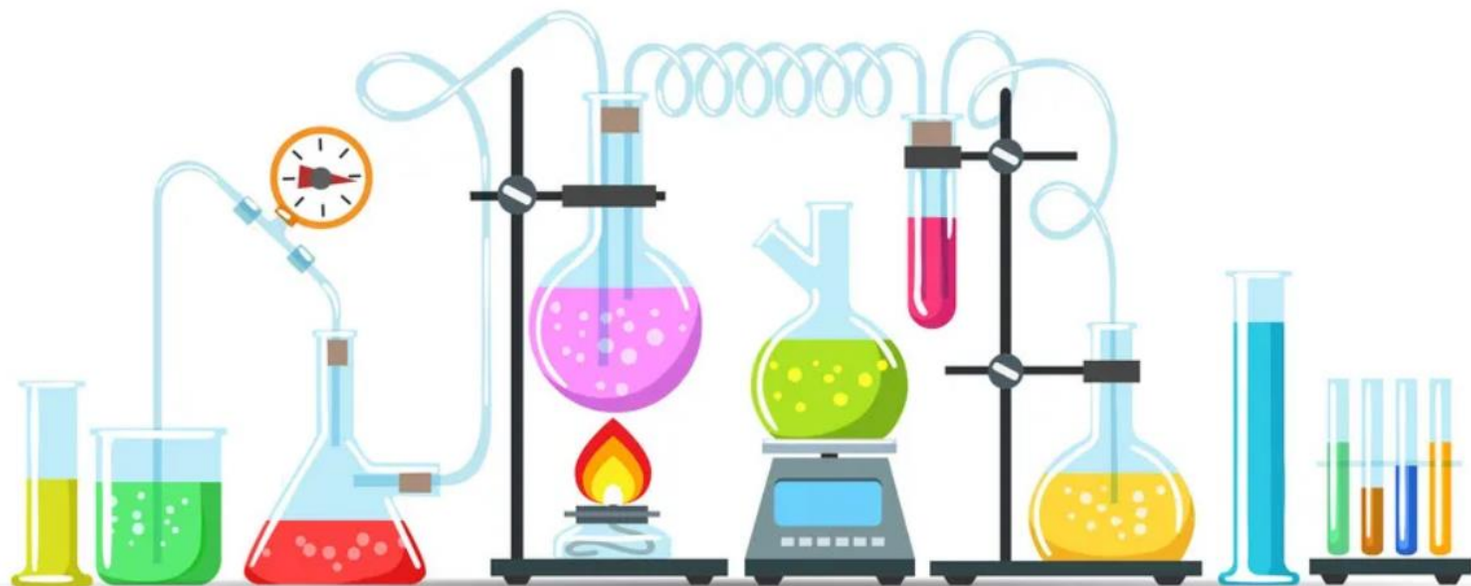


# CHEMISTRY

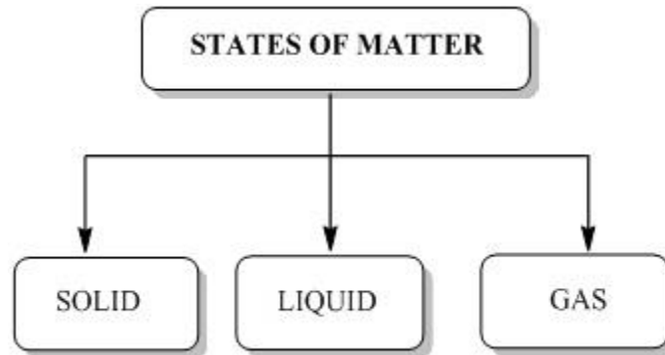
## CHAPTER 1: THE SOLID STATE



## THE SOLID STATE

### Introduction

Everything in our surroundings is known as matter that can be categorized into three states.



In our day to day life solids plays a crucial role to pursue different purposes. Different types of solids with different properties are required for different purposes. The constituent particles and the type of bonds between the particles determine the nature of a specific solid. For eg. Bucket or a container used to carry water, utensils used for cooking food, computer, vehicles, electronic gadgets, notebooks, pencils, papers etc. are all solid substances used in our day to day life.



Liquids and gases on the other hand are another state of matter and are also known as fluids due to their ability to flow. They attain the ability to flow due to the free movement of molecules.

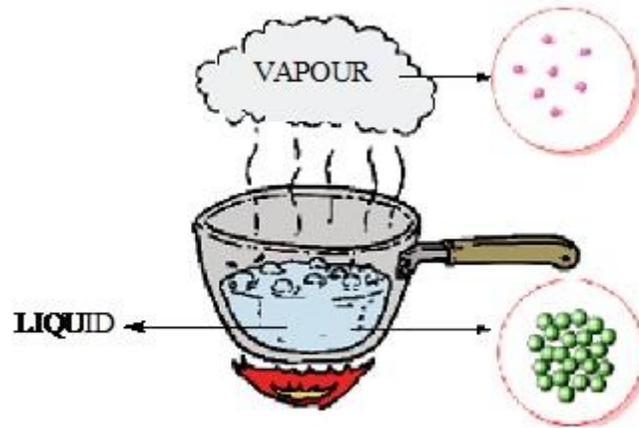
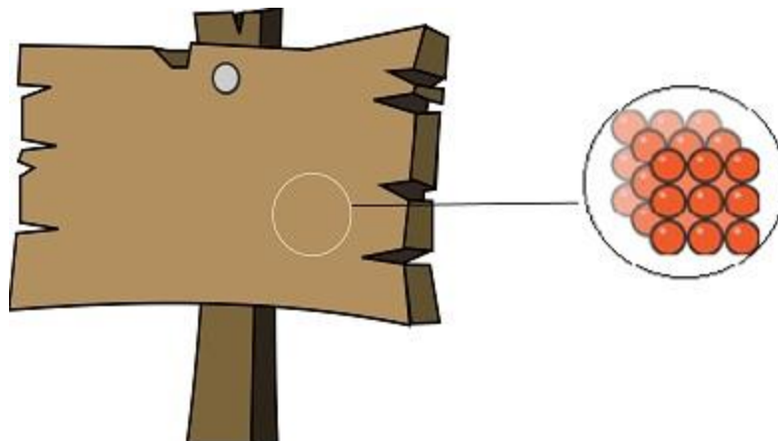


Fig. Particles of liquid (water) are loosely packed than solid and have space between them whereas particles of gas are loosely packed and have excess space between the particles

### Characteristics of solid state

- They have definite shape due to strong Intermolecular forces of attraction.
- They have distinct boundaries.
- They have a fixed volume.
- They cannot flow.
- They have negligible compressibility due to negligible distance between the neighbouring molecules.
- They possess a tendency to uphold their shape when exposed to external force.
- They break under force but it is difficult to change their shape so they are rigid.
- They have high density and do not diffuse at all.



**Fig.** Particles of solid (Wooden plank) are compact together and have less space between them

## Classification of Solids

Solids are classified on the basis of two different parameters i.e.,

1. Packing of constituent particles
2. Forces of attraction among constituent particles.

## Classification on the basis of packing

### 1. Crystalline Solids

These are the solids in which the constituent particles (atoms, ions or molecules) are arranged in a regular, three dimensional orderly arrangement, which gets repeated throughout. Hence, these are also called as long-range order solids. Quartz, diamond, Boron Nitride, NaCl, ZnS, CsCl etc.



### 2. Amorphous Solids

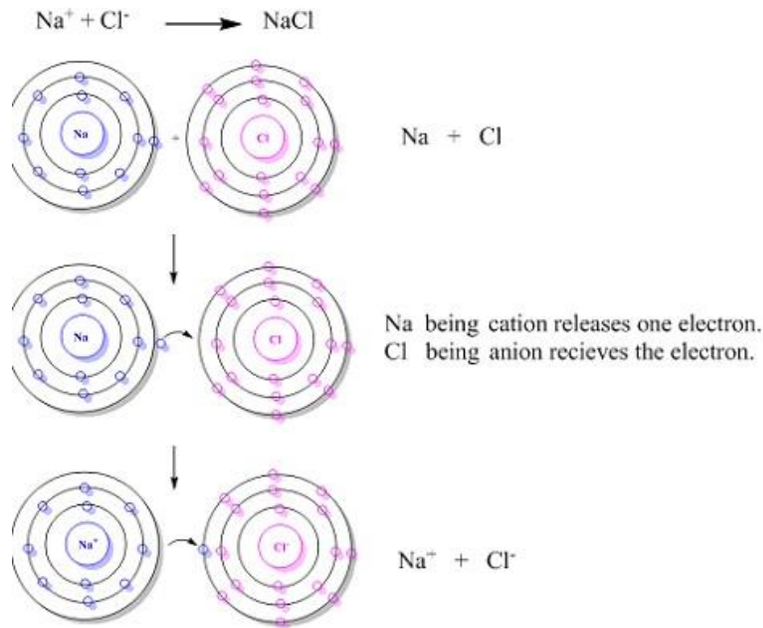
Constituent particles are randomly scattered with, no regular arrangement of particles. Therefore they do not have any definite shape or form. Rubber, glass, plastic etc. are commonly known as amorphous solids.



## Classification on the basis of Nature of Intermolecular Forces

### (i) Ionic Solids

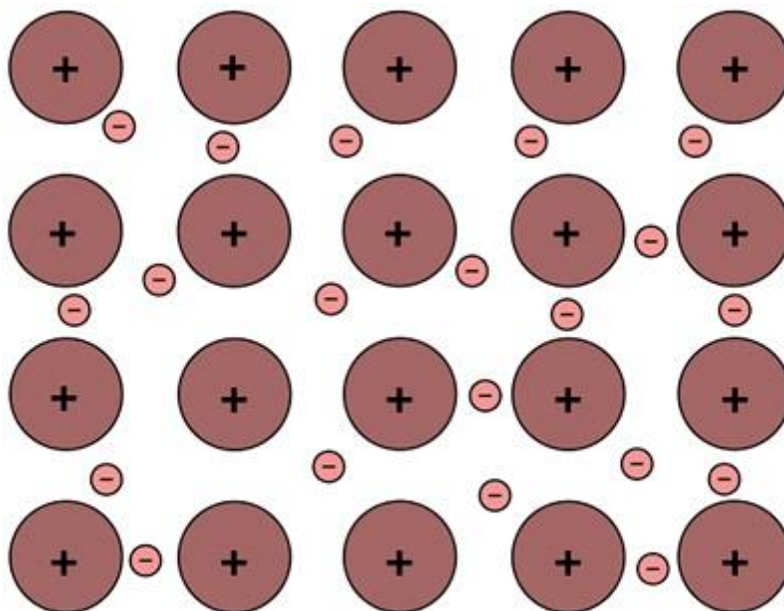
There is a regular arrangement of positively and negatively charged ions throughout the solid where ions are held together by strong coulombic or electrostatic forces. These solids are very hard and brittle and have very high melting points. In solid state, as ions are not free to move, hence they are insulators but in molten state or in aqueous state, it's ions become free to move and it becomes conductor. Ionic solids have high enthalpies of vaporisation. Ex-  $\text{LiF}$ ,  $\text{NaCl}$ ,  $\text{KNO}_3$ ,  $\text{MgO}$ , etc.



## (ii) Metallic Solids

Metal cores and a sea of mobile electrons are the constituents of metallic solids. Each metal atom contributes one or more electrons towards sea of electrons. These electrons are evenly spread out throughout the crystals and weak forces of attraction or metallic bond binds together kernels and sea of electrons.

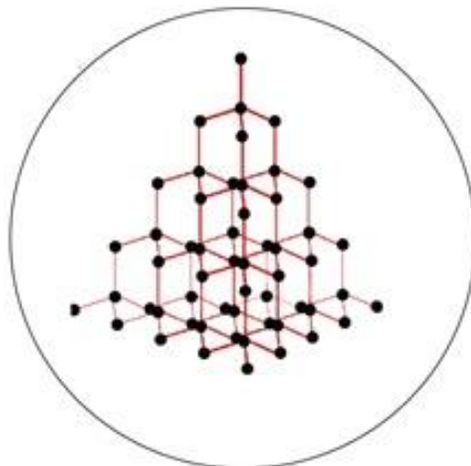
Metallic crystals may be hard as well as soft having moderate enthalpies of fusion. Mobile sea of electrons is responsible for many properties of metals such as malleability (can be beaten into thin sheets), ductility (can be drawn into wires), metallic lustre, thermal conductivity and electrical conductivity etc. Ex- Copper, Iron, Nickel etc.



Metallic Solids

### (iii) Covalent Solids (Network Solid)

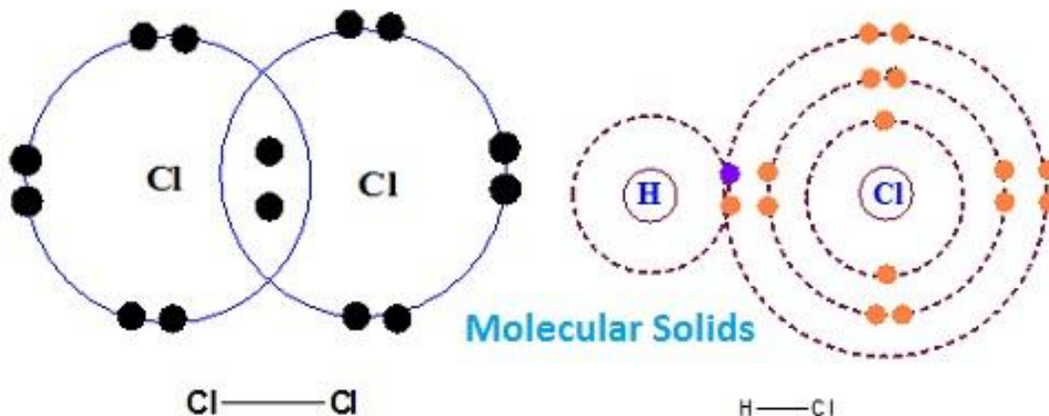
In these atoms are bonded together by covalent bond formation throughout the crystal. It means there is a continuous network of covalent bonds forming a giant three dimensional structure or giant molecule. Covalent bonds are strong and directional in nature. These solids are very hard, brittle and very high melting point. Due to absence of any free electrons or ions they are insulators. Their enthalpies of fusion are very high. Ex- Diamond, Graphite, Boron Nitride, Silicon Carbide (SiC) etc.



Covalent Solids

### (iv) Molecular Solids

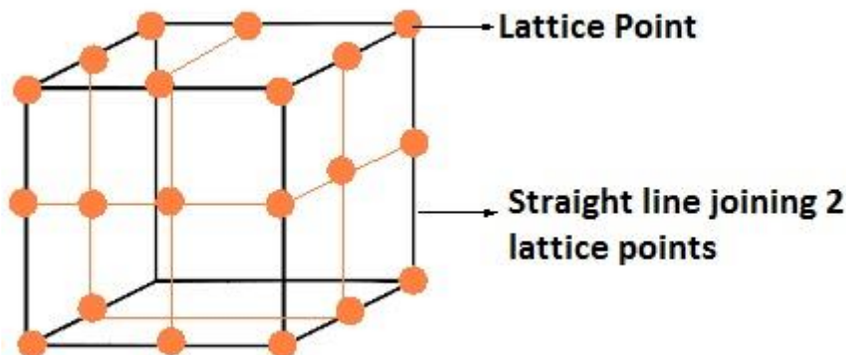
Their molecules are held together by dispersion forces, London forces, dipole-dipole forces or hydrogen bonds. Either atoms or molecules are bonded together by weak dispersion forces or London forces. These are non-conductor soft solids with low m.p. and low enthalpies of vaporisation. They are volatile in nature hence, at room temperature and pressure they are available in liquid or gaseous state. Ex- Iodine, Solid  $H_2$  and  $CO_2$  (dry ice). Naphthalene, Camphor etc.



## Crystal Lattices and Unit Cells

### (i) Crystal Lattices

In a crystalline solid, constituent particles are arranged in a definite, three-dimensional regular geometrical order along all the three axes, in which each particle is depicted as a lattice point. A three-dimensional, regular arrangement of lattice points in space or in a crystal is called a crystal lattice or space lattice.

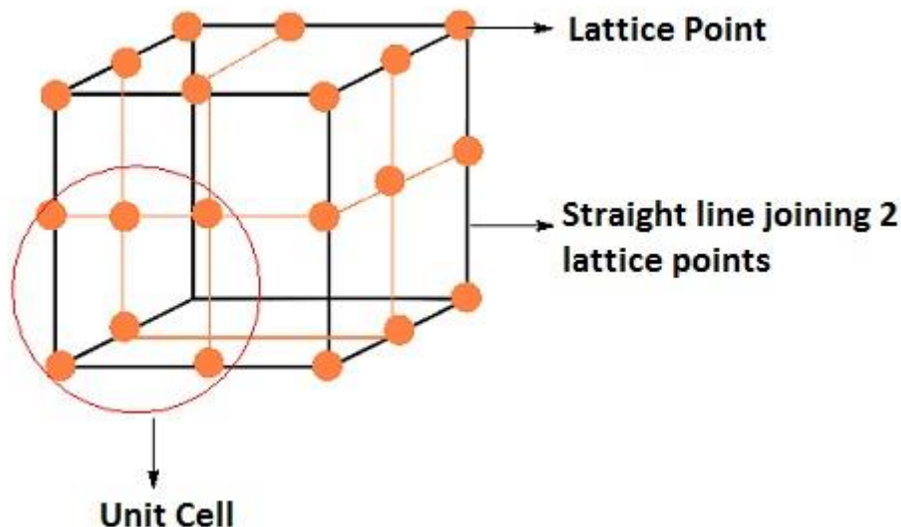


Crystal lattice have following characteristics

- Each point in lattice is called lattice site or lattice point.
- Each lattice point represents one constituent particle i.e. atom, ion or molecule.
- We join lattice points by straight lines to show geometry of the lattice.

### (ii) UNIT CELL

The smallest repeating unit in space lattice which when repeated over and over again in different directions produce complete crystal lattice. There are two important parameters of a unit cell.



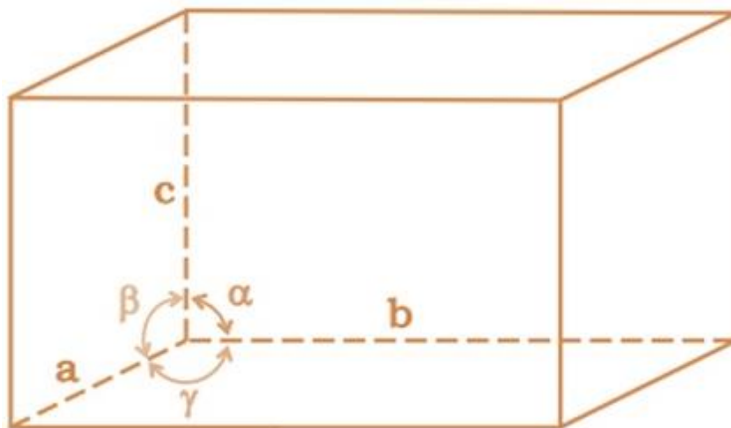
**Edge length or Axial Distance:** Lengths or dimensions along the three edges  $a$ ,  $b$  and  $c$  which may or may not be mutually perpendicular.

**Interaxial Angle:** Angles  $\alpha$ ,  $\beta$  and  $\gamma$  between pair of edges are interaxial angle.

$\alpha$  : between axis B and C

$\beta$  : between axis A and C

$\gamma$  : between axis A and B



### Distinction Between Crystalline and Amorphous Solids

| S.No | Crystalline solid | Amorphous solids |
|------|-------------------|------------------|
|------|-------------------|------------------|

|   |   |   |
|---|---|---|
| 1 | These have definite and regular arrangement of the constituent particles in space.                      | These doesn't have any regular arrangement of the constituent particles in space.             |
| 2 | These are true solids.  | These are super cooled liquids or pseudo solids.  |
| 3 | These have long order arrangement of the particles.   | These have short order arrangement of particle.   |
| 4 | These are anisotropic in nature, i.e., their physical properties are different in different directions. | These are isotropic in nature i.e., their physical properties are same in all the directions. |
| 5 | They have sharp melting points.   | They melt over a certain range of temperature.  |
| 6 | They undergo a clean cleavage when cut.   | They undergo irregular cleavage when cut.   |

### Three-Dimensional Packing

- To understand the packing of constituent particles in a crystal, particles are assumed to be hard spheres of identical size.
- The packing of these hard spheres takes place in such a way that they cover maximum available space and minimum space should be left behind.
- Because of this, the crystal has maximum density. This type of packing is known as close packing.

### Close Packing in One Dimension

- Spheres are arranged in such a way that they should touch each other in the row.
- In this arrangement, each sphere touches two neighbouring spheres. Hence, the coordination number is 2 in this arrangement.



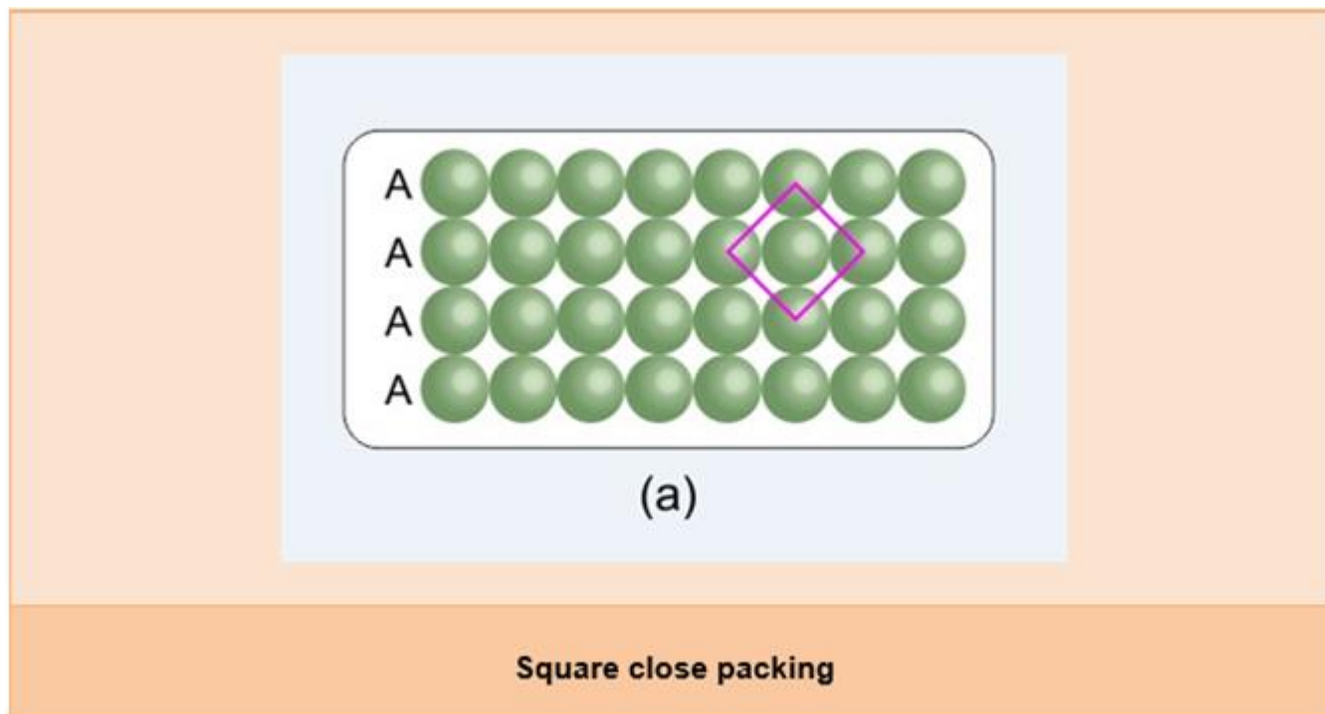
Close packing of spheres in one dimension

## Close Packing in Two Dimensions

- When the rows of one dimension packing are stacked over each other, a two-dimensional closepack structure is formed.
- This stacking is done in two ways:

### (a) Square close packing in two dimensions:

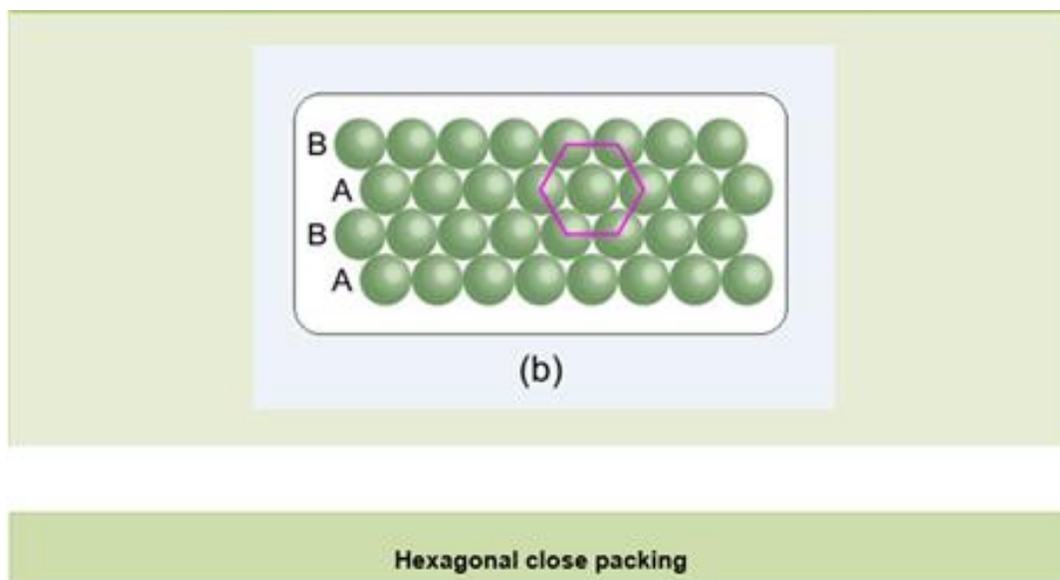
- The spheres in the second row are arranged in such a way that they touch the spheres of the first row and are exactly above the first row.
- If the first row is called the 'A' type row, then the second row will also be the 'A' type as both rows are identical.
- These arrangements are continuous; hence, we can call this arrangement as the AAA type arrangement.
- In this arrangement, each sphere touches four other spheres; hence, the coordination number is four.
- Also, if we join the centres of four spheres touching one particular sphere, then it forms a square; hence, it is called square close packing in two dimensions.



### (b) Hexagonal close packing in two dimensions:

- Spheres in the second row are placed in depressions of the first row. This gives two rows.

- Hence, if we call the first row 'A' type, then the second row will be the 'B' type.
- If a third row is placed in the depression of the second row, then it will be identical to the first row, i.e. the 'A' type.
- The fourth row will be the 'B' type and so on. Hence, this arrangement is called the ABAB type of arrangement.
- In this arrangement, each sphere touches six neighbouring spheres; hence, the coordination number is six.
- Also, if we join the centres of the spheres which touch one particular sphere, then it will give a hexagonal structure; therefore, it is known as hexagonal close packing into two dimensions.
- This type of packing is more effective because maximum space is covered by particles and the space is minimum.

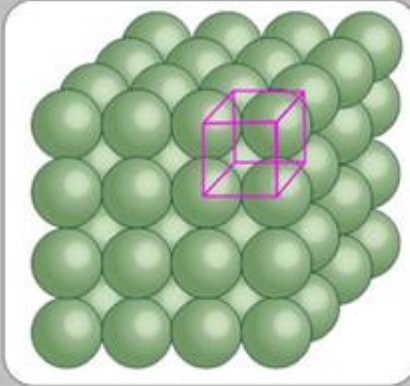


### Close Packing in Three Dimensions

- Three-dimensional packing can be obtained by square close packing and hexagonal close packing.

#### (a) Three-dimensional close packing from two-dimensional square close packing:

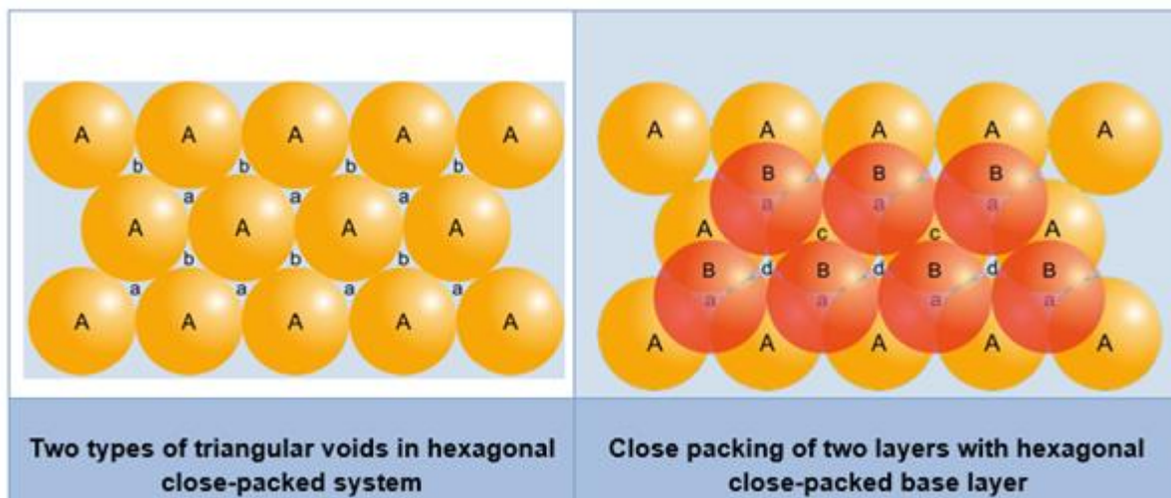
1. The second layer and all further layers are arranged in such a way that they are horizontal and vertically aligned with each other.
2. Hence, if we call the first layer as the 'A' type, then the lattice will be the AAA type.
3. This will give simple cubic lattice, and its unit cell will be a primitive cubic unit cell.



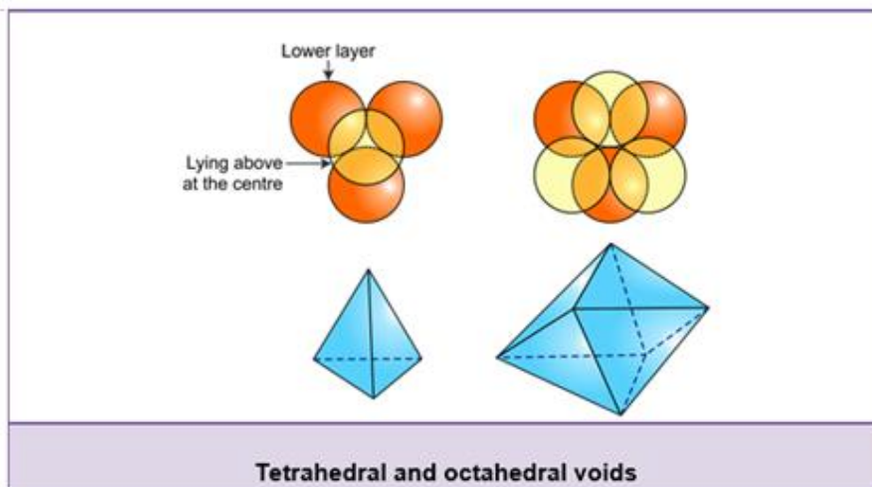
Three-dimensional packing of square close packed layers forming simple cubic lattice with AAA arrangement

**(b) Three-dimensional packing from two-dimensional hexagonal close packing:**

1. We know that more effective packing is given by hexagonal close packing.
2. Assume three-dimensional packing with a hexagonal close packed system. The base layer is called 'A' and the voids between the spheres are named 'a' and 'b' alternately.
3. Both 'a' and 'b' voids are triangular in shape. The only difference is that the apices of voids 'a' point downwards and those of 'b' point upwards.
4. The second layer is placed in such a way that its spheres find place in the 'a' voids of the first layer. The 'b' voids are left unoccupied because no spheres can be placed in them.
5. There are two new types of voids in the second layer—'c' and 'd'. The voids 'c' lie above the spheres of the first layer and the voids 'd' lie on the voids of the first layer.



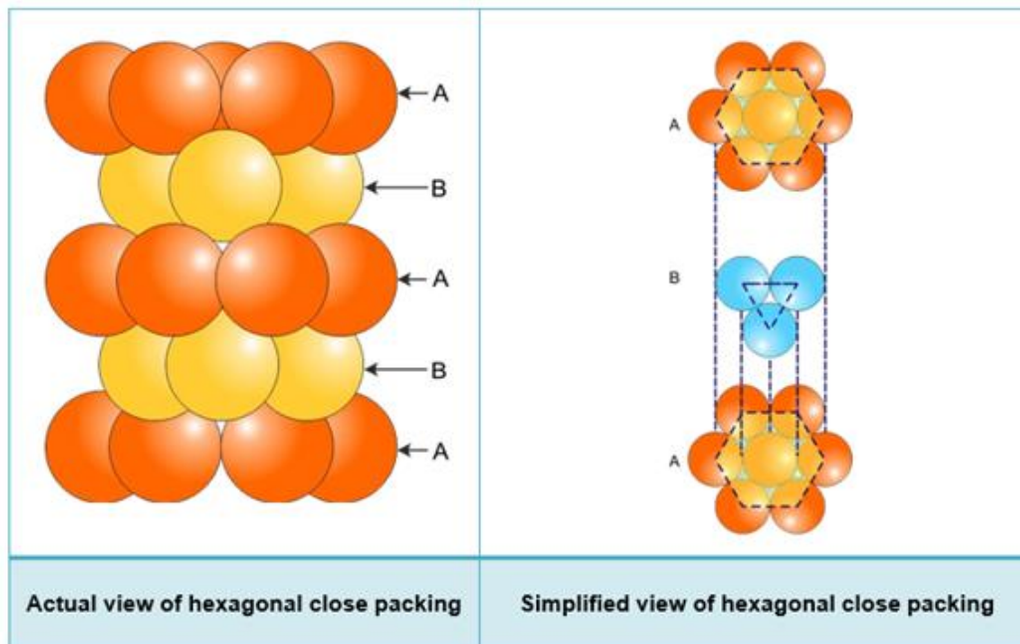
6. A simple triangular void 'c' which is surrounded by four spheres is called a tetrahedral void. The double triangular void (like 'd') which is surrounded by six spheres is called an octahedral void.
7. The voids or holes in the crystals are also called interstices.



8. There are two ways of building the third layer.

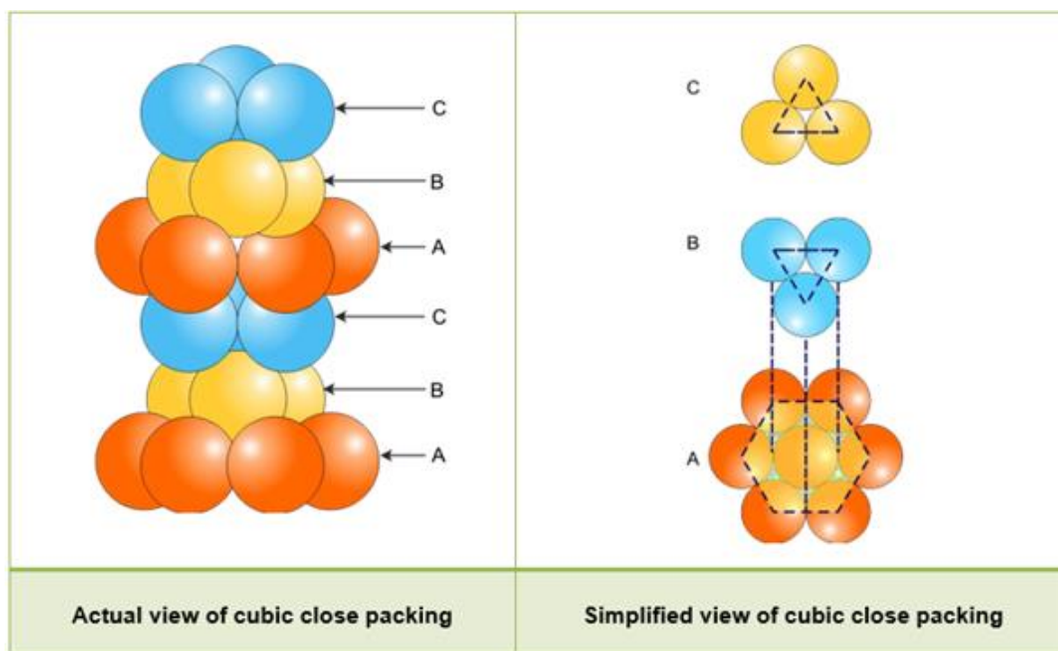
**hcp structure:**

- When the third layer is placed on the second layer in such a way that the spheres cover the tetrahedral voids, three-dimensional closed packing is obtained.
- Consider the first layer as 'A' and the second layer as 'B', the arrangement is of the ABAB type or hexagonal closed packing.
- Molybdenum, magnesium and beryllium crystallise in the hcp structure.



**ccp structure:**

- i. When the third layer is placed on the second layer in such a way that the spheres cover the octahedral voids, two layers 'A' and 'B' are formed. Assume that the new layer be 'C'.
- ii. On continuing, a packing is obtained where the spheres in every fourth layer will be vertically aligned. This pattern is called the ABCABC pattern or cubic close packing.
- iii. It is the same as face-centred cubic close packing.
- iv. Iron, nickel, copper, silver, gold and aluminium crystallise in the ccp structure.

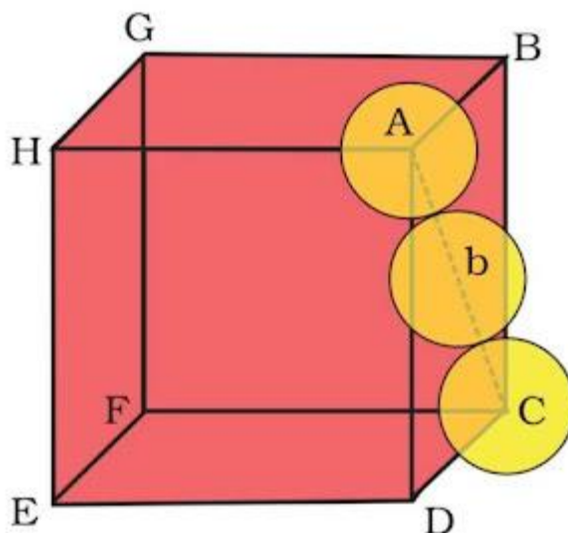


### Packing Efficiency

In all the type of packings there is always some free space in the form of voids or vacant spaces. Packing efficiency is the percentage of total space filled by the particles.

Packing efficiency = Volume of total lattice points / Total volume of unit cell

#### (i). Packing efficiency in fcc or ccp structures



suppose edge length of unit cell =  $a$

And radius of sphere =  $r$

$$\text{In } \triangle ABC, AC^2 = BC^2 + AB^2$$

$$AC^2 = a^2 + a^2$$

$$AC^2 = 2a^2$$

$$\text{Hence, face diagonal } AC = \sqrt{2}a$$

$$\text{But } AC = 4r \therefore 4r = \sqrt{2}a \therefore a = \frac{4r}{\sqrt{2}}$$

$$\text{Hence, volume of unit cell} = a^3 \text{ or}$$

$$a^3 = \left( \frac{4r}{\sqrt{2}} \right)^3 = 16\sqrt{2}r^3$$

As there are total 4 lattice points per unit cell.

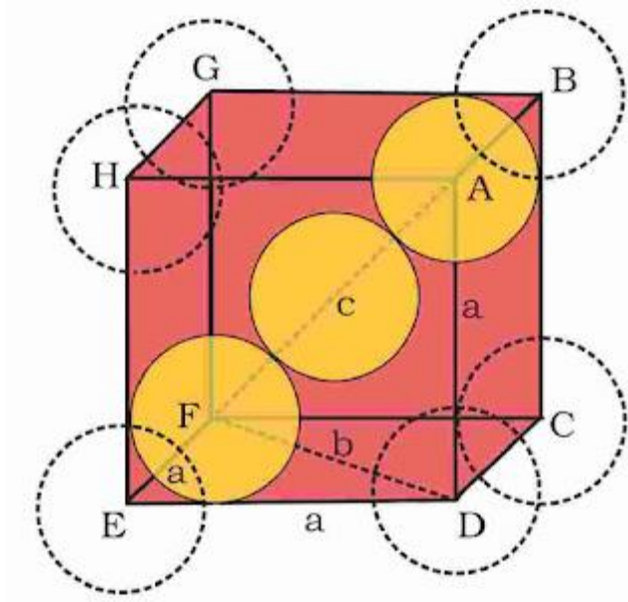
$$\text{Packing efficiency} = \frac{4 \times \frac{4}{3}\pi r^3}{16\sqrt{2}r^3}$$

$$= \frac{16\pi r^3}{3 \times 16\sqrt{2}r^3}$$

$$= \frac{\sqrt{2}\pi}{6} = 0.74$$

Hence, % Packing efficiency = 74%

## (ii). Packing Efficiency in bcc structures



Suppose the edge length =  $a$

Radius of each sphere =  $r$

Number of lattice points per unit cell = 2

$$\text{In } \triangle ABC, AC^2 = BC^2 + AB^2 = a^2 + a^2 = 2a^2$$

Similarly in  $\triangle ACD$ ,

$$AD^2 = AC^2 + CD^2 = 2a^2 + a^2 = 3a^2$$

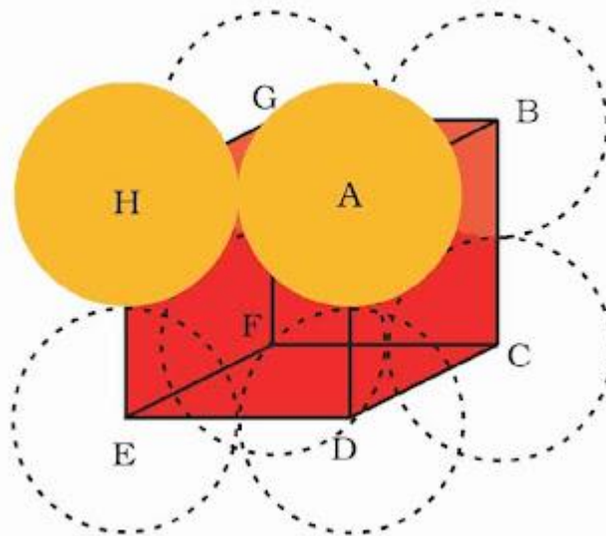
$$AD = \sqrt{3}a$$

$$\text{Now, } AD = 4r \therefore \sqrt{3}a = 4r \text{ or } a = \frac{4r}{\sqrt{3}}$$

$$\begin{aligned} \text{Packing efficiency} &= \frac{2 \times \frac{4}{3} \pi r^3}{\left(\frac{4r}{\sqrt{3}}\right)^3} \\ &= \frac{8\pi r^3 \times 3\sqrt{3}}{3 \times 64r^3} \\ &= \frac{\sqrt{3}\pi}{8} = 0.68 \end{aligned}$$

Hence, % of Packing efficiency = 68%

### (iii). Packing Efficiency in Simple Cubic Unit Cell



Let the edge length of unit cell =  $a$

Radius of sphere =  $r$

As two spheres touch each other at an edge

$$\therefore a = 2r$$

Number of lattice points per unit cell = 1

$$\text{Packing efficiency} = \frac{\frac{4}{3} \pi r^3}{(2r)^3} = \frac{\pi}{6} = 0.524$$

Hence, % Packing efficiency = 52.4%

## Coordination Number

- In both hcp and ccp structures, a sphere is in contact with six other spheres in its own layer.
- It also directly touches three spheres above and three spheres below. Thus, the sphere has 12 close neighbours. Hence, it is said to have coordination number 12.

**Coordination number:** Number of closest neighbours of any constituent particles in the crystal lattice

- The common coordination numbers in different types of crystals are 4, 6, 8 and 12.
- In crystals with directional bonds, the coordination number is lower than that of crystals with a non-directional bond.
- In addition to the above two types, there is another type called body-centred cubic close packing (bcc) in which the space occupied is 68%.
- The coordination number of each atom in the bcc structure is 8.

## Point defects:

Point defects explain about the imperfections of solids along with the types of point defects. Crystalline solids are formed by joining many small crystals. Different types of defects are found in crystals after the process of crystallization.

Point defects are accounted for when the crystallization process occurs at a very fast rate. These defects mainly happen due to deviation in the arrangement of constituting particles. In a crystalline solid, when the ideal arrangement of solids is distorted around a point/atom it is called a point defect.

Defects or Imperfections in crystalline solid can be divided into four groups namely line defects, point defects, volume defects and surface defects. Historically, crystal point defects were first regarded in ionic crystals, not in metal crystals that were much simpler.

### There are 3 types of point defects:

1. Stoichiometric defect
2. Frenkel defect
3. Schottky defect

### 1. Stoichiometric Defect:

In this kind of point defect, the ratio of positive and negative ions (Stoichiometric) and electrical neutrality of a solid is not disturbed. Sometimes it is also known as intrinsic or thermodynamic defects.

**Fundamentally, they are of two types:**

**Vacancy defect:** When an atom is not present at their lattice sites, then that lattice site is vacant and it creates a vacancy defect. Due to this, the density of a substance decreases.

**Interstitial defect:** It is a defect in which an atom or molecule occupies the intermolecular spaces in crystals. In this defect, the density of the substance increases.

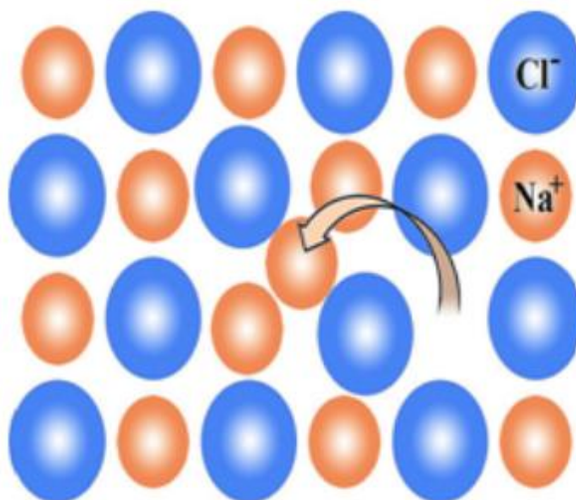
A non-ionic compound mainly shows vacancy and interstitial defects. An ionic compound shows the same in Frenkel and Schottky defect.

## 2. Frenkel Defect:

In ionic solids generally, the smaller ion (cation) moves out of its place and occupies an intermolecular space. In this case, a vacancy defect is created on its original position and the interstitial defect is experienced at its new position.

It is also known as dislocation defect, The density of a substance remains unchanged, It happens when there is a huge difference in the size of anions and cations.

Example: ZnS and AgCl.

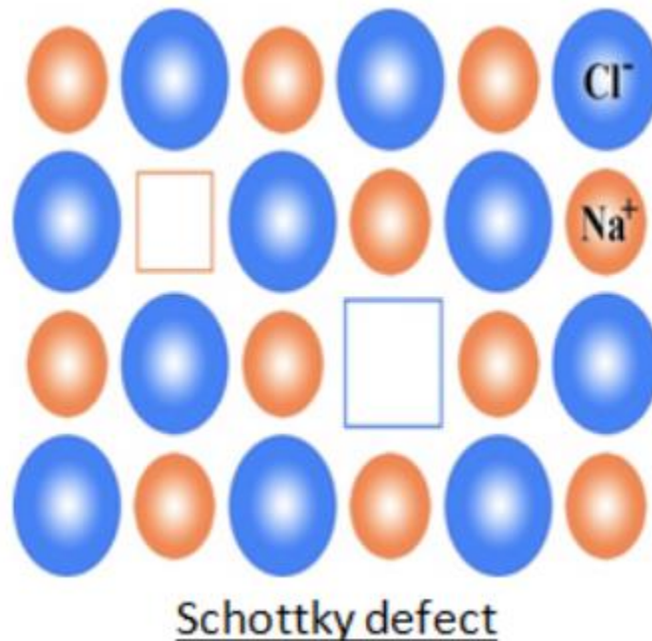


Frenkel defect

## 3. Schottky Defect

This kind of vacancy defects is found in Ionic Solids. But in ionic compounds, we need to balance the electrical neutrality of the compound so an equal number of anions and cations will be missing from the compound. It reduces the density of the substance. In this, the size

of cations and anions are of almost the same.



**Impurity Defect:** Let's understand the impurity defect by an example. If molten NaCl is crystallized with  $\text{SrCl}_2$  compound then the  $\text{Sr}^{2+}$  ions replace two  $\text{Na}^+$  ions and occupy the place of one  $\text{Na}^+$ . In this way the lattice site of one  $\text{Na}^+$  is vacant and it creates an impurity defect.

**Non-Stoichiometric Defect:** In this defect, the cations and anions ratio is disturbed either because of adding or removing of ions.

**Types of Non-Stoichiometric Defect:**

**Metal deficiency defect:** In this, the solids have less number of metals relative to the described Stoichiometric proportion.

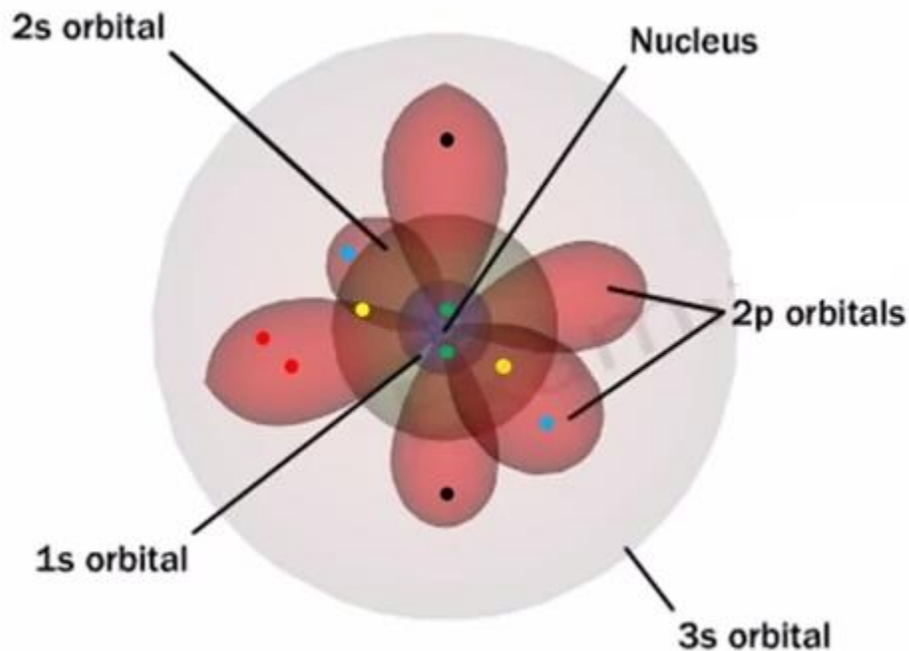
**Metal excess defect:**

There are two types of metal excess defect:

**Metal excess defect due to anionic vacancies:** This occurs due to the absence of anions from its original lattice site in crystals. Therefore, instead of anions, electrons occupy their position

**Metal excess defect due to the presence of extra cations at interstitial sites:** Here, on heating the compound, it releases extra cations. These cations occupy the interstitial sites in crystals and the same number of electrons goes to neighbouring interstitial sites.

**Magnetic Properties:**



- Every substance possesses magnetic properties originated from the electrons present in them.
- Each electron in an atom behaves like a tiny magnet, The magnetic moment of these substances originates from two types of motions its orbital motion around the nucleus and its spin around its own axis.
- Electron being a charged particle undergoes these motions and can be considered as a small loop of current possessing a magnetic moment, Therefore, each electron has a permanent spin and an orbital magnetic moment associated with it.
- Magnitude of this magnetic moment is very small and is measured in the unit called Bohr magneton,  $\mu_B$  and is equal to  $9.27 \times 10^{-24} \text{ A m}^2$ .

On the basis of their magnetic properties, substances can be classified into five categories:

- 1. Paramagnetic:** Paramagnetic materials are materials that tend to get weakly magnetized in the direction of the magnetizing field when placed in a magnetic field. Paramagnetic materials have a permanent dipole moment or permanent magnetic moment. However, if we remove the applied field the materials tend to lose their magnetism. This is because thermal motion randomizes the spin orientations of the electrons.
- 2. Diamagnetic:** The spin motion of electrons and their interaction with one another is what is responsible for the origin of magnetism. The best way to introduce different types of magnetic materials is by describing how materials respond to magnetism. It may come as a surprise to you, but all matter is magnetic. The only difference is that some materials are more magnetic than others. The level of interactions between the magnetic moments is what distinguishes them. In a few materials, there is no collective interaction of atomic

magnetic moment while other materials exhibit strong atomic magnetic moment interaction.

- 3. Ferromagnetic:** There are various kinds of magnetism, out of which ferromagnetism is the strongest type. Ferromagnetic materials are those materials which exhibit a spontaneous net magnetization at the atomic level, even in the absence of an external magnetic field.

When placed in an external magnetic field, ferromagnetic materials are strongly magnetized in the direction of the field. Ferromagnetic materials are strongly attracted to a magnet. These materials will retain their magnetization for some time even after the external magnetizing field is removed. This property is called hysteresis.

- 4. Anti-ferro magnetic:** The materials that exhibit the anti-ferro magnetism are known as antiferromagnetic material. When these materials are kept in the presence of the strong magnetic field, they get magnetized weakly in the direction of the magnetic field. This is known as anti-ferro magnetism.

The antiferromagnetic materials are commonly found among the transition metal compounds. Hematite, chromium, alloys of iron manganese and oxides of nickel are the examples of antiferromagnetic material

- 5. Ferrimagnetic:** Ferrimagnetic materials definition is, in which the magnetic dipoles of the atoms on various subset are opposed, as in anti-ferro magnetism, but the opposing moments are unequal in ferrimagnetic materials, leaving a random net magnetization. Crystal ferrimagnetic materials, including antiferromagnetic materials, have populations of atoms with contrasting magnetic moments. Since the magnitudes of these moments are unequal in ferrimagnet compounds, a random magnetization exists. A mixture of dipole-dipole interactions and exchange interactions arising from the Pauli exclusion theory induce magnetization of ferrimagnetic materials. The key distinction is that in a ferrimagnetic substance, the unit cell contains various groups of atoms.

## Semiconductor

Semiconductors are substances with properties somewhere between them. ICs (integrated circuits) and electronic discrete components such as diodes and transistors are made of semiconductors. Common elemental semiconductors are silicon and germanium. Silicon is well-known of these. Silicon forms most of ICs.

**Conduction of Electricity in Semiconductors** In semiconductors, the gap between the valence band and conduction band is small.

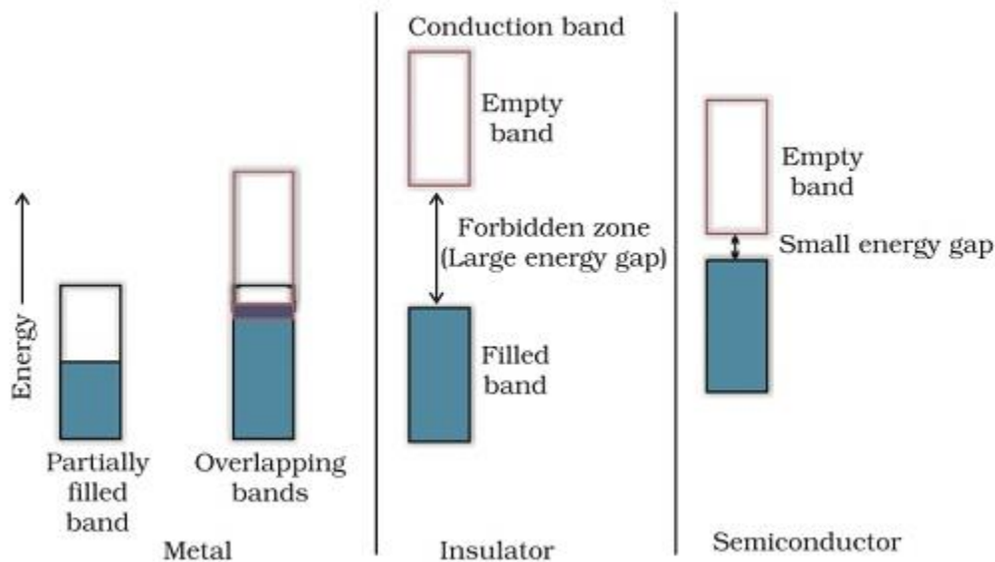
This enables some electrons to jump to conduction band and exhibit their conductivity.

Electrical conductivity of semiconductors increases with increase in temperature, since more electrons can jump to the conduction band due to small gap between the valence band and conduction band.

Silicon and germanium exhibit this behavior and are called intrinsic semiconductors. The conductivity of these intrinsic semiconductors is too low to be practically used.

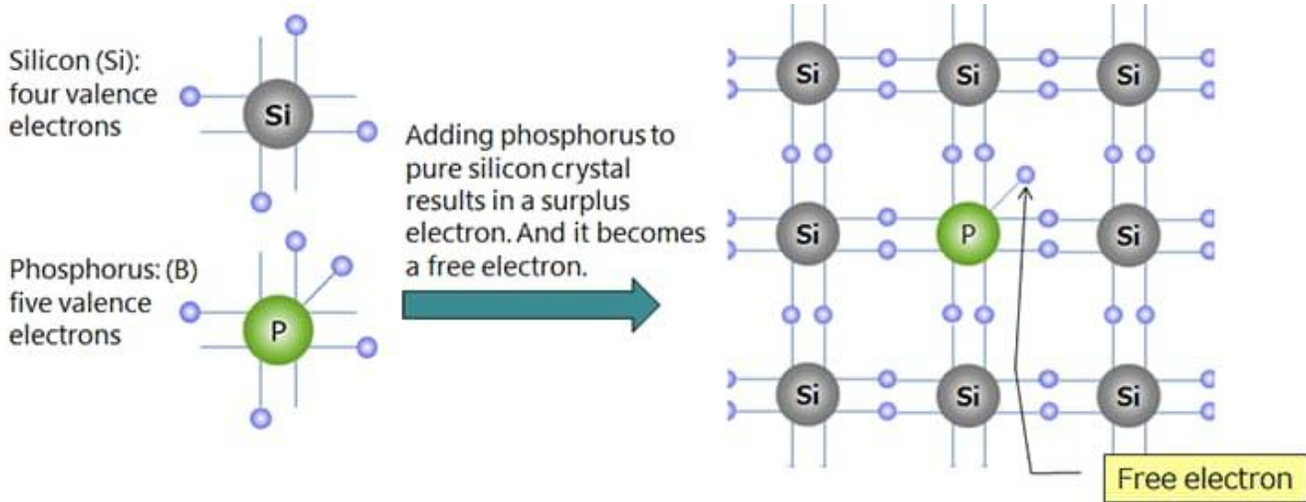
### Semiconductor materials

Solid-state materials are commonly grouped into three classes: insulators, semiconductors, and conductors. (At low temperatures some conductors, semiconductors, and insulators may become superconductors.) The figure shows the conductivities  $\sigma$  (and the corresponding resistivities  $\rho = 1/\sigma$ ) that are associated with some important materials in each of the three classes. Insulators, such as fused quartz and glass, have very low conductivities, on the order of  $10^{-18}$  to  $10^{-10}$  siemens per centimeter; and conductors, such as aluminum, have high conductivities, typically from  $10^4$  to  $10^6$  siemens per centimeter. The conductivities of semiconductors are between these extremes and are generally sensitive to temperature, illumination, magnetic fields, and minute amounts of impurity atoms. For example, the addition of about 10 atoms of boron (known as a dopant) per million atoms of silicon can increase its electrical conductivity a thousand fold



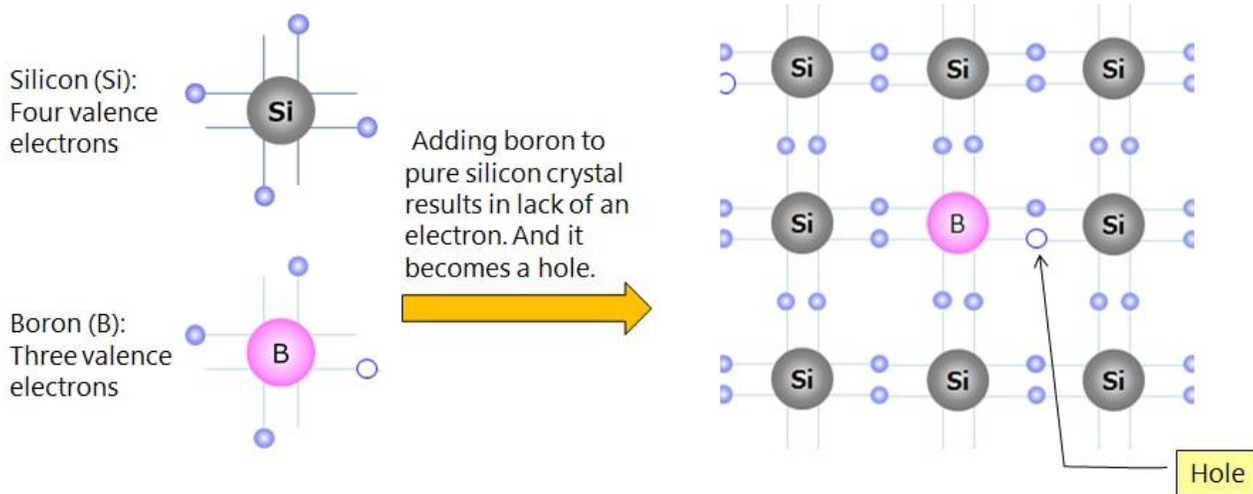
### n-type Semiconductor:

An n-type semiconductor is an intrinsic semiconductor doped with phosphorus (P), arsenic (As), or antimony (Sb) as an impurity. Silicon of Group IV has four valence electrons and phosphorus of Group V has five valence electrons. If a small amount of phosphorus is added to a pure silicon crystal, one of the valence electrons of phosphorus becomes free to move around (free electron\*) as a surplus electron. When this free electron is attracted to the “+” electrode and moves, current flows.



**p-type Semiconductor:**

A p-type semiconductor is an intrinsic semiconductor doped with boron (B) or indium (In). Silicon of Group IV has four valence electrons and boron of Group III has three valence electrons. If a small amount of boron is doped to a single crystal of silicon, valence electrons will be insufficient at one position to bond silicon and boron, resulting in holes\* that lack electrons. When a voltage is applied in this state, the neighboring electrons move to the hole, so that the place where an electron was present becomes a new hole, and the holes appear to move to the "-" electrode in sequence.



**Applications of n-type and p-type semiconductors:**

n-type and p-type semiconductors finds a great use in manufacturing electronic components. Diode is a combination of n-type and p-type semiconductors extensively used as a rectifier. Transistors are manufactured by keeping a layer of one type of semiconductor between two layers of another type of semiconductor. npn and pnp type of transistors are used to detect or amplify radio or audio signals. The solar cell is an efficient photo-diode used for conversion of light energy into electrical

energy.

Gallium arsenide (GaAs) semiconductors have very fast response and have transformed the design of semiconductor devices.

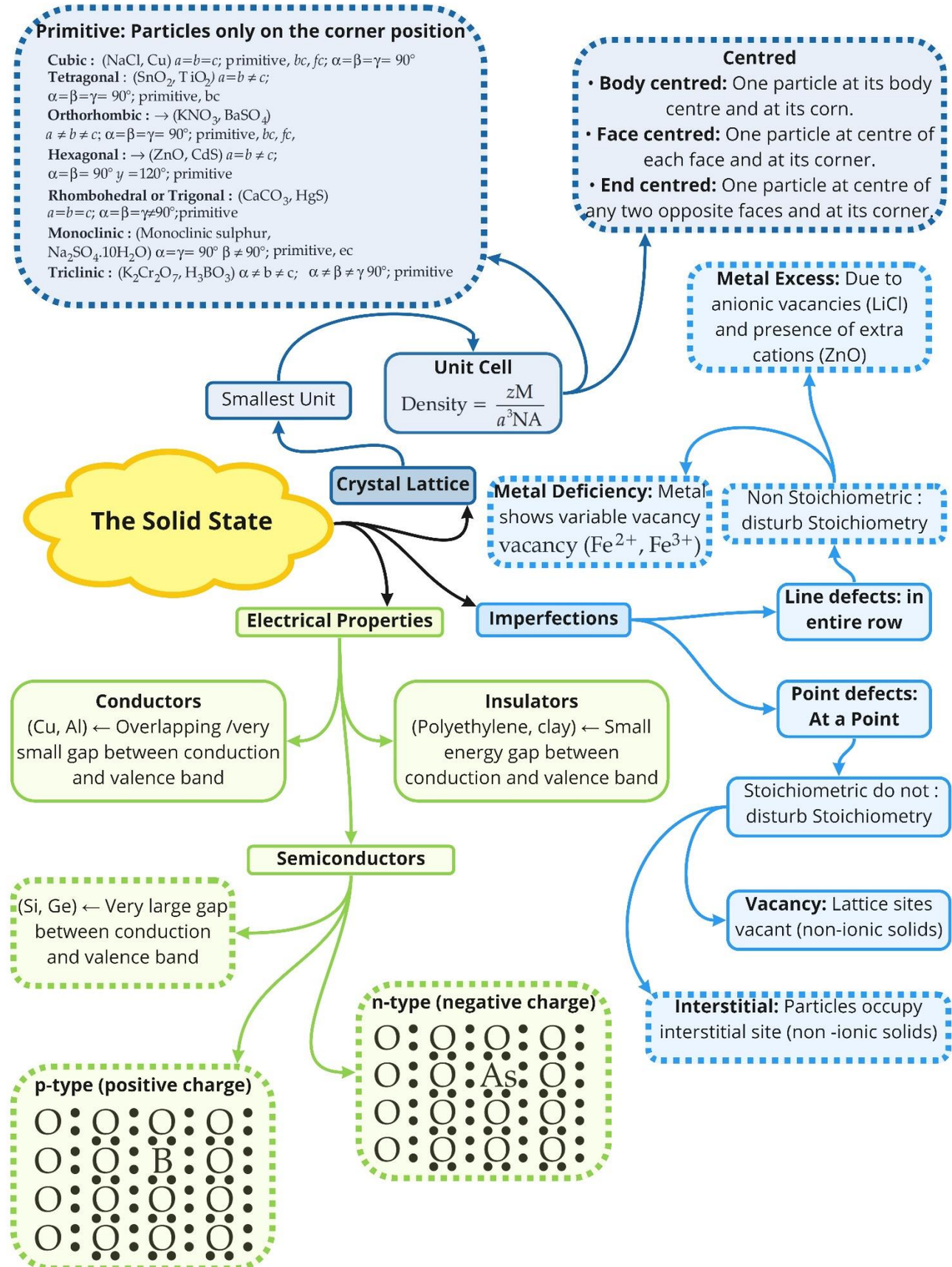
Transition metal oxides show marked differences in electrical properties.

TiO, CrO<sub>2</sub> and ReO<sub>3</sub> behave like metals.

Rhenium oxide, ReO<sub>3</sub> resembles metallic copper in terms of its conductivity and appearance.

Certain other oxides like VO, VO<sub>2</sub>, VO<sub>3</sub> and TiO<sub>3</sub> exhibit metallic or insulating properties depending on temperature.

Class : 12th Chemistry  
Chapter-1 : The Solid State (Part\_1)



Class : 12th Chemistry  
Chapter-1 : The Solid State (Part\_2)

### Magnetic Properties

- **Paramagnetic:** weakly attracted → Unpaired electrons → ( $O_2$ ,  $Cu^{2+}$ )
- **Diamagnetic:** weakly repelled → Paired electrons → ( $H_2O$ ,  $NaCl$ )
- **Ferromagnetic:** strongly attracted → domains in same direction → ( $Fe$ ,  $Co$ )
- **Antiferromagnetic:** domain opposite and equal → ( $MnO$ )
- **Ferromagnetic:** domains unequal → ( $Fe_3O_4$ ,  $MgFe_2O_4$ )

**Amorphous solids:**  
Particles with irregular shape

### Packing Efficiency

- scp → AAA....type → 52.4% → ( $Po$ )
- hcp → ABABAB.....type → 74% → ( $Mg$ ,  $Zn$ )
- ccp/ fcc → ABCABC....type → 74% → ( $Cu$ ,  $Ag$ )
- bcc → square close → 68% → ( $Li$ ,  $Na$ )

## The Solid State

**Voids:** Empty space between spheres

**Crystalline Solids:** Large Number of small crystal with definite shape

### Trigonal

Three spheres in contact →  
0.155 - 0.225

### Tetrahedral

Four spheres at the vertices of  
tetrahedron → 0.225 - 0.414

### Octahedral

Six spheres at vertices of  
octahedron → 0.414 - 0.732

### Metallic solids

Metal atoms held by  
metallic bond → ( $Fe$ ,  $Cu$ )

### Molecular solids

Molecules held by  
intermolecular forces

### Polar

Molecules held by dipole-dipole  
interactions → ( $HCl$ ,  $SO_2$ )

### Non-Polar

Atoms /molecules held by weak  
dispersion forces /London  
forces → ( $Ar$ ,  $He$ )

### Ionic solids

Ions held by strong coulombic  
forces → ( $NaCl$ ,  $MgO$ )

### Hydrogen bonded

Molecules held by hydrogen  
bonding → ( $H_2O$ (ice))

### Covalent solids

Non-metals held by  
covalent bond → ( $SiC$ ,  $C$ )

## Important Questions

### Multiple Choice questions-

- Close packing is maximum in the crystal which is
  - bcc
  - fcc
  - simple cubic
  - end centred cubic
- In a solid lattice, the cation has left a lattice site and is located at an interstitial position. The lattice defect is
  - n-type
  - p-type
  - Frenkel defect
  - Schottky defect
- The coordination number of metal crystallizing in a hexagonal close packing structure are
  - 12
  - 4
  - 8
  - 10
- In a body centred unit cell, the number of atoms present is
  - 1
  - 2
  - 3
  - 4
- In a trigonal crystal
  - $a = b = c, \alpha = \beta = \gamma \neq 90^\circ$
  - $a = b \neq c, \alpha = \beta = \gamma = 90^\circ$
  - $a \neq b \neq c, \alpha = \beta = \gamma = 90^\circ$
  - $a = b \neq c, \alpha = \beta = 90^\circ, \gamma = 120^\circ$
- The appearance of colour in solid alkali metal halides is generally due to
  - Schottky defect
  - Frenkel defect

- (c) F-centre  
(d) Interstitial position
7. In a body centred cubic structure, the space occupied is about
- (a) 74%  
(b) 20%  
(c) 68%  
(d) 52.4%
8. Some polar crystals when heated produce small electrical current. The phenomenon is called
- (a) Ferroelectricity  
(b) Anti-ferroelectricity  
(c) Pyroelectricity  
(d) Piezoelectricity.
9. The empty space within hcp arrangement is
- (a) 34%  
(b) 47.6%  
(c) 32%  
(d) 26%
10. A crystal system with axes  $a \neq b \neq c$  and angles  $\alpha \neq \beta \neq \gamma \neq 90^\circ$  corresponds to
- (a) monoclinic  
(b) triclinic  
(c) cubic  
(d) tetragonal
11. The number of  $\text{Cl}^-$  ions present around each  $\text{Na}^+$  ion in NaCl crystal lattice is
- (a) 3  
(b) 4  
(c) 8  
(d) 6
12. To get n-type semiconductor from silicon, it should be doped with an element having valence electrons of
- (a) 2  
(b) 1  
(c) 3

(d) 5

13. Volume occupied by atoms in fcc is

- (a) 74%
- (b) 68%
- (c) 52.4%
- (d) 75%

14. Percentage empty space in a bcc arrangement is

- (a) 74%
- (b) 68%
- (c) 32%
- (d) 26%

15. To get p-type semiconductor, impurity to be added to silicon should have which of the following number of valence electrons?

- (a) 2
- (b) 3
- (c) 1
- (d) 5

### Very Short Question:

1. Which point defect in crystals does not alter the density of the relevant solid?
2. Which point defect in its crystal units alters the density of a solid?
3. Which point defect in its crystal units increases the density of a solid?
4. How do metallic and ionic substances differ in conducting electricity?
5. Which point defect of its crystals decreases the density of a solid?
6. What is the number of atoms in a unit cell of a face-centred cubic crystal?
7. Write a feature which will distinguish a metallic solid from an ionic solid?
8. What type of semiconductor is obtained when silicon is doped with arsenic?
9. Write a distinguishing feature of metallic solids.
10. What type of interactions hold the molecules together in a polar molecular solid?

### Short Questions:

1. Iron has a body centred cubic unit cell with a cell edge of 286.65 pm. The density of iron is  $7.87 \text{ g cm}^{-3}$ . Use this information to calculate Avogadro's number (At. mass of Fe = 56 g

mol<sup>-1</sup>).

- Silver crystallises with face-centred cubic unit cells. Each side of the unit cell has a length of 409 pm. What is the radius of an atom of silver?
- The well known mineral fluorite is chemically calcium fluoride. It is known that in one unit cell of this mineral there are 4 Ca<sup>2+</sup> ions and 8 F<sup>-</sup> ions and that Ca<sup>2+</sup> ions are arranged in a fee lattice. The F<sup>-</sup> ions fill all the tetrahedral holes in the face centred cubic lattice of Ca<sup>2+</sup> ions. The edge of the unit cell is  $5.46 \times 10^{-8}$  cm in length. The density of the solid is 3.18 g cm<sup>-3</sup>. Use this information to calculate Avogadro's number (Molar mass of CaF<sub>2</sub> = 78.08 g mol<sup>-1</sup>).
- The density of copper metal is 8.95 g cm<sup>-3</sup>. If the radius of copper atom is 127.8 pm, is the copper unit cell a simple cubic, a body-centred cubic or a face centred cubic structure?
- Silver crystallises in face-centred cubic unit cells. Each side of the unit cell has a length of 409 pm. What is the radius of silver atom?
- Silver crystallizes in face-centered cubic unit cell. Each side of this unit cell has a length of 400 pm. Calculate the radius of the silver atom. (Assume the atoms just touch each other on the diagonal across the face of the unit cell. That is each face atom is touching the four comer atoms.)
- The density of lead is 11.35 g cm<sup>-3</sup> and the metal crystallizes with fee unit cell. Estimate the radius of lead atom.  
(At. Mass of lead = 207 g mol<sup>-1</sup> and  $N_A = 6.02 \times 10^{23}$  mol<sup>-1</sup>)
- Tungsten crystallizes in body centred cubic unit cell. If the edge of the unit cell is 316.5 pm, what is the radius of tungsten atom?
- Iron has a body centred cubic unit cell with a cell dimension of 286.65 pm. The density of iron is 7.874 g cm<sup>-3</sup>. Use this information to calculate Avogadro's number. (At mass of Fe = 55.845 u)
- Copper crystallises with face centred cubic unit cell. If the radius of copper atom is 127.8 pm, calculate the density of copper metal.  
(Atomic mass of Cu = 63.55 u and Avogadro's number  $N_A = 6.02 \times 10^{23}$  mol<sup>-1</sup>)

### Long Questions:

- (a) An element has an atomic mass 93 g mol<sup>-1</sup> and density 11.5 g cm<sup>-3</sup>. If the edge length of its unit cell is 300 pm, identify the type of unit cell.  
(b) Write any two differences between amorphous solids and crystalline solids.
- (a) Calculate the number of unit cells in 8.1 g of aluminium if it crystallizes in a f.c.c. structure. (Atomic mass of Al = 27 g mol<sup>-1</sup>)  
(b) Give reasons:

- (i) In stoichiometric defects, NaCl exhibits Schottky defect and not Frenkel defect.
- (ii) Silicon on doping with Phosphorus form n-type semiconductor.
- (iii) Ferrimagnetic substances show better magnetism than antiferromagnetic substances.

### Assertion and Reason Questions:

1. In these questions, a statement of assertion followed by a statement of reason is given. Choose the correct answer out of the following choices.
  - a) Assertion and reason both are correct statements, and reason is correct explanation for assertion.
  - b) Assertion and reason both are correct statements, but reason is not correct explanation for assertion.
  - c) Assertion is correct statement, but reason is wrong statement.
  - d) Assertion is wrong statement, but reason is correct statement.

**Assertion:** The number of tetrahedral voids is double the number of octahedral voids.

**Reason:** The size of the tetrahedral voids is half of that of the octahedral void.

2. In these questions, a statement of assertion followed by a statement of reason is given. Choose the correct answer out of the following choices.
  - a) Assertion and reason both are correct statements, and reason is correct explanation for assertion.
  - b) Assertion and reason both are correct statements, but reason is not correct explanation for assertion.
  - c) Assertion is correct statement, but reason is wrong statement.
  - d) Assertion is wrong statement, but reason is correct statement.

**Assertion:** bcc and hcp has same packing efficiency

**Reason:** bcc arrangement has 2 atoms per unit cell, while fcc has 4 atoms per unit cell.

### Case Study Questions:

1. Read the passage given below and answer the following questions:

The idealized ionic solid consists of two interpenetrating lattices of oppositely-charged point charges that are held in place by a balance of coulombic force of long range. But real ions occupy space, no such "perfect" ionic solid exists in nature. Chemists usually apply the term "ionic solid" to binary compounds of the metallic elements of groups 1 - 2 with one of the halogen elements or oxygen. The most well known ionic solid is sodium chloride, also known by its geological names as rock-salt or halite. Structurally, each ion in sodium chloride is surrounded and held in tension by six neighbouring ions of opposite charge; this is known as (6,

6) coordination. The resulting crystal lattice is of a type known as simple cubic. There are many other fundamental ionic structures (not all cubic) and these are: Zinc blende structure (ZnS) : having ccp arrangement of  $S^{2-}$  and  $Zn^{2+}$  in alternate tetrahedral voids; Wurtzite structure (ZnS) having hcp arrangement of  $S^{2-}$  and  $Zn^{2+}$  in alternate tetrahedral voids; Fluorite structure ( $CaF_2$ ) having ccp arrangement of  $Ca^{2+}$  and  $F^-$  in all tetrahedral voids Antifluorite structure ( $Na_2O$ ) having ccp arrangement of  $O^{2-}$  and  $Na^+$  in all tetrahedral voids. These solids tend to be quite hard and have high melting points.

The following questions are multiple choice questions. Choose the most appropriate answer.

- (i) In NaCl crystal, each  $Cl^-$  ion is surrounded by.
- 4  $Na^+$  ions
  - 6  $Na^+$  ions
  - 1  $Na^+$  ion
  - 2  $Na^+$  ions
- (ii) In an antifluorite structure, cations occupy.
- Tetrahedral voids.
  - Centre of cube
  - Octahedral voids
  - Corners of cube.
- (iii) Anti fluorite structure is derived from fluorite structure by.
- Heating fluorite crystal lattice.
  - Subjecting fluorite structure to high pressure.
  - Interchanging the positions of positive and negative ions in the lattice.
  - None of these.
- (iv) In crystal structure of sodium chloride, the arrangement of  $Cl^-$  ion is.
- fcc
  - both fcc and bcc
  - bcc
  - None of these
- (v) Ionic solid  $BaF_2$  has which kind of structure?
- Fluorite
  - Antifluorite
  - Wurtzite
  - Rock-salt

2. Read the passage given below and answer the following questions:

In contrast to the disorders of gases and liquids, there is translational order in crystals. However, disordered or amorphous solids also exist which lack such order, they are really

highly viscous liquids. In translational order entire structure or lattice, can be generated by repeated replication of a small regular figure, termed as unit cell. The planes of any crystalline structure can be specified using Miller indices, which is also serve to identify single crystal faces. The ordered structure, or lattice, of a solid can be determined by X-ray or neutron diffraction studies, in which a beam of X-rays or neutrons is scattered from the sample to produce a diffraction pattern which can be analyzed to reveal the crystal structure of the sample. All crystal lattices can be classified into 14 Bravais lattices belonging to 7 systems. For example, the simple cubic, face-centred cubic and body-centred cubic lattices are the 3 lattices of the cubic system. Cubic and hexagonal close-packed structures have the structure of tightly packed spheres, where each sphere touches 12 neighbours, 6 in the same plane and 3 above and 3 below. These two close-packed structures differ in the placement of successive planes or layers. For the hexagonal close packing, a third layer is laid down to reproduce the first layer, so that the structure could be represented by ABABAB .... For cubic close packing, third layer is again displaced, corresponding to ABCABC.

The following questions are multiple choice questions. Choose the most appropriate answer:

- (i) In hexagonal close packing, a sphere has coordination number of.
- 4
  - 6
  - 8
  - 12
- (ii) Which of the following arrangements correctly represents hexagonal and cubic close packed structure respectively?
- ABCABC...and ABAB...
  - ABAB...and ABCABC...
  - Both have ABAB ... arrangement.
  - Both have ABCABC... Arrangement.
- (iii) The arrangement of the first two layers, one above the other, in hcp and ccp arrangements is.
- Exactly same in both cases
  - Partly same and partly different
  - Different from each other
  - Nothing definite.
- (iv) Which of the following statements is not correct?
- The amorphous solids have a random, disordered arrangement of constituents.

- b) The simple cubic, face-centred and body-centred are the three lattices of the cubic system.
  - c) The number of Bravais lattice in which a crystal can be categorized is 7.
  - d) A metal that crystallizes in hcp structure has coordination number 12.
- (v) Which of the following statements about amorphous solids is incorrect?
- a) They melt over a range of temperature.
  - b) There is no orderly arrangement of particles.
  - c) They are anisotropic.
  - d) They are rigid and incompressible.

## Answers key

### MCQ answers:

1. (b) fee
2. (c) Frenkel defect
3. (a) 12
4. (b) 2
5. (a)  $a = b = c, \alpha = \beta = \gamma \neq 90^\circ$
6. (c) F-centre
7. (c) 68%
8. (c) Pyroelectricity
9. (d) 26%
10. (b) triclinic
11. (d) 6
12. (d) 5
13. (a) 74%
14. (c) 32%
15. (b) 3

### Very Short Answers:

1. Frenkel defect.

2. Schottky defect.
3. Metal excess defect increases the density of a solid. It is due to presence of extra cations in the interstitial sites.
4. The electrical conductivity in metallic substances is due to free electrons while in ionic substances the conductivity is due to presence of ions.
5. Schottky defect.
6. The number of atoms in a unit cell of fcc-crystal is 4 atoms.
7. The electrical conductivity in metallic solid is due to free electrons while in ionic solid the conductivity is due to presence of ions.
8. n-type semiconductor.
9. Metallic solids possess high electrical and thermal conductivity due to presence of free electrons.
10. Dipole-dipole forces of attractions hold the molecules together in a polar molecular solid.

### Short Answers:

#### 1. Answer

$$a = 286.65 \text{ pm} = 286.65 \times 10^{-10},$$

$$d = 7.87 \text{ g cm}^{-3}, M = 56 \text{ g mol}^{-1}$$

$$Z = 2, N_A = ?$$

$$\text{Using formula : } d = \frac{ZM}{a^3 N_A} \text{ or } N_A = \frac{ZM}{a^3 d}$$

$$\text{or } N_A = \frac{2 \times 56}{(286.65 \times 10^{-10})^3 \times 7.87}$$

$$\text{or } N_A = \frac{112}{(2.87 \times 10^{-8})^3 \times 7.87}$$

$$\text{or } N_A = \frac{112}{23.63 \times 7.87 \times 10^{-24}}$$

$$\text{or } N_A = \frac{112}{185.97 \times 10^{-24}}$$

$$\text{or } N_A = \frac{112}{18.597} \times 10^{23}$$

$$\therefore \text{Avogadro's number } N_A = 6.022 \times 10^{23}$$

#### 2. Answer

$$\text{Given : } a = 409 \text{ pm } r = ?$$

For fcc unit cell, the formula is a

$$r = \frac{a}{2\sqrt{2}}$$

$$\text{or } r = \frac{409}{2\sqrt{2}} = \frac{409}{2 \times 1.414} = \frac{409}{2.828}$$

$$\therefore r = 144.62$$

$\therefore$  Radius of an atom of silver = 144.62 pm

### 3. Answer

Edge of the unit cell (a) =  $5.46 \times 10^{-8}$  cm

Density (P) =  $3.18 \text{ g cm}^{-3}$

According to the formula

$$N_A = \frac{Z \times M}{a^3 \times P} \quad \therefore N_A = \frac{4 \times 78.08}{(5.46 \times 10^{-8})^3 \times 3.18}$$

$$\therefore N_A = \frac{3123.2}{517.612 \times 10^{-23}} = 6.033 \times 10^{23}$$

### 4. Answer:

If copper atom were simple cubic :

$$a = 2 \times r = 2 \times 127.8 \text{ pm} = 255.6 \text{ pm}$$

$$= 255.6 \text{ pm} = 255.6 \times 10^{-10} \text{ cm}$$

$$Z = 1$$

$$\rho = \frac{Z \times M}{a^3 \times N_A} = \frac{1 \times 63.54}{(255.6 \times 10^{-10})^3 \times (6.02 \times 10^{23})}$$

$$\therefore P = 6.34 \text{ g cm}^{-3}$$

Actual density =  $8.95 \text{ g cm}^{-3}$

Hence copper atom is not simple cubic.

If copper atom were body-centred :

$$a = \frac{4r}{\sqrt{3}} = \frac{4 \times 127.8}{1.732} \text{ pm} = 295.15 \text{ pm}$$

$$Z = 2$$

$$P = \frac{Z \times M}{a^3 \times N_A} = \frac{2 \times 63.54}{(295.15 \times 10^{-10})^3 \times 6.02 \times 10^{23}}$$

$$\therefore P = 8.21 \text{ g cm}^{-3}$$

Hence, copper atom is not body centered

If copper atom were face-centered

$$a = 2\sqrt{2}r$$

$$\text{or } a = 2 \times 1.414 \times 127.8 \text{ pm}$$

$$= 361.4 \text{ pm} = 361.4 \times 10^{-10} \text{ cm}$$

$$P = \frac{Z \times M}{a^3 \times N_A} = \frac{4 \times 63.54}{(361.4 \times 10^{-10})^3 \times 6.02 \times 10^{23}}$$

$$\therefore P = 8.94 \text{ cm}^{-3}$$

Hence, copper is face-centred cubic.

### 5. Answer :

Given :  $a = 409 \text{ pm}$   $r = ?$

For fcc unit cell, the formula is

$$r = \frac{a}{2\sqrt{2}}$$

$$\text{or } r = \frac{409}{2\sqrt{2}} = \frac{409}{2 \times 1.414} = \frac{409}{2.828}$$

$$\therefore r = 144.62$$

$\therefore$  Radius of an atom of silver = 144.62 pm

### 6. Answer

Given :  $a = 400 \text{ pm}$ ,  $r = ?$

$$\text{For fcc unit cell : } r = \frac{a}{2\sqrt{2}}$$

$$\text{or } r = \frac{400}{2 \times 1.4142} = \frac{400}{2.828} \therefore r = 141.4$$

$\therefore$  Radius of the silver,  $r = 141.4 \text{ pm}$

### 7. Answer

Given :  $d = 11.35 \text{ g cm}^{-3}$

According to the formula

$$d = \frac{Z \times M}{a^3 \times N_A} \quad \text{or} \quad a^3 = \frac{Z \times M}{d \times N_A}$$

For fcc lattice,  $Z = 4$

$$a^3 = \frac{4 \times 207 \text{ g mol}^{-1}}{11.35 \text{ g cm}^{-3} \times 6.02 \times 10^{23} \text{ mol}^{-1}}$$

$$\text{or } a^3 = \frac{828}{68.327 \times 10^{23}} = 12.118 \times 10^{-23}$$

$$\text{or } a^3 = 1.212 \times 10^{-22} \text{ cm}^3$$

$$\text{or } a = \sqrt[3]{1.212 \times 10^{-22}} \text{ cm}^3$$

$$\therefore a = 4.95 \times 10^{-8} \text{ cm}$$

$$\text{For fcc unit cell, } r = \frac{a}{2\sqrt{2}}$$

$$\text{or } r = \frac{4.95 \times 10^{-8}}{2 \times 1.414} = \frac{4.95 \times 10^{-8}}{2.828}$$

$$\therefore \text{Radius, } r = 1.75 \times 10^{-8} \text{ cm} = 175 \text{ pm}$$

### 8. Answer

$$\begin{aligned} \text{For bcc, unit cell : radius, } r &= \frac{\sqrt{3}a}{4} \quad (\because a = 316.5 \text{ pm}) \\ &= \frac{\sqrt{3} \times 316.5 \text{ pm}}{4} = \frac{1.732 \times 316.5 \text{ pm}}{4} = 137 \text{ pm} \end{aligned}$$

### 9. Answer :

$$\text{Formula : } d = \frac{Z \times M}{a^3 \times N_A}$$

For bcc, lattice  $Z = 2$  [bcc = body centred cubic]

$$7.874 \text{ g cm}^{-3} = \frac{2 \times 55.845 \text{ g mol}^{-1}}{(286.65 \times 10^{-10} \text{ cm})^3 \times N_A}$$

$$N_A = \frac{2 \times 55.845 \text{ g mol}^{-1}}{(286.65 \times 10^{-10} \text{ cm})^3 \times 7.874 \text{ g cm}^{-3}}$$

$$\therefore \text{Avogadro's number, } N_A = 6.02 \times 10^{23} \text{ mol}^{-1}$$

### 10. Answer :

In fcc, lattice,  $Z = 4$  atoms [fcc = face centred cubic]

$$\text{Formula : } d = \frac{Z \times M}{a^3 \times N_A}$$

fcc lattice for copper  $a = 2\sqrt{2}r$

$$\therefore a^3 = (2\sqrt{2}r)^3$$

$$\Rightarrow a^3 = 8 \times 2\sqrt{2} (1.278 \times 10^{-8} \text{ cm})^3$$

$$\Rightarrow a^3 = 4.723 \times 10^{-23} \text{ cm}^3$$

$$d = \frac{4 \times 63.55 \text{ g mol}^{-1}}{4.723 \times 10^{-23} \text{ cm}^3 \times 6.02 \times 10^{23} \text{ mol}^{-1}}$$

$$\Rightarrow d = \frac{254.2 \text{ g mol}^{-1}}{28.43} = 8.94 \text{ g cm}^{-3}$$

## Long Answers:

### 1. Answer

(a) Given:

$$M = 93 \text{ g mol}^{-1}; \rho = 11.5 \text{ g cm}^{-3};$$

$$a = 300 \text{ pm} = 300 \times 10^{-10} \text{ cm} = 3 \times 10^{-8} \text{ cm}$$

Using formula,

$$Z = \frac{\rho \times a^3 \times N_A}{M}$$

$$= \frac{11.5 \times (3 \times 10^{-8})^3 \times 6.022 \times 10^{23}}{93}$$

$$= 2.01 \text{ (approx.)}$$

As the number of atoms present in given unit cells are coming nearly equal to 2, hence the given units cell is body centered cubic unit cell (BCC).

| Amorphous solids   | Crystalline solids   |
|--|--|
| (i) They are isotropic, i.e., they will show same value of physical perties in directions. | (i) They are anisotropic, i.e., tropic, i.e., value of physical properties will be different when measured along different directions. |
| (ii) They have short range order.  | (ii) They have long range order.   |

### 2. Answer

(a) Given:

$$\text{Mass of Al} = 8.1,$$

$$\text{Atomic mass of Al} = 27 \text{ g mol}^{-1}$$

$$\text{No. of atoms} = \eta \times 6.022 \times 10^{23}$$

$$= \frac{8.1}{27} \times 6.022 \times 10^{23}$$

$$= 0.3 \times 6.022 \times 10^{23}$$

$$= 1.8066 \times 10^{23}$$

Since one f.c.c. unit cell has 4 atoms

$$\therefore \text{No. of unit cells} = 1.8066 \times 10^{23} / 4$$

$$= 4.5 \times 10^{22} \text{ unit cells}$$

(b) (i) Schottky defect is shown by the ionic solids having very small difference in their cationic and anionic radius whereas Frenkel defect is shown by ionic solids having large difference in their cationic and anionic radius. NaCl exhibits Schottky defect because radius of both  $\text{Na}^+$  and  $\text{Cl}^-$  have very small difference.

(ii) Phosphorus is pentavalent that is it has 5 valence electrons, an extra electron results in the formation of n-type semi conductors on doping with Silicon. The conductivity is due to presence of extra electrons.

(iii) In antiferromagnetic substances the magnetic moments of domains are half aligned in one direction and remaining half in opposite direction in the presence of magnetic field so magnetic moment will be zero while in ferrimagnetic substances the magnetic moments of domains are aligned in parallel and anti-parallel directions in unequal numbers, hence shows some value of magnetic moment.



## Assertion and Reason Answers:

1. (c) Assertion is correct statement, but reason is wrong statement.

### Explanation:

The size of tetrahedral voids is smaller but not half of the octahedral voids.

2. (d) Assertion is wrong statement, but reason is correct statement.

### Explanation:

bcc and hcp have different arrangements of particles. bcc has 68% and hcp has 74% packing efficiency.

## Case Study Answers:

1. Answer :

(i) (b)  $6 \text{ Na}^+$  ions

**Explanation:**

In NaCl crystal, each  $\text{Cl}^-$  ion is surrounded by 6  $\text{Na}^+$  ions.

(ii) (a) Tetrahedral voids.

**Explanation:**

Anti-fluorite structure (NaCl) have ccp arrangement of  $\text{O}^{2-}$  and  $\text{Na}^+$  in all tetrahedral voids.

(iii) (c) Interchanging the positions of positive and negative ions in the lattice.

**Explanation:**

Antifluorite structure is derived from fluorite structure by interchanging the positions of positive and negative ions.

(iv) (a) fcc

**Explanation:**

In NaCl crystal structure,  $\text{Cl}^-$  ions form fcc.

(v) (a) Fluorite

**Explanation:**

$\text{BaF}_2$  has fluorite structure.

**2. Answer :**

(i) (d) 12

**Explanation:**

For hcp packing, the coordination number of each sphere is 12 (6 in its own layer, 3 in the upper layer and 3 in the lower layer).

(ii) (b) ABAB .... and ABCABC...

**Explanation:**

In hexagonal close packing, each sphere of third layer lies exactly above the sphere of first layer. Thus, hcp is abbreviated as ABAB .... In cubic close packing, the spheres of third layer do not come over those of first layer and spheres of fourth layer correspond with those in first layer. Thus, it is represented as AB CAB C...

(iii) (a) Exactly same in both cases.

(iv) (c) The number of Bravais lattice in which a crystal can be categorized is 7.

**Explanation:**

The number of Bravais lattice in which a crystal can be categorized is 14.

(v) (c) They are anisotropic.

**Explanation:**

They are isotropic as they show same properties in all directions.