Respiration is a biochemical process by which organic compounds are oxidized to liberate chemical energy from food in step-wise process. The organic compounds are carbohydrates, fats and proteins and the energy released is stored as the ATP molecules.

Cellular respiration is the use of oxygen and production of carbon dioxide at cellular level.

**TYPES OF RESPIRATION**

1. **Anaerobic respiration**
   - When food is oxidized without the use of molecular oxygen, it is called anaerobic respiration. The organism undergoing this type of respiration are termed as anaerobes.
   - Examples are anaerobic bacteria, yeasts, many parasitic animals such as Taenia, Fasciola and Ascaris. In microorganisms, this respiration is termed as fermentation and this is termed after the name of the product they form, such as alcoholic fermentation and lactic acid fermentation.
     - Alcoholic fermentation occurs in yeasts, where they oxidize glucose to ethyl alcohol and carbon dioxide:
       \[ C_6H_{12}O_6 \rightarrow 2C_2H_5OH + 2CO_2 + \text{energy} \]
     - Lactic acid fermentation occurs in some bacteria where glucose is metabolized to lactic acid.
     - Anaerobic respiration occurs in cytoplasm and provides less energy (2ATP molecules).
     - In muscles and erythrocytes, glucose is metabolized to form lactic acid which enters the blood and reaches the liver, where it is converted to glycogen aerobically for further reuse. Accumulation of lactic acid in muscles causes fatigue.

2. **Aerobic respiration**
   - When oxygen is used for the oxidation of food it is termed as aerobic respiration and the organisms undergoing this process are termed as aerobes. It is a high energy yielding process. It is of two types:
     a) Direct respiration: It is the exchange of environmental molecular oxygen with the carbon dioxide of the body cells without any special respiratory organ and blood. It is found in aerobic bacteria, protists, plants, sponges, coelenterates, flatworms, roundworms and most arthropods.
     b) Indirect respiration: In this, exchange of gases takes place through special respiratory organs such as skin, gills, bucco-pharyngeal cavity and lungs. It needs blood for transporting oxygen and carbon dioxide after the exchange.
       - The respiration through organs are termed according to their names. Examples skin – cutaneous gills- brachial, buccopharyngeal cavity – buccopharyngeal and lung-pulmonary respiration.
The indirect respiration occurs in two phases: external respiration and internal respiration. These are preceded by a preliminary phase called breathing (ventilation).

Aerobic respiration occurs both in cytoplasm (glycolysis) and in mitochondria (Krebs cycle and electron transport chain) and provides much more energy (38 ATP molecules).

Breathing refers to the movement that sends fresh air to the respiratory organs and removes foul air from them.

External respiration: It is the intake of oxygen by the blood from air or water in the respiratory organs and the elimination of carbon dioxide.

Internal respiration—It involves 4 processes:
- Uptake of oxygen by tissue cells from blood via blood tissue.
- Oxidation of food in the tissue cells by the action of oxidizing enzymes producing carbon dioxide, water, and energy. This is also termed as cell respiration.
- Storage of energy from oxidation in the phosphate bonds of ATP.
- Release of carbon dioxide by tissue cells into the blood via tissue fluid.

RESPIRATORY SURFACE

- The surface at which exchange of gases (CO₂ and O₂) occurs is termed respiratory surface. This surface must have enough area of gas exchange to meet the metabolic needs of the organism.
- For the exchange to be efficient, respiratory surface should have the following features.
  i) It should be thin, large, and moist.
  ii) It should be permeable to respiratory gases.
  iii) It should be highly vascular.
  iv) It must be directly or indirectly in contact with the source of oxygen.

RESPIRATORY STRUCTURE FOR THE EXCHANGE OF GASES IN DIFFERENT GROUPS OF ANIMALS

1. Protozoans (e.g. amoeba, paramecium): Plasma membrane
2. Sponges (e.g. Sycon): Cell’s plasma membrane
3. Cnidarians (e.g. Hydra): Body surface
4. Platyhelminthes
   i) Free living (planaria): Body surface
   ii) Parasites (e.g. Tapeworm): No exchange of gases
5. Nemathelminthes
   i) Free living (e.g. Rhabditis): Body surface
   ii) Parasites (e.g. Ascaris): No exchange of gases
6. Annelids (e.g. Earthworm): Skin (cutaneous respiration)
7. Arthropods
   i) Prawn, crayfish : Gills (Branchial respiration)
   ii) Insects, centipedes, millipedes, ticks: Tracheae (Tracheal respiration)
   iii) Scorpions, Spiders : Book lungs
   iv) King crab (Limulus) Book gills

8. Molluscs
   i) Unio : Two ctenidia (gills)
   ii) Pila : One ctenidium (gill) and one pulmonary sac (lung)

9. Echinoderms (e.g. starfish) : Dermal branchiae, tube feet.

10. Hemichordata (e.g. Balanoglossus): Pharyngeal wall.

11. Chordata
   i) Urochordata (e.g. Herdmania): Pharyngeal wall
   ii) Cephalochordata (e.g. Branchiostoma) : Pharyngeal wall
   iii) Vertebrata
        a) Cyclostomes, fish : gills
        b) Amphibians : Skin, lungs, buccopharyngeal lining.
        c) Reptiles, birds mammals : Lungs.

PROBLEMS OF WATER – BREATHING

- Water-breathing animals face the following problems in ventilating their respiratory surfaces (gills)
  - Water contains much less oxygen than air.
  - Oxygen diffuses through water far more slowly than through air. Therefore, a large quantity of water is required to be passes over the gills to fulfill the oxygen need.
  - Water is absorb 800 times denser than air so that the fish has make a great muscular effort to maintain water flow.
  - At a higher temperature an animal needs more supply of oxygen because rise in temperature increases their metabolic rate, but less oxygen is available to them in warmer water holds less oxygen.

PROBLEMS OF AIR BREATHING.

- Land animals for breathing air face following difficulties:
  - They have to protect their respiratory surface from drying out.
  - Respiratory surface must be kept moist as gases pass through liquid medium.
  - They lose their precious water through evaporation from their respiratory surface.
HUMAN RESPIRATORY SYSTEM.

- The human respiratory system is divided into upper respiratory tract and lower respiratory tract. The upper respiratory tract includes external nostrils, nasal passage, internal nostril and pharynx. The lower respiratory system includes larynx, trachea, bronchi and bronchioles.

- The special mammalian features of respiratory system are:
  1. Presence of nose
  2. Elongation of nasal passage and its complete separation from buccal passage through palate. So, that internal nostrils open deep into nasopharyngeal part of pharynx.
  3. Long wind pipe due to presence of well defined neck
  4. Spongy, solid lungs

EXTERNAL NARES (NOSTRILS)

They are a pair of slit-like opening present on the lower end of nose.

NASAL CAVITY

It occurs between palate and cranium. Nasal cavity is divisible into two nasal chambers by a nasal septum. Each nasal chamber has three parts.

a) Vestibule
   It is a lower smaller part just above external naris which is lined by skin and bears hair as well as oil glands. Hair help in filtering out dust particles from incoming air.

b) Conditioner (Respiratory region)
   It is middle part of nasal chamber. There are three bony projections called nasal conchae or turbinates (superior, middle, inferior) and some sinuses (maxillary, frontal, sphenoid and ethmoid)
   - The conditioner part is reddish pinkish in coloured by ciliated pseudostratified columnar epithelium with mucous and serous glands. The inhaled air is moistened warmed and cleaned.

c) Olfactory region
   Upper part of nasal chamber and superior nasal concha are yellowish brown. They are covered by olfactory epithelium which perceives sensation of smell.

INTERNAL NARES (CHOANAE)

The two nasal chambers open into nasopharynx through internal nares or choanae.
PHARYNX

- Nasopharynx occurs at the base of skull and has lining of ciliated stratified squamous epithelium.
- Nasopharynx leads to oropharynx or common pathway of respiratory and digestive system.
- Oropharynx passes into laryngopharynx which contains epiglottis and passes into larynx.

LARYNX

- Larynx or voice box opens into laryngo-pharynx through a slit-like glottis which can be widened by intrinsic muscles. Glottis can be closed by a large leaf-like cartilaginous flap called epiglottis.
- Larynx has C-shaped thyroid cartilage (on sides and in front where it can be felt as Adam’s apple), a pair of triangular arytenoids (arytaenoid) cartilages (on back), a ring like cricoids cartilages and a pair of nodule like cartilages of santorini (upper end of arytenoids cartilages). Internally larynx has ciliated columnar mucous epithelium and a pair of vocal cord (attached to thyroid and arytenoids cartilages).
- Vocal cords become thickened in adult males. Vocal cord are shorter and thinner in women produced by passage of air between vocal cords and modulations created by tongue, teeth, tips and nasal cavity.

TRACHEA (WIND PIPE)

- It is 10-12 cm long tube with 2-3 cm diameter which arises from larynx and passes upto middle of thorax. Trachea is supported by 16-20 C-shaped incompletely cartilaginous rings and lined by ciliated pseudostratified mucous epithelium.

BRONCHI

- Trachea divides into right and left primary bronchi. Left bronchus is about 5 cm long while right bronchus is only 2.5 cm long. Right bronchus almost directly enters the right lung. Infection of right lung is more common due to this.
- Inside the lung, the primary bronchus divides into secondary bronchi, secondary bronchi into segmental bronchi and latter into bronchioles. All bronchi are lined by ciliated and mucus secreting pseudostratified epithelium and supported by incomplete cartilaginous rings.
- Bronchioles divides into terminal bronchioles, respiratory bronchioles, alveolar ducts, air sacs and alveoli. Mucus secreting cells are absent from terminal bronchioles and their branches. Epithelium is ciliated in bronchioles and terminal bronchioles. It is non-ciliated in respiratory bronchioles and their branches.
RESPIRATORY GAS EXCHANGE

LUNGS

- A pair of conical spongy elastic lunges of pinkish to slate grey colour occur inside air tight thoracic cavity. A small space called mediastinum lies in between the two lungs (especially due to concavity called cardiac notch of left lump). It encloses heart. Each lung is covered by a pleural sac made of an outer parietal pleuron in contact with wall of thoracic cavity and inner visceral pleura in contact with the surface of lung. A narrow pleural cavity (0.02 mm) occurs between them. It contains pleural fluid. It allows frictionless sliding of pleura during inspiration and expiration. Protection and moistening of lungs are also provided. Pleurisy is painful infection involving inflammation of pleura and over-production of pleural fluid. Normally pleural fluid is under negative pressure due to its formation from the membranous covering.

- Left lung is slightly narrower and longer than the right one. Right lung has three lobes – right superior, right middle and right inferior. Left lung has two lobes – left superior and left inferior. It contains a cardiac notch in antero-median region for accommodating heart. Each lobe is divided internally into segments and segments into lobules. A lobule receives a terminal bronchiolo.

- Terminal bronchiolo produces a few respiratory bronchiolo produces a few respiratory bronchiol. A respiratory bronchiolo give rise to 2 – 11 alveolar sac or infundibulum. The latter has a number of small pouches named alveoli or air sacs. Number of alveoli in human pulmonary system is 300 – 400 million with a surface area 100 m². Each alveolus is polyhedral in outline with a thin wall made of non-ciliated squamous epithelium with a few cubical cells that secrete a lipoprotein surfactant to prevent collapse and sticking of alveolar walls during expiration. Life span of epithelial cells is about 3 weeks so that alveolar wall is being continuously replace. Blood capillaries occur on the surface of alveoli for gaseous exchange.

DIAPHRAGM

- It is a membranous musculo – tendinous partition between thorax and abdomen. Normally it is convex with convexity towards thorax.

MUSCLES

- Phrenic muscles attach diaphragm to ribs and vertebral column. Contraction of muscles straighten the diaphragm to increase thoracic cavity

- Intercoastal muscles: There are
  i) External intercoastal
  ii) Internal intercoastal
  iii) External oblique
iv) Internal oblique muscles
- Abdominal muscles: Relaxation allows compression of abdominal organs when diaphragm straightens. Contraction presses the abdominal viscera against diaphragm to bulge it more upwardly (for expiration).

MECHANISM OF BREATHING

INSPIRATION

- It is the process by which the fresh atmospheric air enters into the alveoli of the lungs. It is an active process and is brought about by activity of inspiratory muscles. The main muscles of inspiration in normal quiet breathing are the external intercostal muscles and phrenic or radial muscles of diaphragm. During difficult or deep breathing (forced inhalation) they are assisted by the muscles of abdomen.
  i) Diaphragm: When relaxed the diaphragm is dome-shaped structure which separates the thoracic cavity from the abdominal cavity. Phrenic or radial muscles extend from diaphragm to ribs and vertebral column. When these muscles contract diaphragm becomes flat, thus increases the thoracic cavity antero posteriorly. These are the principle inspiratory muscles and play about 75% role in inspiration, other muscles play 25% role in inspiration.
  ii) External intercostal muscles: They occur between the ribs. These are 11 pairs of muscles extending between 12 pairs of ribs. Their contraction pulls the ribs and sternum upward and outward there by increasing the thoracic cavity dorso-ventrally and laterally.
  iii) Abdominal muscles: These muscles relax and allow compression of abdominal organs by diaphragm.
- Due to simultaneous contraction of inspiratory muscles, volume of thoracic cavity increases in all directions.
- As the lungs are held tightly against thoracic wall, enlargement of thoracic cavity results in expansion of lungs
- This decreases the intrapulmonary pressure than atmospheric pressure by -2 to -6 mmHg.
- As it is property of gases, that they move from the place from higher pressure to place of lower pressure, fresh air rushes through respiratory passage into the lungs to equalize the pressure.
- The movement of fresh air into lungs is called inspiration
EXPIRATION

- It is the process by which foul air is expelled out of the lungs. Expiration is normally a passive process and involves the relaxation of inspiration, expiratory muscles become active, making expiration an active energy consumed process.
  i) Diaphragm: When muscles of diaphragm relax it again becomes dome-shaped, decreasing the thoracic cavity.
  ii) External intercostal muscles: When these muscles relax, sternum and ribs come to their original position. This also decreases thoracic cavity.
  iii) Abdominal muscles: Contraction of abdominal muscles presses the abdominal viscera against the diaphragm, bulging it further upward and thus decreasing the thoracic cavity more vertically.
  iv) Internal intercostal muscles: Contraction of these muscles moves the ribs downward and inward and reduces the thoracic cavity laterally and dorsoventrally. The abdominal and external intercostal muscles are called expiratory muscles.
- Due to the action of above muscles, the overall volume of thoracic cavity decreases and the intra pleural pressure increase by +3 to +4 mmHg. Due to this increased pressure in lungs, foul air is given out of them.
- One breath includes one inspiration and one expiration.
- The respiratory rate is the number of breaths taken per minute. For a person breathing normally at rest, it is equal to 12-14 breath per minute.
- Breathing through nose is healthier as it get filtered and conchal of nose warm up the air.
- Mammals have a negative pressure breathing as it allows them to eat and breath at the same time and in human female thoracic breathing is more predominant.

![Diagram showing the changes in capacity of the thoracic cavity during breathing](image)
PULMONARY AIR VOLUMES AND CAPACITIES

- Spirometry is the process of recording the changes in the volume movement of air into and out of lungs and the instrument used for this purpose is called spirometer or respirometer. The graph showing the changes in the pulmonary volumes and capacities under different conditions of birthing is called spirogram.
- The quantity of air the lungs can receive, hold or expel under different conditions are called pulmonary volumes.
- Combinations of two or more pulmonary volumes are called pulmonary capacities.
- Tidal volume (T.V.): Volume of air inspired or expired in relaxed or resting position -500ml. It consists of 150 ml of dead space volume and 350 ml of alveolar volume.
- Dead space: Part of inspiratory tract not involved in gaseous exchange. (Nose to terminal bronchi, volume 150 ml)
- Residual volume (R.V): Air left in lungs and dead space after forceful expiration. 1.1-1.2 litres. The air left in lungs is useful in uninterrupted gaseous exchange.
- Inspiratory reserve volume (I.R.V = complemental air): Volume of air in excess of tidal volume which can be inhaled due to forceful inspiration.
- Expiratory reserve volume (E.R.V = Supplemental air): Volume of air in excess of tidal volume which can be exhaled due to forceful expiration 1 – 1.1 litres.
- Vital capacity (v.C): It is the total volume of air inspired and expired to a maximum level. It is the sum total of tidal volume, inspiratory reserve volume and expiratory reserve volume
  \[ VC = T.V. + I.R.V. + E.R.V. \]
  It is 3.5 to 4.5 litres
  i) The vital capacity is higher in athletes mountaineers or mountain – dwellers and lower in non-athletes, people living in plains, women, old individuals, cigarette smoker.
  ii) Higher the vital capacity, higher is the amount of air exchanged in each breath.
- Inspiratory capacity (IC): It is the total volume of air that can be inhaled after normal expiration. It includes tidal volume and inspiratory reserve volume.
  \[ IC = T.V. + I.R.V \]
  It is 2.5 to 3.0 litres
- Expiratory capacity (E.C.): It is the total volume of air that can be exhaled after normal inspiration. It includes tidal volume and expiratory reserve volume.
  \[ EC = T.V. + E.R.V \]
  Its value is 1.5 to 1.6 litres
- Functional residual capacity (FRC): It is the sum total of residual volume and the expiratory reserve volume.
  \[ F.R.C. = R.V. + E.R.V. \]
 RESPIRATORY GAS EXCHANGE

Its value is 2.3 to 2.7 litre

- **Total lung capacity (TLC):** It is the total amount of air present in the lungs and the respiratory passage after maximum inspiration. It is the sum total of vital capacity and residual volume

  \[ \text{TLC} = \text{VC} + \text{RV} \]
  
  Its value is 5 to 6 litre

- **Alveolar ventilation:** It is the rate at which the fresh air reaches the alveoli and adjoining areas like alveolar ducts, alveolar sacs and respiratory bronchioles. It is calculated as.

  \[ \text{Alveolar ventilation per minute} = \text{Rate of respiration} \times (\text{TV} - \text{dead s pace volume}) \]
  
  \[ = 12 \times (500 -150) \]
  
  \[ = 12 \times 350 \]
  
  \[ = 4.2 \text{ litres / minute} \]

EXCHANGE OF GASES

- **Alveolar air** is separated from blood present in surrounding blood capillaries by very thin partition of 0.2 μm thickness. It is called respiratory membrane. The membrane consists of alveolar surfactant, alveolar epithelium, epithelial basement membrane, a thin interstitial space, capillary basement membrane and capillary endothelial membrane.

- **Diffusing capacity of a gas across a membrane** is the volume of gas that diffuses per minute for a pressure difference of 1mmHg. The rate of diffusion of CO₂ is 20 times faster than that of oxygen while oxygen diffuses faster (twice) than nitrogen. Partial pressure of O₂ in alveolar air (PO₂) is about 100-104 mmHg while that of deoxygenated blood in alveolar capillary is 40mmHg. Therefore, oxygen diffuses into blood and combines with haemoglobin to form oxyhaemoglobin. Pressure of (PO₂) oxygenated blood is 95 mmHg. PCO₂ (pressure of carbon dioxide) of alveolar capillary blood is 46 mmHg, while in fresh alveolar air it is 40mmHg. As the diffusing capacity of CO₂ is 20 times higher than that of O₂, CO₂ of blood rapidly passes out into alveolar air. Its partial pressure in oxygenated blood is 40 mmHg.

- **Gaseous exchange** occurs again in the tissues cells and capillary blood through the interstitial fluid. Partial pressure of oxygen, PO₂ in respiring cells is 20mmHg, tissue fluid is 40mmHg, while it is 95mmHg in capillary blood. Therefore, O₂ diffuses from blood into tissue fluid and from there into cells. Blood leaving the tissue capillaries has PO₂ of about 40mmHg. PCO₂ of blood capillaries is 40mmHg, tissue fluid 45mmHg and that of cells 52mmHG. Therefore, carbon dioxide diffuses out of cells into tissue fluid and form tissue fluid into blood. Blood leaving the tissue capillaries has a PCO₂ of 46mmHg.

TRANSPORT OF OXYGEN

- Oxygen is carried by blood in two forms in solution and as oxyhaemoglobin as RBCs
RESPIRATORY GAS EXCHANGE

1. IN SOLUTION
   - Oxygen is soluble in plasma to a small extent under normal conditions of temperature and pressure. Hence most of it is carried by red blood cells. About 3% of oxygen is transported by blood in dissolved form in plasma of blood. Example, out of about 4.6ml of oxygen entering each 100ml of blood in lungs only 0.17 ml travels in solution form in the plasma.

2. AS OXYHAEMOGLOBIN
   - RBC contain a protein called haemoglobin. It has four polypeptide chains and four haem groups attached to it or 4 atom of iron in ferrous form (Fe^{2+}), thus it can react with 4 molecules of oxygen to form Hb_{4}O_{8}. This is called oxyhaemoglobin. This combination process is called oxygenation.
   - On an average 15gm of haemoglobin (Hb) is present in 100ml of blood. 1gm of Hb combines with 1.34ml of O_{2}. Thus 100ml of blood carries approximately 20ml of O_{2} (19.4 ml to be exact)
   - But when blood reaches the tissues, its O_{2} concentration reduces gradually to 14.4 ml which is collected by veinules and vein. Thus 5ml of O_{2} is transported by 100ml of blood under normal condition.
   - Haemoglobin has higher affinity for oxygen and this affinity is increased by fall in PCO_{2} of blood.
   - At the alveoli, venous blood has low oxygen and is exposed to low PCO_{2} of alveolus, thus oxygen diffuses into red blood cells and form oxyhaemoglobin (bright red). As CO_{2} diffuses from blood to alveolus, blood PCO_{2} falls increasing further uptake of oxygen.
   - Oxyhaemoglobin remains unchanged till it reaches the tissues where it dissociates readily to release oxygen.

OXYGEN – HAEMOGLOBIN DISSOCIATION CURVE (OXYGEN DISSOCIATION CURVE)

- The percentage of haemoglobin that is bound with O_{2} is called percentage saturation of haemoglobin.
- The relationship between the partial pressure of oxygen (PO_{2}) and percentage of saturation of the haemoglobin with oxygen (O_{2}) is graphically illustrated by a curve called oxygen haemoglobin dissociation curve.
- Under normal conditions, the oxygen haemoglobin dissociation curve is sigmoid shaped or ‘S’ shaped. The lower part of the curve indicates dissociation of oxygen from haemoglobin. The upper part of the curve indicates the acceptance of oxygen by haemoglobin. When the partial pressure of oxygen is 255mmHg the haemoglobin gets saturated to about 50%. It means blood contains 50% oxygen. The partial pressure at which the haemoglobin
saturation is 50% is called $P_{50}$. At 40mmHg of partial pressure of oxygen the saturation is 75%. It becomes 95% when the partial pressure of oxygen is 100mmHg.

- Haemoglobin does not take up oxygen are low $P_O$, but as oxygenation of pigment occurs, its affinity for more $O_2$ increases. In haemoglobin where 4 sub-units are present, acquisition of one molecule of oxygen increases the affinity of neighbouring haems for oxygen. This is known as co-operativity between active sites.

![Oxygen Dissociation Curve Graph](image)

**FACTORS AFFECTING OXYGEN DISSOCIATION CURVE**

i) **Temperature**: At higher temperature haemoglobin gives up oxygen more readily and dissociation curve shits to the right. This is of physiological importance because increased temperature means higher metabolic rate or higher oxygen requirement.

ii) **pH**: Increase in $CO_2$ or other acids lower the pH of plasma and shifts the dissociation curve to the right. At higher $CO_2$ concentration more $O_2$ is given up at any oxygen pressure.

iii) **PCO$_2$**: $CO_2$ lowers the oxygen affinity of haemoglobin even if the pH is kept constant. Oxygen dissociation curve shifts to the right and release more $O_2$ with increase in $PCO_2$

iv) **2,3-diphosphoglyceric acid (2,3-DPG)**: It is present in the red blood cells of adult, formed from 3-phosphoglyceric acid. It competes for oxygen binding sites in haemoglobin molecule. As it binds to the $\beta$-chain of HbA, it causes right shift of dissociation curve resulting in higher $P_{50}$.

v) **Lower CO$_2$ concentration, lower body temperature lower 2,3-DPG lower the $P_{50}$ and the curve moves to the left.**
RESPIRATORY GAS EXCHANGE

BOHR EFFECT

- Shifting of the oxygen haemoglobin dissociation curve to the right by increasing carbon
dioxide partial pressure is known as Bohr effect. It is named after Danish physiologist
Christian Bohr.

- The presence of carbon dioxide decreases the affinity of haemoglobin for oxygen and
increases release of oxygen to the tissues.

- The pH of the blood falls as its CO₂ content increases so that when PCO₂ rises the curve to
the right and P₅₀ rises.

- In the tissue, PO₂ is between 10 to 40 mmHg and PCO₂ is high around 45mmHg. So, an
active tissue will have high PCO₂, low pH and raised temperature leading to the dissociation
of oxygen. Oxygenated blood passing through inactive cells does not given up oxygen even
if its PCO₂ is low but active cells readily gives oxygen as PCO₂ is very high.

TRANSPORT OF CARBON DIOXIDE

I. IN DISSOLVED STATE

Because of its high solubility, about 7% carbon dioxide gets dissolved in the blood
plasma and is carried in solution to lungs. Deoxygenated (venous) blood and oxygenated
(arterial) blood carry about 2.7ml and 2.4ml of CO₂ per 100ml of blood in dissolved state
in plasma respectively.

II. IN THE FORM OF BICARBONATE

- The dissolved carbon dioxide in the blood reacts with water to form carbonic
acid. This reaction is very slow in the blood plasma, but occurs very rapidly inside
RBCs because a zinc containing enzyme, the carbonic anhydrase, present in RBCs,
accelerates its rate about 5000 times.

- Due to this, about, 70% of CO₂, received by blood from the tissues, enters the
RBCs where it reacts with water to form carbonic acid (H₂CO₃).

- Carbonic anhydrase is exclusively found in R.B.Cs. All other tissue contains it in
traces except stomach and pancreas in which have considerable amount. This
enzyme not only speeds up the formation of carbonic acid (H₂CO₃) but also
rapidly converts it back to carbon dioxide and water when blood reaches the
lungs.

- Almost as rapidly as formed all carbonic acid of RBCs dissociates into hydrogen
(H⁺) and bicarbonate ions (HCO₃⁻).

- The most of bicarbonate ions (HCO₃⁻) formed with RBCs diffuse out into blood
plasma along the concentration gradient.

- When the whole blood is saturated with carbon dioxide, the following changes
are seen.
(i) The bicarbonate content of plasma and corpuscles increase.
(ii) The chloride content of plasma is diminished and that of the cells is increased.
(iii) The total base (cations) of both plasma and corpuscles remain unchanged.
(iv) The water content and the volume of corpuscles increase.

- When carbon dioxide is removed from a sample of blood, reverse changes take place. From these observations, it is evident that, when carbon dioxide enters blood, chlorine from plasma enters the RBCs, while the base (NA) is left behind. When carbon dioxide escapes the plasma and combines with the base (Na) again. Due to this alternate movement of chlorine ions, this phenomenon is called chlorine shift or Hamburger phenomenon.

III. AS CARBAMINOHAEMOGLOBIN

- In addition to reacting with water, carbon dioxide also reacts directly with amine radicals (NH$_2$) of haemoglobin to form an unstable compound carbamino-haemoglobin. This is a reversible reaction.

- A small amount of carbon dioxide also reacts in this same way with the plasma proteins. About 23% CO$_2$ is transported in combination with haemoglobin and plasma proteins.

$$\text{HHbNH}_2 + \text{CO}_2 \rightleftharpoons \text{HbNCOOH} + \text{H}^+$$

Reduced \hspace{1cm} Carbamino
haemoglobin \hspace{1cm} haemoglobin
HAEMOGLOBIN AS BUFFER

- Addition of hydrogen ions would make the blood acidic. However, most of the hydrogen ions are neutralized by combination with haemoglobin, which is negatively charged, forming acid haemoglobin. This reduces the acidity of the blood and also releases additional oxygen.
- If the blood becomes too basic, acid haemoglobin dissociates, releasing hydrogen ions.

\[
\text{HHb} \rightarrow \text{H}^+ + \text{Hb}
\]

Thus, the haemoglobin also acts as buffer, a substance that keeps the pH from fluctuating. The haemoglobin of the foetus has a higher affinity for oxygen than the mother’s haemoglobin. After birth, the foetal haemoglobin is gradually replaced by adult haemoglobin.

RESPIRATORY PIGMENTS AND ANIMALS CONTAINING IT

1. Haecocyanin
   A copper containing blue pigment occur in plasma of crustaceans, snails and cephalopods.
2. Chlorocruorin
   It is an iron containing green pigment, occur in plasma of annelids polychaete.
3. Pinnaglobin
   It is a manganese containing brown pigment occurs in blood fluid of some mollusks (pinna).
4. Echinochrome
   Contains iron and occur in the coelomic fluid of sea urchin (echinoderm)
5. Vanadium
   Contains vanadium. Present in the blood of tunicates (urochordates). Ciona contains vanadium in plasma and Ascidia contains in green blood corpuscles (vanadocytes)
6. Myoglobin
   Haemoglobin of the muscle
7. Molpadin
   Occur in molpadia (echinodermata)

RESPIRATORY CENTRE

- It controls the rate of respiration. Respiratory centre is located in medulla oblongata and pons. It has the following well dispersed components.
  i) Dorsal Respiratory Group
     Located dorsally along length of medulla with neurons interconnected to sensory termination of glossopharyngeal (sensory signals from peripheral chemoreceptors) and vagus (sensory signals from lungs and stretch receptors of bronchi) nerves. The area is connected through nerves to phrenic muscles of diaphragm. Nervous signals
from this group brings about normal resting inspiration. Expiration occurs through elastic recoil of thoracic wall and lungs

ii) Pneumotaxic Area
It occurs in pons and is meant for switching off normal inspiration when the limit of lung filling is reached. The latter is however also dependent upon the strength of signal. 0.5 sec when signal is strong and 5 sec when signal is poor.

iii) Ventral Respiratory Group
It occurs ventrolaterally anterior to dorsal respiratory group. The group has two types of neurons, inspiratory and expiratory. They are normally inactive but when the respiratory drive is greater than normal, the group is activated. It results in deeper and quicker inspiration and expiration.

iv) Chemosensitive Area
It lies in the medulla near the place of entry of glossopharyngeal and vagus nerves. It is sensitive to blood carbon dioxide and hydrogen ion concentration. Chemosensitive areas are connected to other areas of respiratory centre.

RESTORATORY DISORDERS

1. TUBERCULOSIS
Bacterial disease caused by Mycobacterium tuberculosis. Infection of several parts but common of lungs. Vaccination with B.C.G. (Bacillus – Calmette – Guerin)

2. PLEURISY ( Pleuritis)
Inflammation of pleura or accumulation of pleural fluid. Presence of excess fluid in the pleural cavity is called hydrothorax. Presence of air in pleural space is called pneumothorax.

3. EMPHYSEMA
It is a permanent abnormal pathological inflation of air spaces distal to terminal bronchioles due to destruction of pulmonary tissues especially alveolar septa and flattening of alveolar ducts. There is little alveolar elasticity. Lung size increases but ventilation is poor. Emphysema develops due to infection, smoking and chronic bronchitis. The disease cannot be cured completely because it involves irreversible change in the alveoli. Bronchodilators, antibiotics and O\textsubscript{2} therapy are used to provide relief and retard progression of disease. Emphysema is preventable if care is taken to reduce exposure to smoke and air pollutants.

4. ASPHIXIA
Paralysis of respiratory centre due to excessive carbon dioxide commonly due to irreversible combination of carbon monoxide with haemoglobin to form carboxyhaemoglobin. It results in death. Common in closed rooms with coal burning, kerosene lamp.

5. PNEUMONIA
- It is a disease of lungs with an incubation period of 1-3 days and characterized by accumulation of mucus/ fluid with dead WBCs in alveoli and bronchioles so that
breathing becomes difficult. It is of several types. Common pneumonia is caused by gram(+) nonmetal paired bacterium called Streptococcus or Diplococcus pneumonia. Other bacterium, fungi, virus, mycoplasma and even some protozoans also produce the disease.

- Three types of individual are more susceptible to disease; elders, infants, immune-compromised. The disease is of two types – bronchopneumonia (young children, elderly person) and lobar pneumonia (10-50 years)
- The disease is transmitted through droplets. There is sudden chill, chest pain, cough with rusty mucoid sputum, rise in temperature, rapid shallow breathing and reduced oxygen level of blood due to poor gaseous exchange. Abdominal distension is also common. Useful drugs are erythromycin, tetracycline, sulphonamide. Bronchiodilator drugs provide some relief untreated pneumonia leads to death.

6. HYPOXIA (Anoxia)
   Shortage of oxygen supply to the body due to:
   a) Normal shortage in air as on high mountain.
   b) Anaemia.
   c) Histotoxicity or poisoning of electron transport system.

7. HICCOUGH (Hiccup)
   Inspiratory spasm caused by sudden contraction of diaphragm accompanied by loud closure of glottis.

8. COUGH
   Violent expiration for expulsion of mucus and particles.

9. WHOOPING COUGH (Pertussis)
   Cough with inspiratory whoop caused by Bordetella (Haemophilus) pertussis.

10. BRONCHIAL ASTHMA
    Due to narrowing of bronchi and spams in bronchial muscles. The disorder is generally due to hypersensitivity of bronchioles to foreign substances. There is intense coughing and difficulty in exhalation. Mucous glands becomes over active producing a lot of mucus that clogs bronchioles and bronchi. Exposure to allergens should be avoided. Bronchodilators, inhalers and antibiotics are given for relief and protection against infection.

11. HAY FEVER
    It is an allergic disorder of nasal lining. It develops due to hypersensitivity of the lining to pollen grains or any other foreign particles.

12. ATELECTASIS
    It is an inability of lungs to expand at birth. This is mainly due to deficiency of surfactants.
13. **SILICOSIS**

   It is due to long exposure to dust containing silicon compounds. Workers of glass industry, potters, gold and copper miners develop progressive fibrosis in the liver.

14. **ASBESTOSIS**

   It is due to inhalation of asbestos – fibres, which may result in cancer of pleura.

15. **DIPHTHERIA**

   Infection of bacterium, corynebacterium diphtheria of upper respiratory tract that produces pseudomembrane in throat. Pseudomembrane obstructs breathing causing hypoxia.

16. **BRONCHITIS**

   Inflammation of bronchi and bronchioles due to hypertrophy and hyperplasia and seromucous gland and goblet cells. There is a regular coughing with thick greenish yellow sputum indicating infection and excessive secretion of mucus. It is commonly caused by viral infection of nasal tract followed by bacterial infection. The disorder is common in smokers and persons exposed to CO rich polluted air. Persons suffering from bronchitis should avoid smoke, irritating chemicals and pollutants. Bronchodilators provide symptomatic relief. Antibiotics are used to cure infection.

17. **CYANOSIS**

   Bluish colouration of skin and mucous membranes due to reduced haemoglobin in blood.

18. **EPISTAXIS**

   Nose bleed. Quite common due to any scratching of nasal membranes. Nasal membrane is highly vascular. Epistaxis can also occur due to hypertension.

19. **PHARYNGITIS**

   Inflammation of Pharynx

20. **LARYNGITIS**

   Inflammation of larynx.

21. **SNEEZING**

   An involuntary, sudden, violent and audible expulsion of air through mouth and nose.

22. **YAWNING**

   A deep involuntary inspiration with mouth open, often accompanied by act of stretching.

23. **SARS**

   It is a killer atypical pneumonia called severe acute respiratory syndrome. The disease is caused by variant of common cold corona virus which spreads by droplet and other methods. There is an initial fever (100.4°F), headache, body aches, dry cough and then difficult breathing.

24. **OCCUPATIONAL LUNG DISEASES**

   i) Black lung: Affects coal workers

   ii) Chronic Beryllium disease (CBD): It affects workers in a variety of metallurgical occupations.
RESPIRATORY GAS EXCHANGE

iii) Byssinosis brown lung disease: It often affects cotton and textile workers when bacteriua released from cotton or other material are inhaled and grow in lungs.

iv) Occupational asthma: It can affect people who work with variety of materials, like dyes, resins, leather, latex, rubber, etc.

- It is always advisable to undertake preventive measures in work place involving pollution risk by:
  (i) Reducing emission of harmful dust and chemicals
  (ii) Using protective gear and clothing.
  (iii) Short duties.
  (iv) Informing workers about risks and preventive measures.
  (v) Regular health check up.

MOUTH SICKNESS

- It is hypoxial or oxygen shortage syndrome, which occurs at altitude of 3500 meters and above. There is decrease in atmospheric pressure as well as oxygen content of atmosphere. Reduced atmospheric pressure, reduces the amount of air taken into the lungs during inspiration. Reduced oxygen contents reduces its partial pressure and rate of diffusion into the blood. Hypoxia increases. As a result, body obtains less O₂ and therefore, produces lesser energy. However, requirement of energy at high altitude is higher due to low temperature and increased physical strain. Effects of deficient availability of energy begins to appear within 8-24 hours.

- It is characterized by breathlessness, fast breathing, nausea, vomiting, cyanosis, headache, muscular weakness and mental fatigue. After sometimes, the symptoms subside due to increased concentration of 2,3-diphosphoglycerate in erythrocytes that attracts more oxygen to form HbO₂ even PO₂ is lower. Soon rise in haemoglobin and erythrocytes count shall start.

TERMS RELATED TO BREATHING

(i) Euponea: Normal breathing
(ii) Hypoponea: Slower breathing
(iii) Hyperponea: Rapid breathing
(iv) Apnoea: No breathing
(v) Dyspnoea : Painful breathing
(vi) Orthopnoea: Difficult breathing in horizontal position.
(vii) Tachypnoea: Rapid shallow breathing.
(viii) Polypnoea: Rapid deep breathing
(ix) Hypercapnia: Excess of CO₂ in blood.
(x) Hypocapnia: Low CO₂ concentration in blood.