CBSE Class-12 Chemistry Quick Revision Notes Chapter-06: General Principles and Processes of Isolation of Elements

Minerals:

The naturally occurring chemical substances in the earth's crust which are obtained by mining are known as minerals.

- Metals may or may not be extracted profitably from them.
- Ores:

The rocky materials which contain sufficient quantity of mineral so that the metal can be extracted profitably or economically are known as ores.

• Gangue:

The earthy or undesirable materials present in ore are known as gangue.

- Metallurgy:
- The entire scientific and technological process used for isolation of the metal from its ores is known as metallurgy.
- Chief Ores and Methods of Extraction of Some Common Metals:

Sodium metal

- a) Occurrence: Rock salt (NaCl), Feldspar (Na₃AlSi₃O₈)
- b) Extraction method: Electrolysis of fused NaCl or NaCl/ CaCl₂
- c) Inference: Sodium is highly reactive and hence, it reacts with water.

Copper metal

- a) Occurrence: Copper pyrites (CuFeS₂), Malachite (CuCO₃.Cu(OH)₂), Cuprite (Cu₂O) Copper glance (Cu₂S)
- b) Extraction method: Roasting of sulphide partially and reduction.

$$2 Cu_2O + Cu_2S \rightarrow 6 Cu + SO_2$$

c) Inference: It is self-reduction in a specially designed converter. Sulphuric acid leaching is also employed.

Aluminium metal

- a) Occurrence: Bauxite:(AlO_x(OH)_{3-2x} where 0 < x < 1), Cryolite (Na₃AlF₆), Kaolinite (Al₂(OH)₄Si₂O₅)
- b) Extraction method: Electrolysis of Al₂O₃ dissolved in molten cryolite or in Na₃AlCl₆
- c) Inference: A good source of electricity is needed in the extraction of Al

Zinc metal

- a) Occurrence: Zinc blende or Sphalerite (ZnS), Zincite (ZnO), Calamine (ZnCO₃)
- b) Extraction method: Roasting and then reduction with carbon.
- c) Inference: The metal may be purified by fractional distillation.

Lead metal

- a) Occurrence: Galena (PbS)
- b) Extraction: Roasting of the sulphide ore and then reduction of the oxide.
- c) Inference: Sulphide ore is concentrated by froth floatation process.

Silver metal

- a) Occurrence: Argentite (Ag₂S)
- b) Extraction method: Sodium cyanide leaching of the sulphide ore and finally replacement of Ag by Zn.
- c) Inference: It involves complex formation and displacement.

Gold metal

- a) Occurrence: Native, small amounts in many ores such as those of copper and silver
- b) Extraction method: Cyanide leaching, same as in case of silver
- c) Inference: Gold reacts with cyanide to form complex

Iron metal

- a) Occurrence: Haematite (Fe₂O₃), Magnetite (Fe₃O₄), Siderite (FeCO₃), Iron pyrites (FeS₂)
- b) Extraction method: Reduction with the help of CO and coke in blast furnace.
- c) Inference: Limestone is added as flux which removes SiO_2 as calcium silicate (slag) floats over molten iron and prevents its oxidation. Temperatures approaching 2170 K is required.

Steps of metallurgy:

- a) Concentration of ore
- b) Conversion of concentrated ore to oxide
- c) Reduction of oxide to metal
- d) Refining of metal

Concentration of ore:

The process of removal unwanted materials like sand, clay, rocks etc from the ore is known as concentration, ore – dressing or benefaction. It involves several steps which depend upon physical properties of metal compound and impurity (gangue). The type of metal, available facilities and environmental factors are also taken into consideration.

• Hydraulic washing (or gravity separation):

It is based on difference in densities of ore and gangue particles. Ore is washed with a stream of water under pressure so that lighter impurities are washed away whereas heavy ores are left behind.

• Magnetic separation:

This method is based on the difference in magnetic and non – magnetic properties of two components of ore (pure and impure). This method is used to remove tungsten ore particles from cassiterite (SnO_2). It is also used to concentrate magnetite (Fe_3O_4), chromite ($FeCr_2O_4$) and pyrolusite (MnO_2) from unwanted gangue.

Froth floatation process:

It is based on the principle that sulphide ores are preferentially wetted by the pine oil or fatty acids or xanthates etc., whereas the gangue particles are wetted by the water.

Collectors are added to enhance the non-wettability of the mineral particles.

Froth stabilizers such as cresols, aniline etc., are added to stabilize the froth.

If two sulphide ores are present, it is possible to separate the two sulphide ores by adjusting proportion of oil to water or by adding depressants.

For example, in the case of an ore containing ZnS and PbS, the depressant used is NaCN. It selectively prevents ZnS from coming to froth but allows PbS to come with the froth.

Leaching (Chemical separation):

It is a process in which ore is treated with suitable solvent which dissolves the ore but not the impurities.

• Purification of Bauxite by leaching (Baeyer's process):

a) Step 1:

$$Al_2O_3(s) + 2NaOH(aq) + 3H_2O(l) \rightarrow 2Na[Al(OH)_4](aq)$$

b) Step 2:

$$2Na[Al(OH)_{A}](aq) + CO_{2}(g) \rightarrow Al_{2}O_{3}XH_{2}O(s) + 2NaHCO_{3}(aq)$$

c) Step 3:

$$Al_2O_3.XH_2O(s) \xrightarrow{Heat \text{ at } 1470\text{K}} Al_2O_3(s) + XH_2O(g)$$

d) Concentration of Gold and Silver Ores by Leaching:

$$4M(s) + 8CN^{-}(aq) + 2H_{2}O(aq) + O_{2}(g) \rightarrow 4[M(CN)_{2}]^{-}(aq) + 4OH^{-}(aq)$$
$$2[M(CN)_{2}]^{-}(aq) + Zn(s) \rightarrow [Zn(CN)_{4}]^{2-}(aq) + 2M(s)$$

Where [M = Ag or Au]

Conversion of ore into oxide:

It is easier to reduce oxide than sulphide or carbonate ore. Therefore, the given ore should be converted into oxide by any one of the following method:

- a) roasting
- b) calcination

Roasting:

- a) It is a process in which ore is heated in a regular supply of air at a temperature below melting point of the metal so as to convert the given ore into oxide ore.
- b) Sulphide ores are converted into oxide by roasting
- c) It is also used to remove impurities as volatile oxides
- d) example $2ZnS + 3O_2 \rightarrow 2ZnO + 2SO_2$

Calcination

- a) It is a process of heating ore in limited supply of air so as to convert carbonate ores into oxides.
- b) Carbonate ores are converted into oxide by roasting
- c) It is also used to remove moisture and volatile impurities.
- d) Example $CaCO_3 \xrightarrow{Heat} CaO + CO_2$

• Reduction of oxide to metal:

The process of converting metal oxide into metal is called reduction. It needs a suitable reducing agent depending upon the reactivity or reducing power of metal. The common reducing agents used are carbon or carbon monoxide or any other metals like Al, Mg etc.

• Thermodynamic principles of metallurgy:

Some basic concepts of thermodynamics help in understanding the conditions of temperature and selecting suitable reducing agent in metallurgical processes:

- a) Gibbs free energy change at any temperature is given by $\Delta G = \Delta H T\Delta S$ where ΔG is free energy change, ΔH is enthalpy change and ΔS is entropy change.
- b) The relationship between ΔG^{θ} and K is $\Delta G^{\theta} = -2.303$ RT log K where K is equilibrium constant. R = 8.314 JK-¹ mol-1, T is temperature in Kelvin.
- c) A negative ΔG means +ve value of K i.e., products are formed more than the reactants. The reaction will proceed in forward direction.
- d) If ΔS is +ve, on increasing temperature the value of $T\Delta S$ increases so that $T\Delta S > \Delta H$ and ΔG will become negative.

Coupled reactions:

If reactants and products of two reactions are put together in a system and the net ΔG of two possible reactions is –ve the overall reaction will take place. These reactions are called coupled reactions.

Ellingham diagrams:

The plots between ΔfG^{θ} of formation of oxides of elements vs. temperature are called Ellingham diagrams. It provides a sound idea about selecting a reducing agent in reduction of oxides. Such diagrams help in predicting the feasibility of a thermal reduction of an ore. ΔG must be negative at a given temperature for a reaction to be feasible.

• Limitations of Ellingham Diagrams:

It does not take kinetics of reduction into consideration, i.e., how fast reduction will take place cannot be determined.

Reduction of iron oxide in blast furnace:

Reduction of oxides takes place in different zones.

a) At 500 – 800 K (lower temperature range in blast furnace)

$$3Fe_2O_3 + CO \rightarrow 2Fe_3O_4 + CO_2$$

$$Fe_3O_4 + 4CO \rightarrow 3Fe + 4CO_2$$

$$Fe_2O_3 + CO \rightarrow 2FeO + CO_2$$

b) At 900 – 1500 K (higher temperature range in blast furnace)

$$C + CO_2 \rightarrow 2CO$$

 $FeO + CO \rightarrow Fe + CO_2$

c) Limestone decomposes to CaO and CO₂

$$CaCO_3 \xrightarrow{Heat} CaO + CO_2$$

e) Silica (impurity) reacts with CaO to form calcium silicate which forms slag. It floats over molten iron and prevents oxidation of iron.

$$CaO + SiO_2 \longrightarrow CaSiO_3$$
Calcium Silicate (slag)

Types of iron:

- a) Pig iron: The iron obtained from blast furnace is called pig iron. It is impure from of iron contains 4% carbon and small amount of S,.P, Si and Mn. It can be casted into variety of shapes.
- b) Cast iron: It is made by melting pig iron with scrap iron and coke using hot air blast. It contains about 3% of carbon content. It is extremely hard and brittle.
- c) Wrought iron: It is the purest form of commercial iron. It is also called malleable iron. It is prepared by oxidative refining of pig iron in reverberatory furnace lined with haematite which oxidises carbon to carbon monoxide.

$$Fe_2O_3 + 3C \rightarrow 2Fe + 3CO$$

The substance which reacts with impurity to form slag is called flux e.g. limestone is flux.

$$S + O_2 \rightarrow SO_2$$

 $4P + 5O_2 \rightarrow 2P_2O_5$
 $Si + O_2 \rightarrow SiO_2$
 $CaO + SiO_2 \rightarrow CaSiO_3(slag)$
 $3CaO + P_2O_5 \rightarrow Ca_3(PO_4)_2(slag)$

The metal is removed and freed from slag by passing through rollers.

• Electrolytic Reduction (Hall - Heroult Process):

Purified bauxite ore is mixed with cryolite (Na₃AlF₆) or CaF₂ which lowers its melting point and increases electrical conductivity. Molten mixture is electrolysed using a number of graphite rods as anode and carbon lining as cathode.

The graphite anode is useful for reduction of metal oxide to metal.

$$2Al_2O_3 + 3C \rightarrow 4Al + 3CO_2$$

$$Al_2O_3 \xrightarrow{Electrolysis} 2Al^{3+} + 3O^{2-}$$

At cathode: $Al^{3+}(melt) + 3e^{-} \rightarrow Al(l)$

At anode: $C(s) + O^{2-}(melt) \to CO(g) + 2e^{-}$ $C(s) + 2O^{2-}(melt) \to CO_2(g) + 4e^{-}$

Graphite rods get burnt forming CO and CO₂. The aluminium thus obtained is refined electrolytically using impure Al as anode, pure Al as cathode and molten cryolite as electrolyte.

At anode:
$$Al \rightarrow Al^{3+} + 3e^{-}$$

At cathode: $Al^{3+} + 3e^{-} \rightarrow Al(pure)$

• Electrolysis of molten NaCl:

$$NaCl \rightarrow Na^+ + Cl^-$$

At cathode: $Na^+ + e^- \rightarrow Na$ At anode: $2Cl^- \rightarrow Cl_2 + 2e^-$

Thus sodium metal is obtained at cathode and Cl₂ (g) is liberated at anode.

Refining:

It is the process of converting an impure metal into pure metal depending upon the nature of metal.

• Distillation:

It is the process used to purify those metals which have low boiling points, e.g., zinc, mercury, sodium, potassium. Impure metal is heated so as to convert it into vapours which changes into pure metal on condensation and is obtained as distillate.

• Liquation:

Those metals which have impurities whose melting points are higher than metal can be purified by this method. In this method, Sn metal can be purified. Tin containing iron as impurities heated on the top of sloping furnace. Tin melts and flows down the sloping surface where iron is left behind and pure tin is obtained.

• Electrolytic refining:

In this method, impure metal is taken as anode, pure metal is taken as cathode, and a soluble salt of metal is used as electrolyte. When electric current is passed, impure metal forms metal ions which are discharged at cathode forming pure metal.

At anode:
$$M_{\text{(Impure)}} \rightarrow M^{n+} + ne^{-}$$

At cathode:
$$M^{n+} + ne^- \rightarrow M_{(Pure)}$$

• Zone refining:

It is based on the principle that impurities are more soluble in the melt than in the solid state of the metal. The impure metal is heated with the help of circular heaters at one end of the rod of impure metal. The molten zone moves forward along with the heater with impurities and reaches the other end and is discarded. Pure metal crystallizes out of the melt. The process is repeated several times and heater is moved in the same direction. It is used for purifying semiconductors like B, Ge, Si, Ga and In.

• Vapour phase refining:

Nickel is purified by Mond's process. Nickel, when heated in stream of carbon monoxide forms volatile Ni(CO)₄ which on further subjecting to higher temperature decomposes to give pure metal.

$$Ni + 4CO \xrightarrow{330-350k} Ni(CO)_4 \xrightarrow{450-470k} Ni + 4CO$$
Impure

• Van- Arkel method:

It is used to get ultra pure metals. Zr and Ti are purified by this process. Zr or Ti are heated in iodine vapours at about 870 K to form volatile ZrI4 or TiI4 which are heated over tungsten filament at 1800K to give pure Zr or Ti.

$$\begin{split} & Ti + 2I_2 \rightarrow TiI_4 \rightarrow \underset{Pure}{Ti} + 2I_2 \\ & Zr + 2I_2 \rightarrow ZrI_4 \rightarrow \underset{Pure}{Zr} + 2I_2 \end{split}$$

Chromatographic method:

It is based on the principle of separation or purification by chromatography which is based on differential adsorption on an adsorbent. In column chromatography, Al_2O_3 is used as adsorbent. The mixture to be separated is taken in suitable solvent and applied on the column. They are then eluted out with suitable solvent (eluent). The weakly adsorbed component is eluted first. This method is suitable for such elements which are available only in minute quantities and the impurities are not very much different in their chemical behaviour from the element to be purified.