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శ 8929803804 (MON-FRI: 9am-6pm) support@misostudy.com
MISO STUDY INDIA PVT. LTD.
2ND FLOOR 65-A, OMPRO TOWER, KALU SARAI, NEW DELHI, DELHI 110016

## Class 11 | Physics

## 01 Units and Measurement

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## 01. Physical Quantities

All the quantities which are used to describe the laws of physics are known as physical quantities.
Classification : Physical quantities can be classified on the following bases :
(A) Based on their directional properties
I. Scalars : The physical quantities which have only magnitude but no direction are called scalar quantities.
e.g. mass, density, volume, time, etc.
II. Vectors : The physical quantities which both magnitude and direction and obey laws of vector algebra are called vector quantities.
e.g. displacement, force, velocity, etc.
(B) Based on their dependency
I. Fundamental or base quantities : The quantities which do not depend upon other quantities for their complete definition are known as fundamental or base quantities.
e.g. length, mass, time, etc.
II. Derived quantities : The quantities which can be expressed in terms of the
fundamental quantities are known as derived quantities.
e.g. Speed (=distance/time), volume, acceleration, force, pressure, etc.

Example Classify the quantities displacement, mass, force, time, speed, velocity, acceleration, pressure and work under the following categories:
(a) base and scalar
(b) base and vector
(c) derived and scalar
(d) derived and vector

Solution
(a) mass, time
(b) displacement
(c) speed, pressure, work
(d) force, velocity, acceleration

## 02. Units of Physical Quantities

The chosen reference standard of measurement in multiples of which, a physical quantity is expressed is called the unit of that quantity.

## System of Units

(i) FPS or British Engineering system : In this system length, mass and time are taken as fundamental quantities and their base units are foot (ft), pound (lb) and second (s) respectively.
(ii) CGS or Gaussian system : In this system the fundamental quantities are length, mass and time and their respective units are centimeter (cm), gram (g) and second (s).
(iii) MKS system : In this system also the fundamental quantities are length, mass and time but their fundamental units are metre ( m ), kilogram ( kg ) and second (s) respectively.
(iv) International system (SI) of units : This system is modification over the MKS system and so it is also known as Rationalised MKS system. Besides the three base units of MKS system four fundamental and tow supplementary units are also included in this system.

| SI BASE QUANTITIES AND THEIR UNITS |  |  |  |
| :---: | :--- | :---: | :---: |
| S. No. | Physical quantity | Unit | Symbol |
| 1 | Length | metre | m |
| 2 | Mass | kilogram | kg |
| 3 | Time | second | s |
| 4 | Temperature | kelvin | K |
| 5 | Electric current | ampere | A |
| 6 | Luminous intensity | candela | cd |
| 7 | Amount of substance | mole | mol |

## 03. Classification of Units

The units of physical quantities can be classified as follows :
(i) Fundamental or base units

The units of fundamental quantities are called base units. In SI there are seven base units.
(ii) Derived units

The units of derived quantities or the units that can be expressed in terms of the base units are called derived units.
e.g. unit of speed $=\frac{\text { unit of distance }}{\text { unit of time }}=\frac{\text { metre }}{\text { second }}=\mathrm{m} / \mathrm{s}$

Some derived units are named in honour of great scientists.
e.g. unit of force - newton ( N ), unit of frequency - hertz ( Hz ), etc.
(iii) Supplementary units

In International System (SI) of units two supplementary units are also defined viz. radian (rad) for plane angle and steradian (sr) for solid angle.

- radian : 1 radian is the angle subtended at the centre of a circle by and arc equal in length to the radius of the circle.
- steradian : 1 steradian is the solid angle subtended at the centre of a sphere, by the surface of the sphere which is equal in area to the square of the radius of the sphere.


## (iv) Practical units

Due to the fixed sizes of SI units, some practical units are also defined for both fundamental and derived quantities. e.g. light year (ly) is a practical unit of distance (a fundamental quantity) and horse power (hp) is a practical unit of power (a derived quantity).
Practical units may or may not belong to a particular system of units but can be expressed in any system of units.
e.g. 1 mile $=1.6 \mathrm{~km}=1.6 \times 10^{3} \mathrm{~m}=1.6 \times 10^{5} \mathrm{~cm}$.

## Conversion factors

To convert a physical quantity from one set of units to the other, the required multiplication factor is called conversion factor.
Magnitude of a physical quantity $=$ numeric value ( n ) $\times$ unit ( $u$ )
While conversion from one set of units to the other the magnitude of the quantity must remain same. Therefore

$$
\mathrm{n}_{1} \mathrm{u}_{1}=\mathrm{n}_{2} \mathrm{u}_{2} \quad \text { or } \quad \mathrm{nu}=\text { constant } \quad \text { or } \quad \mathrm{n} \propto \frac{1}{\mathrm{u}}
$$

This is the numeric value of a physical quantity is inversely proportional to the base unit. $\begin{array}{ll}\text { e.g. } & 1 \mathrm{~m}=\underset{(\mathrm{CGS})}{100 \mathrm{~cm}}=\underset{(\mathrm{FPS})}{3.28 \mathrm{ft}}=39.4 \text { inch } \\ & \\ \text { (SI) }\end{array}$

Example The acceleration due to gravity is $9.8 \mathrm{~m} \mathrm{~s}^{-2}$. Given its value in $\mathrm{ft} \mathrm{s}^{-2}$
Solution $\quad$ As $1 \mathrm{~m}=3.2 \mathrm{ft}$

$$
\therefore \quad 9.8 \mathrm{~m} / \mathrm{s}^{2}=9.8 \times 3.28 \mathrm{ft} / \mathrm{s}^{2}=32.14 \mathrm{ft} / \mathrm{s}^{2} \approx 32 \mathrm{ft} / \mathrm{s}^{2}
$$

## 04. Dimensions

Dimensions of a physical quantity are the powers for exponents to which the base quantities are raised to represent that quantity.

## Dimensional formula

The dimensional formula of any physical quantity is that expression which represents how and which of the base quantities are included in that quantity.
It is written by enclosing the symbols for base quantities with appropriate powers in square brackets i.e. [ ]
e. g. Dimensional formula of mass in $\left[\mathrm{M}^{1} \mathrm{~L}^{0} \mathrm{~T}^{0}\right]$ is the dimensional formula of the force and the dimensions of force are 1 in mass, 1 in length and -2 in time

## 05. Applications of Dimensional Analysis

(i) To convert a physical quantity from one system of units to the other :

This is based on a fact that magnitude of a physical quantity remains same whatever system is used for measurement i.e. magnitude $=$ numeric value $(\mathrm{n}) \times$ unit $(u)=$ constant or $n_{1} u_{1}=n_{2} u_{2}$

So if a quantity is represented by $\left[\mathrm{M}^{a} \mathrm{~L}^{\mathrm{b}} \mathrm{T}^{\mathrm{c}}\right.$ ]
Then $n_{2}=n_{1}\left(\frac{u_{1}}{u_{2}}\right)=n_{1}\left(\frac{M_{1}}{M_{2}}\right)^{\mathrm{a}}\left(\frac{L_{1}}{L_{2}}\right)^{\mathrm{b}}\left(\frac{\mathrm{T}_{1}}{\mathrm{~T}_{2}}\right)^{\mathrm{c}}$

Here
$\mathrm{n}_{2}=$ numerical value in II system
$\mathrm{n}_{1}=$ numerical value in I system
$\mathrm{M}_{1}=$ unit of mass in I system
$\mathrm{M}_{2}=$ unit of mass in II system
$\mathrm{L}_{1}=$ unit of length in I system
$\mathrm{L}_{2}=$ unit of length in II system
$\mathrm{T}_{1}=$ unit of time in I system
$\mathrm{T}_{2}=$ unit of time in II system
Example Convert 1 newton (SI unit of force) into dyne (CGS unit of force)
Solution
The dimensional equation of force is $[F]=\left[\mathrm{M}^{1} \mathrm{~L}^{1} \mathrm{~T}^{-2}\right]$
Therefore if $n_{1}, u_{1}$, and $n_{2}, u_{2}$, corresponds to SI \& CGS units respectively, then $\mathrm{n}_{2}=\mathrm{n}\left[\frac{\mathrm{M}_{1}}{\mathrm{M}_{2}}\right]^{1}\left[\frac{\mathrm{~L}_{1}}{\mathrm{~L}_{2}}\right]^{1}\left[\frac{\mathrm{~T}_{1}}{\mathrm{~T}_{2}}\right]^{-2}=1\left[\frac{\mathrm{~kg}}{\mathrm{~g}}\right]\left[\frac{\mathrm{m}}{\mathrm{cm}}\right]\left[\frac{\mathrm{s}}{\mathrm{s}}\right]^{-2}=1 \times 1000 \times 100 \times 1=10^{5} \quad \therefore$
1 newton $=10^{5}$ dyne.
(ii) To check the dimensional correctness of a given physical relation

If in a given relation, the terms on both the sides have the same dimensions, then the relation is dimensionally correct. This is known as the principle of homogeneity of dimensions.

Example Check the accuracy of the relation $\mathrm{T}=2 \pi \sqrt{\frac{\mathrm{~L}}{\mathrm{~g}}}$ for a simple pendulum using
Solution
The dimensions of LHS $=$ the dimension of $T=\left[M^{0} L^{0} T^{1}\right]$
The dimensions of RHS $=\left(\frac{\text { dimensions of } \leq \text { ngth }}{\text { dimensions of acceleration }}\right)^{1 / 2} \quad(\because 2 \pi$ is a dimensionless constant)

$$
=\left[\frac{\mathrm{L}}{\mathrm{LT}^{-2}}\right]^{1 / 2}=\left[\mathrm{T}^{2}\right]^{1 / 2}=[\mathrm{T}]=\left[\mathrm{M}^{0} \mathrm{~L}^{0} \mathrm{~T}^{1}\right]
$$

Since the dimensions are same on both the sides, the relation is correct.
(iii) To derive relationship between different physical quantities

Using the same principle of homogeneity of dimensions new relations among physical quantities can be derived if the dependent quantities are known.

Example It is known that the time of revolution T of a satellite around the earth depends on the universal gravitational constant $G$, the mass of the earth $M$, and the radius of the circular orbit R. Obtain an expression for T using dimensional analysis.

$$
\text { We have } \quad[\mathrm{T}]=[\mathrm{G}]^{\mathrm{a}}[\mathrm{M}]^{\mathrm{b}}[\mathrm{R}]^{\mathrm{c}}
$$

Solution

$$
[\mathrm{M}]^{0}[\mathrm{~L}]^{0}[\mathrm{~T}]^{1}=[\mathrm{M}]^{-\mathrm{a}}[\mathrm{~L}]^{3 \mathrm{a}}[\mathrm{~T}]^{-2 \mathrm{a}} \times[\mathrm{M}]^{\mathrm{b}} \times[\mathrm{L}]^{\mathrm{c}}=[\mathrm{M}]^{\mathrm{b}-\mathrm{a}}[\mathrm{~L}]^{\mathrm{c}+3 \mathrm{a}}[\mathrm{~T}]^{-2 \mathrm{a}}
$$

Comparing the exponents
For $[\mathrm{T}]: 1=-2 \mathrm{a} \Rightarrow \mathrm{a}=-\frac{1}{2} \quad$ For $[\mathrm{M}]: 0=\mathrm{b}-\mathrm{a} \Rightarrow \mathrm{b}=\mathrm{a}=-\frac{1}{2}$
For $[L]: 0=c+3 a \Rightarrow c=-3 a=\frac{3}{2}$

Putting the values we get $T \propto G^{-1 / 2} M^{-1 / 2} R^{3 / 2} \Rightarrow T \propto \sqrt{\frac{R^{3}}{G M}}$
The actual expression is $\mathrm{T}=2 \pi \sqrt{\frac{\mathrm{R}^{3}}{\mathrm{GM}}}$
Dimensions of trigonometric, exponential, logarithmic function etc.
All trigonometric, exponential and logarithmic functions and their arguments are dimensionless.

NOTE Trigonometric function $\sin \theta$ and its $\operatorname{argument} \theta$ are dimensionless.

## 06. Limitations of this Method

- In Mechanics the formula for a physical quantity depending one more than three physical quantities cannot be derived. It can only be checked.
- This method can be used only if the dependency is of multiplication type. The formulae containing exponential, trigonometrical and logarithmic functions cant't be derived using this method. Formulae containing more than one term which are added or subtracted like s $=u t+\mathrm{at}^{2} / 2$ also can't be derived.
- The relation derived from this method gives no information about the dimensionless constants.
- If dimensions are given, physical quantity may not be unique as many physical quantities have the same dimensions.
- It gives no information whether a physical quantity is a scalar or a vector.


## 07. Significant Figures or Digits

The significant figure (SF) in a measurement are the figure or digits that are known with certainity plus one that is uncertain.
Significant figures in a measured value of a physical quantity tell the number of digits in which we have confidence. Larger the number of significant figure obtained in a measurement, greater is its accuracy and vice versa.

## Rules to find out the number of significant figures

I Rule : All the non-zero digits are significant e.g. 1984 has 4 SF.
II Rule : All the zeros between two non-zero digits are significant. e.g. 10806 has 5 SF
III Rule : All the zeros to the left of first non-zero digit are not significant. e.g. 00108 has 3 SF .
IV Rule : If the number is less than 1, zeros on the right of the decimal point but to the left of the first non-zero digit are not significant. e.g. 0.002308 has 4 SF.
$\mathbf{V}$ Rule : The trailing zeros (zeros to the right of the last non-zero digit) in a number with a decimal point are significant. e.g. 01.080 has 4 SF .

VI Rule : The trailing zeros in a number without a decimal point are not significant e.g. 010100 has 3 SF. But if the number comes from some actual measurement then the trailing zeros become significant. e.g. $\mathrm{m}=100 \mathrm{~kg}$ has 3 SF .
VII Rule : When the number is expressed in exponential form, the exponential term does not affect the number of S.F. For example in $\mathrm{x}=12.3=1.23 \times 10^{1}=$ $0.123 \times 10^{2}=0.0123 \times 10^{3}=123 \times 10^{-1}$ each term has 3 SF only.

## Rules for arithmetical operations with significant figures

I Rule : In addition or subtraction the number of decimal places in the result should be equal to the number of decimal places of that term in the operation which contain lesser number of decimal places. e.g. $12.587-12.5=0.087=$ 0.1 ( $\because$ second term contain lesser i.e. one decimal place)

II Rule : In multiplication or division, the number of SF in the product or quotient is same as the smallest number of SF in any of the factors. e.g. $4.0 \times 0.12=$ $0.484=0.48$

- To avoid the confusion regarding the trailing zeros of the numbers without the decimal point the best way is to report every measurement in scientific notation (in the power of 10 ). In this notation every number is expressed in the form a $\times 10 \mathrm{~b}$, where a is the base number between 1 and 10 and b is any positive or negative exponent of 10 . The base number (a) is written in decimal form with the decimal after the first digit. While counting the number of SF only base number is considered (Rule VII).
- The change in the unit of measurement of a quantity does not effect the number of SF. For example in $2.308 \mathrm{~cm}=23.08 \mathrm{~mm}=0.02308 \mathrm{~m}=23080 \mu \mathrm{~m}$ each term has 4 SF .

Example Write down the number of significant figures in the following.
(a) 165
(b) 2.05
(c) 34.000 m
(d) 0.005
(e) $0.02340 \mathrm{~N} \mathrm{~m}^{-1}$
(f) 26900
(g) 26900 kg

Solution
(a) 165
(b) 2.05
(c) 34.000 m
(d) 0.005
(e) $0.02340 \mathrm{~N} \mathrm{~m}^{-1}$
(f) 26900
(g) 26900 kg

3 SF (following rule I)
3 SF (following rule I \& II)
5 SF (following rule I \& V)
1 SF (following rules I \& IV)
4 SF (following rule I, IV \& V)
3 SF (see rule VI)
5 SF (see rule VI)

## 08. Rounding Off

To represent the result of any computation containing more than one uncertain digit, it is rounded off to appropriate number of significant figures.

## Rules for rounding off the numbers :

I Rule : If the digit to be rounded off is more than 5, then the preceding digit is increased by one. e.g. $6.87 \approx 6.9$
II Rule : If the digit to be rounded off is less than 5, than the preceding digit is unaffected and is left unchanged. e.g. $3.94 \approx 3.9$
III Rule : If the digit to be rounded off is 5 than the preceding digit is increased by one if it odd and is left unchanged if it is even. e.g. $14.35 \approx 14.4$ and $14.45 \approx 14.4$

Example The length, breadth and thickness of a metal sheet are $4.234 \mathrm{~m}, 1.005 \mathrm{~m}$ and 2.01 cm respectively. Give the area and volume of the sheet to correct number of significant figures.
Solution
length $(l)=4.234 \mathrm{~m} \quad$ breadth $(b)=1.005 \mathrm{~m}$
thickness $(\mathrm{t})=2.01 \mathrm{~cm}=2.01 \times 10^{-2} \mathrm{~m}$
Therefore area of the sheet $=2(l \times \mathrm{b}+\mathrm{b} \times \mathrm{t}+\mathrm{t} \times \mathrm{l})$

$$
=2(4.234 \times 1.005+1.005 \times 0.0201+0.0201 \times
$$

4.234) $\mathrm{m}^{2}$

$$
=2(4.3604739) \mathrm{m}^{2}=8.720978 \mathrm{~m}^{2}
$$

Since area can contain a max ${ }^{\mathrm{m}}$ of 3 SF (Rule II of article 4.2) therefore, rounding off, we get
Area $=8.72 \mathrm{~m}^{2}$
Like wise volume $=l \times \mathrm{b} \times \mathrm{t}=4.234 \times 1.005 \times 0.0201 \mathrm{~m}^{3}=0.0855289 \mathrm{~m}^{3}$
Since volume can contain 3 SF , therefore, rounding off, we get
Volume $=0.0855 \mathrm{~m}^{3}$

## 09. Order of Magnitude

Order of magnitude of a quantity is the power of 10 required to represent that quantity. This power is determined after rounding off the value of the quantity properly. For rounding off, the last digit is simply ignored if it is less than 5 and, is increased by one if it is 5 or more than 5 .

- When a number is divided by $10^{x}$ (where x is the order of the number) the result will always lie between 0.5 and 5 i.e. $0.5 \leq \mathrm{N} / 10^{\mathrm{x}}<5$

Example Order of magnitude of the following values can be determined as follows :
(a) $49=4.9 \times 10^{1} \approx 10^{1}$

Solution
Solution
Solution
Solution
Solution
$\therefore \quad$ Order of magnitude $=1$
(b) $51=5.1 \times 10^{1} \approx 10^{2}$
$\therefore$ Order of magnitude $=2$
(c) $0.049=4.9 \times 10^{-2} \approx 10^{-2}$
$\therefore$ Order of magnitude $=-2$
(d) $0.050=5.0 \times 10^{-2} \approx 10^{-1}$
$\therefore \quad$ Order of magnitude $=-1$
(e) $0.051=5.1 \times 10^{-2} \approx 10^{-1}$
$\therefore \quad$ Order of magnitude $=-1$

## - Accuracy, Precision of Instruments and Errors in Measurement Accuracy and Precision

The result of every measurement by any measuring instrument contains some uncertainty. This uncertainty is called error. Every calculated quantity which is based on measured value, also has an error. Every measurement is limited by the reliability of the measuring instrument and skill of the person making the measurement. If we repeat a particular measurement, we usually do not get precisely the same result as each result is subjected to some experimental error. This imperfection in measurement can be described in terms of accuracy and precision. The accuracy of a measurement is a measure of how close the measured value is to the true value of the quantity. Precision tells us to what resolution or limit the quantity is measured, we can illustrate the difference between accuracy and precision with help of a example. Suppose the true value of a certain length is 1.234 cm . In one experiment, using a measuring instrument of resolution 0.1 cm , the measured value is found to be 1.1 cm , while in another experiment using a measuring device of greater resolution of 0.01 m , the length is determined to be 1.53 cm . The first measurement has move accuracy (as it is closer to the true value) but less precision (as resolution is only 0.1 cm ), while the second measurement is less accurate but more precise.

## 10. Errors

The difference between the true value and the measured value of a quantity is known as the error of measurement.
Errors may arise from different sources and are usually classified as follows

## Systematic or Controllable Errors

Systematic errors are the errors whose causes are known. They can be either positive or negative. Due to the known causes these errors can be minimised. Systematic errors can further be classified into three categories
(i) Instrumental errors :- These errors are due to imperfect design or erroneous manufacture or misuse of the measuring instrument. These can be reduced by using more accurate instruments.
(ii) Environmental errors :- These are due to the changes in external environmental conditions such as temperature, pressure, humidity, dust vibrations or magnetic and electrostatic fields.
(iii) Observational errors :- These errors arise due to improper setting of the apparatus or carelessness in taking observations.

## Random Errors

These errors are due to unknown causes. Therefore they occur irregularly and are variable in magnitude and sign. Since the causes of these errors are not known precisely they can not be eliminated completely. For example, when the same person repeats the same observation in the same conditions, he may get different readings different times.
Random errors can be reduced by repeating the observation a large number of times and taking the arithmetic mean of all the observations. This mean value would be very close to the most accurate reading.

NOTE If the number of observations is made $n$ times then the random error reduces to $\left(\frac{1}{n}\right)$ times.

Gross Errors : Gross errors arise due to human carelessness and mistakes in reading the instruments or calculating and recording the measurement results.
For example :-
(i) Reading instrument without proper initial settings.
(ii) Taking the observations wrongly without taking necessary precautions.
(iii) Exhibiting mistakes in recording the observations.
(iv) Putting improper values of the observations in calculations.

These errors can be minimised by increasing the sincerity and alertness of the observer.

## 11. Representation of Errors

Errors can be expressed in the following ways
Absolute Error ( $\Delta \mathbf{a}$ ) : The difference between the true value and the individual measured value of the quantity is called the absolute error of the measurement.
Suppose a physical quantity is measured $n$ times and the measured values are $a_{1}, a_{2}, a_{3}$ $\ldots . . . . . . . . a_{n}$. The arithmetic mean $\left(a_{m}\right)$ of these values is

$$
\begin{equation*}
\mathrm{a}_{\mathrm{m}}=\frac{\mathrm{a}_{1}+\mathrm{a}_{2}+\mathrm{a}_{3}+\ldots \ldots \ldots . \mathrm{a}_{\mathrm{n}}}{\mathrm{n}}=\frac{1}{\mathrm{n}} \sum_{\mathrm{i}=1}^{\mathrm{n}} \mathrm{a}_{\mathrm{i}} \tag{i}
\end{equation*}
$$

If the true value of the quantity is not given then mean value $\left(a_{m}\right)$ can be taken as the true value. Then the absolute errors in the individual measured values are

$$
\begin{aligned}
& \Delta \mathrm{a}_{1}=\mathrm{a}_{\mathrm{m}}-\mathrm{a}_{1} \\
& \Delta \mathrm{a}_{2}=\mathrm{a}_{\mathrm{m}}-\mathrm{a}_{2} \\
& \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \\
& \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots \\
& \Delta \mathrm{a}_{\mathrm{n}}=\mathrm{a}_{\mathrm{m}}-\mathrm{a}_{\mathrm{n}}
\end{aligned}
$$

The arithmetic mean of all the absolute errors is defined as the final or mean absolute error $(\Delta \mathrm{a})_{\mathrm{m}}$ or $\Delta \mathrm{a}$ of the value of the physical quantity a

$$
\begin{equation*}
\left(\Delta a_{\mathrm{m}}\right)=\frac{\left|\Delta \mathrm{a}_{1}\right|+\left|\Delta \mathrm{a}_{2}\right|+\ldots \ldots \ldots \ldots+\left|\Delta \mathrm{a}_{\mathrm{n}}\right|}{\mathrm{n}}=\frac{1}{\mathrm{n}} \sum_{\mathrm{i}=1}^{\mathrm{n}}\left|\Delta \mathrm{a}_{\mathrm{i}}\right| \tag{ii}
\end{equation*}
$$

So if the measured value of a quantity be ' $a$ ' and the error in measurement be $\Delta \mathrm{a}$, then the true value $\left(a_{t}\right)$ can be written as

$$
\begin{equation*}
\mathrm{a}_{\mathrm{t}}=\mathrm{a} \pm \Delta \mathrm{a} \tag{iii}
\end{equation*}
$$

Relative or Fractional Error : It is defined as the ratio of the mean absolute error $\left((\Delta a)_{m}\right.$ or $\overline{\Delta a}$ ) to the true value or the mean value ( $a_{m}$ or $\bar{a}$ ) of the quantity measured.
Relative or fractional error $=\frac{\text { Mean absolute error }}{\text { Mean value }}=\frac{(\Delta \mathrm{a})_{\mathrm{m}}}{\mathrm{a}_{\mathrm{m}}}$ or $\frac{\overline{\Delta \mathrm{a}}}{\overline{\mathrm{a}}}$
When the relative error is expressed in percentage, it is known as percentage error, percentage

$$
\begin{equation*}
\text { error }=\text { relative error } \times 100 \tag{v}
\end{equation*}
$$

or percentage error $=\frac{\text { mean absolute error }}{\text { true value }} \times 100 \%=\frac{\overline{\Delta a}}{a} \times 100 \%$

## 12. Propagation of Errors in Mathematical Operations

Rule I : The maximum absolute error in the sum or difference of the two quantities is equal to the sum of the absolute errors in the individual quantities.
If $X=A+B$ or $X=A-B$ and if $\pm \Delta \mathrm{A}$ and $\pm \Delta \mathrm{B}$ represent the absolute errors in A and B respectively, then the maximum absolute error in $X=\triangle X=\triangle A+\triangle B$ and
Maximum percentage error $\quad=\frac{\Delta X}{X} \times 100$
The result will be written as $X \pm \Delta X \quad$ (in terms of absolute error)
or

$$
X \pm \frac{\Delta X}{X} \times 100 \% \text { (in terms of percentage error) }
$$

Rule II : The maximum fractional or relative error in the product or division of quantities is equal to the sum of the fractional or relative errors in the individual quantities.
If $\quad \mathrm{X}=\mathrm{A} \times \mathrm{B} \quad$ or $\quad \mathrm{X}=\mathrm{A} / \mathrm{B}$
then $\quad \frac{\Delta X}{X}= \pm\left(\frac{\Delta A}{A}+\frac{\Delta B}{B}\right)$
Rule III : The maximum fractional error in a quantity raised to a power (n) is $n$ times the fractional error in the quantity itself, i.e.
If
$X=A^{n}$
then
$\frac{\Delta X}{X}=n\left(\frac{\Delta \mathrm{~A}}{\mathrm{~A}}\right)$
If $\quad X=A^{p} B^{q} C^{r}$
then
$\frac{\Delta X}{X}=\left[p\left(\frac{\Delta A}{A}\right)+q\left(\frac{\Delta B}{B}\right)+r\left(\frac{\Delta C}{C}\right)\right]$
If $\quad X=\frac{A^{p} B^{q}}{C^{r}} \quad$ then

$$
\frac{\Delta X}{X}=\left[p\left(\frac{\Delta A}{A}\right)+q\left(\frac{\Delta B}{B}\right)+r\left(\frac{\Delta C}{C}\right)\right]
$$

## IMPORTANT POINTS

- Systematic errors are repeated consistently with the repetition of the experiment and are produced due to improper conditions or procedures that are consistent in action whereas random errors are accidental and their magnitude and sign cannot be predicated from the knowledge of the measuring system and conditions of measurement.
Systematic errors can therefore be minimised by improving experimental techniques, selecting better instruments and improving personal skills whereas random errors can be minimised by repeating the observation several times.
- Mean absolute error has the units and dimensions of the quantity itself whereas fractional or relative error is unitless and dimensionless.
- Absolute errors may be positive in certain cases and negative in other cases.

Example The initial and final temperatures of water as recorded by an observer are (40.6 $\pm 0.2)^{\circ} \mathrm{C}$ and $(78.3 \pm 0.3)^{\circ} \mathrm{C}$. Calculate the rise in temperature with proper error limits.
Solution
Given $\theta_{1}=(40.6 \pm 0.2)^{\circ} \mathrm{C}$ and $\theta_{2}=(78.3 \pm 0.3)^{\circ} \mathrm{C}$
Rise in temp. $\theta=\theta_{2}-\theta_{1}=78.3-40.6=37.7^{\circ} \mathrm{C}$.
$\Delta \theta= \pm\left(\Delta \theta_{1}+\Delta \theta_{2}\right)= \pm(0.2+0.3)= \pm 0.5^{\circ} \mathrm{C} \quad \therefore$ rise in temperature $=$ $(37.7 \pm 0.5)^{\circ} \mathrm{C}$

## 13. Least Count

The smallest value of a physical quantity which can be measured accurately with and instrument is called the least count (L. C.) of the measuring instrument.

## Least Count of Vernier Callipers

Suppose the size of one main scale division (M.S.D.) is $M$ units and that of one vernier scale division (V. S. D.) is V units. Also let the length of ' $a$ ' main scale divisions is equal to the length of ' $b$ ' vernier scale divisions.


$$
\begin{aligned}
& a M=b V \Rightarrow V=\frac{a}{b} M \\
& \therefore \quad M-V=M-\frac{a}{b} M \text { or } \quad M-V=\left(\frac{b-a}{b}\right) M
\end{aligned}
$$

The quantity ( $\mathrm{M}-\mathrm{V}$ ) is called vernier constant (V. C.) or least count (L. C.) of the vernier callipers.

$$
\text { L.C. }=\mathrm{M}-\mathrm{V}=\left(\frac{\mathrm{b}-\mathrm{a}}{\mathrm{~b}}\right) \mathrm{M}
$$

## Least Count of screw gauge or spherometer



$$
\text { Least Count }=\frac{\text { Pitch }}{\text { Total number of divisions on the circul scale }}
$$

where pitch is defined as the distance moved by the screw head when the circular scale is given one complete rotation. i.e.

$$
\text { Pitch }=\frac{\text { Distance moved by the screw on the linear scale }}{\text { No. of full rotations given }}
$$

NOTE With the decrease in the least count of the measuring instrument, the accuracy of the measurement increases and the error in the measurement decreases.

## Example

One cm on the main scale of vernier callipers is divided into ten equal parts. If 20 divisions of vernier scale coincide with 8 small divisions of the main scale. What will be the least count of callipers ?

Solution

$$
20 \text { div. of vernier scale }=8 \text { div. of main scale } \Rightarrow 1 \mathrm{~V} . \mathrm{S} . \mathrm{D} .=\left(\frac{8}{20}\right) \mathrm{M} . \mathrm{S} . \mathrm{D} .=
$$ $\left(\frac{2}{5}\right)$ M. S. D.

Least count $=1$ M. S. D. -1 V. S. D. $=1$ M. S. D. $-\left(\frac{2}{5}\right)$ M. S. D. $=$

$$
\begin{aligned}
& \text { ( } 1-\frac{2}{5} \text { ) M. S. D. } \\
& =\frac{3}{5} \mathrm{M} . \mathrm{S} . \text { D. }=\frac{3}{5} \times 0.1 \mathrm{~cm}=0.06 \mathrm{~cm} \\
& \left(\because 1 \text { M. S. D. }=\frac{1}{10} \mathrm{~cm}=0.1 \mathrm{~cm}\right)
\end{aligned}
$$

NOTE The final absolute error in this type of questions is taken to be equal to the least count of the measuring instrument.

## NEET <br> Exercise (1)

1. A physical quantity of the dimension of length that can be formed out of $c, G$ and $\frac{e^{2}}{4 \pi \varepsilon_{0}}$ is [c is velocity of light, $G$ is universal constant of gravitation and $e$ is charge]
(a) $\frac{1}{c^{2}}\left[G \frac{e^{2}}{4 \pi \varepsilon_{0}}\right]^{1 / 2}$
(c) $\frac{1}{c^{2}}\left[\frac{e^{2}}{G 4 \pi \varepsilon_{0}}\right]^{1 / 2}$
(b) $c^{2}\left[G \frac{e^{2}}{4 \pi \varepsilon_{0}}\right]^{1 / 2}$
(d) $\frac{1}{c} G \frac{e^{2}}{4 \pi \varepsilon_{0}}$
2. If dimensions of critical velocity $\mathrm{v}_{c}$ of a liquid flowing through a tube are expressed as [ $\eta^{x} \rho^{y}$ $r^{2}$ ], where $\eta, \rho$ and $r$ are the coefficient of viscosity of liquid, density of liquid and radius of the tube respectively, then the values of $x, y$ and $z$ are given by
(a) $1,-1,-1$
(b) $-1,-1,1$
(c) $-1,-1,-1$
(d) $1,1,1$
3. In an experiment, four quantities $a, b, c$ and $d$ are measured with percentage error $1 \%, 2 \%, 3 \%$ and $4 \%$ respectively. Quantity $P$ is calculated $P=\frac{a^{3} b^{2}}{c d} \%$. Error in $P$ is
(a) $14 \%$
(b) $10 \%$
(c) $7 \%$
(d) $4 \%$
4. If energy ( $E$ ), velocity (v) and time ( $T$ ) are chosen as the fundamental quantities, the dimensional formula of surface tension will be
(a) $\left[\mathrm{Ev}^{-2} \mathrm{~T}^{-1}\right]$
(b) $\left[\mathrm{Ev}^{-1} \mathrm{~T}^{-2}\right]$
(c) $\left[\mathrm{Ev}^{-2} \mathrm{~T}^{-2}\right]$
(d) $\left[\mathrm{E}^{-2} \mathrm{v}^{-1} \mathrm{~T}^{-3}\right]$
5. Dimensions of resistance in an electrical circuit, in terms of dimension of mass $M$, of length $L$, of time $T$ and of current $I$, would be
(a) $\left[\mathrm{ML}^{2} \mathrm{~T}^{-3} \mathrm{I}^{-1}\right]$
(b) $\left[\mathrm{ML}^{2} \mathrm{~T}^{-2}\right]$
(c) $\left[\mathrm{ML}^{2} \mathrm{~T}^{-1} \mathrm{I}^{-1}\right]$
(d) $\left[\mathrm{ML}^{2} \mathrm{~T}^{-3} \mathrm{I}^{-2}\right]$
6. Which two of following five physical parameters have the same dimensions?
(i) Energy density
(ii) Refractive index
(iii) Dielectric constant
(iv) Young's modulus
(v) Magnetic field
(a) (ii) and (iv)
(b) (iii) and (v)
(c) (i) and (iv)
(d) (i) and (v)
7. In a vernier callipers $N$ divisions of vernier scale coincide with $N-1$ divisions of main scale (in which length of one division is 1 mm ). The least count of the instrument should be
(a) $N$
(b) $N-1$
(c) $\frac{1}{10 N}$
(d) $\frac{1}{(N-1)}$
8. The dimensional formula for permeability of free space, $\mu_{0}$ is
(a) $\left[\mathrm{MLT}^{-2} \mathrm{~A}^{-2}\right]$
(b) $\left[\mathrm{ML}^{-1} \mathrm{~T}^{2} \mathrm{~A}^{-2}\right]$
(c) $\left[\mathrm{ML}^{-1} \mathrm{~T}^{-2} \mathrm{~A}^{2}\right]$
(d) $\left[\mathrm{MLT}^{-2} \mathrm{~A}^{-1}\right]$
9. If $p$ represents radiation pressure, $c$ represents speed of light and $S$ represent radiation energy striking unit area per sec. The non-zero integers $\mathrm{x}, \mathrm{y}, \mathrm{z}$ such that $p^{\mathrm{x}} S^{\mathrm{y}} c^{\mathrm{z}}$ is dimensionless are
(a) $\mathrm{x}=1, \mathrm{y}=1, \mathrm{z}=1$
(b) $\mathrm{x}=-1, \mathrm{y}=1, \mathrm{z}=1$
(c) $\mathrm{x}=1, \mathrm{y}=-1, \mathrm{z}=1$
(d) $\mathrm{x}=1, \mathrm{y}=1, \mathrm{z}=-1$
10. A certain body weighs 22.42 g and has a measured volume of 4.7 cc . The possible error in the measurement of mass and volume are 0.01 g and 0.1 cc . Then, maximum error in the density will be
(a) $22 \%$
(b) $2 \%$
(c) $0.2 \%$
(d) $0.02 \%$

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## Class 11 | Chemistry

## 02 Mole Concept



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## 01. Classification of matter

Chemistry deals with the composition, structure and properties of matter. These aspects can be best described and understood in terms of basic constituents of matter: atoms and molecules. That is why chemistry is called the science of atoms and molecules.


## Matter

The thing which occupy space and have mass, which can be felt by our five sense is called as matter. Matter is further classified into two categories :
a. Physical classification
b. Chemical classification

## 02. Prefixed Used With Units

The S.I. system recommends the multiples such as $10^{3}, 10^{6}, 10^{9}$ etc. and fraction such as $10^{-3}, 10^{-6}, 10^{-9}$ etc. i.e. the powers are the multiples of 3 . These are indicated by special prefixes. These along with some other fractions or multiples in common use, along with their prefixes are given below in Table and illustrated for length (m).
※TABLE : SOME COMMONLY USED PREFIXES WITH THE BASE UNIT

| Prefix | Symbol | Multiplication Factor | Example |
| :--- | :---: | :---: | :--- |
| deci | $\mathbf{d}$ | $10^{-1}$ | 1 decimetre $(\mathrm{dm})=10^{-1} \mathrm{~m}$ |
| centi | $\mathbf{c}$ | $10^{-2}$ | 1 centimetre $(\mathrm{cm})=10^{-2} \mathrm{~m}$ |
| milli | $\mathbf{m}$ | $10^{-3}$ | 1 millimetre $(\mathrm{mm})=10^{-3} \mathrm{~m}$ |
| micro | $\mu$ | $10^{-6}$ | 1 micrometre $(\mu \mathrm{m})=10^{-6} \mathrm{~m}$ |
| nano | $\mathbf{n}$ | $10^{-9}$ | 1 nanometre $(\mathrm{nm})=10^{-9} \mathrm{~m}$ |
| pico | $\mathbf{p}$ | $10^{-12}$ | 1 picometre $(\mathrm{pm})=10^{-12} \mathrm{~m}$ |
| femto | $\mathbf{f}$ | $10^{-15}$ | 1 femtometre $(\mathrm{fm})=10^{-15} \mathrm{~m}$ |
| atto | $\mathbf{a}$ | $10^{-18}$ | 1 attometre $(\mathrm{am})=10^{-18} \mathrm{~m}$ |
| deka | $\mathbf{d a}$ | $10^{1}$ | 1 dekametre $(\mathrm{dam})=10^{1} \mathrm{~m}$ |
| hecto | $\mathbf{h}$ | $10^{2}$ | 1 hectometre $(\mathrm{hm})=10^{2} \mathrm{~m}$ |
| kilo | $\mathbf{k}$ | $10^{3}$ | 1 kilometre $(\mathrm{km})=10^{3} \mathrm{~m}$ |
| mega | $\mathbf{M}$ | $10^{6}$ | 1 megametre $(\mathrm{Mm})=10^{6} \mathrm{~m}$ |
| giga | $\mathbf{G}$ | $10^{9}$ | 1 gigametre $(\mathrm{Gm})=10^{9} \mathrm{~m}$ |
| tera | $\mathbf{T}$ | $10^{12}$ | 1 teremetre $(\mathrm{Tm})=10^{12} \mathrm{~m}$ |
| peta | $\mathbf{P}$ | $10^{15}$ | 1 petametre $(\mathrm{Pm})=10^{15} \mathrm{~m}$ |
| exa | $\mathbf{E}$ | $10^{18}$ | 1 exametre $(\mathrm{Em})=10^{18} \mathrm{~m}$ |

As volume is very often expressed in litres, it is important to note that the equivalence in S.I. units for volume is as under: 1 litre $(1 \mathrm{~L})=1 \mathrm{dm}^{3}=1000 \mathrm{~cm}^{3}$ and 1 millilitre $(1 \mathrm{ml})=1 \mathrm{~cm}^{3}=1 \mathrm{cc}$

Example Convert 2 atm into cm of Hg .
Solution $\quad 2 \mathrm{~atm}=2 \times 76 \mathrm{~cm}$ of $\mathrm{Hg}=152 \mathrm{~cm}$ of $\mathrm{Hg} \quad\{1$ atmosphere $=76 \mathrm{~cm}$ of Hg

## 03. Different types of masses

## One mole

Avogadro's Number $\left(\mathrm{N}_{\mathrm{A}}\right)=6.023 \times 10^{23}$. It is the number of atoms present in exactly 12 g of $\left(\mathrm{C}^{12}\right)$ isotope.

## Atomic Weight (A)

Atomic weight is the relative weight of one atom of an element with respect to a standard weight.
$A=\frac{\text { Weight of one atom of an element }}{\frac{1}{12} \text { th part by weight of an atom of }\left(\mathrm{C}^{12}\right) \text { isotope }}$
amu (atomic mass unit)
$1 \mathrm{amu}=\frac{1}{12}$ th part by weight of an atom of $\left(\mathrm{C}^{12}\right)$ isotope

$$
=\frac{1}{N_{A}} g=1.66 \times 10^{-24} g
$$

Atomic weight $(A) \times \mathrm{amu}=$ Absolute atomic weight.

NOTE Atomic weight is a relative weight that indicates the relative heaviness oof one atom of an element with respect to amu weight. Atomic weight has no unit because it is the ratio of weights. One mole of an amu $=1.00 \mathrm{~g}$.

## Change of Scale for Atomic Weight

If an amu is defined differently as $(1 / \mathrm{x})$ th part by weight of an atom of $\left(\mathrm{C}^{12}\right)$ isotope rather $(1 / 12)$ th part then the atomic weight $\left(A^{\prime}\right)$ can be derived as:
$A^{\prime}=A\left(\frac{x}{12}\right)$
Where, $A=$ conventional atomic weight

## Molecular Weight (MW)

Like atomic weight, it is the relative weight of a molecule or a compound with respect to amu weight.
Molecular Weight $=\frac{\text { Weight of one molecule of a compound }}{\frac{1}{12} \text { th part by weight of an atom of } \mathrm{C}^{12} \text { isotope }}$
Gram Atomic, Gram Molecular Weight ( $M$ )
It is the weight of 1.0 mole (Avogadro's numbers) of atoms, molecules or ions in gram unit. $M=A$ amu $\times$ Avogadro number $=A$ gram Hence, gram molecular weight $(M)$ is numerically equal to the atomic weight or (molecular weight) in gram unit because

$$
1.0 \text { mole of amu is } 1.0 \mathrm{~g} \text {. }
$$

Example A piece of Cu contain $6.022 \times 10^{24}$ atoms. How many mole of Cu atoms does it contain?

Solution

$$
\text { No. of mole }=\frac{6.022 \times 10^{24}}{\mathrm{~N}_{\mathrm{A}}}=\frac{6.022 \times 10^{24}}{6.022 \times 10^{23}}=10 \mathrm{~mole}
$$

## 04. Law of conservation of mass (Lavoisier-1774):

In any physical or chemical change, mass can neither be created nor be destroyed.

## It means:

Total mass of the reactants = total mass of the products.
This relationship holds good when reactants are completely converted into products.
In case the reacting material are not completely consumed the relationship will beTotal mass of the reactants $=$ Total mass of the products + mass of unreacted reactants.

Example $\quad 1.7$ gram of silver nitrate dissolved in 100 gram of water is taken. 0.585 gram of sodium chloride dissolved in 100 gram of water is added it and chemical reaction occurs. 1.435 gm of AgCl and 0.85 gm NaNO 3 are formed. Show that these results illustrate the law of conservation of mass.

Solution
Total masses before chemical change
$=$ mass of $\mathrm{AgNO}_{3}+$ mass of $\mathrm{NaCl}+$ mass of water
$=1.70+0.585+200 \mathrm{~g}$
$=202.285 \mathrm{~g}$
Total masses after the chemical reaction
$=$ mass of $\mathrm{AgCl}+$ mass of $\mathrm{AgNO}_{3}+$ mass of water
$=1.435+0.85+200$
$=202.258 \mathrm{~g}$
Then, in this chemical change
Total masses of reactants $=$ Total masses of product

## 05. Law of constant composition : [proust 1799]

A chemical compound always contains the same element combined together in fixed proportion by mass.

Example $\quad 1.08$ gram of Cu wire was allowed to react with nitric acid. The resulting solution was dried and ignited when 1.35 gram of copper oxide was obtained. In another experiment 1.15 gram of copper oxide was heated in presence of Hydrogen yielding 0.92 gram of copper. Show that the above data are in accordance with law of constant composition?
Solution

$$
\begin{aligned}
\% \text { of "Cu" in copper oxide in } 1^{\text {st }} \text { case } & =\frac{1.08}{1.35} \times 100 \\
& =80 \% \\
\% \text { of oxygen } & =20 \% \\
\% \text { of "Cu" in copper oxide in } 2^{\text {nd }} \text { case } & =\frac{0.92}{1.15} \times 100 \\
& =80 \% \\
\% \text { of oxygen } & =20 \%
\end{aligned}
$$

## 06. Law of multiple proportion : [Dalton 1806]

When two elements combine to form two or more compounds, the different masses of one element which combine with a fixed mass of the other element, bear a simple ratio to one another.

Example Two compounds each containing only tin and oxygen had the following composition.

|  | Mass \% of Tin | Mass \% of oxygen |
| :--- | :--- | :--- |
| Compound A | 78.77 | 21.23 |
| Compound B | 88.12 | 11.88 |

Show that these data illustrate the law of multiple proportion?
Solution In compound A
21.23 parts of oxygen combine with 78.77 parts of tin.

1 part of oxygen combine with $\frac{78.77}{21.23}=3.7$ parts of Sn .

## In compound $B$

11.88 parts of oxygen combine with 88.12 parts of tin.

1 part of oxygen combine with $\frac{88.12}{11.88}=7.4$ parts of tin.
Thus the mass of Tin in compound A and B which combine with a fixed mass of oxygen are in the ratio $3.7: 7.4$ or $1: 2$. This is a simple ratio. Hence the data illustrate the law of multiple proportion.

## 07. Law of reciprocal proportion : [Richter 1794]

When two different elements combine with the same mass of a third element, the ratio on which the do so will be same or simple multiple if both directly combined with each other.

Example The \% composition of $\mathrm{NH}_{3}, \mathrm{H}_{2} \mathrm{O}$ and $\mathrm{N}_{2} \mathrm{O}_{3}$ is as given below:
$\mathrm{NH}_{3} \rightarrow 82.35 \% \mathrm{~N}$ and 17.65 H .
$\mathrm{H}_{2} \mathrm{O} \rightarrow 88.9 \% \mathrm{O}$ and 11.1 H
$\mathrm{N}_{2} \mathrm{O}_{3} \rightarrow 63.15 \% \mathrm{O}$ and $36.85 \% \mathrm{~N}$
On the basis of above data prove the law of reciprocal proportion?
Solution $\quad 63.15$ parts of oxygen combine with 36.85 parts of "N"
Therefore, 88.9 part of oxygen combine with $\left(\frac{36.85}{63.15} \times 88.9\right)=51.87$ part of
"Nitrogen"
Therefore ratio is $\frac{11.1}{11.1}: \frac{51.87}{11.1}=1: 4.67$
Now compare with the ratio of Nitrogen and Hydrogen in $\mathrm{NH}_{3}$

$$
\frac{17.65}{17.65}: \frac{82.35}{17.65}=1: 4.67
$$

Hence the Law of reciprocal proportion is verified

## 08. Gay- Lussac's law of gaseous volumes [Gay-Lussac-1808]

When gases combined or produced in a chemical reaction, they do so in a simple ratio by volume provided all the gases are at same temperature and pressure.

## 09. Limiting Reagent

It is the reagent that is consumed completely during a chemical reaction. If the supplied mass ratio of reactants are not stoichiometric ratio, one of the reagent is consumed completely leaving parts of others unreacted. One that is consumed completely is known as limiting reagent.
'Limiting reagent determine the amount of product in a given chemical reaction'
Example If 20 gm of $\mathrm{CaCO}_{3}$ is treated with 20 gm of HCl , how many grams of $\mathrm{CO}_{2}$ can be generated according to following reaction?
$\mathrm{CaCo}_{3}(\mathrm{~g})+2 \mathrm{HCl}(\mathrm{aq}) \rightarrow \mathrm{CaCl}_{2}(\mathrm{aq})+\mathrm{H}_{2} \mathrm{O}(\ell)+\mathrm{CO}_{2}(\mathrm{~g})$
Solution
$\mathrm{CaCO}_{3}+2 \mathrm{HCl} \rightarrow \mathrm{CaCl}_{2}+\mathrm{H}_{2} \mathrm{O}+\mathrm{CO}_{2}$
1 mole $\mathrm{CaCO}_{3} \rightarrow 2$ mole HCl
$\therefore 100 \mathrm{~g} \mathrm{CaCO}_{3}$ combine $\rightarrow 2 \times 36.5 \mathrm{~g} \mathrm{HCl}$
$\therefore 20 \mathrm{~g} \rightarrow \frac{2 \times 36.5 \times 20}{100}=14.6 \mathrm{HCl}$
$\mathrm{CaCO}_{3}$ completely consumes in the reaction and HCl is in excess.
Therefore,
$\mathrm{CaCO}_{3} \rightarrow$ Limiting reagent
$\mathrm{HCl} \rightarrow$ Excess reagent
Now
$100 \mathrm{~g} \mathrm{CaCO}_{3}$ given 1 mole $\mathrm{CO}_{2}\left(44 \mathrm{~g} \mathrm{CO}_{2}\right)$
$20 \mathrm{~g} \mathrm{CaCO}_{3}$ will give $\frac{20 \times 40}{100}=8 \mathrm{~g} \mathrm{CO}_{2}$

## 10. Percentage yield

In general, when a reaction is carried out on the laboratory we do not obtain the theoretical amount of product. The amount of product that is actually obtained is called the actual yield. Knowing the actual yield and theoretical yield, the \% yield can be calculated by the following formula-

$$
\text { Percentage yield }=\frac{\text { Actual yield }}{\text { Theoritical yield }} \times 100 \%
$$

Example $\quad$ For the reaction
$\mathrm{CaO}+2 \mathrm{HCl} \rightarrow \mathrm{CaCl}_{2}+\mathrm{H}_{2} \mathrm{O}$
1.12 gram of CaO is reacted with excess of hydrochloric acid and 1.85 gm $\mathrm{CaCl}_{2}$ is formed. What is the $\%$ yield of the reaction?
Solution 1 mole CaO gives 1 mole $\mathrm{CaCl}_{2}$ 56 g CaO gives $111 \mathrm{~g} \mathrm{CaCl}_{2}$ 1.12 g CaO will give $\frac{111 \times 1.12}{56} \mathrm{~g} \mathrm{CaCl}_{2}=2.22 \mathrm{~g} \mathrm{CaCl}_{2}$

Now

$$
\% \text { yield }=\frac{\text { Actual yield }}{\text { Theoretical yield }} \times 100
$$

Actual yield $=1.85 \mathrm{gm}$
Theoretical yield $=2.22 \mathrm{gm}$
$\%$ yield $=\frac{1.85}{2.22} \times 100=83.33 \%$

## 11. Percentage Purity

Depending upon the mass of the product, the equivalent amount of reactant present can be determined with the help of given chemical equation. Knowing the actual amount of the reactant taken and the amount calculated with the help of a chemical equation, the purity van be determined, as
Percentage purity $=\left[\frac{\text { Amount of reactant calculated from the chemical equation }}{\text { Actual amount of reactant taken }}\right] \times 100 \%$

Example $\quad$ Calculate the amount of $(\mathrm{CaO})$ in kg that can be produced by heating 200 kg lime stone that is $90 \%$ pure $\mathrm{CaCO}_{3}$.

## Solution

$\mathrm{CaCO}_{3} \rightarrow \mathrm{CaO}+\mathrm{CO}_{2}$
1 mole $\mathrm{CaCO}_{3}$ gives 1 mole CaO
$100 \mathrm{~g} \mathrm{CaCO}_{3}$ gives 56 g CaO

$$
\begin{aligned}
\left(\frac{200 \times 90 \times 1000}{100}\right) \mathrm{g} \mathrm{CaCO}_{3} \text { gives } & =\frac{180000 \times 56}{100} \mathrm{~g} \mathrm{CaO} \\
& =\frac{1800 \times 56}{100} \mathrm{~g} \mathrm{CaO} \\
& =\frac{1008}{10} \mathrm{~g} \mathrm{CaO} \\
& =100.8 \mathrm{~g} \mathrm{CaO}
\end{aligned}
$$

## 12. Types of Average masses

## Average Atomic Mass

Average atomic mass $=$
Let a sample contains $n_{1}$ mole of atomic mass $\mathrm{M}_{1}$ and $\mathrm{n}_{2}$ mole of atoms with atomic mass $\mathrm{M}_{2}$ then

$$
M_{a v}=\frac{n_{1} M_{1}+n_{2} M_{2}}{n_{1}+n_{2}}
$$

## Average Molecular Mass

Average molecular mass $=$
Let a sample contains $n_{1}$ mole of molecules with molecular mass $M_{1}$ and $n_{2}$ mole of molecules with molecular mass $\mathrm{M}_{2}$, then

$$
\mathrm{M}=\frac{\mathrm{n}_{1} \mathrm{M}_{1}+\mathrm{n}_{2} \mathrm{M}_{2}}{\mathrm{n}_{1}+\mathrm{n}_{2}}
$$

Example Find the average atomic mass of a mixture containing $25 \%$ by mole $\mathrm{Cl}^{37}$ and $75 \%$ by mole $\mathrm{Cl}^{35}$ ?
Solution

$$
\mathrm{n}_{1}=25 \mathrm{n}_{2}=75 \mathrm{M}_{1}=37 \quad \mathrm{M}_{2}=35
$$

$$
M_{a v}=\frac{25 \times 37+75 \times 35}{25+75}=35.5
$$

## 13. Empirical \& molecular formula

The empirical formula of a compound is a chemical formula showing the relative number of atoms in the simplest ratio. An empirical formula represents the simplest. whole number ratio of various atoms present in a compound.
The molecular formula gives the actual number of atoms of each element in a molecule. The molecular formula shows the exact number of different types of atoms present in a molecule of a compound. The molecular formula is an integral multiple of the empirical formula.

$$
\text { i.e. molecular formula=empirical formula } \times \mathrm{n} \quad \text { where } \mathrm{n}=\frac{\text { molecular formula mass }}{\text { empirical formula mass }}
$$

Example An organic substance containing carbon, hydrogen and oxygen gave the following percentage composition.
$\mathrm{C}=40,684 \%$; $\mathrm{H}=5,085 \%$; and $\mathrm{O}=54,228 \%$
The molecular weight of the compound is 118 g . Calculate the molecular formula of the compound.
Solution Step-1 : To calculate the empirical formula of the compound.

| Element | Sym <br> bol <br> percent <br> age of <br> element | At. <br> mass of <br> element | Relative no. of <br> atoms= <br> Percentage | Simplest <br> atomic <br> ratio | Simplest <br> whole no. <br> atomic ratio |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Carbon | C | 40.678 | 12 | $\frac{40.678}{12}=3.390$ | $\frac{3.390}{3.389}=1$ | 2 |
| Hydrogen | H | 5.085 | 1 | $\frac{5.085}{1}=5.085$ | $\frac{5.085}{3.389}=1.5$ | 3 |
| Oxygen | O | 54.228 | 16 | $\frac{54.228}{16}=3.389$ | $\frac{3.389}{3.389}=1$ | 2 |

$\therefore$ Empirical Formula is $\mathrm{C}_{2} \mathrm{H}_{3} \mathrm{O}_{2}$
Step-2 : To calculate the empirical formula mass. The empirical formula of the compound is $\mathrm{C}_{2} \mathrm{H}_{3} \mathrm{O}_{2}$
Empirical formula mass $=(2 \times 12)+(3 \times 1)+(2 \times 16)=59$.
Step-3 : To calculate the value of ' $n$ '

$$
\mathrm{n}=\frac{\text { Molecular mass }}{\text { Empirical formula mass }}=\frac{118}{59}=2
$$

Step-4 : To calculate the molecular formula of the salt.
Molecular formula $=\mathrm{n}=($ Empirical formula $)=2 \times \mathrm{C}_{2} \mathrm{H}_{3} \mathrm{O}_{2}=\mathrm{C}_{4} \mathrm{H}_{6} \mathrm{O}_{4}$
Thus the molecular formula is $\mathrm{C}_{4} \mathrm{H}_{6} \mathrm{O}_{4}$

## 14. Vapour Density

Some times in numericals molecular mass of volatile substance is not given, instead vapour density is given. Vapour density van be defined as

$$
\text { V.D. }=\frac{\text { Density of gas at a given } \mathrm{T} \text { and } \mathrm{P}}{\text { Density of } \mathrm{H}_{2} \text { at same } \mathrm{T} \text { and } \mathrm{P}}
$$

or, V.D. $=\frac{M_{g a s}}{2}$

$$
\mathrm{M}_{\mathrm{gas}}=2 \times \mathrm{V} . \mathrm{D} .
$$

## 15. Eudiometry - Gas Analysis

The study of gaseous reactions is done in a eudiometer tube with the help of Gay-Lussac's law and Avogadro's law. Eudiometer tube is a closed graduated tube open at one end. The other end is a closed one which is provided with platinum terminals for passing electricity for electric spark, through the known volume of mixture of gases and known volume of oxygen gas. Volume of $\mathrm{CO}_{2}$ formed is determined by absorbing in KOH solution, $\mathrm{O}_{2}$ is determined by dissolving unreacted $\mathrm{O}_{2}$ in alkaline pyrogallol and water vapours formed are determined by nothing contraction in volume caused due to cooling.

## 16. Avogadro's Law

In 1812, Amadeo Avogadro stated that samples of different gases which contain the same number of molecules (any complexity, size, shape) occupy the same volume at the same temperature and pressure.
For ideal gas at constant Temperature \& Pressure, pressure is directely proportional to no. of moles

## 17. Some Absorbents of Gases

The absorbent which is used for specific gas is listed below

| Absorbent | Gas or gases absorbed |
| :--- | :--- |
| Turpentine oil | $\mathrm{O}_{3}$ |
| Alkaline pyrogallol | $\mathrm{O}_{2}$ |
| Ferrous sulphate solution | NO |
| Heated magnesium | $\mathrm{N}_{2}$ |
| Heated palladium | $\mathrm{H}_{2}$ |
| Ammonical coprous chloride | $\mathrm{O}_{2}, \mathrm{CO}, \mathrm{C}_{2} \mathrm{H}_{2}$ or $\mathrm{CH} \equiv \mathrm{CH}$ |
| Copper sulphate solution | $\mathrm{H}_{2} \mathrm{~S}, \mathrm{PH}, \mathrm{AsH}_{3}$ |
| Conc. $\mathrm{H}_{2} \mathrm{SO}_{4}$ | $\mathrm{H}_{2} \mathrm{O}$ I.e., moisture, $\mathrm{NH}_{3}$. |
| NaOH or KOH solution | $\mathrm{CO}_{2}, \mathrm{NO}_{2}, \mathrm{SO}_{2}, \mathrm{X}$, all acidic oxides |

## 18. Volume Expansion and Contraction In The Eudiometer Tube

$\mathrm{aA}(\mathrm{g})+\mathrm{bB}(\mathrm{g}) \quad \mathrm{cC}(\mathrm{g})+\mathrm{dD}(\mathrm{g})$
$\triangle_{\mathrm{n}_{\mathrm{g}}}=\mathrm{No}$. of gaseous products -No . of gaseous reactants $=(\mathrm{c}+\mathrm{d})-(\mathrm{a}+\mathrm{b})$
(a) If $\triangle \mathrm{n}_{\mathrm{g}}>0$, then expansion will occur
(b) If $\triangle \mathrm{n}_{\mathrm{g}}=0$, No contraction/expansion (volume remains constant)
(c) If $\triangle \mathrm{n}_{\mathrm{g}}<0$, then contraction will occur

## 19. Assumptions

(i) All gases are assumed to be ideal.
(ii) Nitrogen gas formed during reaction will not react with any other gas.
(iii) The volume of solids and liquids are negligible in comparision to the volume of gas.

## 20. General Reactions for Combustion of Organic Compounds

(i) When an organic compound is hydrocarbon :

$$
\mathrm{C}_{\mathrm{x}} \mathrm{H}_{\mathrm{Y}}+\left(\mathrm{x}+\frac{\mathrm{y}}{4}\right) \mathrm{O}_{2} \rightarrow \mathrm{xCO}_{2}+\frac{\mathrm{y}}{2} \mathrm{H}_{2} \mathrm{O}
$$

(ii) When an organic compound contain carbon, hydrogen and oxygen :

$$
\mathrm{C}_{\mathrm{x}} \mathrm{H}_{\mathrm{y}} \mathrm{O}_{z}+\left(\mathrm{x}+\frac{\mathrm{y}}{4}-\frac{z}{2}\right) \mathrm{O}_{2} \rightarrow \mathrm{xCO}_{2}+\frac{\mathrm{y}}{2} \mathrm{H}_{2} \mathrm{O}
$$

(iii) When an organic compound contain carbon, hydrogen and nitrogen :

$$
\mathrm{C}_{\mathrm{x}} \mathrm{H}_{\mathrm{y}} \mathrm{~N}_{\mathrm{z}}+\left(\mathrm{x}+\frac{\mathrm{y}}{4}\right) \mathrm{O}_{2} \rightarrow \mathrm{xCO}_{2}+\frac{\mathrm{y}}{2} \mathrm{H}_{2} \mathrm{O}+\frac{\mathrm{z}}{2} \mathrm{~N}_{2}
$$

Example $\quad 10 \mathrm{ml}$ of a mixture of $\mathrm{CO}, \mathrm{CH}_{4}$ and $\mathrm{N}_{2}$, exploded with excess of oxygen, gave a contraction of 6.5 ml . There was a further contraction of 7 ml . when the residual gas was treated with KOH . What is the composition of the original mixture?
Solution

$$
\begin{array}{lc}
\mathrm{CH}_{4}+2 \mathrm{O}_{2} \rightarrow \mathrm{CO}_{2}+2 \mathrm{H}_{2} \mathrm{O} \\
x \mathrm{ml} & \frac{x}{2} \mathrm{ml}
\end{array} \quad x \mathrm{ml},
$$

$$
\text { volume of oxygen used }=2 y+\frac{x}{2} \mathrm{ml}
$$

Total volume of all gases before combustion $=10+2 y+x / 2$

## 21. Percentage

Concentration of solution is the amount of solute dissolved in a known amount of the solvent or solution. The concentration of solution can be expressed in various ways as discussed below.

It refers to the amount of the solute per 100 parts if the solution. It can also be called as parts per hundred (pph). It can be expressed by any of following four methods:
(i) Weight by weight percentage $(\mathbf{\%} \mathbf{w} / \mathbf{w})=\frac{\text { Wt. of solute }(\mathrm{g})}{\text { Wt. of solution }(\mathrm{g})} \times 100$ e.g., $10 \% \mathrm{Na}_{2} \mathrm{CO}_{3}$ solution $\mathrm{w} / \mathrm{w}$ means 10 g of $\mathrm{Na}_{2} \mathrm{CO}_{3}$ is dissolved in 100 g of the solution. (It means $10 \mathrm{~g} \mathrm{Na}_{2} \mathrm{CO}_{3}$ is dissolved in 90 of solvent)
(ii) Weight by volume percent (\%w/v) $=\frac{\text { Wt. of solute }(\mathrm{g})}{\text { Wt. of solution }\left(\mathrm{cm}^{3}\right)} \times 100$ e.g., $10 \% \mathrm{Na}_{2} \mathrm{CO}_{3}(\mathrm{w} / \mathrm{v})$ means $10 \mathrm{~g} \mathrm{Na}_{2} \mathrm{CO}_{3}$ is dissolved in $100 \mathrm{~cm}^{3}$ of solution
(iii) Volume by volume percent $(\% \mathbf{v} / \mathbf{v})=\frac{\text { Volume of solute }\left(\mathrm{cm}^{3}\right)}{\text { Volume of solution }\left(\mathrm{cm}^{3)}\right.} \times 100$ e.g., $10 \%$ ethanol ( $\mathrm{v} / \mathrm{v}$ ) means $10 \mathrm{~cm}^{3}$ of ethanol dissolved in $100 \mathrm{~cm}^{3}$ of solution.
(iv) Volume by volume percent $(\mathbf{\%} \mathbf{v} / \mathbf{v})=\frac{\text { Vol. of solute }}{\text { Wt. of solution }} \times 100$ e.g., $10 \%$ ethanol ( $\mathrm{v} / \mathrm{w}$ ) means $10 \mathrm{~cm}^{3}$ of ethanol dissolved in 100 g of solution.

Example Concentrated nitric acid used as laboratory reagent is usually $69 \%$ by mass of nitric acid. Calculate the volume of the solution which contains 23 g nitric acid. The density of concentrated acid is $1.41 \mathrm{~g} \mathrm{~cm}^{-3}$.
Solution $\quad 69 \mathrm{~g}$ of $\mathrm{HNO}_{3}$ in 100 g solution given density $=1.41 \mathrm{~g} / \mathrm{cc}$

$$
\begin{aligned}
\text { density } & =\frac{\text { mass }}{\text { volume }} \\
\text { volume } & =\frac{\text { mass }}{\text { volume }}=\frac{10000}{1.41} \mathrm{cc}
\end{aligned}
$$

Now,
$69 \mathrm{~g} \mathrm{HNO}_{3}$ is in $\frac{10000}{141}$ volume solution
$23 \mathrm{~g} \mathrm{HNO}_{3} \rightarrow \frac{23}{69} \times \frac{10000}{141}$ volume solution $=\mathbf{2 3 . 6 4} \mathbf{c m}^{\mathbf{3}}$

## 22. Molarity (M)

The number of moles of solute dissolved in one litre solution is called its molarity.

$$
\begin{aligned}
& \qquad \begin{aligned}
& \text { Molarity }=\frac{\text { Number of moles of solute }}{\text { volume of solution in litres }}=\frac{\mathrm{n}}{\mathrm{~V}} \\
& \text { weight of solute in gram }=\frac{\text { Molarity } \times \text { Volume of solution in } \mathrm{mL} \times \text { molecular weight }}{1000} \\
& \text { Molarity }=\frac{\text { Numbers of moles of solute } \times 1000}{\text { Volume of solution in } \mathrm{mL}} \\
&=\frac{\text { weight of solute in grams } \times 1000}{\text { Molecular weight } \times \text { Volume of solution in mL }} \\
&=\frac{\text { strength of solution in gram/litre }}{\text { Molecular weight of solute }}=\text { Normality } \times \frac{\text { Equivalent weight of solute }}{\text { Molecular weight of solute }} .
\end{aligned}
\end{aligned}
$$

Example $\quad$ A bottle of commercial sulphuric acid (density $1.787 \mathrm{~g} \mathrm{ml}^{-1}$ ) is labelled as $86 \%$ by weight. What is the molarity of acid?
Solution $\quad 86 \mathrm{~g} \mathrm{H}_{2} \mathrm{SO}_{4}$ is in 100 g solution
Now

$$
\begin{aligned}
\text { density } & =\frac{\text { mass }}{\text { volume }} \\
1.78 & =\frac{100}{\mathrm{~V}} \\
\mathrm{~V} & =\frac{10000}{1.78}=56.18 \mathrm{ml} .
\end{aligned}
$$

Again

$$
\begin{aligned}
\text { molarity }(M) & =\frac{\text { no. of moles of } \mathrm{H}_{2} \mathrm{SO}_{4} \times 1000}{\text { Volume of sol. in ml. }} \\
& =\frac{\left(\frac{86}{98}\right) \times 1000}{56.18}=\mathbf{1 5 . 6 2} \mathbf{~ M}
\end{aligned}
$$

## 23. Molarity (m)

The number of moles or gram molecules of solute dissolved in 1000 gram of the solvent is called molality of the solution.
Molality of a solution $=\frac{\text { Number of moles of solute }}{\text { Amount of solvent in kg }}=\frac{\text { Number of moles of solute } \times 1000}{\text { Amount of solved in gram }}$ It is independent of temperature.

## 24. Parts per million (ppm) and parts per billion (ppb)

When a solute is present in very small quantity, it is convenient to express the concentration in parts per million and parts per billion. It is the number of parts of solute per million $\left(10^{6}\right)$ or per billion $\left(10^{9}\right)$ parts of solution. It is independent of the temperature.

$$
\begin{aligned}
& \mathrm{ppm}=\frac{\text { Mass of solute component }}{} \\
& \mathrm{ppb}=\underline{\text { Mass of solute component }}
\end{aligned}
$$

Example Calculate the parts per million of $\mathrm{SO}_{2}$ gas in 250 ml water (density $1 \mathrm{~g} \mathrm{~cm}^{-3}$ ) containing $5 \times 10^{-4} \mathrm{~g}$ of $\mathrm{SO}_{2}$ gas.

Solution $\quad$ Mass of $\mathrm{SO}_{2}$ gas $=5 \times 10^{-4} \mathrm{~g}$; Mass of $\mathrm{H}_{2} \mathrm{O}=$ Volume $\times$ Density $=250 \mathrm{~cm}^{3} \times 1 \mathrm{~g} \mathrm{~cm}^{3}=250 \mathrm{~g}$
$\therefore$ Parts per million of $\mathrm{SO}_{2}$ gas $=\frac{5 \times 10^{-4}}{250 \mathrm{~g}} \times 10^{6}=\mathbf{2}$

## 25. Formality (F)

Formality of solution may be defined as the number of gram formula units of the ionic solute dissolved per litre of the solution. It is represented by $\mathbf{F}$. Commonly, the term formality is used to express the concentration of the ionic solids which do not exist as molecules but exist as network of ions. A solution containing one gram formula mass of solute per liter of the solution has formality equal to one and os called Formal solution. It may be mentioned here that the formality of a solution changes with change in temperature.
Formality $(\mathrm{F})=\frac{\text { Number of gram formula units of solute }}{\text { Volume of solution in litres }}$

$$
=\frac{\text { Mass of ionic solute (g) }}{\text { gram formula unit mass of solute } \times \text { Volume of solution (1) }}
$$

Example What will be the formality of $\mathrm{KNO}_{3}$ solution having strength equal to 2.02 g per litre?
Solution Strength of $\mathrm{KNO}_{3}=2.02 \mathrm{gL}^{-1}$ and g formula weight of $\mathrm{KNO}_{3}=101 \mathrm{~g}$

$$
\therefore \text { Formality of } \mathrm{KNO}_{3}=\frac{\text { strength in } \mathrm{g} l^{-1}}{\mathrm{~g} . \text { formula wt. of } \mathrm{KNO}_{3}}=\frac{2.02}{101}=\mathbf{0 . 0 2 F}
$$

## 26. Mole fraction (x)

The ratio of moles of one component to the total number of moles of all the components present in the solution, is called the mole fraction of that component.
Mole fraction of solute $X_{A}$ is given by $X_{A}=\frac{n_{A}}{n_{A}+n_{B}}$
Mole fraction of solute $X_{B}$ is given by $X_{B}=\frac{n_{B}}{n_{A}+n_{B}}$
where $n_{A}$ is moles of solute $A$ and $n_{B}$ is moles of solvent $B$.

## 27. Mass Fraction

Mass fraction of a component in a solution is the mass of the component divided by the total mass of the solution. For a solution containing $w_{A} g m$ of $A$ and $w_{B} g m$ of $B$.

$$
\text { Mass fraction of } A=\frac{W_{A}}{W_{A}+W_{B}} \quad \text { Mass fraction of } B=\frac{W_{B}}{W_{A}+W_{B}}
$$

NOTE It may be notes that molarity, mole fraction, mass fraction etc. are preferred to molarity, normality, formality etc. Because the former involve the weights of the solute and solvent where as later involve volumes of solutions. Temperature has no effect on weights but it has significant effect on volumes.

## 28. Equivalent Weight

Equivalent weight of an element is that part by weight which combines with 1.0 g of hydrogen or 8.0 g of oxygen or 35.5 g of chlorine.
(i) Equivalent weight of a salt $(\mathrm{EW})=\frac{\text { Molar mass }}{\text { Net positive (or nagative) valency }}$
e.g. Equivalent weight $\mathrm{CaCl}_{2}=\frac{\mathrm{M}}{2}, \mathrm{Alcl}_{3}=\frac{\mathrm{M}}{3}, \mathrm{Al}_{2}\left(\mathrm{So}_{4}\right)_{3}=\frac{\mathrm{M}}{6}$
(ii) Equivalent weight of acids $=\frac{\text { Moller mass }}{\text { Basicity }}$
e.g. Equivalent weight $\quad \mathrm{HCl}=\mathrm{M}($ basicity $=1) ; \mathrm{H}_{2} \mathrm{SO}_{4}=\frac{\mathrm{M}}{2}($ basicity $=2)$
$\mathrm{H}_{3} \mathrm{PO}_{4}=\frac{\mathrm{M}}{3}($ basicity $=3)$
(iii) Equivalent weight of bases $=\frac{\text { Moller mass }}{\text { Acidity }}$

$$
\text { e.g. Equivalent weight } \quad \mathrm{NaOH}=\mathrm{M}, \mathrm{Ca}(\mathrm{OH})_{2}=\frac{\mathrm{M}}{2}, \mathrm{Al}(\mathrm{OH})_{3}=\frac{\mathrm{M}}{3}
$$

The number of gram-equivalents (Eq)

$$
\text { Equivalent }=\frac{\text { Weight of compound }}{\equiv \text { valent weight }}=\frac{\mathrm{W}}{\text { Equivalent weight }}
$$

Mole Equivalent Relationship In a given weight (w) of sample, number of moles (n) and number of equivalents (eq) are related as
$\mathrm{n}=\frac{\mathrm{w}}{\mathrm{m}}$ and $\mathrm{Eq}=\frac{\mathrm{w}}{\text { Equivalent weight }}$

$$
\frac{\mathrm{Eq}}{\mathrm{n}}=\frac{\mathrm{M}}{\text { Equivalent weight }}=\mathrm{n}-\text { factor }
$$

$\mathbf{n}$-factor For salt, it is valency, for acid it is basicity, for base it is acidity.

## Normally/Molarity Relationship

$$
\mathrm{N}=\frac{\mathrm{Eq}}{\mathrm{~V}} \text { and } \mathrm{M}=\frac{\mathrm{n}}{\mathrm{~V}} \Rightarrow \frac{\mathrm{~N}}{\mathrm{M}}=\frac{\mathrm{Eq}}{\mathrm{n}}=\frac{\mathrm{MW}}{\mathrm{EW}}=\mathrm{n}-\text { factor }
$$

## 29. Relation Between Molarity And Normality

$S=$ Molarity $\times$ molecular weight of solute and $S=$ Normality $\times$ equivalent weight of solute.

## So we can write

Molarity $\times$ molecular weight of solute $=$ Normality $\times$ equivalent weight of solute.
Normality $=\frac{\text { molarity } \times \text { molecular weight of solute }}{\text { equivalent weight of solute }}=\frac{\text { molarity } \times \text { molecular weight of solute }}{(\text { moleculer weight of solute } / \text { valency factor }}$
Normality = molarity $\times$ valency factor

$$
\mathrm{N}=\mathrm{M} \times \mathrm{n} \quad ; \quad \mathrm{N}>\mathrm{M}
$$

Example Calculate the molarity and molality of a solution of $\mathrm{H}_{2} \mathrm{SO}_{4}$ (sp. gr.=1.98) containing $27 \% \mathrm{H}_{2} \mathrm{SO}_{4}$ by mass.

$$
3.3 \mathrm{M}, 3.77 \mathrm{M}
$$

Solution

$$
\begin{aligned}
& \text { Vol of } 100 \mathrm{~g} \text { of } 27 \% \mathrm{H}_{2} \mathrm{SO}_{4}=\frac{\mathrm{wt} .}{\mathrm{d}}=\frac{100}{1.098} \mathrm{ml} \\
& \mathrm{M}_{\mathrm{H}_{2} \mathrm{SO}_{4}}=\frac{\text { wt./mol.wt. }}{\text { vol, of solution (litre) }}=\frac{27 \times 1.198 \times 1000}{98 \times 100}=\mathbf{3 . 3 m o l} L^{-1} \\
& \mathrm{M}_{\mathrm{H}_{2} \mathrm{SO}_{4}}=\frac{\text { wt. } / \mathrm{mol} . \mathrm{wt} .}{\text { vol, of solvent }(\mathrm{kg})}=\frac{27 \times 1000}{(100-27) \times 98}=\mathbf{3 . 7 7 m o l ~ \mathbf { K g } ^ { - 1 }}
\end{aligned}
$$

## 30. Dilution Formula

If a concentrated solution is diluted, following formula work $M_{1} V_{1}=M_{2} V_{2}$
( $M_{1}$ and $V_{1}$ are the molarity and volumes before dilution and $M_{2}$ and $V_{2}$ are molarity and volumes after dilution)

## 31. Mixing of two or more solutions of different molarities

If two or more solutions of molarities $\left(M_{1}, M_{2}, M_{3}, \ldots\right)$ are mixed together, molarity of the resulting

$$
M=\frac{M_{1} V_{1}+M_{2} V_{2}+M_{3} V_{3} \cdots}{V_{1}+V_{2}+V_{3} \cdots}
$$

solution can be worked out as :

## 32. Strength of $\mathbf{H}_{2} \mathrm{O}_{2}$ solution

The strength of $\mathrm{H}_{2} \mathrm{O}_{2}$ is aqueous solution is expressed in the following two ways:
(i) Percentage strength

The mass of $\mathrm{H}_{2} \mathrm{O}_{2}$ present in 100 ml of the aqueous solution is termed as percentage strength. For example, a $25 \%$ solution (w/v) of $\mathrm{H}_{2} \mathrm{O}_{2}$ means that 25 grams of $\mathrm{H}_{2} \mathrm{O}_{2}$ are present in 100 ml of the solution.
(ii) Volume strength

Strength of the sample of $\mathrm{H}_{2} \mathrm{O}_{2}$ is generally expressed in terms of the volume of oxygen at $0^{\circ} \mathrm{C}$ and 1 atm that one volume of the sample of hydrogen peroxide gives on heating. The commercial samples are marked as ' 10 volume'. ' 15 volume' or ' 20 volume'. 10 volume means that one volume of the sample of hydrogen peroxide gives 10 volumes of oxygen at $0^{\circ} \mathrm{C}$ and 1 atm .1 ml of a 10 volume solution of $\mathrm{H}_{2} \mathrm{O}_{2}$ will liberate 10 ml of oxygen at $0^{\circ} \mathrm{C}$ and 1 atm .

## 33. Percentage labelling of oleum

Oleum is fuming sulphuric acid which contains extra $\mathrm{SO}_{3}$ dissolved in $\mathrm{H}_{2} \mathrm{SO}_{4}$. To convert this extra $\mathrm{SO}_{3}$ into $\mathrm{H}_{2} \mathrm{SO}_{4}$, water has to be added $\left(\mathrm{SO}_{3}+\mathrm{H}_{2} \mathrm{O} \rightarrow \mathrm{H}_{2} \mathrm{SO}_{4}\right)$. The amount of sulphuric acid obtained when just sufficient water is added into 100 g of oleum so that all $\mathrm{SO}_{3}$ present in it is converted into $\mathrm{H}_{2} \mathrm{SO}_{4}$ is called percentage labelling of oleum.

## 34. Relationship Between Different Concentration Terms

(i) $\mathrm{N}=\mathrm{M} \times \mathrm{n}$ factor
(ii) $\mathrm{M}=\frac{\mathrm{md}}{1+\mathrm{mM}_{2} / 1000}$
(iii) $\mathrm{m}=\frac{1000 \times \mathrm{x}_{2}}{\mathrm{x}_{1} \mathrm{M}_{1}}$
(iv) $\mathrm{M}=\frac{1000 \times \mathrm{dx}_{2}}{\mathrm{x}_{1} \mathrm{M}_{1}+\mathrm{x}_{1} \mathrm{M}_{1}}$
(v) $\quad d=M\left(\frac{1}{m}+\frac{M_{2}}{1000}\right)$
(vi) Volume strength of $\mathrm{H}_{2} \mathrm{O}_{2}=5.6 \times \mathrm{N}=\frac{5.6 \times \text { Percentagestrength }}{\text { Eq. wt. of } \mathrm{H}_{2} \mathrm{O}_{2}(17)} \times 10$
(vii) Volume strength of $\mathrm{H}_{2} \mathrm{O}_{2}=11.2 \times \mathrm{M}=\frac{11.2 \times \text { Percentagestrength } \times 10}{\text { Mol. wt. of } \mathrm{H}_{2} \mathrm{O}_{2}(34)}$
(viii) In oleum labelled as $(100+x) \%$
$\%$ of free $\mathrm{SO}_{3}=\left(\frac{80 \times \mathrm{x}}{18}\right)(\mathrm{w} / \mathrm{w})$
where $\mathrm{N}=$ Normality

| $\mathrm{M}=$ Molarity | $\mathrm{m}=$ molarity |
| :--- | :--- |
| $\mathrm{d}=$ density of solution | $\mathrm{M}_{2}=$ Molecular mass of solute |
| $\mathrm{x}_{2}=$ Mole fraction of solute | $\mathrm{x}_{1}=$ Mole fraction of solvent |
| $\mathrm{M}_{1}=$ Molecular mass of solvent | $\mathrm{d}=$ Density of solution |

## NEET Pattern Exercise (1)

1. Which has maximum molecules?
(a) $7 \mathrm{~g} \mathrm{~N}_{2} \mathrm{O}$
(b) $20 \mathrm{~g} \mathrm{H}_{2}$
(c) $16 \mathrm{~g} \mathrm{NO}_{2}$
(d) $16 \mathrm{~g} \mathrm{SO}_{2}$
2. Haemoglobin contains $0.334 \%$ of iron by weight. The molecular weight of haemoglobin is approximately 67200. The number of iron atoms (Atomic weight of Fe is 56 ) present in one molecule of haemoglobin is
(a) 4
(b) 6
(c) 3
(d) 2
3. If the weight of metal chloride is $x$ gram containing $y$ gram of metal, the equivalent weight of metal will be
(a) $E=\frac{X}{y} \times 35.5$
(b) $E=\frac{8(y-x)}{x}$
(c) $E=\frac{y}{(x-y)} \times 35.5$
(d) $E=\frac{8(x-y)}{y}$
4. An organic compound containing C and H gave the following analysis $\mathrm{C}=40 \%, \mathrm{H}=6.7 \%$. Its empirical formula would be
(a) $\mathrm{CH}_{4}$
(b) $\mathrm{CH}_{2} \mathrm{O}$
(c) $\mathrm{C}_{2} \mathrm{H}_{4} \mathrm{O}_{2}$
(d) $\mathrm{C}_{2} \mathrm{H}_{4}$
5. 2.5 litre of 1 M NaOH solution mixed with another 3 litre of 0.5 M NaOH solution. Then find out molarity of resultant solution.
(a) 0.80 M
(b) 1.0 M
(c) 0.73 M
(d) 0.50 M
6. The total number of electrons in 2.0 g of $\mathrm{D}_{2} \mathrm{O}$ to that in 1.8 g of $\mathrm{H}_{2} \mathrm{O}$
(a) Double
(b) Same
(c) Triple
(d) One fourth
7. If the weight of metal oxide is x g containing of $y \mathrm{~g}$ of oxygen, the equivalent weight of metal will be
(a) $E=\frac{8 x}{y}$
(b) $E=\frac{8(y-x)}{x}$
(c) $\mathrm{E}=\frac{\mathrm{y}}{8}$
(d) $E=\frac{8(x-y)}{y}$
8. The number of significant figures in $2.653 \times 10^{4}$ is
(a) 8
(b) 4
(c) 7
(d) 1
9. Mole fraction of solute in aqueous solution of $30 \% \mathrm{NaOH}$.
(a) 0.16
(b) 0.05
(c) 0.25
(d) 0.95

## Assertion-Reason Type Questions

10. In the following questions, a statements of assertion (A) is followed by a statement of reason (R)
(i) If both Assertion \& Reason are true and the reason is the correct explanation of the assertion, then mark (a).
(ii) If both Assertion \& Reason are true but the reason is not the correct explanation of the assertion, then mark (b).
(iii) If Assertion is true statement but Reason is false, then mark (c).
(iv) If both Assertion and Reason are false statement, then mark (d).

A: In the reaction
$2 \mathrm{NaOH}+\mathrm{H}_{3} \mathrm{PO}_{4} \longrightarrow \mathrm{Na}_{2} \mathrm{HPO}_{4}+2 \mathrm{H}_{2} \mathrm{O}$,
equivalent weight of $\mathrm{H}_{3} \mathrm{PO}_{4}$ is $\frac{\mathrm{M}}{2}$, where M is its molecular weight.
R : Equivalent weight $=\frac{\text { Molecular weight }}{n-\text { factor }}$

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## 01. Introduction

- Biology; Bios $=$ life, Logos $=$ Study, means study of life is biology.
- "Biology is the science of life forms and living processes"
- Systematic and monumental description of life forms made human to make a detailed system of identification, nomenclature and classification of organisms i.e. Taxonomy.
- Hence the study of identification, nomenclature and identification is called taxonomy.
- All the organisms have been evolved by a very long and complex process of evolution, so they all are related to each other by sharing of some common genetic material but up to varying degrees. This sharing may be less or more among different cases..
- When human came to know this fact then he humbled and led to cultural movements for conservation of biodiversity.
- Sharing of common characters was also proved when human studied the similarities among living organisms both horizontally and vertically.
- The living world is full of amazing diversity of living organisms.
- The diversity of habitats of organisms is also very vast and amazing.
- This diversity makes us deeply reflect on "What indeed is life" ? This question actually asks to solve two problems.
(i) First is a technical $\rightarrow$ What living is as opposed to the non living means Living v/s Non living.
(ii) Second is a philosophical one $\rightarrow$ what the purpose of life is? As scientists we will try to solve the first question, because the second question is more related to philosophy rather science.


## 02. Characters of Living Organism

Following are the main characters of "living"-
$\left.\begin{array}{l}\text { GROWTH } \\ \text { REPRODUCTION }\end{array}\right\}$ Not the defining properties/characters/features
METABOLISM
$\left.\begin{array}{l}\text { CELLULARORGANISATION } \\ \text { CONSCIOUSNESS }\end{array}\right\}$ Defining properties/characters/features
The character which has no exception is called as defining property of life.

## Growth

- Increase in mass or overall size of a tissue or organism or its parts is called growth.
- Increase in mass and increase in number of individuals are twin characters of growth.
- Growth is an irreversible permanent increase in size of an organ or its parts or even of an individual cell.
- Growth is of two types :
(a) Intrinsic growth :- This growth is from inside of the body of living organisms.
(b) Extrinsic growth :- This growth is from outside i.e. accumulation of material on any body surface Non living exhibts this type of growth.
- Growth is of two types :
(a) Indeterminate growth $=$ Unlimited growth $\rightarrow$ Growth which occurs continuously throughout their life span is indeterminate growth or unlimited growth. It occurs in plants and not in animals.
(b) Determinate growth $=$ Limited growth $\rightarrow$ Growth which occurs only up to a certain age is determinate growth or Limited growth. It occurs in animals. However, cell division occurs in certain tissues to replace lost cells.
- In majority of higher plants and animals, growth and reproduction are mutually exclusive events.
- Because both living and nonliving exhibit growth so it can not be taken as defining property.
- Growth from inside (intrinsic growth) can be taken as defining property.


## Reproduction

Reproduction is one of the fundamental characteristics of living organisms. It can be defined as the production of new individuals of same kind by the grown up individuals. It is characteristic exhibited by living organisms which can produce new young ones of their own kind. There are two modes of reproduction - asexual and sexual.

- In lower organisms like yeasts and Hydra, budding takes place in which new individuals are produced by the formation of an outgrowth known as 'bud'.
- Fragmentation is another mode of asexual reproduction, as in this, body of an organism (parent body) breaks up into two or more parts (known as fragments) each of which grows into a new individual. It is also quite common in filamentous algae, fungus, bryophytes (at protonema stage which occurs during life cycle in mosses).
- Planaria (flat worms) exhibit an extraordinary ability to regenerate its lost body parts completely (which is known as true regeneration).
- This is a method of reproduction as new planarians develop by splitting of parent planarian body either lengthwise or transversely. In higher organisms like plants, animals sexual mode of reproduction is quite common which involves formation of gametes (i.e., sex cells) from two parents of opposite sexes but same species. These gametes then fuse to form zygote ( 2 n ) which develops to form a new organism of same kind.
- Hence, reproduction is shown by all living organisms except a few which are either sterile or infertile, like mule, worker-bees, infertile human couples, etc. do not reproduce at all.
- Hence, reproduction can be regarded as characteristic of living organisms but it is not their exclusive defining characteristic.


## Metabolism :

- The sum total all the chemical reactions occuring in our body is metabolism.
- All living organisms, both unicellular and multicellular exhibit metabolism.
- No non-living object shows metabolism.
- In this way metabolism is a defining character of living organisms because it has no exceptions.
- Now we have known most of the chemical or metabolic reactions occuring in our body so we can demonstrate many of them in a cell free medium or in a test tube in lab.
- The isolated metabolic reaction outside the body of an organism, performed in a test tube (in-vitro) is neither living nor nonliving.
- These isolated reaction can be regarded as living things, but they are definitely living reactions because they are similar to the reactions performing in our body.
- Here we should not forget the fact that metabolism is the total sum of all the chemical reactions performing in our body, it is not the sum of few or more living reactions.
- All organisms are made of small or big chemicals perform thousands of reactions and form some other chemicals also in the bodies of living organisms.
- All plants, animals, fungi and microbes exhibit metabolism.


## Consciousness

- Most obvious and technically complicated feature of all living organism. All living organisms are able to detect changes, i.e., sense their surroundings and can also respond to them. This is known as sensitivity which is defined as the ability to detect changes in the environment and to give response towards it accordingly. Any change that can be detected by an organism is called stimulus. This can be physical (like intensity, duration, direction of light, sound, change in temperature, duration of day length, i.e., photoperiod, etc.), chemical (like acids, pollutants, etc.) or biological (like other organisms).
- Besides, human being is the only organism, who is aware of himself. He has self-consciousness too with awareness of the surroundings. He relate his mind to the changes taking place in the world. He is an intelligent animal with thoughts, feelings and self-hood. sensitivity or awareness is regarded as defining property of living organisms as non-living things do not have power of sensing their surroundings and give response according to it. However, patients lying in coma in hospitals virtually supported by machines which replace heart and lungs are neither living nor dead otherwise brain-dead.


## 03. Diversity In The Living World

- We consider vast areas like forest, desert, plateau etc. we find that number and kinds of living organisms increase many folds. These different kinds of plants, animals and other organisms are referred to as 'Biodiversity' of this earth.
- Biodiversity is the number and various kinds of organisms found on earth. It stands for the variability found among living organisms inhabiting this world.
- Biodiversity (Greek word bios = life; diversity $=$ forms) or biological diversity can be defined as the vast array of species of microorganisms, algae, fungi, plants, animals occuring on the earth either in the terrestrial or aquatic habitats and the ecological complexes of which they are a part.
- According to IUCN (International Union of Conservation of Nature and Natural resources), currently known and described species of all organisms are between 1.7-1.8 million. These millions of plants, animals and other organisms in the world cannot be recognised, studied or described by an individual on its own.
- There is need to standardize the names of all living organisms after proper identification, in order to study such diverse life forms. Therefore, for better understanding of biodiversity scientists have established a definite system of principles, procedures and terms which identifies, categories and assigns specific name to each and every organism known to us. Such systems are acceptable to all biologists all over the world.
- The scientific need for simple, stable and internationally accepted systems for naming the living organisms of the world has generated, a process called "Nomenclature". And, before assigning a specific name to an organism, one should determine or know its kind or features correctly, so that one can identify it in each every part of the world. This is known as "Identification"


## 04. Taxonomy

- This word was proposed by A.P. De Candolle in his book "Theories elementaire de la botanique" (Theory of elementary botany)
- Taxonomy includes study of following 4 points

Characterization - Identification of character
Identification - Identification of living organisms
Nomenclature - Nomenclature of living organisms
Classification - Classification of living organisms in groups

## 05. Systematics

(i) The term "Systematics" was pro posed by Linnaeus. The word systematics is derived from the latin word "systema" which means systematic arrangement of organisms.
(ii) According to G. Simpson : Systematics is the study of diversity of organisms and all their comparative and evolutionary relationship.

NOTE It includes description of external morphological characters of plants of living organisms.
eg. Morphological characters of Root, Stem, Leaves, Flowers.

## 06. New Systematics or Neosystematics or Biosystematics

- Neo-systematics - A new branch - Name given by Julian Huxley (1940)
- It includes description of all the characters (external + internal)
e.g. Anatomical characters, Histological characters, Embryological characters, Identical characters
- It is used to know the inter-relationship between living organisms.

NOTE New systematics is mainly based on evolutionary as well as genetic relationship as compared to morphological characters.

## 07. Types of Taxonomy

- Alpha taxonomy or classical taxonomy : It is based on external morphology of plants.
- Bete taxonomy or Explorative taxonomy : Besides external morphology it also includes internal characters like embryological, cryological, anatomical characters etc.
- Omega taxonomy or Encyclopaedic taxonomy : Omega taxonomy have widest scope. It is based on all the information or data available about plants.
- Chemotaxonomy : The uses of chemical characters of plants in classification or in solving taxonomic problems is called chemotaxonomy or chemical taxonomy. It is based on the chemical constituents of plants. The fragrance and taste vary from species to species.

The basic chemical compounds used in chemotaxonomy are alkaloids, carotenoides, tannins, polysaccharide, nucleic acids, fatty acids, amino acids, aromatic compounds etc.

- Cytotaxonomy : The use of cytological characters of plants in classification or in solving taxonomic problems is called cytotaxonomy.
- Following characters are used in cytotaxonomy
* Chromosome number.
* Chromosome morphology - including chromosome size, total length of chromatin, arms ratio, primary and secondary constriction.
* Heterochromatin.
* Chromosome behaviour at meiosis.
* Polyploidy - autopolyploidy, allopolyploidy and aneuploidy.
* Chromosomal aberrations.
* Different type of cytochromes.

Cytological data have been used in many cases to find out the affinities among genera.

- Karyotaxonomy : Based on characters of nucleus and chromosomes. Pattern of chromosomal bands (dark bands and light bands) is most specific character.
- Adansonian system or Phenetic Classification or Numerical Classification.
* Proposed by "Sokel and Sneath"
* In it plants are classified on the basis of number of similarities and dissimilarities.
* In this, importance to any one character is not given, all characters have same importance.
While in natural classification floral characters have importance than morphological characters.


## 08. Significance of Taxonomy

- Most significant feature is identification of living organism.
- With the help of taxonomy diversity of living being can be studied easily.
- Maximum diversity of living beings are found in tropical rain forests.
- These forest have heavy rain fall through out the year.
- In India maximum tropical rain forest are found in Andaman \& Nicobar and in all eastern states (Assam, Meghalaya, West Bengal etc.)
- At present, 300 lakh ( $\mathbf{3 0}$ million) type of living organism are found on our earth.
- 17 lakh ( $\mathbf{1 . 7}$ million) type of living organism have been discovered till now.
- Out of them $\mathbf{1 2}$ lakh types are animal and $\mathbf{5}$ lakh types are plants.


## 09. Taxonomic Category

Classification is not a single step process but involves heirarchy of steps in which each step represent a rank or category.

## - Species :

Taxonomic studies consider a group of individual organism with fundamental similarities as a species. One should be able to distinguish one species from the other closely related species based on the distinct morphological difference.

- Genus :

Genus comperises a group of related species which has more characters in common in comparison to species of other genera.

- Family :

Family has a group of related genera with still less number of similarities as compared to genus and species. Families are characterized on the basis of both vegetative and reproductive feature of plant species.
For example :Three different genera Solanum, Petunia and Datura are included in family solanaceae.

- Order :

Order being a higher category is the assemblage of families which exhibit a few similar character.
For example : Plant families like convolvulaceae, Solanaceae are included in the order polymoniales mainly based on the floral characters.

- Class :

Class includes organism of related orders having less similarities than orders.

- Division :

Division includes all organisms belonging to different classes having a few common characters. There 7 main taxonomic categories. They are obligate or essential or broad categories i.e. they are strictly used at the time of any plant classification. There are some extra or sub categories, like sub division, sub order, sub family, etc. They are used only when they are needed.

| Biggest group $\rightarrow$ | - Kingdom |
| ---: | :--- |
|  | - Division/Phylum |
|  | - Class |
|  | - Order - Cohort |
|  | - Family |
|  | - Genus |
|  | - Species |

* The classification of any plant or animal is written in descending or ascending order.
* Hierarchy - Descending or ascending arrangement of taxonomic categories is known as hierarchy.
$\star$ Species : Smallest taxonomic category $\rightarrow$ It is basic unit of classification.

NOTE As we go higher from species to kingdom, number of common characters decreases. Lower the taxa more are the characteristics that the members with in the taxon share. Higher the category, greater is the difficulty of determining the relationship to other taxa at the same level.

| Suffix for taxa | (Taxon) |  |
| :--- | :--- | :--- |
| Division | - | phyta |
| Sub div | - | phytina |
| Class | - | opsida, phyceae, ae |
| Order | - | ales |
| Sub-order | - | ineae |
| Family | - | aceae |
| Sub Family | - | oideae |

NOTE There is no suffix for Genus, Species and Kingdom.

## 10. Nomenclature

- Polynomical system :
* According to this system name of any plant consists of many words.
* For eg. Caryophyllum $\rightarrow$ Caryophyllum saxatilis folis gramineus umbellatis corymbis
* Based of morphology mainly
- Trinomical system :
* Proposed by Lamarck.
* According to this system name of any plant or species is composed of three names-
- Generic name
- Specific epithet
- Subspecific name (Name of variety)
* When members of any species have large variations then trinomial system is used.
* On the basis of dissimilarities this species is classified into sub species. eg.

Brassica oleracea var. botrytis (Cauliflower)
Brassica oleracea var. capitata (Cabbage)
Brassica oleracea var. caulorapa (Knol-Khol)

## - Binomial system :

* Binomial system was first proposed by Gaspard Bauhin in his book -"Pinax Theatre Botanica"
* Carolus Linnaeus : Linnaeus used this nomenclature system for the first time on large scale and proposed scientific name of all the plants and animals.
* Linnaeus is the founder of binomial system.
* Linnaeus proposed scientific name of plants in his book "Species Plantarum"
* It was published on 1 May, 1753. So this was the initiation of binomial system for plants.
* So any name proposed (for plants) before this date is not accepted today.
$\not$ * Linnaeus proposed scientific name of animals in his book "Systema Naturae" $\left(10^{\text {th }}\right.$ edition).
* This $10^{\text {th }}$ edition of Systema Naturae was first published on 1 August, 1758.
* So initiation of binomial system for animals is believed to be started on 1 August, 1758.


## 11. ICBN

- Main rules of ICBN :
* According to binomial system name of any species consists of two components or words.
- Generic name Name of genus
- Specific epithet
$\begin{array}{ccccc}\text { e.g. } & \text { Solanum tuberosum (Potato) } & \text { Mangifera indica (Mango) } \\ & \downarrow & \downarrow & \downarrow & \downarrow \\ & \text { Generic name } & \text { Specific epithet } & \text { Generic name } & \text { Specific epithet }\end{array}$
* In plant nomenclature (ICBN) tautonyms are not valid i.e. generic name and specific epithet should not be same in plants.
eg. Mangifera mangifera
But tautonyms are valid in animal nomenclature (ICZN-International Code of Zoological Nomenclature)
eg. Naja naja (Indian cobra), Rattus rattus (Rat)
$\star$ First letter of generic name should be in capital letter and first letter of specific epithet should be in small letter.
eg. Mangifera indica
* But if specific epithet is based on the name of any person, its first letter should be in capital letter.
eg. Isoetes Pantii
* When written with free hand or typed, then generic name and specific epithet should be separately underlined.
But during printing name should be in italics to indicate their latin orgin.
* Name of scientist (who proposed nomenclature) should be written in short after the specific epithet.
eg. Mangifera indica Linn.
* Name of scientist should be neither underlined nor in italics, but written in Roman letters (simple alphabets)
* Sciectific names should be derived from Latin (usually) or Greek languages because they are dead languages.
* Type specimen (Herbarium Sheet) of newly discovered plant should be placed in herbarium (Dry garden).


## 12. Classification

According to "A.P. de Candolle", Classification is of two types

## (i) Empirical Classification <br> (ii) Rational Classification

- Empirical Classification :
* In this type, the actual nature or character of plants is not considered.
* Plants are classified on the basis of their alphabetical order.
* In this way plants are classified into 26 groups.
- Rational Classification

In this classification, plants are classified on the basis of their actual character or nature i.e. by viewing the characters.

Types of rational classification
(iii) Practical classification

In this type of classification, plants are classified on the basis of their economic importance. e.g. Oil yielding plants $\rightarrow$ Coconut, Walnut, Soyabean

Fibre yielding plants $\rightarrow$ Jute, Cotton
Medicinal plants $\rightarrow$ Rauwolfia, Cinchona, Eucalyptus

NOTE In this classification, any one plant can be a member of more than one group. eg. Turmeric : Multi uses plant, it gives both medicines and spices.

## (iv) Artificial classification :

In this type of classification plants are classified on the basis of one or two morphological characters. i.e. over all morphology is not considered.
for e.g. - Classification proposed by Linnaeus is Artificial
Linnaeus classified plant kingdom on the basis stamen into 24 classes.

NOTE Linnaeus classification is also called sexual classification.
NOTE Linnaeus divided flowering plants into 23 classed starting with class monandria with a single stamen (eg. Canna) and plants with twenty or more stamens attached with calyx were assigned to class Icosandria. He also included all non-flowering plants such as algae, fungi, mosses and ferns in a separate class called cryptogamia.
(v) Natural classification:

In this type, plants are classified on the basis of their complete morphological characters of stem, root, leaves, flowers etc.
Importance -
Natural classification is believed to be the best classification, because it represents the natural similarities and dissimilarities of plants i.e. it represents the interrelationship among plants.
In this classification, the plants belonging to the same group shows many similarities, while in artificial classification, the plants belonging to the same group shows only, 1 or 2 similar characters. They have many dissimilarities.
Natural classification is of two types

- Natural formal
- Natural phylogenetic
(a) Natural formal $\rightarrow$ In this classification, the phylogeny of the plant is not considered i.e. only the morphology of the plant is considered.
(b) Natural phylogenetic $\rightarrow$ In this classification, both morphology and phylogeny are considered. In phylogenetic classification, the plants are arranged on the basis of their evolution.

Thallophyta $\rightarrow$ Bryophyta $\rightarrow$ Pteridophyta $\rightarrow$ Gymnosperm $\rightarrow$ Angiosperm (most advanced plants)
Note : Phylogenetic classification also known as cladistic classification
(vi) Adansonian system or phenetic classification or Numerical classification :

In it plants are classified on the basis of numbers of similarities and dissimilarities. This classification is easily carried out by using computers and it is based on all observable characteristics. In this classification number and codes are assigned to all the characters and the data are prepared and then processed. Those organism which have maximum similarities are placed in same group. In this way each character is given equal importance and at the same hundreds of characters can be considered.

NOTE In this classification importance to any one character is not given, all characters have same importance. While in natural classification floral (reproductive) characters have more importance than vegetative (root, stem and leaves) characters.

## 13. Taxonomical Aids

- Accurate classification and identification of organisms is required which needs field studies and intensive laboratory work. This is done after collection of actual specimens of plants and animal species which is the primary source of all taxonomic studies.
- Hence, these taxonomical studies help in

Fundamental study of different living organisms.
Also aid in their systematic study.
Information gathered is stored with specimens for future studies.

## Herbarium

- It is defined as "store house of collected plant specimens that are dried, pressed and preserved on sheets". Further, these sheets are arranged in the sequence of a universally accepted system of classification.
- Such herbaria serve as quick source of reference in taxonomical studies. It also provides information about the local flora as well as flora of distant areas. This information is also useful in locating wild varieties and relatives of economically important plants.
List of some Herbaria of the world :
(i) Royal Botanical Gardens, Kew (England)
(ii) Central National Herbarium, Calcutta


## 14. Botanical Gardens

Botanical garden is an institution located in an enclosed piece of land which grows numerous kinds of plants obtained from different places for botanical studies. Each plant is first identified and then labelled indicating its botanical/scientific name and its family.

## List of some Botanical Gardens

(i) Royal Botanical Garden, Kew (England)
(ii) National Botanica Garden, Lucknow
(iii) Indian Botanical Garden, Howrah
(iv) Lloyd Botanical Garden, Darjeeling

## 15. Museum

- Museum is a place used for storing, preservation and exhibition of both plants and animals. All educational institutes and universities maintain museums in their Botany and Zoology departments.
- A commonly used preservative solutions is "Formalin". Plants and animals specimen may also be preserved as dry specimens. For instances, insects are collected, killed and pinned before preserving them in special insect boxes while larger animals like reptiles, birds and mammals are usually stuffed and then preserved.
- National Museum of Natural History (NMNH) in Delhi is important from natural science point of view.


## 16. Zoological Parks

- Zoological parks commonly known as zoos are the places where live wild animals are kept in protected environment which is made similar to their natural habitats as much as possible. Here, they are provided with protection and care by human beings.
- These parks serve as ideal means to study and learn different food habits and behaviour of variety of animals. So, students should visit nearby zoos for knowledge and entertainment both.
- National Zoological Park (Delhi) is one of the finest zoos of Asia.


## 17. Key

- Key is an important taxonomic aid used for identification of plants and animals based on the similarities and dissimilarities. Actually, it is a set of alternate characters of different types arranged sequence wise in such a fashion that by selection and elimination one can quickly find out the name of the organism.
- The keys are based on the set of contrasting characters generally in a pair known as "couplet". Each character of the couplet or statement in the key is called as "lead".
- One has to choose correct option between two statements of characters of definite species so that the animal or plant is identified accurately. Keys are generally analytical in nature.


## 18. Flora, Manuals, Monographs and Catalogues

- These flora, manuals, monographs, etc. are recorded descriptions of plants, animals and other organisms. They provide correct identification and description of variety of living organisms.
* Flora : It is a book containing information about plants found in a particular area. It gives the actual account of habitat and distribution of various plants of a given area. These provide the index to the plant species found in a particular area. For example, Flora of Delhi by J.K. Maheshwari.
* Manual : It is a book containing complete listing and description of the plants growing in a particular area. They provide useful information for identification of names of various species found in an area.
* Monograph : It contains information of any one taxon.
* Catalogue : It include the alphabetical arrangement of species of a particular place describing their features.


## 19. Species concept

John Ray : Proposed the term and concept of species.

## - Biological concept of species

* Enst Mayr (Darwin of $20^{\text {th }}$ century) proposed the biological concept of species.
* Mayr defined the "species" in the form of biological concept.
* According to Mayr "All the members that can interbreed among them self and can produce fertile offsprings are the members of same species"

But this definition of Mayr was incomplete because this definition is applicable to sexually reproducing living beings because there are many organisms that have only asexual mode of reproduction.
eg. Bacteria, Mycoplasma, BGA

* The main character in determination of any species is interbreeding. But this character is not used in taxonomy. In taxonomy, the determination of species is mainly based on morphological characters.
* In higher plants, the determination of species is mainly based on the morphology of flower (floral morphology). Because floral (reproductive) characters are more conservative as compared to vegetative (Root, Stem, Leaf) characters i.e. they do not show major changes.
- Typological Concept
* It was proposed by "Aristotle" and "Plato".
$\div$ This concept is based on single individual of a species.
* According to this concept, "There is a definite type or pattern of characters in the each species and every living organisms and all the members of species shows maximum resemblance with this pattern".
* The species in which one foxed pattern of characters is present are called as monotypic species.
eg. Bacteria, Blue green algae
* In many species, more than one type of pattern of characters are present. These are called "Polytypic species" or "Macrospecies".
eg. Brassica oleracea $\rightarrow$ Cauliflower, Cabbage, Knol-Khol.


## NEET Pattern Exercise (1)

1. Which one of the following scientist's name is correctly matched with the theory put forth by him?
(a) Weismann - Theory of continuity of germplasm
(b) Pasteur - inheritance of acquired characters
(c) De Vries - Natural selection
(d) Mendel - Theory of pangenesis
2. Carbohydrates the most abundant biomolecules on earth, are produced by
(a) all bacteria, fungi and algae
(b) fungi, algae and green plant cells
(c) some bacteria, algae and green plant cells
(d) viruses, fungi and bacteria
3. Age of fossils in the past was generally determined by radio-carbon method and other methods involving radioactive elements found in the rocks. More precise methods, which were used recently and led to the revision of the evolutionary periods for different groups of organism, include
(a) study of carbohydrates/proteins in fossils
(b) study of the conditions of fossilisation
(c) Electron Spin Resonance (ESR) and fossil DNA
(d) study of carbohydrates/proteins in rocks
4. According to Oparin, which one of the following was not present in the primitive atmosphere of the earth?
(a) Methane
(b) Oxygen
(c) Hydrogen
(d) Water vapour
5. There is no life on moon due to the absence of
(a) $\mathrm{O}_{2}$
(b) water
(c) light
(d) temperature
6. First life on earth was
(a) cyanobacteria
(b) chemoheterotrophs
(c) autotrophs
(d) photoautotrophs
7. If there was no $\mathrm{CO}_{2}$ in the earth's atmosphere the temperature of earth's surface would be
(a) higher than the present
(b) less than the present
(c) the same
(d) dependent on the amount of oxygen in the atmosphere
8. The $\mathrm{CO}_{2}$ content by volume, in the atmospheric air is about
(a) $0.0314 \%$
(b) $0.34 \%$
(c) $3.34 \%$
(d) $4 \%$
9. Glycogen is a polymer of
(a) galactose
(b) glucose
(c) fructose
(d) sucrose
10. A nucleotide is formed of
(a) purine, pyrimidine and phosphate
(b) purine, sugar and phosphate
(c) nitrogen base, sugar and phosphate
(d) pyrimidine, sugar and phosphate

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Q1 (a)
Pasteur gave the Theory about the Germ Plasm, De Vries give the Theory about Mutation and mendel works on Theory of Inheritance. So the Weismann give the Theory of Continuity of germplasm, according to the germ-plasm theory, inheritance in a multicellular organism only takes place by means of the germ cells: the gametes, such as egg cells and sperm cells. Genetic information cannot pass from soma to germ-plasm and on to the next generation.

Q2 (c)
Some Photosynthetic Bacteria such as Rhodopseudomonas can prepare carbohydrates. but during this type of food synthesis $\mathrm{O}_{2}$ is not evolved because in this case hydrogen donor is other than $\mathrm{H}_{2} \mathrm{O}$. Algae (blue and green-blue) and all green plant cells prepare their food (carbohydrate) through photosynthesis. Here, hydrogen ions are donated by water molecules by the process of photolysis of water, i.e. $\mathrm{O}_{2}$ is released during this type of food synthesis.

Q3 (c)
Electron Spin Resonance (ESR) is a technique used to date newly formed materials, which Radiocarbon dating cannot, like carbonates, tooth enamel, or materials that have been previously heated like igneous rock. The age of substance can be determined by measuring the dosage of radiation since the time of its formation. DNA in ices and DNA in hydrated solids give nearly identical results, suggesting that the DNA strands in ices are as closely packed as those in the hydrated solid DNA samples. Our results suggest that previous reports of extensive electron-transfer distances for DNA in icy media are found to be better explained by substantial inter-double-strand electron transfer. this was best method which were used recently for study the evolutionary periods for different groups of organism.

Q4 (b)
According to Oparin, in the primitive Earth's surface, carbon, hydrogen, water vapour, and ammonia reacted to form the first organic compounds. He believed that organic molecules could be formed from abiogenic materials in the presence of an external energy source (e.g., ultraviolet radiation) and that the primitive atmosphere was reducing (having very low amounts of free oxygen) and contained ammonia and water vapour, among other gases. Both also suspected that the first life-forms appeared in the warm, primitive ocean and were heterotrophic (obtaining preformed nutrients from the compounds in existence on early Earth) rather than autotrophic (generating food and nutrients
from sunlight or inorganic materials). So in the primitive environment Oxygen is absent.
Q5 (b)
Water is an essential constituent of cytoplasm of all living organisms it helps in distribution of substances within the organism elimination of waste products, body temperature maintenance etc. It is absent on moon. Anaerobic organisms can live in the absence of $\mathrm{O}_{2}$. Light and temperature are already known to exist on moon. Lack of liquid water: Life, as we know it, requires liquid water to thrive. The lack of atmosphere means no greenhouse effect and very low pressure. The surface temperature is either extremely cold, or extremely hot- water cannot exist in a liquid state on the lunar surface.
Q6 (b)
Chemoautotrophs does carbon fixation on their own and chemoautotrophs gets the carbon from outside. If you consider the process of evolution, traits evolve over long periods of time and applying the principle of incremental complexity, the complex will follow simpler. Since autotrophs need more time to evolve complex mechanisms to synthesize carbon on their own compared to get it somewhere else, heterotrophs would have evolved first on the earth.

Q7 (b)
$\mathrm{CO}_{2}$ keeps our earth warm by not letting earth's terrestrial radiation escape into space. It acts like a greenhouse. Remove this layer of insulation, out earth will become really cold. The amount of atmospheric $\mathrm{CO}_{2}$, or carbon dioxide, is inextricably linked to climate. Average global carbon dioxide levels and average global temperatures go hand in hand, and temperature changes cause changes in precipitation. Concentrations of carbon dioxide in the atmosphere tend to fluctuate in a cyclical pattern over tens of thousands of years, as do worldwide periods of cold including Ice Ages and warmth.

Q8 (a)
Carbon dioxide $\left(\mathrm{CO}_{2}\right)$ is the $5^{\text {th }}$ most abundant gas in the atmosphere and, together with water vapor, probably the best known of the greenhouse gases. Carbon dioxide levels in the atmosphere are not constant -- they have risen by nearly 40 percent since the Industrial Revolution, according to climate scientist Todd Sanford. They are small compared to the main atmospheric components of nitrogen and oxygen. Scientists express them as parts per million, or ppm. In March 2014, carbon dioxide levels were at 314 ppm , which is 0.0314 percent of the atmosphere. This roughly corresponds to a mass of 3 trillion tons. After nitrogen, oxygen, water vapor and argon, carbon dioxide is the fifth most abundant gas in the atmosphere.

Q9 (b)
Glycogen is a readily mobilized storage form of glucose. It is a very large, branched polymer of glucose residues that can be broken down to yield glucose molecules when energy is needed. Most of the glucose residues in glycogen are linked by $\alpha-1,4$-glycosidic bonds. Branches at about every tenth residue are created by $\alpha-1,6$-glycosidic bonds. Recall that $\alpha$-glycosidic linkages form open helical polymers, whereas $\beta$ linkages produce nearly straight strands that form structural fibrils, as in cellulose.

Q10 (c)
DNA is just a pattern made up of four different nucleotides. Each chain of the double helix is made up of repeating units called nucleotides. Each nucleotide consists of a sugar (deoxyribose) in
the middle of a phosphate group and a nitrogenous base. Nucleotides form a pair in a molecule of DNA where two adjacent bases form hydrogen bonds. The nitrogenous bases of the DNA always pair up in specific way, purine with pyrimidine (A with $T$, $G$ with $C$ ), held together by weak hydrogen bonds. A single nucleotide is composed of three functional groups: a sugar, a triphosphate, and a nitrogenous (nitrogen-containing) base.


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## Class 12 | Physics

## 01 Electrostatics-I

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## 01. Introduction

Electrostatics, deals with the study of charges in rest. These stationary charges occurs due to friction of two insulting bodies, therefore it is often called frictional electricity.

## Important points

(i) Gravitational force is the weakest while nuclear force is the strongest force of the nature
(ii) Nuclear force does not depend upon charge, it acts equally between proton-proton, proton neutron and neutron-neutron.
(iii) There are weak forces acting in $\beta$-degradiation in radio-activity.
(iv) A stationary charge produces electric filed while a moving charge produce electric as well as magnetic field.
(v) Moving charge produce electric field as well as magnetic field but does not radiate energy while uniform acceleration.
(vi) Accelerated charge produce electric field as well as magnetic field and radiate energy.

## 02. Charge

Property of a substance by virtue of which it can repel or attract another charged substance.
Charges are of two types
(a) Positive charge : Lesser number of electrons than number of protons.
(b) Negative charge : More number of electrons than number of protons

Importants Points : Only, electron is responsible for a substance to be charged and not the proton.

## Properties of Charge

(i) Like charges repel while unlike charges attract each other.
(ii) Charge is quantized in nature i.e. The magnitude of charge possessed by different objects is always and integral multiple of charge of electron (or proton) i.e. $q= \pm$ ne where $n$ $=1,2,3$ $\qquad$
(iii) The minimum possible charge that can exist in nature is the charge of electron which has a magnitude of $\mathrm{e}=1.60207 \times 10^{-19}$ coulomb. This is also known as quantum of charge or fundamental charge.
(iv) In an isolated system the algebraic sum of total charge remains constant. This is the law of 'Conservation of charge'.

## 03. Coulomb's Las

The force of attraction or repulsion between two stationary point charges is directly proportional to the product of charges and inversely proportional to the square of distance between them. This force acts along the line joining the two. If $q_{1} \& q_{2}$ are charges in consideration r , the distance between them and F , the force acting between them


Then, $\mathrm{F} \propto \mathrm{q}_{1} \mathrm{q}_{2}$

$$
\begin{aligned}
& \mathrm{F} \propto 1 / \mathrm{r}^{2} \\
\therefore \quad & \mathrm{~F} \propto \frac{\mathrm{q}_{1} \mathrm{q}_{2}}{\mathrm{r}^{2}} \\
\Rightarrow \quad & \mathrm{~F}=\mathrm{K} \frac{\mathrm{q}_{1} \mathrm{q}_{2}}{\mathrm{r}^{2}}, \text { where } \mathrm{k}=\text { constant. } \\
& \mathrm{K}=\frac{1}{4 \pi \varepsilon_{0} \varepsilon_{\mathrm{r}}}=\frac{9 \times 10^{9}}{\varepsilon_{\mathrm{r}}} \mathrm{Nm}^{2} \mathrm{C}^{-2}
\end{aligned}
$$

where,
$\varepsilon_{0}=$ Electric permittivity of vacuum or air $=8.85 \times 10^{-12} \mathrm{C}^{2} \mathrm{~N}^{-1} \mathrm{~m}^{-2}$ and
K or $\varepsilon_{\mathrm{r}}=$ Relative permittivity or Dielectric constant or Specific inductive capacity
$\varepsilon_{\mathrm{r}}=\frac{\varepsilon}{\varepsilon_{0}} \Rightarrow \varepsilon=\varepsilon_{0} \varepsilon_{\mathrm{r}}$
[Newton's law for particles is analogous to coulomb's law for rest charge. The difference is that Newton's law gives attraction force while coulomb's law gives attraction as well as repulsion force]

NOTE (i) Coulomb's law is applicable to point charges only. But it can be applied for distributed charges also
(ii) This law is valid only for stationary charges and cannot be applied for moving charges.
(iii) This law is valid only if the distance between two charges is not less than $10^{-15}$ m

## Direction

Direction of the force acting between two charges depends upon their nature and it is along the line joining two charges.

$\overrightarrow{\mathrm{F}}_{21}=$ force on $\mathrm{q}_{2}$ due to $\mathrm{q}_{1}$

$$
\begin{equation*}
\overrightarrow{\mathrm{F}}_{21}=\frac{\mathrm{q}_{1} \mathrm{q}_{2}}{4 \pi \varepsilon_{0} \varepsilon_{\mathrm{r}} \mathrm{r}_{12}^{2}} \hat{\mathrm{r}}_{12} \tag{A}
\end{equation*}
$$

(where $\hat{r}_{12}$ is a unit vector pointing from $\mathrm{q}_{1}$ to $\mathrm{q}_{2}$ )
$\overrightarrow{\mathrm{F}}_{12}=$ Force on $\mathrm{q}_{1}$ due to $\mathrm{q}_{2}$
$\overrightarrow{\mathrm{F}}_{12}=\frac{\mathrm{q}_{1} \mathrm{q}_{2}}{4 \pi \varepsilon_{0} \varepsilon_{\mathrm{r}} \mathrm{r}_{12}^{2}} \hat{\mathrm{r}}_{21}$
(where $\hat{\mathrm{r}}_{21}$ is a unit vector pointing from $\mathrm{q}_{2}$ to $\mathrm{q}_{1}$ )
$\Rightarrow$ Electric force between two charges not depends on neighbouring charges.
$\Rightarrow$ If a dielectric slab ( $\epsilon_{\mathrm{r}}$ ) of thickness ' t ' is placed between two charges (distance d), force decreases.

$$
\mathrm{F}=\frac{\mathrm{Q}_{1} \mathrm{Q}_{2}}{4 \pi \in_{0} \mathrm{r}^{2}} \quad \text { where } \mathrm{r}=\mathrm{d}-\mathrm{t}+\mathrm{t} \sqrt{\epsilon_{\mathrm{r}}}
$$

## 04. Electric Field

A charge produces something called and electric field in the space around it and this electric field exerts a force on any charge placed in it.

NOTE The electric field does not exert force on source charge.

## Electric field Intensity

Force experienced by a unit positive charge placed in and electric field at a point is called electric field intensity at that point. It is also known as electric field simply. Let $\mathrm{q}_{0}$ be the positive test charge placed in an electric field. If $\overrightarrow{\mathrm{F}}$ is the force experienced by this charge, then

$$
\vec{E}=\lim _{\mathrm{q}_{0} \rightarrow 0} \frac{\overrightarrow{\mathrm{~F}}}{\mathrm{q}_{0}}
$$

(i) Unit : N/C or volt/metre
(ii) This is a vector quantity and its direction is the same as force on the positive test charge.

(iii) Since $\vec{E}$ is the force on unit charge, force on charge $q$ is. $\vec{F}=q \vec{E}$.
(iv) Dimension is $\left[\mathrm{M}^{1} \mathrm{~L}^{1} \mathrm{~T}^{-3} \mathrm{~A}^{-1}\right]$
(v) Electric field due to a point charge is

$$
\overrightarrow{\mathrm{E}}=\frac{\mathrm{kq}}{\mathrm{r}^{2}} \cdot \hat{\mathrm{r}}
$$

(vi) Direction of electric field due to positive charge is away from charge while direction of electric field due to negative charge is towards the charge.


## Special point

(a) If $\mathrm{q}_{1}$ and $\mathrm{q}_{2}$ are at a distance r and both have the same type of charge, then the distance ' $d$ ' of the point from $q_{1}$ where electric field is zero is given by $d=\frac{\sqrt{q_{1}} r}{\left(\sqrt{q_{1}}+\sqrt{q_{2}}\right)}$. This point will lie between line joining $\mathrm{q}_{1} \& \mathrm{q}_{2}$.

(b) If $\mathrm{q}_{1}$ and $\mathrm{q}_{2}$ have opposite charges then distance ' d ' of the point ' p ' from $\mathrm{q}_{1}$ where electric field is zero is given by

$$
\mathrm{d}=\frac{\sqrt{\mathrm{q}_{1}} \mathrm{r}}{\sqrt{\mathrm{q}_{1}}-\sqrt{\mathrm{q}_{2}}}, \quad\left[\left|\mathrm{q}_{1}\right|>\left|\mathrm{q}_{2}\right|\right]
$$

(c) There charges $+\mathrm{Q}_{1},+\mathrm{Q}_{2}$ and q are placed on a straight line. If this system of charges is in equillibrium, charge q should be a given

$$
\mathrm{q}=\frac{\mathrm{Q}_{1} \mathrm{Q}_{2}}{\left(\sqrt{\mathrm{Q}_{1}}+\sqrt{\mathrm{Q}_{2}}\right)^{2}}
$$

$\Rightarrow$ For measuring $\overrightarrow{\mathrm{E}}$ practically a test charge ( +ve ) of magnitude much less than the source charge should be used.
$\Rightarrow$ Electric force on a charge in uniform $E$ is constant and hence acceleration is constant, so equations of motion can be used

$$
\text { (acceleration } \mathrm{a}=\frac{\mathrm{qE}}{\mathrm{~m}} \text { ) }
$$

$\Rightarrow$ Electric field due to linear charge distribution (a) Finite wire
(d) Finite wire


$$
\mathrm{E}_{\mathrm{p}}=\frac{2 \mathrm{k} \lambda}{\mathrm{R}} \sin \frac{\alpha}{2}
$$

(e) Infinite wire $\left(\alpha=180^{\circ}\right)$


$$
\mathrm{E}_{\mathrm{P}}=\frac{2 \mathrm{k} \lambda}{\mathrm{R}}
$$

(f) Charged arc


$$
\mathrm{E}_{\mathrm{P}}=\frac{2 \mathrm{k} \lambda}{\mathrm{R}} \sin \left(\frac{\alpha}{2}\right)
$$

(g) Charged ring of radius R

at and axial point $\mathrm{E}_{\mathrm{P}}=\frac{\mathrm{kQx}}{\left(\mathrm{R}^{2}+\mathrm{x}^{2}\right)^{3 / 2}}$;

$$
x \gg R \Rightarrow E_{P}=\frac{k Q}{x^{2}}
$$



$$
\mathrm{x} \ll \mathrm{R} \Rightarrow \mathrm{E}_{\mathrm{P}}=\frac{\mathrm{kqx}}{\mathrm{R}^{2}}
$$

As x is increases: $\overrightarrow{\mathrm{E}}$ due to ring first $\uparrow$ then $\downarrow$ and at $\mathrm{X}=\frac{\mathrm{R}}{\sqrt{2}}$ it is maximum.

## Electric lines of forces : (ELF)

The electric field in a region can be represented by drawing certain curves known as electric lines of force.
An electric line of force is that imaginary smooth curve drawn in an electric field along which a free isolated unit positive charge moves.

## Properties

(i) Imaginary
(ii) Can never cross each other
(iii) Can nerver be closed loops
(iv) The number of lines originating or terminating on a charge is proportional to the magnitude of charge. In rationalised $\mathrm{MKS}\left(\frac{1}{\varepsilon_{0}}\right)$ system electric lines are associated with unit charge, so if a body encloses q , total lines of force associated with it (called flux) will be $\mathrm{q} / \varepsilon_{0}$.
(v) Total lines of force may be fractional as lines of force are imaginary.

(vi) Lines of force ends or starts normally at the surface of a conductor.
(vii) If there is no electric field there will be no lines of force.
(viii) Lines of force per unit area normal to the area at a point represents magnitude of intensity, crowded lines represent strong field while distant weak field.
(ix) Tangent to the line of force at a point in an electric field gives the direction of intensity. So a positive charge free to move follow the line of force.

## 05. Electric Dipole

(i) An system consisting of two equal and opposite charges separated by a small distance is termed and electric dipole.


Example : $\mathrm{Na}^{+} \mathrm{Cl}^{-}, \mathrm{H}^{+} \mathrm{Cl}^{-}$etc.
(ii) An isolated atom is not a dipole because centre of positive charge coincides with centre of negative centres. But if atom is placed in an electric field, then the positive and negative centres are displaced relative to each other and atom become a dipole.
(iii) DIPOLE MOMENT: The product of the magnitude of charges and distance between them is called the dipole moment.
It is denoted by $\vec{p}$ and $|\vec{p}|=q \times 2 \ell$
(a) This is a vector quantity which is directed from negative to positive charge.
(b) Unit : C-m
(c) Dimension : $\left[\mathrm{M}^{0} \mathrm{~L}^{1} \mathrm{~T}^{1} \mathrm{~A}^{1}\right]$

## Electric field due to a dipole

(i) There are two components of electric field at any point
(a) $\mathrm{E}_{\mathrm{r}} \rightarrow$ in the direction of $\overrightarrow{\mathrm{r}}$
(b) $\mathrm{E}_{\theta} \rightarrow$ in the direction perpendicular to $\overrightarrow{\mathrm{r}}$

$$
\begin{aligned}
& \mathrm{E}_{\mathrm{r}}=\frac{1}{4 \pi \varepsilon_{0}} \cdot \frac{2 \mathrm{p} \cos \theta}{\mathrm{r}^{3}} \\
& \mathrm{E}_{\theta}=\frac{1}{4 \pi \varepsilon_{0}} \cdot\left(\frac{\mathrm{p} \sin \theta}{\mathrm{r}^{3}}\right)
\end{aligned}
$$


(i) Resultant

$$
\mathrm{E}=\sqrt{\mathrm{E}_{\mathrm{r}}^{2}+\mathrm{E}_{\theta}^{2}}=\frac{\mathrm{p}}{4 \pi \varepsilon_{0} \mathrm{r}^{3}} \sqrt{1+3 \cos ^{2} \theta}
$$

(ii) Angle between the resultant $\overrightarrow{\mathrm{E}}$ and $\overrightarrow{\mathrm{r}}, \alpha$ given
by $\alpha=\tan ^{-1}\left(\frac{\mathrm{E}_{\theta}}{\mathrm{E}_{\mathrm{r}}}\right)=\tan ^{-1}\left(\frac{1}{2} \tan \theta\right)$
(iii) If $\theta=0$, i.e point is on the axis -

$$
\mathrm{E}_{\mathrm{axis}}=\frac{2 \mathrm{kp}}{\mathrm{r}^{3}} \quad(\mathrm{r} \gg \ell)
$$

(iv) If $\theta=90^{\circ}$, i.e. point is on the line bisecting the dipole perpendicularly

$$
\mathrm{E}_{\text {equatorial }}=\frac{\mathrm{kp}}{\mathrm{r}^{3}} \quad(\mathrm{r} \gg \ell)
$$

(v) $\mathrm{So}, \mathrm{E}_{\text {axis }}=2 \mathrm{E}_{\text {equatorial }}$ (for same r )
(vi) $\quad \mathrm{E}_{\text {axis }}=\frac{1}{4 \pi \varepsilon_{0}} \cdot \frac{2 \mathrm{pr}}{\left(\mathrm{r}^{2}-\ell^{2}\right)^{2}}$

$$
\mathrm{E}_{\text {equatorial }}=\frac{1}{4 \pi \varepsilon_{0}} \cdot \frac{\mathrm{p}}{\left(\mathrm{r}^{2}+\ell^{2}\right)^{3 / 2}}
$$

where $\mathrm{p}=\mathrm{q} \cdot(2 \ell)$
(vii)

potential at a general point.

$$
\mathrm{V}=\frac{1}{4 \pi \varepsilon_{0}} \frac{\mathrm{p} \cos \theta}{\mathrm{r}^{2}}
$$

(viii)If $\theta=0^{\circ}, V_{\text {axis }}=\frac{\mathrm{kp}}{\mathrm{r}^{2}}$
(ix) If $\theta=90^{\circ}, \quad \mathrm{V}_{\text {equator }}=0$
(x) Here we see that $\mathrm{V}=0$ but $\mathrm{E} \neq 0$ for points at equatorial position.
(xi) Again, if $r \gg d$ is not true and $d=2 \ell$,

$$
\begin{aligned}
& \mathrm{V}_{\text {axis }}=\frac{1}{4 \pi \varepsilon_{0}} \cdot \frac{\mathrm{p}}{\left(\mathrm{r}^{2}-\ell^{2}\right)} \\
& \mathrm{V}_{\text {equator }}=0
\end{aligned}
$$

NOTE (i) This is not essential that at a point, where $E=0, V$ will also be zero there eg. inside a uniformly charged sphere, $E=0$ but $V \neq 0$
(ii) Also if $V=0$, it if not essential for $E$ to be zero eg. in equatorial position of dipole $V=0$, but $\mathbf{E} \neq 0$

## Electric Dipole In an Electric Field - Uniform Electric Field

(i) When an electric dipole is placed in a uniform electric dipole, A torque acts on it which subjects the dipole to rotatory motion. This $\tau$ is given by $\tau=\mathrm{pE} \sin \theta$ or $\vec{\tau}=\overrightarrow{\mathrm{p}} \times \overrightarrow{\mathrm{E}}$
(ii) Potential energy of the dipole
$\mathrm{U}=-\mathrm{pE} \cos \theta=-\overrightarrow{\mathrm{p}} \cdot \overrightarrow{\mathrm{E}}$


## Cases

(a) If $\theta=0^{\circ}$, i.e. $\vec{p} \| \vec{E}, \tau=0$ and $U=-\mathrm{pE}$, dipole is in the minimum potential energy state and no torque acting on it and hence it is in the stable equilibium state.
(b) For $\theta=180^{\circ}$, i.e. $\vec{p}$ and $\overrightarrow{\mathrm{E}}$ are in opposite direction, then $\tau=0$ but $U=\mathrm{pE}$ which is maximum potential energy state. Although it is in equilibium but it is not a stable state and a slight perturbation can disturb it.
(c) $\theta=90^{\circ}$, i.e. $\overrightarrow{\mathrm{p}} \perp \overrightarrow{\mathrm{E}}$, then
$\tau=\mathrm{pE}$ (maximum) and $\mathrm{U}=0$

## Work done in rotating an electric dipole in and electric field

(i) To rotate the dipole by an angle $\theta$ from the state of stable equilibrium $\mathrm{W}=\mathrm{PE}$ $(1-\cos \theta)$.
(ii) Work done in rotating the dipole from $\theta_{1}$ to $\theta_{2}$ in an uniform electric filed

$$
\mathrm{W}=\mathrm{pE}\left(\cos \theta_{1}-\cos \theta_{2}\right)
$$

(iii) Work done in rotating the dipole through $180^{\circ}$ from stable equilibrium state

$$
\mathrm{W}=2 \mathrm{pE}=2 \text { (potential energy) }
$$

## 06. Electric Flux

(i) It is denoted by ' $\phi$ '.
(ii) It is a scalar quantity.
(iii) It is defined as the total number of lines of force passing normally through a curved surface placed in the field.
(iv) It is given by the dot product of $\overrightarrow{\mathrm{E}}$ and normal infinitesimal $\overrightarrow{\mathrm{ds}}$ area integrated over a closed surface.

$$
\begin{gathered}
\mathrm{d} \phi=\overrightarrow{\mathrm{E}} \cdot \overrightarrow{\mathrm{ds}} \\
\phi=\oint \overrightarrow{\mathrm{E}} \cdot \mathrm{ds}=\oint \mathrm{Eds} \cos \theta
\end{gathered}
$$

where $\theta=$ angle between electric field and normal to the area

(v) Unit: (a) $\mathrm{N}-\mathrm{m}^{2} / \mathrm{C}$
(b) volt - meter
(vi) Dimension : $\left[\mathrm{ML}^{3} \mathrm{~T}^{-3} \mathrm{~A}^{-1}\right]$
(vii) Flux due to a positive charge goes out of the surface while that due to negative charge comes into the surface.
(viii) Value of electric flux is independent of shape and size of the surface.
(ix) If only a dipole is present in the surface then net flux is zero.
(x) Net flux from a surface is zero does not imply that intensity of electric field is also zero.

## 07. Gauss's Law

This law states that electric flux $\phi_{\mathrm{E}}$ through any closed surface is equal to $1 / \varepsilon_{0}$ times the net charge ' $q$ ' enclosed by the surface i.e

$$
\phi_{\mathrm{E}}=\oint \overrightarrow{\mathrm{E}} \cdot \overrightarrow{\mathrm{~d}} \mathrm{~s}=\frac{\mathrm{q}}{\varepsilon_{0}}
$$

## Important point about flux

(i) Independent of distances between charges inside the surface and their distribution.
(ii) Independent of shape, size and nature of surface.
(iii) Net flux due to a charge outside the surface will be zero.
(iv) Gauss law is valid only for the vector fields which obey inverse square law

Example A hemispherical surface of radius R is kept in a uniform electric field E such that E is parallel to the axis of hemi-sphere, Net flux from the surface will be


Solution

$$
\begin{aligned}
& \phi=\oint \overrightarrow{\mathrm{E}} \cdot \overrightarrow{\mathrm{~d} \mathrm{~s}}=\mathrm{E} \cdot \pi \mathrm{R}^{2} . \\
& =(\mathrm{E})(\text { Area of surface perpendicular to } \mathrm{E}) \\
& =\mathrm{E} \cdot \pi \mathrm{R}^{2} .
\end{aligned}
$$

## 08. Application of Gauss's Law

Electric field due to a solid conducting sphere/Hollow conducting sphere.

(i) Case: $1 \mathrm{r}>\mathrm{R} \quad \overrightarrow{\mathrm{E}}=\frac{\mathrm{kq}}{\mathrm{r}^{2}} \hat{\mathrm{r}}=\frac{1}{\varepsilon_{0}} \frac{\sigma \mathrm{R}^{2}}{\mathrm{r}^{2}} \hat{\mathrm{r}}$

Case: $2 \mathrm{r}=\mathrm{R} \quad \overrightarrow{\mathrm{E}}=\frac{\sigma}{\varepsilon_{0}} \hat{\mathrm{r}}$
Case: $3 \mathrm{r}<\mathrm{R} \quad \overrightarrow{\mathrm{E}}=0$
i.e. At point interior to a conducting or a hollow sphere, electric field intensity is zero.
(ii) For points outside the sphere, it behaves like all the charge is present at the centre.
(iii) Intensity of electric field is maximum at the surface

## Imp.

(iv) Electric field at the surface is always perpendicular to the surface.
(v) For points, near the surface of the conductor, $\mathrm{E}=\frac{\sigma}{\varepsilon_{0}}$ perpendicular to the surface

## Electric field due to infinitely long charge

(i) $\underset{\rightarrow}{A}$ long wire is given a line charge density $\lambda$. If wire is positively charged, direction of $\overrightarrow{\mathrm{E}}$ will be away from the wire (outward $\perp$ ) while fro a negatively charged wire, direction of $\overrightarrow{\mathrm{E}}$ will be (inward $\perp$ ) towards the wirde.
(ii) E at point p

$$
\overrightarrow{\mathrm{E}}=\frac{\lambda}{2 \pi \varepsilon_{0} \mathrm{r}} \hat{\mathrm{r}} \quad \text { or } \quad \mathrm{E}=\frac{\lambda}{2 \pi \varepsilon_{0} \mathrm{r}}
$$


(iii) Potential difference between points $A\left(r_{1}\right) \& B\left(r_{2}\right)=V_{2}-V_{B}=\frac{\lambda}{2 \pi \varepsilon_{0}} \ln \left(\frac{r_{2}}{r_{1}}\right)$
(iv) Potential difference between points $A\left(r_{1}\right) \& B\left(r_{2}\right)=V_{2}-V_{B}=\frac{\lambda}{2 \pi \varepsilon_{0}} \ln \left(\frac{r_{2}}{r_{1}}\right)$

## Electric field at a point due to an infinite sheet of charge

(i) If $\sigma=$ surface charge density. Intensity at points near to the sheet $\overrightarrow{\mathrm{E}}=\frac{\sigma}{2 \varepsilon_{0}} \hat{\mathrm{r}}$

(ii) Direction of electric field is perpendicular to the sheet of charge.
(iii) Intensity of electric field does not depend upon the distance of points from the sheet for the points in front of sheet i.e. There is an equipotential region near the charged sheet.
(iv) Potential difference between two points $A \& B$ at distances $r_{1} \& r_{2}$ respectively is

$$
\mathrm{V}_{\mathrm{A}}-\mathrm{V}_{\mathrm{B}}=\frac{\sigma}{2 \varepsilon_{0}}\left(\mathrm{r}_{2}-\mathrm{r}_{1}\right)
$$

## Electric field due to infinite charged metal plate

(i) Intensity at points near the plate $\overrightarrow{\mathrm{E}}=\frac{\sigma}{\varepsilon_{0}} \hat{\mathrm{r}}$
where $\sigma=$ surface charge density
(ii) $\overrightarrow{\mathrm{E}}$ is independent of distance of the point from the plate and also of the area of sheet i.e. There is an equipotential region near the plate.

(iii) Direction of electric field is perpendicular to the palate.
(iv) Potential difference between two point $A\left(r_{1}\right)$ and $B\left(r_{2}\right)\left(r_{1}<r_{2}\right)$ near the plate is

$$
\Delta \mathrm{V}=\mathrm{V}_{\mathrm{A}}-\mathrm{V}_{\mathrm{B}}=\frac{\sigma}{\varepsilon_{0}}\left(\mathrm{r}_{2}-\mathrm{r}_{1}\right)
$$

Electric field due to charged ring : $\mathbf{Q}$ charge is distributed over a ring of radius $\mathbf{R}$.
(i) Intensity of electric field at a distance x from the centre of ring along it's axis-

$$
\begin{aligned}
& \mathrm{E}=\frac{1}{4 \pi \varepsilon_{0}} \frac{\mathrm{Q} \cos \theta}{\mathrm{r}^{2}}=\frac{\mathrm{Qx}}{4 \pi \varepsilon_{0}{ }^{3}} \quad(\because \cos \theta=\mathrm{x} / \mathrm{r}) \\
& =\frac{1}{4 \pi \varepsilon_{0}} \frac{\mathrm{Qx}}{\left(\mathrm{R}^{2}+\mathrm{x}^{2}\right)^{3 / 2}}\left[\because \mathrm{r}=\sqrt{\mathrm{R}^{2}+\mathrm{x}^{2}}\right]
\end{aligned}
$$

and it's direction will be along the axis of the ring.

(ii) Intensity will be zero at the centre of the ring.
(iii) Intensity will be maximum at a distance $\mathrm{R} / \sqrt{2}$ from the centre and

$$
\mathrm{E}_{\max }=\frac{2}{3 \sqrt{3}} \cdot \frac{1}{4 \pi \varepsilon_{0}} \cdot \frac{\mathrm{Q}}{\mathrm{R}^{2}}
$$

(iv) Electric potential at a distance x from centre,

$$
\mathrm{V}=\frac{1}{4 \pi \varepsilon_{0}} \frac{\mathrm{Q}}{\sqrt{\left(\mathrm{x}^{2}+\mathrm{R}^{2}\right)}}
$$

## NEET <br> Exercise (1)

1. The force between two charges situated in air is $F$, The force between the same charges if the distance between them is reduced to half and they are situated in a medium having dielectric constant 4 is
(a) $F / 4$
(b) $4 F$
(c) $16 F$
(d) $F$
2. If charge $q$ is placed at the centre of the line joining two equal charges $Q$, the system of three charges will be in equilibrium if $q$ is
(a) $-Q / 2$
(b) $-Q / 4$
(c) $-4 Q$
(d) $+Q / 2$
3. Two equally charged identical metal spheres $A$ and $B$ repel each other with a force $3 \times 10^{-5} \mathrm{~N}$. Another identical unchanged sphere $C$ is touched with $A$ and then placed at the mid-point between $A$ and $B$. Net force on $C$ is
(a) $1 \times 10^{-5} \mathrm{~N}$
(b) $2 \times 10^{-5} \mathrm{~N}$
(c) $1.5 \times 10^{-5} \mathrm{~N}$
(d) $3 \times 10^{-5} \mathrm{~N}$
4. The electric strength of air is $2 \times 10^{7} \mathrm{~N} / \mathrm{C}$. The maximum charge that a metallic sphere of diameter 6 mm can hold is
(a) $3 n C$
(b) $20 n C$
(c) $1.5 n C$
(d) $2 n C$
5. An electric dipole is placed at an angle of $60^{\circ}$ with an electric field of intensity $10^{5} \mathrm{NC}^{-1}$. It experiences a torque equal to $8 \sqrt{3} \mathrm{Nm}$. Calculate the charge on the dipole, if the dipole length is 2 cm .
(a) $-8 \times 10^{3} \mathrm{C}$
(b) $8.54 \times 10^{-4} \mathrm{C}$
(c) $8 \times 10^{-3} \mathrm{C}$
(d) $0.85 \times 10^{-6} \mathrm{C}$
6. An electric field exists in the space around a point charge $+Q$. A positive charge $+q$ is carried from $A$ to $B$ and $A$ to $C$, where $B$ and $C$ lie on a circle with $+Q$ at the centre, Work done is


Figure
(a) greater along the path $A C$ than along $A B$
(b) greater along the path $A B$ than along $A C$
(c) same in both the cases
(d) zero in both the cases.
7. An $\alpha$-particle and a proton are accelerated through same potential difference from rest. Find the ratio of their final velocity
(a) $\sqrt{2}: 1$
(b) $1: 1$
(c) $1: \sqrt{2}$
(d) $1: 2$
8. $\mathrm{A}, \mathrm{B}$ and C are three points in a uniform electric field, in Figure. The electric potential is :

(a) Same at all the three points $A, B$ and $C$
(b) Maximum at $A$
(c) Maximum at $B$
(d) Maximum at $C$
9. $n$ small drops of same size are charged to $V$ volt each. If they coalesce to form a single large drop, then the potential will be
(a) $V n$
(b) $V n^{-1}$
(c) $V n^{1 / 3}$
(d) $V n^{2 / 3}$
10. A long string with a charge of $\lambda$ per unit length passes through an imaginary cube of edge $a$. The maximum flux of the electric field through the cube will be
(a) $\lambda a / \in_{0}$
(b) $\sqrt{2} \lambda a / \in_{0}$
(c) $6 \lambda a^{2} / \in_{0}$
(d) $\sqrt{3} \lambda a / \in_{0}$

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## 02 Solid State

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Positive


## 01. The Solid State

The solid are characterized by incompressibility, rigidity and mechanical strength. The molecules, atoms or ions in solids are closely packed i.e they are held together by strong forces and can not move about at random. Thus solids have definite volume, shape, slow definite, low vapour pressure and possesses the unique property of being rigid. Such solids are known as true solids e.g. $\mathrm{NaCl}, \mathrm{KCl}, \mathrm{Sugar}, \mathrm{Ag}, \mathrm{Cu}$ etc. On the other hand the solid which loses shapes on long standing, flows under its own weight and easily distorted by even mild distortion forces are called pseudo solids e.g. glass, plastic etc.
Some solids such as NaCl , Sugar, Sulphur etc. have properties not only of rigidity and incompressibility but also of having typical geometrical forms. These solids are called as crystalline solids. In such solids there is definite arrangements of particles (atoms, ions or molecules) throughout the entire three dimensional network of a crystal. This is named as long-range order. This three dimensional arrangement is called crystal lattice or space lattice. Other solids such as glass, rubber, plastics etc. have rigidity and incompressibility to a certain extent but they do not have definite geometrical forms or do not have long range order are known as amorphous solids.

## 02. Differences Between Crystalline and Amorphous Solids

## (i). Characteristic Geometry

In the crystalline solids the particles (atoms, ions, or molecules are definitely and orderly arranged thus these have characteristic geometry while amorphous solids do not have characteristic geometry.

## Melting Points

A crystalling solids has a sharp melting point i.e. it changes into liquid state at a definite temperature. On the contrary an amorphous solid does not has a sharp melting point.
Cooling curve
Amorphous solids show smooth cooling curve while crystalline solids show two breaks in cooling curve. In the case of crystalline solids two breaks points ' $a$ ' and ' $b$ ' are appear. These points indicate the beginning and the end of the process of crystallization. In this time interval temperature remains constant. This is due to the fact that during crystallization process energy is liberated which compensates for the loss of heat thus the temperature remains constant.


## (i). Isotropy and Anisotropy

Amorphous solids differ from crystalline solids and resemble liquids in many respects. The properties of amorphous solids, such as electrical conductivity, thermal conductivity, mechanical strength, refractive index, coefficient of thermal expansion etc. are same in all directions. Such solids are known as isotropic. Gases and liquids are also isotropic. On the other hand crystalline solids show these physical properties different in different directions. Therefore crystalline solids are called anisotropic.

## 03. Crystalline State

"A crystal is a solid composed of atoms (ions or molecules) arranged in an orderly repetitive array"
"The smallest geometrical position of the crystal which can be used as repetitive unit to build up the whole crystal is called a unit cell." The unit cell should have same symmetry elements as the crystal and there should be no gaps between unit cells.
The angle between the two perpendiculars to the two intersecting faces is termed as the interfacial angle which may be same as the angle between the unit cell edges. Goniometer is used to measure the interfacial angle. It is important to note that interfacial angle of a substance remains the same although its shape may be different due to conditions of formation.


## 04. Types of the Crystals

Crystals are divided into four important types on the basis of chemical bonding of the constituent atoms.
(i) lonic Crystals

These are formed by a combination of highly electro-positive ions (cations) and highly electronegative ions (anions). Thus strong electrostatic force of attraction acts with in the ionic crystals. Therefore, a large amount of energy is required to separate ions from one another.
e.g. $\mathrm{NaCl}, \mathrm{KF}, \mathrm{CsCl}$ etc.


## (ii) Covalent Crystals

These are formed by sharing of valence electrons between two atoms resulting in the formation of a covalent bond. The covalent bonds extend in two or three dimensions forming a giant interlocking structure called network. Diamond and graphite are the good examples of this type.
(iii) Molecular Crystals

In these crystals, molecules occupy the lattice points of the unit cells, except in solidified noble gases in which the units are atoms, where the binding is due to vander Waal's' forces and dipole-dipole forces. Since vander Waal's' forces are non-directional hence structure of the crystal is determined by geometric consideration only. Solid $\mathrm{H}_{2}, \mathrm{~N}_{2}, \mathrm{O}_{2}$, $\mathrm{CO}_{2}, \mathrm{I}_{2}$, sugar etc. are well known examples of such crystal in which vander Waal's' forces are acting.

## (iv) Metallic Crystals

These are formed by a combination of atoms of electropositive elements. These atoms are bound by metallic bonds. It may be defined as:
The force that binds a metal ion to a number of electrons within its sphere of influences is known as metallic bond.

## 05. Isomorphism

The occurrence of a given substance in more than one solid crystalline forms have different physical properties is known as polymorphism. This property when occurs in elements is known as allotropy.
Sometimes we come across examples of chemically different solids which crystalline in the crystalline shape. Such substances are said to be Isomorphous (same shape). Their chemical constitutions are very similar and in some cases crystals of one substance may continue to grow when placed in a saturated solution of the other e.g. potash alum and chrome alum crystals have the same shape and can be grown in each other's solutions.

## 06. SPACE LATTICE/CRYSTALLINE LATTICE/3-D LATTICE

Space lattice is a regular arrangement of lattice points showing how the particles are arranged at different sites in 3D-view.
"The three dimensional distribution of component particles in a crystal can be found by X-ray diffraction of different faces of the crystal. On the basis of the classification of symmetry, the crystals have been divided into seven systems. These can be grouped into 32 classes which in turn can be regrouped into 7 crystal systems. These seven systems with the characteristics of their axes (angles and intercepts) along with some examples of each are given in the following table.

## The Seven Crystal System

| Name of system |  | Axes | Angles | Bravais Lattices |
| :---: | :---: | :---: | :---: | :---: |
| 1. | Cubic <br> [Isometric] | $\mathrm{a}=\mathrm{b}=\mathrm{c}$ | $\alpha=\beta=\Upsilon=90^{\circ}$ | Primitive, Face-centred, Body centred $=3$ |
| 2. | Tetragonal | $\mathrm{a}=\mathrm{b} \neq \mathrm{c}$ | $\alpha=\beta=\Upsilon=90^{\circ}$ | Primitive, Body centred $=2$ |
| 3. | Rhombohedral or Trigonal | $\mathrm{a}=\mathrm{b}=\mathrm{c}$ | $\alpha=\beta=\Upsilon \neq 90^{\circ}$ | Primitive $=1$ |
| 4. | Orthorhombic or Rhombic | $\mathrm{a} \neq \mathrm{b} \neq \mathrm{c}$ | $\alpha=\beta=\Upsilon=90^{\circ}$ | Primitive, Face-centred, Body centred End centred $=4$ |
| 5. | Monoclinic | $\mathrm{a} \neq \mathrm{b} \neq \mathrm{c}$ | $\begin{aligned} & \alpha=\Upsilon=90^{\circ} ; \\ & \beta \neq 90^{\circ} \end{aligned}$ | Primitive, End - centred $=2$ |
| 6. | Triclinic | $\mathrm{a} \neq \mathrm{b} \neq \mathrm{c}$ | $\alpha \neq \beta \neq \Upsilon \neq 90^{\circ}$ | Primitive $=1$ |
| 7. | Hexagonal | $\mathrm{a}=\mathrm{b} \neq \mathrm{c}$ | $\begin{gathered} \alpha=\beta=90^{\circ} \\ \Upsilon=120^{\circ} \end{gathered}$ | $\begin{aligned} & \text { Primitive }=1 \\ & \text { Total }=14 \end{aligned}$ |

## 07. Types of Unit Cells

There are four types of unit cells (among the cubic system).
(a) Simple or primitive. Particles are only at the comers of the unit cell.

(b) Face-centred cubic unit cell. A particle is present at the centre of each face of the unit cell.

(c) End-face-centred cubic unit cell. A particle is present at each comer plus particles are present on the end faces of the unit cell.

(d) Body-centred cubic unit cell. In addition to the particles at the corners, there is one particle in the body-centre of the unit cell.


## 08. Coordination Number

If atoms are represented by spheres, the number of spheres which are touching a particular sphere is called coordination number of that sphere. In ionic crystals, the coordination number may be defined as the number of the oppositely charged ions surrounding a particular ion.

## 09. The number of particles per Unit Cell

(i) An atom at the corner is shared by 8 unit cells. Therefore, its contribution is $=1 / 8$.
(ii) An atom at the face is shared by 2 unit cells. Therefore, its contribution is $=1 / 2$.
(iii) An atom present in the body is unshared. Therefore, its contribution is 1 .
(iv) An atom present on the edge is shared by four unit cells. Therefore, its contribution is $=$ $1 / 4$. Thus, in
(v) a primitive or simple cubic unit cell, the total number of particles is $1 / 8 \times 8=1$.
(vi) a face-centred cubic unit cell; the total number of particles is $1 / 8 \times 8+1 / 2 \times 6=4$.
(vii) a body-centred cubic unit cell, the total number of particles is $1 / 8 \times 8+1=2$.

The number of atoms per unit cell are in the same ratio as the stoichiometry of the compound or the ratio of the number of particles $A$ and $B$ present per unit cell gives the formula of the compound formed between A and B .

## 10. LENGTH OF FACE DIAGONAL AND CUBE DIAGONAL



Consider the triangle ABC , with the help of pythagoras theorem $\mathrm{AC}=\sqrt{\mathrm{AB}^{2}+\mathrm{BC}^{2}}=\sqrt{\mathrm{a}^{2}+\mathrm{a}^{2}}=\sqrt{2} \mathrm{a} \quad$ (length of face diagonal.)

Consider the triangle DAC, with the help of pythagoras theorem
$\mathrm{DC}=\sqrt{\mathrm{DA}^{2}+\mathrm{AC}^{2}}=\sqrt{\mathrm{a}^{2}+(\sqrt{2} \mathrm{a})^{2}}=\sqrt{3} \mathrm{a} \quad$ (length of cube diagonal)

## TYPES OF UNIT CELL (BRAVAIS LATTICE)

The distance between successive lattice planes of the same type is called the spacing of planes or inter planar distance between the planes. On the basis of this aspect, the lattices may be divided in following classes.

## 11. Simple/Primitive/Basic Unit cell

A unit cell having lattice point only at corners called as primitive or simple unit cell. i.e. in this case there is one atom at each of the eight corners of the unit cell considering an atom at one corner as the centre, it will be found that this atom is surrounded by six equidistant neighbours (atoms) and thus the co-ordination number will be six. If ' $a$ ' is the side of the unit cell, then the distance between the nearest neighbours shall be equal to ' $a$ '.

(a) Relationship between edge length ' $a$ ' and atomic radius ' $r$ ':


$$
\left.\mathrm{a}=2 \mathrm{r} \quad \text { i.e. } \mathrm{r}=\frac{\mathrm{a}}{2} \text { (One face of } \mathrm{SCC}\right)
$$

(b) Number of atoms present in unit cell

In this case one atom or ion lies at the each corner. Hence simple cubic unit cell contains a total of $\frac{1}{8} \times 8=1$ atom or ion/unit cell.
(c) Packing efficiency(P. E.)

$$
\begin{aligned}
& \text { P.E. }=\quad \begin{array}{l}
\text { Volume occupied by atoms present in unit cell } \\
\quad \text { Volume of unit cell }
\end{array} \\
& =\frac{\mathrm{n} \times \frac{4}{3} \pi \mathrm{r}^{3}}{\mathrm{~V}}\left[\because \text { Volume of atom }=\frac{4}{3} \pi \mathrm{r}^{3}\right] \\
& \text { For SCC }: \quad \text { P.E. }=\frac{1 \times \frac{4}{3} \times \pi \times\left(\frac{\mathrm{a}}{2}\right)^{3}}{\mathrm{a}^{3}} \quad\left[\because \mathrm{r}=\frac{\mathrm{a}}{2} \text { and } \mathrm{V}=\mathrm{a}^{3}, \mathrm{n}=1\right] \\
& =\frac{\pi}{6}=0.524 \quad \text { or } 52.4 \%
\end{aligned}
$$

## 12. Body centred cubic (b.c.c.) cell:

A unit cell having lattice point at the body centre in addition to the lattice point at every corner is called as body centered unit cell.
Here the central atom is surrounded by eight equidistant atoms and hence the co-ordination number is eight. The nearest distance between two atoms will be $\frac{a \sqrt{3}}{2}$

(a) Relationship between edge length ' $a$ ' and atomic radius ' $r$ ' :


In BCC, along cube diagonal all atoms touches each other and the length of cube diagonal is $\sqrt{3} \mathrm{a}$
So, $\sqrt{3} \mathrm{a}=4 \mathrm{r} \quad$ i.e. $\mathrm{r}=\frac{\sqrt{3} \mathrm{a}}{4}$
(b) Number of atom present in unit cell:
$\left(\frac{1}{8} \times 8\right)+(1 \times 1)=1+1=2$ atoms/unit cell
In this case one atom or ion lies at the each corner of the cube. Thus contribution of the 8 corners is $\left(\frac{1}{8} \times 8\right)=1$, while that of thee body centred is 1 in the unit cell. Hence total number of atoms per unit cell is $1+1=2$ atoms (or ions)
(c) Packing efficiency:
P.E. $=\frac{\mathrm{n} \times \frac{4}{3} \pi \mathrm{r}^{3}}{\mathrm{~V}}=\frac{2 \times \frac{4}{3} \times \pi\left(\frac{\sqrt{3} \mathrm{a}}{4}\right)^{3}}{\mathrm{a}^{3}}=\frac{\sqrt{3} \pi}{8}=0.68 \quad\left[\because \mathrm{n}=2, r=\frac{\sqrt{3} \mathrm{a}}{4}, V=\mathrm{a}^{3}\right]$

In B.C.C. $68 \%$ of total volume is occupied by atom or ions.

## 13. Face centred cubic (f.c.c.) cell:

A unit cell having lattice point at every face centre in addition to the lattice point at every corner called as face centred unit cell. i.e. in this case there are eight atoms at the eight corners of the unit cell and six atoms at the centre of six faces. Considering an atom at the face centre as origin, it will be found that this face is common to two cubes and there are twelve points surrounding it situated at a distance which is equal to half the face diagonal of the unit cell. Thus the co-ordination number will be twelve and the distance between the two nearest atoms will be $\frac{\mathrm{a}}{\sqrt{2}}$.

(a) Relationship between edge length ' $a$ ' and atomic radius ' $r$ ' :

In FCC, along the face diagonal all atoms touches each other and the length of face diagonal is $\quad \sqrt{2} \mathrm{a}$.
So $\quad 4 \mathrm{r}=\sqrt{2} \mathrm{a} \quad$ i.e. $\mathrm{r}=\frac{\sqrt{2} \mathrm{a}}{4}=\frac{\mathrm{a}}{2 \sqrt{2}}, \quad \mathrm{r}=\frac{2}{2 \sqrt{2}}$

(b) Number of atoms per unit cell :
$\left(\frac{1}{8} \times 8\right)+\left(6 \times \frac{1}{2}\right)=1+3=4$ atoms /unit cell
In this case one atom or ion lies at the each corner of the cube and one atom or ion lies at the centre of each face of the cube. It may noted that only $\frac{1}{2}$ of each face sphere lie with in the unit cell and there are six such faces. The total contribution of 8 corners is $\left(\frac{1}{8} \times 8\right)=1$, while that of 6 face centred atoms is $\left(\frac{1}{2} \times 6\right)=3$ in the unit cell.
Hence total number of atoms per unit cell is $1+3=4$ atoms (or ions).
(c) Packing efficiency :

$$
\begin{aligned}
& \text { P.E. }=\frac{\mathrm{n} \times \frac{4}{3} \pi \mathrm{r}^{3}}{\mathrm{~V}} \quad\left[\because \text { for } \mathrm{FCC} \mathrm{n}=4, \mathrm{r}=\frac{\mathrm{a}}{2 \sqrt{2}}, \mathrm{~V}=\mathrm{a}^{3}\right] \\
& =\frac{4 \times \frac{4}{3} \pi \times\left(\frac{\mathrm{a}}{2 \sqrt{2}}\right)^{3}}{\mathrm{a}^{3}}=\frac{\pi}{3 \sqrt{2}}=0.74 \text { or } 74 \%
\end{aligned}
$$

i.e. In FCC, $74 \%$ of total volume is occupied by atoms.

## 14. End Centered Unit Cell:

A unit cell having lattice point at the centres of only. one set of opposite faces in addition to the lattice point at every corner called as end centered unit cell.

NOTE This type of Bravais lattice is obtained only in orthorhombic and monoclinic type unit call.

## 15. CRYSTAL DENSITY OF THE CRYSTAL

If the length of edge of the unit cell is known we can calculate the density of the crystal as follow: Let length of edge of the unit cell be $\ell$.
$\therefore$ Volume of the unit cell $=\ell^{3}=\mathrm{V} \mathrm{cm}^{3}$
Density of the unit cell $=\frac{\text { Mass of unit cell }}{\text { Volume of unit cell }}$
Mass of the unit cell $=$ Number of atoms present in a unit cell $\times$ Mass of one atom $=$ $\mathrm{n} \times \mathrm{m} \mathrm{g}$

But mass of one atom $(\mathrm{m})=\frac{\text { Atomic mass }}{\text { Avogadro Number }}=\frac{\mathrm{M}}{\mathrm{N}_{\mathrm{A}}}$
Mass of the unit cell $=\mathrm{n} \times \frac{\mathrm{M}}{\mathrm{N}_{\mathrm{A}}} \mathrm{g}$
Density of the unit cell $=\frac{\mathrm{n} \times \frac{\mathrm{M}}{\mathrm{N}_{\mathrm{A}}}}{\mathrm{V}} \mathrm{gm} \mathrm{cm}^{-3}$

Density of the unit cell $=\frac{\mathrm{n} \times \mathrm{M}}{\mathrm{V} \times \mathrm{N}_{\mathrm{A}}} \mathrm{g} \mathrm{cm}^{-3}=$ Crystal density $(p)$

Example An element (atomic mass $=60$ ) having face centred cubic crystal has a density of $6.23 \mathrm{~g} \mathrm{~cm}^{-3}$. What is the edge length of the unit cell (Avogadro constant, $\mathrm{N}_{\mathrm{A}}=$ $6.02 \times 10^{23} \mathrm{~mol}^{-1}$ ).
Solution

$$
\begin{array}{rlrl}
\text { Density } & =\frac{Z \times M}{a^{3} \times N_{A}} & \mathrm{Z}=4 \text { (for } f c c \text { ) } \\
& & \\
6.23 & =\frac{4 \times 60}{a^{3} \times 6.022 \times 10^{23}} & a^{3}=\frac{4 \times 60}{6.23 \times 6.022 \times 10^{23}} \\
& =64 \times 10^{-24} & & \\
a & =4 \times 10^{-8} \mathrm{~cm} \\
& =\mathbf{4} \times \mathbf{1 0}^{-\mathbf{1 0}} \mathbf{m}=\mathbf{4 0 0} \mathbf{~ p m .} & &
\end{array}
$$

## 16. Close packing

It is type of packing with maximum packing fraction \& space utilization. The constituent particles of a solid are like hard spheres. These spheres can pack in space in various manner to form a packing. To clearly understand the packing of these spheres, the packing can be categorised as :
(i) Close packing in one dimension.
(ii) Close packing in two dimension.
(iii) Close packing in three dimension.

## 17. Close packing in one dimension :

In one dimension, only one arrangement of spheres is possible as shown in fig.


Close packing of spheres in one dimension

## Two dimensional packing of spheres :

Two possible types of two dimensional packing are.
(i) Square close packing in two dimension.
(ii) Hexagonal close packing in two dimension.

## 18. Two dimensional square packing of spheres :

When two rows are placed in such a manner, that spheres of one row are placed immediately below of the other, the resulting packing is called two dimensional square close packing.
(i) Since all the rows are identical the packing is called AAA $\qquad$ type packing.
(ii) Each sphere is touched by four other hence coordination number is four.
(iii) Of centres of spheres are connected, square cells are formed, hence also called two dimensional square packing.
(iv) This type of packing is not very effective in terms of utilization of space.
(v) Packing efficiency in $2-\mathrm{D}=\frac{1 \times \pi \mathrm{r}^{2}}{\mathrm{a}^{2}}=\frac{1 \times \pi(\mathrm{a} / 2)^{2}}{\mathrm{a}^{2}}=\frac{\pi}{4}=0.74$.
(vi) Packing efficiency in $3-\mathrm{D}=\frac{1 \times \frac{4}{3} \pi\left(\frac{\mathrm{a}}{2}\right)^{3}}{\mathrm{a}^{3}}=0.52 \quad$ [In 3-D its unit cell is simple cubic]


## 19. Two dimensional hexagonal packing :

If various one dimensional close pack rows are placed in such a way that spheres of top row fits in depression of bottom row spheres, the resulting packing is called two dimensional close packing structure.
(i) Every third row sphere comes exactly at top of first row sphere, hence the packing is called ABABAB packing.
(ii) If centres are joined, hexagonal unit cells are formed. Hence this is called two dimensional hexagonal close packing.
(iii) This packing is most efficient in utilizing space in two dimensional arrangement.
(iv) Each sphere is touched by six other, hence coordination number is six.
(v) Packing efficiency in $2-\mathrm{D}=\frac{3 \times \pi\left(\frac{\mathrm{a}}{2}\right)^{2}}{\frac{\mathrm{a}^{3} \sqrt{3}}{4} \times 6}=\frac{\pi}{2 \sqrt{3}}=0.91$
(vi) Packing efficiency in $3-\mathrm{D}=\frac{3 \times \frac{4}{3} \pi\left(\frac{\mathrm{a}}{2}\right)^{3}}{\frac{\mathrm{a}^{2} \sqrt{3}}{4} \times 6 \times \mathrm{a}}=\frac{\pi}{3 \sqrt{3}}=0.60$


## 20. Close packing in three dimensions :

When two dimensional packing structure are arranged one above the other, depending upon type of two dimensional arrangement in a layer, and the relative positions of spheres in above or below layer, various types of three dimensional packing results. To define 3-D lattice six lattice parameters are required -3 edge lengths \& 3 angles.
(i) Simple cubic packing (A A A A)
(ii) Hexagonal close packing ( AB AB AB .....)
(iii) Cubic close packing or face centered cubic (... ABC ABC...)

## 21. Simple cubic packing in three dimension :

The two dimensional square close packed layer are placed, in such a manner that spheres in each layer comes immediately on top of below layer, simple cubic packing results. Important points.
(i) Atoms all aligned vertically and horizontally in all directions.
(ii) The unit cell for this packing is simple cubic unit cell.
(iii) In this packing, only $52 \%$ of available space is occupied by atoms.


## 22. Three Dimensionally close packing :

In hexagonal close packing, there are two types of the voids (open space or space left) which are divided into two sets ' $b$ ' and ' $c$ ' for convenience. The spaces marked ' $c$ ' are curved triangular spaces with tips pointing upwards whereas spaces marked ' $b$ ' are curved triangular spaces with tips pointing downwards. Now we extend the arrangement of spheres in three dimensions by placing second close packed layer (hexagonal close packing) (B) on the first layer (A). The spheres of second layer may placed either on space denoted by ' $b$ ' or ' $c$ '. It may be noted that it is not possible to place sphered on both types of voids (i.e. b and c). Thus half of the voids remain unoccupied by the second layer. The second layer also have voids of the types ' $b$ ' and in order to build up the third layer,
there are following two ways :


## 23. Hexagonal close packing (HCP) AB-AB-AB

## Type :

(i) In one way, the spheres of the third layer lie on spaces of second layer (B) in such a way that they lie directly above those in the first layer (A). In other words we can say that the layer becomes identical to the first layer. If this arrangement is continued idefinitely in the same order this represented as A BA BA B .....
This type of arrangement represent hexagonal close packing (hcp) symmetry (or structure), which means that the whole structure has only one 6 -fold axis of symmetry i.e. the crystal has same appearance on rotation through an angle of $60^{\circ}$.
(ii) Every third layer sphere lies on top of first layer sphere. (ABABAB $\qquad$ packing)
(iii) Maximum possible spade is occupied by spheres.
(iv) Each sphere is touched by 12 other spheres in 3D ( 6 is one layer, 3 in top layer and 3 in bottom.)
(v) The unit cell for hexagonal close packing is hexagonal unit cell.
(vi) For every atom forming hcp there are effectively two tetra voids and one octa void. That why this generate ABAB ..... AB pattern. One type of void always remain on unoccupied.
Unit cell : $\mathrm{a}=2 \mathrm{r}=\mathrm{b} ; \quad \Upsilon=120$

(vii) Packing efficiency of HCP units

Relation between $\mathbf{a}, \mathrm{b}, \mathrm{c}$ and $\mathbf{R}$ :
$\mathrm{a}=\mathrm{b}=2 \mathrm{R}$

$\tan 30=\frac{\mathrm{a}}{2 \times \mathrm{y}}$
so

$$
\mathrm{y}=\frac{\mathrm{a} \times \sqrt{3}}{2 \times 1}=\frac{\sqrt{3}}{2} \mathrm{a}
$$

Area $=6\left[\frac{1 \mathrm{a}}{2} \times \frac{\sqrt{3} \mathrm{a}}{2}\right]=\frac{6 \sqrt{3} \mathrm{a}^{2}}{4}$

## Calculation of c :

$\cos 30^{\circ}=\frac{\mathrm{a}}{2 \times \mathrm{x}}$

$$
\mathrm{x}=\frac{2 \mathrm{a}}{2 \times \sqrt{3}}=\frac{\mathrm{a}}{\sqrt{3}}
$$



Applying pythogoras theorem : $\mathrm{x}^{2}+\mathrm{h}^{2}=\mathrm{a}^{2}$
So $\quad h^{2}=a^{2}-x^{2}=a^{2}-\frac{a^{2}}{3}=\frac{2}{3} a^{2}$
$\mathrm{h}=\frac{2}{\sqrt{3}} \mathrm{a} \quad$ so $\quad \mathrm{c}=2 \mathrm{~h}=2 \sqrt{\frac{2}{3}} \mathrm{a}$

## 24. $\mathrm{ABC}-\mathrm{ABC}$ arrangement (Cubic close packing (FCC)

In second way, the spheres of the third layer (C) lie on the second layer (B) in such a way that they lie over the unoccupied spaces ' C ' of the first layer( A ). If this arrangement is continuous in the same order this is represented as $A B C$ ABC $A B C \ldots$. . This type of arrangement represent cubic close packed (ccp) structure. This structure has 3-fold axes of symmetry which pass though the diagonal of the cube. since in this system, there is a sphere at the centre if each face of the unit cell and hence this structure is also known as face-centred cubic (fcc) structure. It may be noted that in ccp (or fcc) structures each sphere is surrounded by 12 spheres hence the coordination number of each sphere is 12 . The spheres occupy $74 \%$ of the total volume and $26 \%$ of is the empty space in both (hcp and ccp) structure.


- III ${ }^{\text {rd }}$ layer will be different from $\mathrm{I}^{\text {st }}$ layer as well as $\mathrm{II}^{\text {nd }}$ layer.
- It is also known as cubical close packing (CCP), unit cell chosen is face centered unit cell (FCC).


## Relation between ' $a$ ' and ' $R$ ' :

$\mathrm{a} \neq 2 \mathrm{R} \quad \sqrt{2} \mathrm{a}=4 \mathrm{R}$ (Sphere are touching along the face diagonal)
Effective no. of atoms per unit cell $(\mathbf{Z})=Z=\frac{1}{8} \times 8+\frac{1}{2} \times 6=4$

## Packing fraction

P.F. $=\frac{4 \times \frac{4}{3} \pi \mathrm{R}^{3}}{4 \times 4 \times 4 \mathrm{R}^{3}} \times \sqrt{2} \times 2=\frac{\pi}{3 \sqrt{2}}=0.74(74 \%)$

Coordination number,$(\mathbf{C N})=12$
Density (d) $=\frac{\mathrm{Z} \times \mathrm{M}}{\mathrm{N}_{\mathrm{A}} \cdot \mathrm{a}^{3}}$

## INTERSTICES OR VOIDS OR HOLES IN CRYSTALS

It has been shown that the particles are closely packed in the crystals even than there is some empty space left in between the spheres. This is known as interstices (or interstitial site or hole empty space or voids). In three dimentional close packing (CCP \& HCP) the interstices are of two types : (i) tetrahedral interstices and (ii) octahedral interstices.


## 25. Tetrahedral Interstices :

We have seen that in hexagonal close packing (hcp) and cubic close packing (ccp) each sphere of second layer touches with three spheres of first layer. Thus they, leave a small space in between which is known as tetrahedral site or interstices. or The vacant space between 4 touching spheres is called as tetrahedral void. Since a sphere touches three spheres in the below layer and three spheres in the above layer hence there are two tetrahedral sites associated with one sphere. It may by noted that a tetrahedral site does not mean that the site is tetrahedral in geometry but it means that this site is surrounded by four spheres and the centres of these four spheres lie at the apices of a regular tetrahedron.


In FCC, one corner and its three face centred atom of faces meeting at that corner form a tetrahedral void.
In FCC, two tetrahedral voids are obtained along one cube diagonal. So in FCC 8 tetrahedral voids are present.
In FCC total number of atoms $=4$
In FCC total number of tetrahedral voids $=8$
So, we can say that, in 3D close packing 2 tetrahedral voids are attached with one atom.


## 26. Octahedral - Interstices :

Hexagonal close packing (hcp) and cubic close packing (ccp) also form another type of interstices (or site) which is called octahedral site (or interstices). or The vacant space between 6 touching spheres is called as octahedral void.
In the figure two layers of close packed spheres are shown. The spheres of first layer are shown by full circles while that of second layer by dotted circles. Two triangles are drawn by joining the centres of three touching spheres of both the layers.
In FCC, 6 face centres form a octahedral void at body centre.
The apices of these triangles point are in opposite directions. On super imposing these triangles on one another a octahedral site is created. It may be noted that an octahedral site does not mean that the hole is octahedral in shape but it means that this site is surrounded by six nearest neighbour lattice points arranged octahedrally.


## 27. Positions of Tetrahedral Voids in an fcc Unit Cell :

The centre of tetrahedral void is located on the centre of body diagonal of each small cube of volume $\left(\frac{a^{3}}{8}\right)$.
Total number of atoms per unit cell $=\frac{1}{2} \times 6+8 \times \frac{1}{8}=4$
Total number of tetra void $=8$
Effective number of tetra void per atom forming close pack=2.
If a fcc unit cell is divided into eight equal mini-cubes (edge $=a / 2$ ) then centre of tetra voids lies at the centres of each of there mini-cubes. In FCC center of tetra void is at perpendicular distance of a/4 from nearest face.

## 28. Positions of Octahedral Voids in an fcc Unit Cell :

Position of octa void is at mid-point of each edge (total 12 edges in a cube) and at the centre of cube. Each octa void located at mid point of edge contributes $1 / 4$ to the unit cell. The octa void situated at the centre contributes 1 .
In FCC, total number of octahedral voids are

$$
(1 \times 1) \quad+\quad\left(12 \times \frac{1}{4}\right)=1+3=4
$$

(Cube centre) (edge)
In FCC, number of atoms $=4$
In FCC, number of octahedral voids $=4$
So, we can say that, in any type of close packing one octahedral void is attached with one atom.

## 29. Limiting Radius Ratios

An ionic crystal contains a large number of cations and anions. Generally cations are smaller in size than anions. The cations are surrounded by anions and they touch each other. These ions are arranged in space in such a way to produce maximum stability. The stability of the ionic crystal may be described in terms of radius ratio i.e. the ratio of the radius of cation (r) to that of anion ( R ) is $(\mathrm{r} / \mathrm{R})$. The range of $(\mathrm{r} / \mathrm{R})$ may be expressed as limiting radius ratio. This value is important to determine the arrangement of the ion in different types of crystals.

Limiting radius ratio for various types of sites

| Limiting radius <br> ratio $=\mathbf{r} / \mathbf{R}$ | Coordination <br> Number of cation | Structural Arrangement <br> (Geometry of voids) | Example |
| :---: | :---: | :--- | :--- |
| $0.155-0.225$ | 3 | Plane Trigonal | Boron Oxide |
| $0.225-0.414$ | 4 | Tetrahedral | $\mathrm{ZnS}, \mathrm{SiO}_{2}$ |
| $0.414-0.732$ | 4 | Square planar | .- |
| $0.414-0.732$ | 6 | Octahedral | $\mathrm{NaCl}, \mathrm{MgO}_{2}$ |
| $0.732-1.000$ | 8 | Cubic | CsCl |

Example $\quad$ A solid $\mathrm{A}^{+} \mathrm{B}^{-}$has NaCl type close packed structure. If the anion has a radius of 250 pm , what should be the ideal radius for the cation ? Can a cation $\mathrm{C}^{+}$having a radius of 180 pm be slipped into the tetrahedral site of the crystal $\mathrm{A}^{+} \mathrm{B}^{-}$? Give reason for tour answer.
Solution $\quad \mathrm{NaCl}=$ FCC Closed packed structure

$$
\frac{r_{+}\left(C^{+}\right)}{r_{-}\left(B^{-}\right)}=\frac{180 p m}{250 p m}=0.72
$$

It does not lie in the range $0.225-0.414$. Hence, $\mathrm{C}^{+}$cannot be slipped into the tetrahedral site.

## 30. STRUCTURE of $\mathbf{N a C l}$ :

The bigger $\mathrm{Cl}^{-}$forms cubic close packing and small $\mathrm{Na}+$ occupy positions of all octahedral voids. The radius ratio $\frac{\mathrm{r}^{+}}{\mathrm{r}^{-}}$lie in the range $0.414-0.73$.
(i) Each $\mathrm{Na}^{+}$is surrounded by six $\mathrm{Cl}^{-}$is surrounded by six $\mathrm{Na}^{+}$ion. [6:6 coordination]
(ii) Total no. of $\mathrm{Na}^{+}$and $\mathrm{Cl}^{-}$in each limit cell is 4 .
(iii) Number of formula units of NaCl per unit cell is equal to 4 .
(iv) The density of NaCl crystal is given by $\mathrm{d}=\left(\frac{4 \times \mathrm{M}_{\mathrm{NaCl}}}{\mathrm{N}_{\mathrm{A}} \times \mathrm{a}^{3}}\right)$ $\mathrm{N}_{\mathrm{A}}=$ Avogadro's number $; \quad \mathrm{a}=$ Edge length The edge length of NaCl unit cell is given by $\left(2 \mathrm{r}^{+}+2 \mathrm{r}^{-}\right) \Rightarrow \frac{\mathrm{a}}{2}=\mathrm{r}^{+}+\mathrm{r}^{-}$(FCC \& Octa void)

## 31. Zinc Blende (Sphalerite) Structure :

Larger anions form ccp arrangement and smaller cations filling half of alternate tetrahedral voids.
(i) C.N. of $\mathrm{Zn}^{2+}=4 \quad$; C.N. of $\mathrm{S}^{2-}=4 \quad[4: 4$ coordination $]$
(ii) Formula units of ${ }^{\mathcal{F}}$ per unit cell $=4$.
(iii) $\mathrm{d}_{\mathrm{ZnS}}=\frac{4 \times \mathrm{M}_{\mathrm{ZnS}}}{\mathrm{N}_{\mathrm{A}} \times \mathrm{a}^{3}}$
(iv) $\mathrm{r}_{\mathrm{Zn}^{2+}}+\mathrm{r}_{\mathrm{S}^{2-}}=\frac{\mathrm{a} \sqrt{3}}{4}$

## 32. Cesium Halide Structure :

$(\mathrm{CsCl}) \mathrm{Cl}^{-}$at the corners of cube and $\mathrm{Cs}^{+}$in the center.
(i) C.N. of $\mathrm{Cs}^{+}=8 \quad ; \quad$ C.N. of $\mathrm{Cl}^{-}=8 \quad[8: 8$ coordination $]$
(ii) Formula units of CsCl per cube $=1$
(iii) $\mathrm{d}_{\mathrm{CsCl}}=\frac{\mathrm{M}_{\mathrm{CsCl}}}{\mathrm{N}_{\mathrm{A}} \times \mathrm{a}^{3}}$
(iv) $\mathrm{r}_{\mathrm{Cs}+}=\mathrm{r}_{\mathrm{Cs}-}=\frac{\mathrm{a} \sqrt{3}}{2} \Rightarrow \mathrm{r}^{+}+\mathrm{r}^{-}=\frac{\mathrm{a} \sqrt{3}}{4}$ (FCC \& tetra void)

## 33. Fluorite Structure :

$\left(\mathrm{CaF}_{2}\right) \mathrm{Ca}^{2+}$ forming ccp arrangement and $\mathrm{F}^{-}$filling all tetrahedral voids.
(i) C.N. of $\mathrm{F}^{-}=4 \quad ; \quad$ C.N. of $\mathrm{Cs}=8 \quad[8: 4$ coordination $]$
(ii) Formula units of $\mathrm{CaF}_{2}$ per unit cell $=4$
(iii) $\mathrm{d}_{\mathrm{CaF}_{2}}=\frac{4 \times \mathrm{M}_{\mathrm{CaF}_{2}}}{\mathrm{~N}_{\mathrm{A}} \times \mathrm{a}^{3}}$
(iv) $\mathrm{r}_{\mathrm{Ca}^{2+}}+\mathrm{r}_{\mathrm{F}^{-}}=\frac{\mathrm{a} \sqrt{3}}{4}$

## 34. Imperfections In Solid

Any deviation from the perfectly ordered arrangement constitutes a defect or imperfection. These defects sometimes called thermodynamic defect because the number of these defects depends on the temperature. Crystals may also possess additional defect due to the presence of impurities. Imperfection not only modify the properties of solids but also give rise to new properties.

## 35. Stoichiometric Defects

Those compounds in which the number of positive and negative ions are exactly in the ratio indicated by their chemical formula are called stoichiometric compounds. eg NaCl (1:1). These solids show following types of defects :

## 36. Schottky Defect :

This type of defect is created when same number of positive ion and negative ion are missing from their respective positions leaving behind a pair of holes. Schottky defect is more common in ionic compounds with high co-ordination number and where the size of positive and negative ions are almost equal. The presence of large number of schottky defects in crystal results in significant decrease in its density. $\mathrm{Eg} . \mathrm{NaCl}, \mathrm{KCl}, \mathrm{CsCl}, \mathrm{KBr}$ etc.


## 37. Interstitial Defect :

This type of defect is caused due to the presence of ions in the normally vacant interstitial sites in the crystal.

## 38. Frenkel Defect :

This type of defect is created when an ion leaves its correct lattice site and occupies an interstitial site. Frenkel defects are common in ionic compounds which have low co-ordination number and in which there is large difference in size between positive and negative ions. Eg. $\mathrm{ZnS}, \mathrm{AgCl}, \mathrm{AgBr}, \mathrm{Ag}$ etc.

## 39. Non Stoichiometric Defect Frenkel Defect :

There are many compound in which the ratio of positive and negative ions present in the compound differs from that required by ideal formula of the compound. Such compounds are called Non stoichiometric compounds. eg. $\mathrm{VO}_{\mathrm{x}}$ (Where x can vary between 0.6 to 1.3.) In these compounds balance of positive and negative charges is maintained by having extra electrons or extra positive charge. So, these defects are of following types:

## 40. Metal Excess Defects Due to Anion Vacancies

A compound may have excess metal ion if a negative ion is absent from its lattice site, leaving a hole which is occupied by electron to maintain electrical neutrality. The holes occupied by electrons are called F-centres and are responsible for the colour if the compound. Eg.

- The excess sodium in NaCl makes the crystal appears yellow.
- Excess potassium in KCl makes it voilet.
- Excess lithium in LiCl makes it Pink.
- Greater the number of F-centres greater is the intensity of colour. This type of defects are found in crystal which are likely to possess schottky defects.



## 41. Metal Excess Defects Due to Interstitial Cations

It may occur if an extra positive ion is present in an interstitial site, Electrically neutrality is maintained by the presence of an electron in the interstitial site. This type of defects are exhibited by the crystals which are likely to exhibit "Frenkel defects". Eg. Yellow colour of ZnS .


## 42. Metal Deficiency Due to Cation Vacancies

The non-stoichiometric compound may have metal deficiency due to the absence of a metal ion from its lattice site. The charge is balanced by an adjacent ion having higher positive charge. This type of defects are generally shown by compounds of transition metals.


Example Analysis shows that nickel oxide has the formula $\mathrm{Ni}_{0.98} \mathrm{O}$. What fractions of the nickel exist as $\mathrm{Ni}^{2+}$ and $\mathrm{Ni}^{3+}$ ?
Solution Let $\mathrm{Ni}^{2+}$ be $x$ so that $\mathrm{Ni}^{3+}$ will be $0.98-\mathrm{x}$. Total charge on the compound must be zero so that
$+2 x+3(0.98-x)-2=0$
$2 x+2.94-3 x-2=0$

$$
-x=-0.94
$$

or

$$
x=0.94
$$

$$
\begin{array}{r}
\% \text { of } \mathrm{Ni}^{2+}=\frac{0.94}{0.98} \times 100=96 \% \\
\% \text { of } \mathbf{N i}^{\mathbf{3 +}}=\mathbf{4 \%}
\end{array}
$$

## NEET Pattern Exercise (1)

Q1. A metal crystallizes with a face-centered cubic lattice. The edge of the unit cell is 408 pm . The diameter of the metal atom is
(a) 144 pm
(b) 204 pm
(c) 288 pm
(d) 408 pm

Q2. Percentage of free space in a body centred cubic unit cell is
(a) $28 \%$
(b) $30 \%$
(c) $32 \%$
(d) $34 \%$

Q3. If a is the length of the side of a cube, the distance between the body centered atom and one corner atom in the cube will be
(a) $\frac{2}{\sqrt{3}}$ a
(c) $\frac{\sqrt{3}}{4}$ a
(b) $\frac{4}{\sqrt{3}}$ a
(d) $\frac{\sqrt{3}}{2} \mathrm{a}$

Q4. Structure of a mixed oxide is cubic close packed (ccp). The cubic unit cell of mixed oxide is composed of oxide ions. One fourth of the tetrahedral voids are occupied by divalent metal A and the octahedral voids are occupied by a monovalent metal B . The formula of the oxide is
(a) $\mathrm{ABO}_{2}$
(b) $\mathrm{A}_{2} \mathrm{BO}_{2}$
(c) $\mathrm{A}_{2} \mathrm{~B}_{3} \mathrm{O}_{4}$
(d) $\mathrm{AB}_{2} \mathrm{O}_{2}$

Q5. The number of octahedral $\operatorname{void}(\mathrm{s})$ per atom present in a cubic close-packed structure is
(a) 2
(b) 4
(c) 1
(d) 3

Q6. Lithium metal crystallizes in a body centred cubic crystal. If the length of the side of the unit cell of lithium is 351 pm , the atomic radius of the lithium will nearly be
(a) 152 pm
(b) 75 pm
(c) 300 pm
(d) 240 pm

Q7. AB crystallizes in a body centred cubic lattice with edge length 'a' equal to 387 pm . The distance between two oppositively charged ions in the lattice is
(a) 335 pm
(b) 250 pm
(c) 200 pm
(d) 300 pm

Q8. If ' $a$ ' stands for the edge length of the cubic systems : simple cubic, body centred cubic and face centred cubic, then the ratio of radii of the spheres in these systems will be respectively
(a) 1a: $\sqrt{3} \mathrm{a}: \sqrt{2} \mathrm{a}$
(b) $\frac{1}{2} \mathrm{a}: \frac{\sqrt{3}}{4} \mathrm{a}: \frac{1}{2 \sqrt{2}} \mathrm{a}$
(c) $\frac{1}{2} \mathrm{a}: \sqrt{3} \mathrm{a}: \frac{1}{\sqrt{2}} \mathrm{a}$
(d) $\frac{1}{2} \mathrm{a}: \frac{\sqrt{3}}{2} \mathrm{a}: \frac{\sqrt{2}}{2} \mathrm{a}$

Q9. A given metal crystallizes out with a cubic structure having edge length of 361 pm . If there are four metal atoms in one unit cell, what is the radius of one atom?
(a) 108 pm
(b) 40 pm
(c) 127 pm
(d) 80 pm

Q10. The packing efficiency of two-dimensional square unit cell shown below is

(a) $39.27 \%$
(b) $68.02 \%$
(c) $74.05 \%$
(d) $78.54 \%$

## ANSWER

| Q1 | Q2 | Q3 | Q4 | Q5 |
| :--- | :--- | :--- | :--- | :--- |
| (c) | (c) | (d) | (d) | (c) |
| Q6 | Q7 | Q8 | Q9 | Q10 |
| (a) | (a) | (b) | (c) | (d) |

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## Class 12 | Biology

## 03 Reproduction in Organisms

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## 01. Introduction

A vast number of plant and animal species have existed on the earth for several thousand of years. The process in living organisms that ensures this continuity is Reproduction. Reproduction is one of the most characteristic feature of living organisms. Life will not exist on the earth if plants and animals do not reproduce to make offsprings.

## 02. Life span

Life span can be defined as the period from birth to the natural death of an organism. It can vary from as short as few days to as long as a number of years.
Maximum Life Span : Maximum life span is the maximum number of years survived or the greatest age reached by any member of a species. The average life span refers to the average number of years survived or age reached by the members of a population. The maximum life span of a domestic dog is about 20 years and that of a laboratory mouse is 4.5 years. The maximum life span of humans has been estimated to be about 121 years. This rests on the fact that a man in Japan, Shirechiyo lzymi, reached the age of 120 years and 237 days in 1986. He died after developing pneumonia. Average life span and life expectancy of humans have grown dramatically. In general the rate of mortality of humans has gone down and the life span has increased. It is 56 in India whereas in the United State, it is 78.

## 03. Reproduction

Reproduction is the means of self perpetuation of a race in which new, young, similar looking individuals are formed by the grown up or adult individuals. The adults which give rise to young ones are called parents.

## Functions of Reproduction :

(i) It replaces the individuals dying due to senescence or ageing.
(ii) Individuals removed from population due to predation or disease are replaced through reproduction.
(iii) It introduces variations essential for adaptability and struggle for existence.

## Basic Features of Reproduction :

(i) Replication of DNA.
(ii) Division of cells. It may or may not involve meiosis.
(iii) Growth due to synthesis of more protoplasm.
(iv) Formation of reproductive units.
(v) Elaboration and development of reproductive units to form new young individuals.

## 04. Types of Reproduction

Broadly speaking, there are two types of reproduction, asexual and sexual. Asexual reproduction does not involve gamete formation and fusion. It is uniparental. On the other hand, sexual reproduction consists of formation and fusion of gametes of opposite sexes. It is mostly biparental with two types o parents of different sexes but can be single/uniparental also, as in case of bisexual or hermaphrodite animals.

## I. Asexual Reproduction

It is the mode of reproduction in which new individuals develop directly from specialised or unspecialised parts of a single parent without involving fusion of gametes or sex cells. Asexual reproduction occurs in both single celled and multicelled individuals. The parent individual splits, buds or fragments to form identical daughter cells or individuals, e.g., Amoeba, Paramoecium, Euglena (acellular protists), Sycon, Hydra, Tubularia, Planaria, Ascidia (metazoans). Asexual reproduction is also called agamogenesis or agamogeny. In this mode of reproduction, somatic cells undergo mitosis during the formation of a new individual. Therefore, it is also called somatogenic reproduction. Young ones resulting from asexual reproduction are exactly identical with the parent except in size and are called clones. Each individual of a clone is referred to as a ramet.

Asexual reproduction occurs by fission, budding and fragmentation.
(A) Fission : It is a mode of asexual reproduction in which the body of a mature individual divides into two or more similar and equal sized daughter individuals. Fission can be binary fission or multiple fission.
(a) Binary Fission : It is the division of the body of an individual into two equal halves, each of which functions as an independent daughter individual. In unicellular organisms, binary fission is accompanied by mitotic division of nucleus followed by cytokinesis. In metazoans. The organisms which undergo binary fission seldom die of senescence or old age because as soon as they mature, they divide into two daughters. They are, therefore, nearly immortal. Depending on the plane of division, binary fission is of the following types:
(i) Simple Binary Fission (Irregular Binary Fission) : Division can occur through any plane e.g., Amoeba.
(ii) Longitudinal Binary Fission : The plane of fission passes along the longitudinal axis of the organism, e.g., Euglena, Vorticella.
(iii) Oblique Binary Fission : The plane of binary fission lies at an angle to the transverse axis e.g., Ceratium, Gonyaulax.
(iv) Transverse Binary Fission : The plane of binary fission runs along the transverse axis of the individual, e.g., Paramoecium, diatoms, bacteria. In Paramoecium, transverse binary fission is preceeded by a mitotic division of meganucleus and mitotic division of micronucleus. In it, binary fission produces two dissimilar daughters, one proter (anterior) and the other opisthe (posterior). Both develop the deficient components and become similar.
(b) Multiple Fission : The nucleus divides several times by amitosis to produce many nuclei, without involving any cytokinesis. Later, each nucleus gathers a small amount of cytoplasm around it and the mother individual splits into many tiny daughter cells (e.g., Amoeba, Plasmodium, Monocystis, etc). In course of time, each of these daughter cells starts a free life and transforms into an adult individual. This kind of fission is called multiple fission.
Cyst formation : In response to unfavourable living conditions, an Amoeba withdrawn its pseudopodia and secretes a three-layered hard covering or cyst around itself. This phenomenon is termed as encystation. During favourable conditions, the encysted Amoeba divides by multiple fission and produces many minute amoubae or pseudopodiospores; the cyst wall bursts out and the spores are liberated in the surrounding medium to grow up into many Amoebae. This phenomenon is known as sporulation. Acellular protists like sporozoans (e.g., Monocystis, Plasmodium, etc.) typically exhibit sporulation in their life cycles.
(B) Budding : In budding, new individuals are formed by mitosis. Initially, a small outgrowth of the parent's body develops into a miniature individual. It then separates from the mother to lead a free life (e.g., Hydra). This type of budding is known as exogenous budding. Sometimes, the buds do not get separated from the mother individual and form a colony. For example, in Obelia, the colony consists of a number of individuals jor zooids that perform different functions. In all fresh water sponges (e.g., Spongilla) and some marine sponges (e.g., Sycon), the parent individual releases a specialized mass of cells enclosed in a common opaque envelope, called the gemmule. On germination, each gemmule gives rise to offspring and the archeocytes present in it give rise to various cells of the body of sponge as they are totipotent. Gemmules are thought to be internal buds.
(C) Fragmentation : The body of the parent breaks into distinct pieces, each of which can produce an offspring (e.g., Hydra, some marine worms, sea-stars).

## Advantages of Asexual Reproduction :

(a) It is uniparental.
(b) It is a rapid mode of reproduction.
(c) The young ones are exact replicas of their parent.
(d) Asexual reproduction is simpler than sexual reproduction.

## Disadvantages of Asexual Reproduction :

(a) As there is rapid multiplication, a large number of young ones are formed which causes overcrowding.
(b) There is no mixing of genetic material, so no new combination or variation takes place.
(c) There is no crossing over, hence hew linkages are not formed.
(d) It has no role in evolution.
(e) Adaptability to changes in environment is low due to absence of new variations.

## 05. Sexual Reproduction

Sexual reproduction involves formation and fusion of gametes to form the zygote which develops to form a new organism.

## Characteristics :

(a) Two fusing gametes can be produced by same individual or different individuals.
(b) Offsprings produced are not identical to parents.
(c) It involves meiosis and syngamy (fusion or gametes).
(d) It is a slow, elaborate or complex process, so multiplication is not so rapid.

## 06. Phases in Life Cycle

(a) Juvenile phase
(b) Reproductive phase
(c) Senescent phase
(a) Juvenile phase/Pre-reproductive phase : During this phase organism will show growth so that it can attain certain maturity to perform the sexual reproduction. This phase is known as vegetative phase in plants. It is of variable durations in different organisms.
(b) Reproductive phase : Reproductive organs develop and mature during this phase. In the higher plants (Angiosperms). end of juvenile phase or onset of reproductive phase is easily marked. In the higher plants during this phase, there is formation of reproductive structures i.e., flowers.
(c) Senescent phase : It is a post-reproductive phase. It involves structural and functional deterioration of body by accumulation of waste metabolites which ultimately leads to death.

## 07. Events in Sexual Reproduction

After attainment of maturity, all sexually reproducing organisms exhibit events and processes that have remarkable fundamental similarity, even though the structures associated with sexual reproduction are indeed very different. These sequential events may be grouped into three distinct stages, namely, the pre-fertilization, fertilization and the post-fertilization events.

## A. Pre-fertilization Events

These are events in sexual reproduction which occur prior to the process of fertilization. The two main pre-fertilization events are gametogenesis and gamete transfer.
(a) Gametogenesis : It refers to the process of formation of gametes - male and female. Categories of Gametes :
(i) Isogametes : When the fusing gametes are morphologically similar they are known as isogametes or homogametes. They are produced in some algae and fungi.

- Algae : Cladophora, Chlamydomonas debaryana, Ulothrix
- Fungi : Synchytrium, Rhizopus
(ii) Heterogametes : When the fusing gametes are morphologically distinct types, they are known as heterogametes. It is the feature of majority of sexually reproducing organisms. e.g.
(a) Algae : Fucus, Volvox, Chara
(b) All Bryophytes, Pteridophytes, Gymnosperms and Angiosperms. In such organisms, male gamete is called antherozoid or sperm and the female gamete is called egg or ovum

Cell Division During Gamete Formation : Gametes are always haploid i.e., they possess only one set of chromosomes or genome though the parent body producing gametes may be either haploid or diploid. As gametes are always haploid so surely in haploid parent, gametes are produced by mitotic division.
In plants belonging to group pteridophytes, gymnosperms and angiosperms and animals the parental body is diploid. Here reductional division occurs before or at the time of gamete formation. The cells which undergo meiosis ar called meiocyte. If meiocyte is indulged in gamete formation, then it is called gamete mother cell.
In haploid organisms, gametes are produced through mitosis but you must not think that meiosis never occurs in life cycle of haploid organisms. This could be made clear from what you have learnt in previous classes. In these organisms like haploid algae and some fungi, meiosis occurs in zygote or zygospore which is called zygotic meiosis.

## 08. Sexuality in Organisms :

Lower Organisms : In most of the lower sexually reproducing organisms, two fusing gametes are morphologically similar. If these gametes belong to the same parent then such organisms are called homothallic, e.g., fungi (Mucor mucedo). When these gametes belong to different parents then these organisms are called heterothallic.

Higher Organisms : In higher plants there are well-developed sex organs and there is clear distinction between male and female sex organs. Angiosperms possess flowers as reproductive structures. The male sex organ is called stamen and female sex organ is carpel or pistil. If male and female sex organs occur in the same flower then these plants are called bisexual, e.g., China rose. If flowers possess only stamen or carpel then these plants are called unisexual. When male flower (staminate) and female flower (pistillate) are present on same plant body such plants are monoecious, e.g., cucurbits, coconut and maize. However, if they are present on separate plant body then these plants are known as dioecious, e.g., date palm and papaya.

## 09. Gamete Transfer :

After the formation of male and female gametes, compatible gametes must be physically brought together to facilitate fusion (fertilisation or syngamy). In few fungi and algae, both types of gametes are motile. But in majority of organisms male gamete is motile and the female gamete is non-motile. So there is a need of a medium through which the male gametes move.
In seed plants both male and female gametes are non-motile. Here pollen grains are the carrier of male gametes and ovule has the egg. As the male gamete is non-motile so it cannot swim through water medium to reach female gamete rather pollen tube serve this purpose.
For this pollen grain produced in anther ( $\widehat{\delta}$ part) are transferred to the stigma of female organ i.e., carpel through the process of pollination. Pollination is of two type i.e., self pollination and cross pollination. Self pollination is the transfer of the pollen grains from anther of a flower to the stigma of same flower or different flower of the same plant. Cross pollination is transfer of the pollen grain from anther of one flower to the stigma of different flower of other plant.

## 10. Fertilization

The most vital event of sexual reproduction is the fusion of gametes. This process is called syngamy or fertilization which results in the formation of a diploid zygote.
(a) External fertilization : Syngamy occurs outside the body of organism in external medium (water). It is shown by majority of aquatic organisms like most of algae, fishes as well as amphibians.
(b) Internal fertilization : Syngamy occurs inside the body of organisms. It is present in majority of plants like bryophytes, pteridophytes, gymnosperms and angiosperms. It occurs in few algae like spirogyra. In all these organisms egg is formed inside the female body where syngamy occurs.

## 11. Post-Fertilization Events

Events in sexual reproduction after the formation of zygote are called post-fertilization events. Zygote : It is the first cell of the new generation in all sexually reproducing organisms. Zygote is always diploid. It is formed in the external aquatic medium in those organisms which perform external fertilization. Zygote is produced inside the body in cases where fertilization in internal.
In many algae and fungi, the zygote secretes a thick wall that is resistant to desiccation and damage, which help organisms to tide over unfavourable conditions. During unfavourable conditions it undergoes a period of rest until a swing back to sustainability occurs.
Embryogenesis : Embryogenesis is the process of development of embryo from zygote. Embryo is a multicellular stage in the life cycle of a plant or animal prior to formation of an independent individual.

In embryogenesis, the zygote undergoes repeated cell divisions through mitosis. Cell differentiation occurs at specific locations resulting in production of different tissues, organs and organ systems. Development of different external and internal structures is called morphogenesis. Embryo formation is present in all plant groups, except algae. In flowering plants, zygote develops into embryo. The food for development of embryo comes from a special tissue known as endosperm. Ultimately, the fertilized ovule matures into a seed. Inside the mature seed is the progenitor of the next generation, the embryo. A number of seeds develop in an ovary depending upon the number of ovules. Meanwhile, wall of the ovary also proliferates. It produces pericarp or fruit wall. The pericarp can be dry or fleshy. The ripened ovary with pericarp and seeds is called fruit.

## Differences between Asexual and Sexual Reproduction

## Asexual Reproduction

i. New individuals are formed from a single parent.
ii. Asexual reproduction does not require the production of sex organs.
iii. It does not involve meiosis. All divisions are mitotic.
iv. Asexual reproduction does not involve fusion of cells or gametes.
v. New individual develops from one cell or a part of one parent.
vi. New individuals are genetically similar to the parents.
vii. It does not introduce variability. Hence, asexual reproduction has no evolutionary importance.
viii. It is quick method of multiplication.
ix. It is simple process.

## Sexual Reproduction

i. Commonly two parents are involved in the formation of new individuals through sexual reproduction.
ii. Formation of sex organs is a pre-requisite for sexual reproduction.
iii. Sexual reproduction involves meiosis at one or the other stage. In higher plants, it occurs at the time of spore formation or sporogenesis.
iv. It involves fusion of gametes.
v. New individual develops from zygote i.e., fusion product of two gametes.
vi. Offspring or new individuals are genetically different from either of the two parents.
vii. It introduces variability and is, hence of evolutionary importance.
viii. Sexual reproduction is a slower method of multiplication.
ix. It is elaborate or complex process.

## NEET Pattern Exercise (1)

1. Which one of the following shows isogamy with non-flagellate gametes?
(a) Sargassum
(b) Ectocarpus
(c) Ulothrix
(d) Spirogyra
2. Select the wrong statement.
(a) Isogametes are similar in structure, function and behaviour
(b) Anisogametes differ either in structure, function and behaviour
(c) In oomycetes female gamete is smaller and motile, while male gamete is larger and non-motile
(d) Chlamydomonas exhibits both isogamy and anisogamy and Fucus shows oogamy
3. Product of sexual reproduction generally generates
(a) longer viability of seeds
(b) prolonged dormancy
(c) new genetic combination leading to variation
(d) large biomass
4. Which one of the following is correctly matched?
(a) Onion
Bulb
(b) Ginger - Sucker
(c) Chlamydomonas - Conidia
(d) Yeast - Zoospores
5. The 'Eyes' of the potato tuber are
(a) flower buds
(b) shoot buds
(c) axillary buds
(d) root buds
6. Why is vivipary an undesirable character for annual crop plants?
(a) It reduces the vigour of the plant
(b) It adversely affects the fertility of the plant
(c) The seeds exhibit long dormancy
(d) the seeds cannot be stored under normal conditions for the next season
7. During regeneration modification of an organ to other organ is known as
(a) morphogenesis
(b) cpimorphosis
(c) morphallcixis
(d) acretopmaruy growth
8. The process of series of changes from larva to adult after embryonic development is called
(a) regeneration
(b) metamorphosis
(c) growth
(d) ageing
9. A population of genetically identical individuals, obtained from asexual reproduction is
(a) callus
(b) clone
(c) deme
(d) aggregate
10. New banana plants develop from
(a) rhizome
(b) sucker
(c) stolon
(d) seed

## : <br> Answer \& Solution

1. (d)

Isogamy with non-flagellated of gametes is seen is Spirogyra can reproduce both by sexual and asexual (vegetative) means.


They reproduce sexually by conjugation in which two non-flagellated morphologically similar but physiologically different gemetes (isogamous) fuse together. One filament acts as male like gamete and passes through the conjugation tube of another filament which acts as female like gamete.
2. (c)

Oomycetes include water moulds, white rusts and downy mildews. In these female gamete is larger and non-motile, wherease, male gamete is smaller and motile. whereas, male gamete is algae like Uiothrix, Chalmydomonas, Spirogyra etc., which are similar in structure, function and behaviour. Anisogametes are found in Chlamydomonas in which one gamete is larger and non-motile and the other one is motile egg with motile sperm. The gametes, differ both morphologically as well as physiologically It occurs in chalamydomonas, Fucus, Chara, volvox etc.
3. (c)

Sexual reproduction leads to new genetic combination leading to variation in new products. Longer viability of seeds prolonged dormancy and large biomass are not related to sexual reproduction.
4. (a)

The correctly matched pair to onion-bulb. Onion is a simple tunicated layered bulb while ginger is a straggling rhizome having uniparous cyme reproduces by budding and Chlamydomonas by zoospores,
5. (c)

Tuber is oval or spherical swollen underground modified stem lacking adventitious roots. It possesses a number of spirally arranged depression called eyes. Each eye represents node and consists of 1-3 axillary bus in the axils of small scally leaves.
6. (d)

Vivipary is the condition when seeds germinate on the plant, It an undesirable character for annual crop plants because germinated seeds cannot be stored under normal condition for the next season.
7. (b)

Epimorphosis is the replacement of a lost organ of the body by proliferating new cells from the surface of the wound or injured part. Morphogenesis (Gr. morhe $=$ form and genesis $=$ origin) is the growth, shaping and arrangement of body parts according to genetically predefined patterns. The extent direction and rate of morphogenesis depend on genetic controls and environmental factors.
8. (b)

Matamorphosis (meta $=$ change, morphe $=$ form $)$ is a process by which an animal undergoes a comparatively rapid change from larval to adult form Regeneration is regrowth of the part of body which has been removed due to the injury or other causes Growth is an increase in dry mass of an organism. Ageing is progressive deteriortion in activity of cell tissues organs, etc.
9. (b)

Clone refers to the population of genetically identical individuals obtained from asexual reproduction or produced vegetatively from single organism. An individual member of a clone is called ramete.
10.(b)

New banana plants develop through sucker. Sucker is the sub-aerial modification of stem which originates from the basal and underground portion of main stem. It also occurs in mint, Chrysanthemum, etc

