl is introduced above the paste through the side tube. Electrical contact is made through Pt wire (dipped in Hg). This glass tube is taken into another glass tube, which has a side tube and a porous plug at the bottom. This porous plug act as salt bridge. The conc of KCl solution used is either decinormal, normal or saturated. Correspondingly, the electrode is known as decinormal, normal or saturated calomel electrode. This constitutes one half cell. Calomel electrode can act as either anode or cathode. It is connected to the other half cell through salt bridge.



The calomel electrode is represented as, Hg  $/Hg_2Cl_2$  (s) Cl (anode) Or Cl /  $Hg_2Cl_2$  (s) Hg (Cathode)

Working: Depending on the potential of electrode with which the calomel electrode is connected, calomel electrode acts as anode or cathode.

(a). When it acts as anode, electrode reaction is

$$
2Hg \longrightarrow Hg_2^{2+} + 2e^-
$$
  
\n
$$
Hg_2^{2+} + 2Cl^- \longrightarrow Hg_2Cl_2
$$
  
\n
$$
2Hg + 2Cl^- \longrightarrow Hg_2Cl_2 + 2e^-
$$

(b) When it acts as cathode, electrode reaction is

 $Hg_2^{2+} + 2e^- \longrightarrow 2Hg$  $\text{Hg}_2\text{Cl}_2 \longrightarrow \text{Hg}_2^{2+} + 2\text{Cl}^ Hg_2Cl_2 + 2e^ \longrightarrow$  2Hg + 2Cl<sup>-</sup>

Thus, Net reversible electrode reaction is

 $Hg_2Cl_2 + 2e$   $\overline{\bullet}$   $2Hg + 2Cl$ 



Electrode potential is calculated using Nernst equation,

$$
E_{cal} = E^{\circ} - \frac{2.303RT}{2F} \quad \log [CI^{-}]^{2}
$$

Applications:1. It is used as secondary reference electrode in all potentiometric determinations. 2. Used in glass or combined electrode to determine the pH of the unknown solution.

**Answer 9b. Principle:** The procedure of using measurement of emf to determine the concentration of ionic species in solution is called as potentiometry also known as potentiometric titration. When a metal M is immersed in a solution containing its own ions  $M_{n+}$ , the electrode potential is

given by Nernst equation.

$$
E = E^{\circ} + \underbrace{0.0591}_{n} \log \left[ M^{n+} \right]
$$

Thus, the concentration can be calculated, provided  $E<sub>o</sub>$ of the electrode is known. The principle involved in potentiometric titration is the measurement of emf between two electrodes, an indicator electrode, (the potential of which is function of the concentration of the ion to be determined) and a reference electrode of constant potential. In this method, the measurement of emf is made while the titration is in progress. The equivalence point of the reaction is revealed by a sudden change in potential in the plot of emf readings against the volume of titrant.

**Instrumentation:** It consists of: (i) Calomel electrode as a reference electrode, (ii) Platinum electrode as an indicator electrode, (iii) a device for measuring the potential and (iv) Magnetic stirrer.

**Application:** Potentiometric estimation of FAS using standard K2Cr2O7 solution: Pipette out 25ml of FAS into a beaker. Add 1 t.t dil H2SO4, immerse calomel electrode + platinum electrode into it. Connect the assembly to a potentiometer and measure the potential by adding K2Cr2O7 in the increments of 0.5ml. Plot graph  $\Delta E/$  $\Delta V$  against volume of K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub>, and determine the equivalence point. From the normality and volume  $K_2Cr_2O_7$ , solutions calculate the normality and the weight of FAS in the given solution.



**Answer 9c. Concentration Cells:** In these cells, the two electrodes of same element is in contact with solution of same metal ion (electrolyte) but of different concentration.

ΔE ΛV

Cell representation: Ag / Ag<sup>+</sup>( $C_1 = 0.001$  M) // Ag<sup>+</sup>( $C_2 = x$  M) / Cu

The Electrode reactions are as follows:



At anode  $\text{Ag} \rightarrow \text{Ag}^{2+} (\text{C}_1=0.001 \text{ M}) + e^{-}$ 

At cathode  $Ag^+(C_2 = x M) + e^- \rightarrow Ag$ 

EMF of cell is given by

$$
E_{\text{cell}} = \frac{2.303RT}{nF} \log \frac{C_2}{C_1}
$$

 $0.059 = 2.303 \times 8.314 \times 298$  log X **1×96500** 0.001 **X = 0.01 M**

**Answer 10a. . Ion Selective electrodes** These electrodes selectively respond to a specific ion in a mixture and potential developed is a function of concentration of that ion in the solution. These electrodes consist of a membrane which is capable of exchanging specific ions with solution with which it is in contact. These are also called as membrane electrodes. Eg. Glass electrode.

**Glass Electrode Construction:** The glass electrode consist of glass tube, the bottom of the glass tube is glass bulb made up of very thin glass membrane. The thickness of glass membrane varies from 0.03 mm to 0.1 mm. The membrane is made up of special glass of low melting point and high electrical conductivity. Its



composition is  $SiO_2 - 72\%$ , Na<sub>2</sub>O- 22%, CaO- 6%. It can sense H<sup>+</sup> ions up to a pH of about 9. Glass bulb contains 0.1 N HCl (Assume concentration is  $C_2$ ). An Ag/AgCl electrode (internal reference electrode) is placed in the solution and connected by a pt wire for electrical contact.

The electrode is represented as, Ag/AgCl (s)/ 0.1N HCl/ Glass

**Working of glass electrode:** When the glass electrode is dipped into any solution containing H + ions, the Na+ ions of the glass membrane are exchanged for H+ ions of the test solution.

 $H^+ + Na^+$  GI  $\longrightarrow Na^+ + H^+$ GI

If a thin walled bulb containing an acid is immersed in another solution containing  $H^+$  ions (fig), a potential is developed across the glass membrane. This is called the boundary potential E<sub>b</sub>. It is a potential developed across the glass membrane when concentration of the solution





inside and outside the glass membrane are different. The  $E<sub>b</sub>$  is due to the difference in potential  $(E_2-E_1)$  developed across the gel layer of the glass membrane between the two liquid.

Mathematically it is represented as,

$$
E_b = E_2 - E_1
$$
 Where,  $E_2$ = Potential due to H<sup>+</sup> present in outside solution (Unknown solution)  
 $E_1$  = Potential due to H<sup>+</sup> present in inside solution (known solution)  
According to Nernst equation

$$
E_b = \frac{2.303RT}{nF} \log \frac{C_2}{C_1}
$$
  
\n
$$
E_b = 0.0591 \log C_2 - 0.0591 \log C_1
$$
 (1)

Where,  $C_2$  is the concentration of H<sup>+</sup> ions of the solution into which glass membrane is dipped. The concentration of  $H^+$  ion inside the bulb (C<sub>1</sub>) is constant i.e.  $C_1 = 0.1$  M.

Thus, Eb = 0.0591 log C2 + K or = K + 0.0591 log C<sup>2</sup> n n

Glass electrode selects only  $H^+$  ions ignoring other ions.

Hence 
$$
C_2 = H^+
$$

 $E_b = K + 0.0591 \log [H^+]$ 

Where,  $log[H^+] = -pH$ 

Thus,  $E_b = K - 0.0591pH$  ----------------------(2)

The combined glass electrode is dipped into acidic solution, then the potential of the glass electrode is given by….

E<sup>G</sup> = E<sup>b</sup> +EAg-AgCl ………………………(3)

From equation 1, theoretically if  $C_1 = C_2$ ,  $E_b$  should be 0, however it has been observed practically that even when  $C_1 = C_2$ , a small potential is developed which is called as asymmetric potential (Easym). Hence equation 3 can be rewritten as

E<sup>G</sup> = E<sup>b</sup> +EAg-AgCl + Easym ………………………(4)

Substituting the value of  $E_b$  from equation (2) in equation (4)

$$
E_G=K-0.0591pH\ + E\ _{Ag/AgCl}+E\ _{assy}
$$

E<sup>G</sup> = Eº<sup>G</sup> 0.0591pH **………………(5)** Where (Eº<sup>G</sup> = K + EAg/AgCl + Eassy)

The above expression (eq 5) indicate that the potential of glass electrode, E<sup>G</sup> varies with the pH of the acidic solution.

.**Answer 10b. Principle:** Colorimetry is a scientific technique that is used to determine the concentration of

colored compounds in solutions.

When a beam of incident light of intensity **I**<sup>0</sup> passes through a solution, a part of the incident light

is relected (**Ir**), a part is absorbed (**Ia**) and rest of the light is transmitted (**It**)

Thus,  $I_0 = I_r + I_a + I_t$ 

In colorimeter, **Ir** is eliminated. For this purpose, the amount of light relected (**Ir**) is kept constant



by using cells that have identical properties. (**I0**) & (**It**) is then measured. Colorimetry measurements are based on Beer-Lambert's law which states that when a monochromatic light passes through a transparent medium, the amount of light absorbed is directly proportional to the concentration and path length of the solution.

**A = εct** Where A is absorbance, ε is the molar extinction coefficient, c is the concentration, t is the path length. If t, the path length is kept constant, then,  $A \propto c$ . Hence a plot of absorbance against concentration gives a straight line.

**Instrumentation:** Photoelectric colorimeter consists of

(i) Tungsten lamp as the light source.

(ii) A filter which provides the desired wavelength range wherein the solution gives the maximum absorbance.

#### (iii) A sample cell

(iv) A photocell detector: Detector are photosensitive elements which converts light energy into electrical energy

#### **Application: Colorimetric estimation**

**of Cu in CuSO4.** Draw out 2, 4, 6, 8, and 10 ml cm3 of the Copper sulphate solution into 50cm3 volumetric flask. Add 5cm3 of ammonia solution to each of them and dilute upto the mark with distilled water and mix well. Prepare a blank solution by diluting 5cm<sup>3</sup> of ammonia solution in 50cm<sup>3</sup> volumetric flasks. For test solution add 5ml of NH<sup>3</sup> and make up to the mark. Measure the absorbance of each of these against blank solution at 620 nm. Plot a graph of absorbance (OD) against volume of







copper sulphate solution and determine the volume of copper sulphate solution in the test sample as shown in the figure and find the amount of copper present in it.

**Answer 10c Principle**: Conductometry is based on Ohm's law which states that the current i (amperes) flowing in a conductor is directly proportional to the applied electromotive force, E (volts), and inversely proportional to the resistance R (ohms) of the conductor. The reciprocal of the resistance is called the conductance (Ease with which electric current flows through a conductor). Specific conductance of a solution is defined as the conductance of a solution present between two parallel electrodes which have 1cm2 area of cross section and which have kept 1 cm apart. The conductance of solution depends on the number and mobility of ions. The substitution of ions with different ionic mobility affects the electrolytic conductivity. Therefore, the equivalence point can be determined by means of conductivity measurement for a neutralization reaction between an acid and a base. Equivalence point is determined graphically by plotting conductance against titer values.



### **Instrumentation:**

Conductometer consists of: (1) conductivity cell having two platinum electrodes; and a (ii) conductometer. A simple arrangement of conductometric titration is depicted in figure. The solution to be titrated is taken in the beaker.



**Application in the estimation of weak:** Pipette out 50ml of sample into a beaker. Immerse the conductivity cell into it. Connect the conductivity cell to a conductivity meter and measure the

conductance by adding NaOH from the burette by increment of 1 ml. Plot a graph of conductance against volume of NaOH. Determine the neutralization point from the graph as shown below**.**

**Weak acid with a strong base: (CH3COOH Vs NaOH):** In the conductometric titration of a weak acid with a strong base, the conductance of the acid will be initially low due to poor dissociation of acetic acid. On complete neutralization of the acid, further addition of base leads to an increase in the number of more mobile OH- ions. Hence conductance increases sharply.