



Colloidal Dispersions

Dispersed Systems

Dispersed systems consist of particulate matter, known as the dispersed phase, distributed throughout a continuous or dispersion medium.

The dispersed material may range in size from particles of atomic and molecular dimensions to particles whose size is measured in millimeters.

Accordingly, a convenient means of classifying dispersed systems is on the basis of the mean particle diameter of the dispersed material.

Based on the size of the dispersed phase, three types of dispersed systems are generally considered: (a) molecular dispersions, (b) colloidal dispersions, and (c) coarse dispersions.

The size ranges assigned to these classes, together with some of the associated characteristics, are shown in the accompanying table

Classification of Dispersed Systems Based on Particle Size

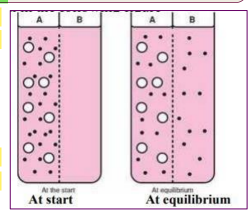
Particle		
Size*	Characteristics of System	Examples
Less than 1 nm	2- Pass through ultrafilter and	Oxygen molecules, ordinary ions, glucose
From 1 nm to 0.5 μm	Visible in electron microscope 2- Pass through filter paper	Colloidal silver sols, natural and synthetic polymers, cheese, butter, jelly, paint, milk, shaving cream, etc.
Greater than 0.5 µm	filter paper 3-Do not pass through	Grains of sand, most pharmaceutical emulsions and suspensions, red blood cells
	Less than 1 nm From 1 nm to 0.5 µm	Size* Characteristics of System 1- Invisible in electron microscope 2- Pass through ultrafilter and semipermeable membrane 3- Undergo rapid diffusion 1- Not resolved by ordinary microscope (although may be detected under ultramicroscope) Visible in electron microscope 2- Pass through filter paper 3- Do not pass semipermeable membrane 4- Diffuse very slowly 1-Visible under microscope 2- Do not pass through normal filter paper 3- Do not pass through normal filter paper 3- Do not pass through semipermeable membrane



Diffusion through a semipermeable membrane

Because the size of colloidal particles, they can be separated from molecular particles by the use of semipermeable membrane. This technique is known as **Dialysis.** As shown in the following figure

Conditions on the two sides, A and B, of the membrane are shown at the start and at equilibrium. The open circles are the colloidal particles that are too large to pass through the membrane.



The solid dots are the electrolyte particles that pass through the pores of the membrane.

- Dialysis occurs in vivo. Thus, ions and small molecules pass readily from the blood, through a natural semipermeable membrane, to the tissue fluids; the colloidal components of the blood remain within the capillary system.
- The principle of dialysis is utilized in the artificial kidney, which removes low—molecularweight impurities from the body by passage through a semipermeable membrane

Types of Colloidal Dispersions*

(According to the type of dispersed phase and dispersion medium)

Dispersed	Dispersion		
Phase	Medium	Colloid Type	Examples
Solid	Solid	Solid sol	Pearls, opals
Liquid	Solid	Solid emulsion	Cheese, butter
Gas	Solid	Solid foam	Pumice, marshmallow
Solid	Liquid	Sol, gel	Jelly, paint
Liquid	Liquid	Emulsion	Milk, mayonnaise
Gas	Liquid	Foam	Whipped cream, shaving cream
Solid	Gas	Solid aerosols	Smoke, dust
Liquid	Gas	Liquid aerosols	Clouds, mist, fog

Note :- a gas in gas always produce a solution





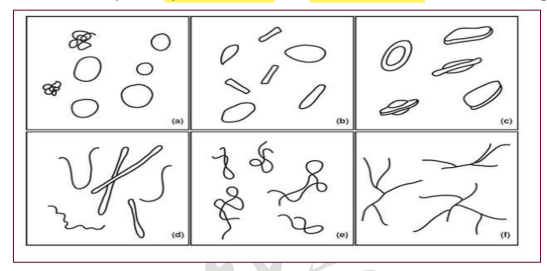
Types of Colloidal Dispersion

PHYSICAL PHARMACY

(According to the shape of dispersed particles)

The shape adopted by particles in dispersion is important because the more extended the particle, the greater is its specific surface and the greater is the opportunity for attractive forces to develop between the particles of the dispersed phase and the dispersion medium.

Some representative shapes of spherocolloids and fibrous colloids are shown in Figure below



Some shapes that can be assumed by colloidal particles: (a) spheres and globules, (b) short rods and prolate ellipsoids, (c) oblate ellipsoids and flakes, (d) long rods and threads, (e) loosely coiled threads, and (f) branched threads.



Types of Colloidal Systems

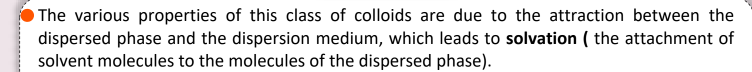
(According to the interaction between the dispersed phase and dispersion medium)

Lyophilic Colloids

Systems containing colloidal particles that interact to an appreciable extent with the dispersion medium are referred to as lyophilic (solvent-loving) colloids. Owing to their affinity for the dispersion medium, such materials form colloidal dispersions, or sols, with relative ease.

Thus, lyophilic colloidal sols are usually obtained simply by dissolving the material in the solvent being used.

For example, the dissolution of acacia or gelatin in water leads to the formation of a sol.



In the case of hydrophilic colloids, in which water is the dispersion medium, this is term is known as hydration.

((One of the most important property of lyophilic colloid is the presence of solvent sheath)) Most lyophilic colloids are organic molecules, for example, gelatin, acacia, insulin, albumin, rubber, and polystyrene.

Of these, the first four produce lyophilic colloids in aqueous dispersion media (hydrophilic sols).

Rubber and polystyrene form lyophilic colloids in nonaqueous, organic solvents. These materials accordingly are referred to as lipophilic colloids.

These examples illustrate the important point that the term lyophilic has meaning only when applied to the material dispersed in a specific dispersion medium. A material that forms a lyophilic colloidal system in one liquid (e.g., water) may not do so in another liquid (e.g., benzene).

Lyophobic Colloids

The second class of colloids is composed of materials that have little attraction, if any, for the dispersion medium. These are the lyophobic (solvent-hating) colloids and, so, their properties differ from those of the lyophilic colloids.

This is primarily due to the absence of a solvent sheath around the particle. Lyophobic colloids are generally composed of inorganic particles dispersed in water. Examples of such materials are gold, silver, sulfur, arsenous sulfide, and silver iodide.

In contrast to lyophilic colloids, it is necessary to use special methods to prepare lyophobic colloids.

♦ Association Colloid

Association or amphiphilic colloids form the third group in this classification. They are certain
organic molecules or ions, termed amphiphiles or surface-active agents, are characterized by
having two distinct regions of opposing solution affinities within the same molecule or ion.



When present in a liquid medium at low concentrations, the amphiphiles exist separately and are of such a size as to be subcolloidal. As the concentration is increased, aggregation occurs over a narrow concentration range.

These aggregates, which may contain 50 or more monomers, are called micelles. Because the diameter of each micelle is of the order of 50 Å, micelles lie within the size range of colloidal particles, such compounds are called association colloidal.

Lyophilic	Lyophobic	Association (Amphiphilic)
Dispersed phase consists	Dispersed phase ordinarily	Dispersed phase consists of
generally of large organic	consists of inorganic particles,	aggregates (micelles) of small
molecules lying within colloidal	such as gold or silver	organic molecules or ions whose
size range		size individually is below the
		colloidal range
Molecules of dispersed phase are	Little if any interaction occurs	Hydrophilic or lipophilic portion
solvated, i.e., they are associated	between particles and dispersion	of the molecule is solvated,
with the molecules comprising	medium (no solvation)	depending on whether the
the dispersion medium		dispersion medium is aqueous or
		nonaqueous
Molecules disperse	Material does not disperse	Colloidal aggregates are formed
spontaneously to form colloidal	spontaneously, and special	spontaneously when the
solution	procedures therefore must be	concentration of amphiphile
	adopted to produce colloidal	exceeds the critical micelle
	dispersion	concentration
Viscosity of the dispersion	Viscosity of the dispersion	Viscosity of the system increases
medium ordinarily is increased	medium is not greatly increased	as the concentration of the
greatly by the presence of the	by the presence of lyophobic	amphiphile increases, as micelles
dispersed phase; at sufficiently	colloidal particles, which tend to	increase in number and become
high concentrations, the sol may	be unsolvated and symmetric	asymmetric
become a gel; viscosity and gel		
formation are related to	armacy 'Ur	110
solvation effects and to the shape	difficulty of	-6
of the molecules, which are		
usually highly asymmetric		
Dispersions are stable generally	Lyophobic dispersions are	In aqueous solutions, the critical
in the presence of electrolytes;	unstable in the presence of even	micelle concentration is reduced
they may be salted out by high	small concentrations of	by the addition of electrolytes;
concentrations of very soluble	electrolytes; effect is due to	salting out may occur at higher
electrolytes; effect is due	neutralization of the charge on	salt concentrations
primarily to desolvation of	the particles; lyophilic colloids	
lyophilic molecules	exert a protective effect	

♦ (Kinetic Properties of Colloids

- There are several properties of colloidal systems that relate to the motion of particles with respect to the dispersion medium. The motion may be:-
 - 1-Thermally induced
 - 2- Gravitationally induced



- 3- Applied externally
- 4- Electrically induced motion



1-Thermally induced (increased by increasing temp)

Brownian motion

- Brownian motion describes the random movement of colloidal particles. The erratic motion, resulting from the bombardment of the particles by the molecules of the dispersion medium.
- The motion of the molecules cannot be observed, of course, because the molecules are too small to see. The velocity of the particles increases with decreasing particle size, while increasing the viscosity of the medium decreases and finally stops the Brownian movement.

Diffusion

- Particles diffuse spontaneously from a region of higher concentration to one of lower concentration until the concentration of the system is uniform throughout.
- Diffusion is a direct result of Brownian movement, therefore (all factors affect Brownian movement will affect diffusion)
- According to **Fick's first law**, the amount (dq) of substance diffusing in time (dt) across a plane of area (S) is directly proportional to the change of concentration, dc, with distance traveled, dx. Fick's law is written as:

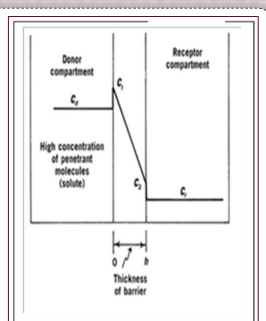
$$dq = -DS \frac{dc}{dx} dt$$

$$g = -D \text{ cm}^2 \frac{g/cm \text{ 3}}{cm} \text{ sec} \longrightarrow D = \text{cm}^2/\text{sec.}$$
 (Units)

- D is the diffusion coefficient, the amount of material diffusing per unit time across a unit area when dc/dx, called the concentration gradient, is unity. D thus has the dimensions of area per unit time.
- Negative sign in the equation, signifies that the diffusion occurs in decreasing concentration of diffusion
- The passage of a substance may be through porous membrane or through tortuous pores or channels (example. Skin)
- Smaller particles diffuse faster in a given media

Molecular > colloid > coarse

 Passive diffusion caused by a concentration gradient and carried out through Brownian motion



It is normal for the concentration curve to increase or decrease sharply at the boundaries of the barrier because, in general, C1 is different from Cd, and C2 is different from Cr

It is important for the release of drug from topical preparation and in the GIT absorption of drugs

Osmosis

- It is the action in which only the solvent is transferred, while the diffusion involve the passage of solute.
- The osmotic pressure, π , of a dilute colloidal solution is described by the van't Hoff equation:

$$\pi = cRT$$

where c is molar concentration of solute, R, gas constant = 0.082 atm.mole/ L.deg This equation can be used to calculate the molecular weight of a colloid in a dilute solution.

2- Gravitationally induced Sedimentation

The velocity, v, of sedimentation of spherical particles having a density ρ in a dispersion medium of density ρ₀ and a viscosity η₀ is given by Stokes's law:

$$V = \frac{2r^2(\rho - \rho_0) g}{9 \eta 0}$$



r = radius of particle, ρ = density of particle, ρ_0 = density of the dispersion medium η_0 = viscosity of the dispersion medium, g = is the acceleration due to gravity Factors affecting sedimentation

- -Particle size (\downarrow particle size $\rightarrow \downarrow$ sedimentation)
- -Viscosity (\uparrow viscosity $\rightarrow \downarrow$ sedimentation)
- -Difference between the densities of the dispersed particles and the dispersion medium (个 difference $\rightarrow \uparrow$ sedimentation)

If the particles are subjected only to the force of gravity, then the lower size limit of particles obeying Stokes's equation is about 0.5 μm. This is because Brownian movement becomes significant and tends to offset sedimentation due to gravity and promotes mixing instead.

Consequently, a stronger force must be applied to bring about the sedimentation of colloidal particles in a quantitative and measurable manner. This is accomplished by use of the ultracentrifuge, which can produce a force one million times that of gravity.

3- Applied externally Viscosity

Viscosity is an expression of the resistance to flow of a system under an applied stress. The more viscous a liquid is, the greater is the applied force required to make it flow at a particular rate.

Viscosity study can give us information about :

- -The shape of particles in dispersion
- -The molecular weight of the dispersed phase



The shape of particles in dispersion

The shapes of particles of the disperse phase affect the viscosity of colloidal dispersions.

Spherocolloids form dispersions of relatively low viscosity, whereas systems containing linear particles are more viscous. The relationship of shape and viscosity reflects the degree of solvation of the particles.

If a linear colloid is placed in a solvent for which it has a low affinity, it tends to "ball up," that is, to assume a spherical shape, and the viscosity falls. This provides a means of detecting changes in the shape of flexible colloidal particles and macromolecules.



The molecular weight of the dispersed phase

The molecular weight of the is measured by using viscometer and the molecular weight obtained by this technique is called viscosity average molecular weight .



The molecular weight of the polymer solution is very high so the viscosity of polymer solution is very high compared to that of pure solvent.

From the Mark-Houwink equation the relationship between the molecular weight and viscosity is given below:

$$[\eta] = KM^{\alpha}$$

Where $[\eta]$ is the intrinsic viscosity, M is the molecular weight, K and α are constants for a particular polymer solvent system.

If we know the k and α values for a given polymer solution, the intrinsic viscosity and molecular weight can be calculated using the above equation.



4- Electrically induced motion

It is the movement of charged surface with respect to an adjacent liquid phase. One of the most important application for such movement is the electrophoresis.



RHEOLOGY

The term rheology, comes from the Greek rheo ("to flow") and logos ("science").

Rheology is used to describe the flow of liquids and the deformation of solids.

Viscosity is an expression of the resistance of a fluid to flow; the higher the viscosity, the greater is the resistance.

The simple liquids can be described in terms of absolute viscosity (single value).

The heterogeneous dispersions rheologic properties are more complex, however, and cannot be expressed by a single value.

Manufacturers of medicinal and cosmetic creams, pastes, and lotions must be capable of producing products with acceptable consistency and smoothness.

Valuable information can be obtained by use of analytic methods of rheology, for formulating better pharmaceutical products.

The rheology of product can affect patient acceptability, physical stability, and even biologic availability.