Membrane separation:

Membrane separations represent a new type of unit operation. The membrane acts as a semi permeable barrier and separation occurs by the membrane controlling the rate of movement of various molecules between two liquid phases, two gas phases, or a liquid and a gas phase. The two fluid phases are usually miscible and the membrane barrier prevents actual, ordinary hydrodynamic flow.

Classification of membrane processes

- Gas diffusion: The rates of gas diffusion depend on the pore sizes and the molecular weights. We may have molecular, transition, and Knudsen diffusion regions depending on the relative sizes of pore and gas molecule.

- Microfiltration (MF): This refers to membranes that have pore m. It is used to filter diameters from 0.1 to 10 filter suspended particulates, bacteria or large colloids from solution.

- Ultrafiltration (UF): This refers to membranes having pore diameters in the range 20-1000 Å. It can be used to filter dissolved macromolecules, such as proteins and polymers, from solution. Reverse osmosis (RO): The membrane pores are in the range of 5-20 Å in diameter, which are within the range of the thermal motion of the polymer chains.

Membrane modules:

The practical equipment where the actual membrane based separation occurs is known as membrane modules. The basic aim of development of these modules is to provide maximum membrane area in relatively smaller volume, so that the permeate flux i.e., the productivity of the system is maximum. These membrane modules are of four types, (i) plate and frame module, (ii) hollow fiber module, (iii) spiral wound and (iv) tubular modules.

Plate and frame modules: The heart of plate-frame module is the support plate that is sandwiched between two flat sheet membranes. The membranes are sealed to the plate, either gaskets with locking devices, glue or directly bonded. The plate is internally porous
and provides a flow channel for the permeate which is collected from a tube on the side of the plate. Ribs or grooves on the face of the plate provide a feed side flow channel. The feed channel can be a clear path with channel heights from 0.3 to 0.75 mm. The higher channel heights are necessary for high viscosity feeds; reduction in power consumption of 20 to 40\% can be achieved by using a 0.6 mm channel compared to a 0.3 mm channel. Alternatively, retentate separator screens (20 or 50 mesh polypropylene) can be used. Commercial plate-frame units are usually horizontal with the membrane plates mounted vertically. They can be run with each plate in parallel plates in two or three series.

**Tubular modules**

In such modules, the membrane is cast on the inside surface of a porous tube. Tubular membranes operate in tangential, or cross-flow, design where process fluid is pumped along the membrane surface in a sweeping type action. The feed solution is pumped through the center of the tube at velocities as high as 6 m/s. These cross-flow velocities minimize the formation of a concentration polarization layer on the membrane surface, promoting high and stable flux and easy cleaning, especially when the objective is to achieve high suspended solids in the MF, UF or NF concentrate. Permeate is driven through the membrane to be directed out of the system or back into the process depending on the application. There are many advantages in tubular membrane configurations. Besides their rugged construction, they have a distinct advantage of being able to process high suspended solids, and concentrate product successfully and repeatedly to relatively high end point concentration levels without plugging. A common objective of an end-of-pipe waste treatment UF system is to reduce waste volume as much as possible to reduce concentrate hauling costs. For juice clarification applications, tubular membrane systems produce the greatest yields and the highest final suspended solids concentration levels. Tubular MF, UF and NF systems do not require significant prefiltration. Some tubular products have the ability to be mechanically cleaned with spongeballs. Spongeballs can be used in process, and are also used to enhance chemical cleaning by reducing time and cleaning chemicals. Tubular membranes are ideally suited to treatment of metalworking oily waste, wastewater minimization and recovery from industrial processes, juice clarification, treatment of pulp and paper industry waste, etc. Tubular membranes typically have life up to 2 to 10 years.

**Hollow fiber module**
In hollow fiber module, lots of hollow fibers (each fiber is a tubular module) are kept in a large pipe. Geometry allows a high membrane surface area to be contained in a compact module. This means large volumes can be filtered, while utilizing minimal space, with low power consumption. Hollow fiber membranes can be designed for circulation, dead end and single pass operation. Some of the many hollow fiber membrane filtration applications include, potable water treatment, juice clarification, wine filtration, dairy processing, etc. The advantages of such modules include reduction in space requirement, lowering in labor cost, lowering in chemical cost, delivery of high quality product water, etc. Hollow fiber membranes offer the unique benefits of high membrane packing densities, sanitary designs and, due to their structural integrity and construction, can withstand permeate back pressure thus allowing flexibility in system design and operation. Most hollow fiber products are available in (i) 1" diameter laboratory test cartridges ranging up to 10" diameter for commercial products, (ii) Standard commercial cartridge lengths of 25", 43", 48", 60" and 72", (iii) Nominal separation ranges from 0.2 micron down to 1,000 MWCO, (iv) Fiber inside diameters from 0.02"(.5mm) up to 0.106"(2.7mm), (v) Various materials of construction including polysulfone and polyacrylonitrile. The following figure (Fig. 3.5c) shows some hollow fiber cartridges of 5, 8 and 10” diameter with endcaps. Benefits of hollow fiber membranes include (i) controlled flow hydraulics, (ii) tangential flow along the membrane surface limits membrane fouling, (iii) membranes can be backflushed to remove solids from the membrane inside surface, thus extending the time between two chemical cleaning cycles, (iv) high membrane packing density resulting in high throughput, (v) modular in structure so that future extension of the plant becomes easier.

**Spiral wound module**

In spiral wound membrane, membrane is cast as a film onto flat sheet. Membranes are sandwiched together with feed spacers (typical thickness 0.03 to 0.1 inch) and permeate carrier. They are sealed at each edge and wound up around a perforated tube. The module diameter ranges from 2.5 to 18 inch and length varies from 30 to 60 inch.

**Driving forces in membrane separation processes:**

Separation in membrane is the result of differences in the transport rates of chemical species through it. Transport rate is determined by the driving force acting on individual components, their mobility, concentration of solute in membrane phase, etc.. Mobility:
Depending on solute size and structure of membrane. Concentration: Chemical compatibility of solute & interface material.

**Fouling of membrane:**

Fouling of membrane is of two types reversible and irreversible.

Reversible fouling: It can be washed away by adopting an appropriate cleaning protocol, like membrane washing. After cleaning, membrane permeability is restored. Concentration polarization is reversible fouling.

Irreversible fouling: In this case, membrane pores are blocked permanently and they cannot be removed, even after proper washing. Permeability is lost permanently.

Fouling results in a continuous decline in membrane permeation rate, an increased rejection of low molecular weight solutes and eventually blocking of flow channels. On start-up of a process, a reduction in membrane permeation rate to 30–10 per cent of the pure water permeation rate after a few minutes of operation is common for ultrafiltration. Such a rapid decrease may be even more extreme for microfiltration. This is often followed by a more gradual decrease throughout processing. Fouling is partly due to blocking or reduction in effective diameter of membrane pores, and partly due to the formation of a slowly thickening layer on the membrane surface. The extent of membrane fouling depends on the nature of the membrane used and on the properties of the process feed. The first means of control is therefore careful choice of membrane type. Secondly, a module design which provides suitable hydrodynamic conditions for the particular application should be chosen. Process feed pretreatment is also important. The type of pretreatment used in reverse osmosis for desalination applications. In biotechnological applications pretreatment might include prefiltration, pasteurisation to destroy bacteria, or adjustment of pH or ionic strength to prevent protein precipitation. When membrane fouling has occurred, backflushing of the membrane may substantially restore the permeation rate. This is seldom totally effective however, so that chemical cleaning is eventually required. This involves interruption of the separation process, and consequently time losses due to the extensive nature of cleaning required. Thus, a typical cleaning procedure might involve: flushing with filtered water at 35–50°C to displace residual retentate; recirculation or back-flushing with a cleaning agent, possibly at elevated temperature; rinsing with water to remove the cleaning agent; sterilisation by
recirculation of a solution of 50–100 ppm of chlorine for 10–30 minutes (600–1800s) at (293–303 K) (20–30 °C); and flushing with water to remove sterilising solution. More recent approaches to the control of membrane fouling include the use of more sophisticated hydrodynamic control effected by pulsed feed flows or non-planar membrane surfaces, and the application of further perturbations at the membrane surface, such as continuous or pulsed electric fields.

**What is membrane?**

A membrane may be defined as “an interphase separating two phases and selectively controlling the transport of materials between those phases”. A membrane is an interphase rather than an interface because it occupies a finite, though normally small, element of space.

**What is mean by membrane separation?**

The use of thin barriers (membranes) between miscible fluids for separating a mixture; a suitable driving force across the membrane, for example concentration or pressure differential, leads to preferential transport of one or more feed components.

Separations of liquids and gases are commonly accomplished using membrane separation methods which include dialysis, reverse osmosis, and ultrafiltration Hybrid and more exotic membrane methods that have also proven effective are electrodialysis, helium separation through glass, hydrogen separation through Palladium and alloy membranes, immobilized solvent and liquid-surfactant membranes.

Permeation of liquids and gases through polymeric membranes occurs where a constituent passes through the membrane by diffusion and sorption by the fluid on the other side of the membrane. The driving force is achieved either by pressure or concentration difference across the membrane.

**Membranes modules can be configured in three ways**

1. single stage feed and bleed
2. multistage feed and bleed
3. continuous single pass as shown in figures
Principle of Operation

When a solution and water are separated by a semi-permeable membrane, the water will move into the solution to equilibrate the system. This is known as osmotic pressure. If a mechanical force is applied to exceed the osmotic pressure (up to 700 psi), the water is forced to move down the concentration gradient i.e. from low to high concentration. Permeate designates the liquid passing through the membrane, and retentate (concentrate) designates the fraction not passing through the membrane.

Advantage of membrane over other process

- A higher overall liquid removal rate is achieved by prevention of the formation of an extensive filter cake.
- The process feed remains in the form of mobile slurry suitable for further processing.
- The solids content of the product slurry may be varied over a wide range.
- It may be possible to fractionate particles of different sizes.

Limitation of membrane processes

The specification and the characteristics of the feed is important in membrane process, cleaning of membranes at regular interval is necessary, concentration polarization, membrane fouling are the two usual phenomena in membranes.

Example for membrane separation process

Some of the membrane processes are dialysis, reverse osmosis, ultrafiltration, and electrodialysis.

Membrane materials for various application

Initially most of the membranes were cellulosic in nature. These are now being replaced by polyamide, polysulphone, polycarbonate and a number of other advanced polymers. These synthetic polymers have improved chemical stability and better resistance to microbial degradation. Membranes have most commonly been produced by a form of phase inversion known as immersion precipitation.
Uses of membrane separation

In general, membrane separation techniques are especially useful in separating:

- Mixtures of similar chemical compounds,
- Mixtures of thermally unstable components (since no heating is needed), and in conjunction with conventional separation methods (such as using membranes to break azeotropic mixtures before feeding them)
- Proteins can be separated in whey for the production of whey protein concentrate (WPC)
- Milk can be concentrated prior to cheese making at the farm level. Apple juice and wine can be clarified.
- Waste treatment and product recovery is possible in edible oil, fat, potato, and fish processing
- Fermentation broths can be clarified and separated
- Whole egg and egg white ultrafiltration as a pre-concentration prior to spray

Distinguish between osmosis & reverse osmosis

When two solutions of differing concentrations of dissolved materials are separated by a semi-permeable membrane, the liquid component will tend to flow from the lower to the more highly concentrated side. In a sense, the concentration difference will tend to equilibrate across the membrane. This process is called osmosis.

If the liquid on the more concentrated side is maintained at a higher pressure, however, this process can be reversed: the solvent will flow from the concentrated side to the less concentrated side. Since the membrane blocks the passage of the dissolved waste constituents, the concentrated solution becomes even more concentrated. This process is called reverse osmosis.
Define thermal diffusion

It involves the formation of a concentration difference within a single liquid or gaseous phase by imposition of a temperature gradient upon the fluid, thus making a separation of the components of the solution possible.

Diffusion is the movement, under the influence of a physical stimulus, of an individual component through a mixture. The term thermal itself denotes that the diffusion is caused due to temperature gradient. Molecular diffusion which is induced by temperature is referred to as Thermal Diffusion.

Example: – Helium-3 separation from Helium-4

Sweep diffusion

If a condensable vapor (steam) is allowed to diffuse through a gas mixture, it carries one of the components along with it, thus making a separation by the operation known as Sweep diffusion.

What is ultra filtration?

Ultrafiltration (UF) is a membrane separation process, driven by a pressure gradient, in which the membrane fractionates components of a liquid as a function of their solvated size and structure.

Define electro dialysis

A ionic mobility under the influence of an electric field through the anionic and cationic membranes is known as electrodialysis. Charged ions can be removed from a solution by synthetic polymer membranes containing ion exchange groups. Anion exchange membranes carry cationic groups which repel cations and are permeable to anions, and cation exchange membranes contain anionic groups and are permeable only to cations. Electrodialysis membranes are comprised of polymer chains - styrene-divinyl benzene made anionic with quaternary ammonium groups and made cationic with sulphonic groups. 1-2V is then applied across each pair of membranes.
APPLICATIONS OF ELECTRODIALYSIS

- Cheese whey demineralization
- Brackish water desalination
- Nitrate removal for drinking water
- Food/sugar products desalting
- Tartaric wine stabilization
- NaCl removal from amino acid salts
- Acid removal from organic product
- Conversion of organic salts into acid and base
- Desalting of amines.
- De-acidification of fruit juices
- Metals removal from ethylene glycol
**Ion exchange.**

The ion exchange is the exchange of equivalent numbers of similarly charged ions, between an immobile phase, which may be a crystal lattice or a gel, and a liquid surrounding the immobile phase. If the exchanging ions are positively charged, the ion exchanger is termed cationic, and anionic if they are negatively charged. The rate at which ions diffuse between an exchanger and the liquid is determined, not only by the concentration differences in the two phases, but also by the necessity to maintain electroneutrality in both phases.

Ion exchange is not a membrane process but it is used for product of protein isolates of higher concentration than obtainable by membrane concentration. Fractionation may also be accomplished using ion exchange processing. It relies on inert resins (cellulose or silica based) that can adsorb charged particles at either end of the pH scale. The design can be a batch type, stirred tank or continuous column. The column is more suitable for selective fractionation. Whey protein isolate (WPI), with a 95% protein content, can be produced by this method. Following adsorption and draining of the deproteined whey, the pH or charge properties are altered and proteins are eluted. Protein is recovered from the dilute stream through UF and drying. Selective resins may be used for fractionated protein products or enriched in fraction allow tailoring of ingredients.

Application examples of ion exchange membrane

* Production of High Purity Chemicals
* Production of Ultra Pure Water
* Battery Diaphragm
* Recovered developing solution.
* Food and pharmaceutical
* Demineralization of Cheese Whey
* Demineralization of Organic Acids and Amino Acids
* Desalination of Soy-Sauce
* Stabilization of Wine
* Demineralization, Deacidification of Fruit Juice

* Demineralization of Natural Extract

**Environmental conservation**

* Desalination of Leachate

* Removal of Nitrate from Under-Ground Water

**Others**

* Production of Salt from Sea water

* Production of Drinking Water from Brackish Water

* Desalination of Deep Sea Water

* Acid Recovery from Waste Acid

* Recovery of Valuable Metals

* Recovery of Acid from Pickling Waste

**Principles of ion exchange**

The ion-exchange solids first used were porous, natural or synthetic minerals containing silica, the zeolites, such as the mineral Na$_2$O.A1$_2$0$_3$.4Si0$_2$.2H$_2$0, for example. Positively charged ions (cations) of a solution which are capable of diffusing through the pores will exchange with the Na$^+$ ions of such a mineral, and the latter is therefore called a cation exchanger. For example,

\[
Ca^{++} + Na_2R \rightarrow CaR + 2Na^+
\]

where R represents the residual material of the zeolite. In this manner "hard" water containing Ca$^{++}$ may be softened by contact with the zeolite, the less objectionable Na$^+$ replacing the C$^{++}$ in solution and the latter becoming immobilized in the solid. The reaction is reversible, and after saturation with Ca$^{++}$ the zeolite may be regenerated by contact with a solution of salt,
\[ \text{CaR} + 2\text{NaCl} \rightarrow \text{Na}_2\text{R} + \text{CaCl}_2 \]

Later certain carbonaceous cation exchangers were manufactured by treating substances such as coal with reagents such as fuming sulfuric acid, and the like. The resulting exchangers can be regenerated to a hydrogen form, HR, by treatment with acid rather than salt. Thus, hard water containing \( \text{Ca(HCO}_3\text{)}_2 \) would contain \( \text{H}_2\text{CO}_3 \) after removal of the \( \text{Ca}^{++} \) by exchange, and since the carbonic acid is readily removed by degasification procedures, the total solids content of the water may be reduced in this manner. Early applications of ion exchangers using these principles were largely limited to water-softening problems.

Synthetic, insoluble polymeric resins containing sulfonic, carboxylic, .or phenolic groups can be considered as consisting of an exceedingly large anion and a replaceable or exchangeable cation. These make exchanges of the following type possible,

\[ \text{Na}^+ + \text{HR} \leftrightarrow \text{NaR} + \text{H}^+ \]

and different cations will exchange with the resin with different relative ease. The \( \text{Na}^+ \) immobilized in the resin may be exchanged with other cations or with \( \text{H}^+ \), for example, much as one solute may replace another adsorbed upon a conventional adsorbent. Similarly synthetic, insoluble polymeric resins containing amine groups and anions may be used to exchange anions in solution. The mechanism of this action is evidently not so simple as in the case of the cation exchangers, but for present purposes it may be considered simply as an ion exchange. For example,

\[ \text{RNH}_3\text{OH} + \text{Cl}^- \leftrightarrow \text{RNH}_3\text{Cl} + \text{OH}^- \]
\[ \text{H}^+ + \text{OH}^- \rightarrow \text{H}_2\text{O} \]

where \( \text{RNH}_3 \) represents the immobile cationic portion of the resin. Such resins may be regenerated by contact with solutions of sodium carbonate or hydroxide. The synthetic ion-exchange resins are available in a variety of formulations of different exchange abilities, usually in the form of fine, granular solids or beads, \( 6 \) to 325 mesh. The individual beads are frequently nearly perfect spheres.
Ion exchange resins

Ion-exchange operations are essentially chemical reactions between an electrolyte in solution and an insoluble electrolyte with which the solution is contacted. The mechanisms of these reactions and the techniques used to bring them about resemble those of adsorption so closely that for most engineering purposes ion exchange may be simply considered as a special case of adsorption.

There are two general types of ion exchange resins: those that exchange positive ions, called cation resins, and those that exchange negative ions, called anion resins. A cation is an ion with a positive charge. Common cations include Ca, Mg, Fe, and H. A cation resin is one that exchanges positive ions. An anion is an ion with a negative charge. Common anions include Cl, SO, and OH. An anion resin is one that exchanges negative ions. Chemically, both types are similar and belong to a group of compounds called polymers, which are extremely large molecules that are formed by the combination of many molecules of one or two compounds in a repeating structure that produces long chains. A mixed-bed demineralizer is a vessel, usually with a volume of several cubic feet, that contains the resin. Physically, ion exchange resins are formed in the shape of very small beads, called resin beads, with an average diameter of about 0.005 millimeters. Wet resin has the appearance of damp, transparent, amber sand and is insoluble in water, acids, and bases. Retention elements or other suitable devices in the top and bottom have openings smaller than the diameter of the resin beads. The resin itself is a uniform mixture of cation and anion resins in a specific volume ratio depending on their specific gravities. The ratio is normally 2 parts cation resin to 3 parts anion resin. In some cases, there may be chemical bonds formed between individual chain molecules at various points along the chain.

Such polymers are said to be cross-linked. This type of polymer constitutes the basic structure of ion exchange resins. In particular, cross-linked polystyrene is the polymer commonly used in ion exchange resins. However, chemical treatment of polystyrene is required to give it ion exchange capability, and this treatment varies depending on whether the final product is to be an anion resin or a cation resin. All of the resin, except the exchangeable ion, is inert in the exchange process.
Define resin capacity

Various measures of the capacity of a resin for ion exchange are in common use. The maximum capacity measures the total number of exchangeable ions per unit mass of resin, commonly expressed in milliequivalents per gram (meq/g). It is the number of fixed ionic groups which determines the maximum exchange capacity of a resin although the extent to which that capacity may be exploited depends also on the chemical nature of those groups.

Osmosis And Reverse Osmosis

When two solutions of differing concentrations of dissolved materials are separated by a semi-permeable membrane, the liquid component will tend to flow from the lower to the more highly concentrated side. In a sense, the concentration difference will tend to equilibrate across the membrane. This process is called osmosis.

If the liquid on the more concentrated side is maintained at a higher pressure, however, this process can be reversed: the solvent will flow from the concentrated side to the less concentrated side. Since the membrane blocks the passage of the dissolved waste constituents, the concentrated solution becomes even more concentrated. This process is called reverse osmosis.

Reverse osmosis (RO) designates a membrane separation process, driven by a pressure gradient, in which the membrane separates the solvent (generally water) from other components of a solution. The membrane configuration is usually cross-flow. With reverse osmosis, the membrane pore size is very small allowing only small amounts of very low molecular weight solutes to pass through the membranes. It is a concentration process using a 100 MW cutoff, 700 psig, temperatures less than 40°C with cellulose acetate membranes and 70-80°C with composite membranes.

Reverse osmosis is often used to remove dissolved organics and metals where concentrations are less than 300 parts per million. However, special care and testing must be performed to assure that the wastes don't dissolve or clog the membrane. Low solubility salts are also prone to precipitate on the membrane surface.
**Electrodialysis**

Ions in aqueous solution can be separated using a direct current electrical driving force on an ion-selective membrane. Electrodialysis usually uses many thin compartments of solution separated by membranes that permit passage of either positive ions (cations) or negative ions (anions) and block passage of the oppositely charged ion.

Cation-exchange membranes are alternatively stacked with anion-exchange membranes placed between two electrodes. The solution to be treated is circulated through the compartments and a direct current power source is applied. All cations gravitate toward the cathode (negatively terminal) and transfer through one membrane, while anions move in the opposite direction, thereby concentrating in alternative compartments.

Electrodialysis is commonly used to recover spent acid and metal salts from plating rinse. It obviously is not effective for non-polar solutions. Electrodialysis is used for demineralization of milk products and whey for infant formula and special dietary products. Also used for desalination of water.

**Principles of operation:**

Under the influence of an electric field, ions move in an aqueous solution. The ionic mobility is directly proportioned to specific conductivity and inversely proportioned to number of molecules in solution. ~3-6 x 10² mm/sec. Charged ions can be removed from a solution by synthetic polymer membranes containing ion exchange groups. Anion exchange membranes carry cationic groups which repel cations and are permeable to anions, and cation exchange membranes contain anionic groups and are permeable only to cations. Electrodialysis membranes are comprised of polymer chains - styrene-divinyl benzene made anionic with quaternary ammonium groups and made cationic with sulphonic groups. 1-2V is then applied across each pair of membranes.

**Electrodialysis process:**

Anion and cation exchange membranes are arranged alternately in parallel between an anode and a cathode (see schematic diagram). The distance between the membranes is 1mm or less.
Categories of membrane separation

Membrane separation processes are classified under different categories depending on the materials to be separated and the driving force applied:

1. In ultrafiltration, liquids and low-molecular-weight dissolved species pass through porous membranes while colloidal particles and macromolecules are rejected. The driving force is a pressure difference.

2. In dialysis, low-molecular-weight solutes and ions pass through while colloidal particles and solutes with molecular weights greater than 1000 are rejected under the conditions of a concentration difference across the membrane.

3. In electrodialysis, ions pass through the membrane in preference to all other species, due to a voltage difference.

4. In reverse osmosis, virtually all dissolved and suspended materials are rejected and the permeate is a liquid, typically water.

5. For gas and liquid separations, unequal rates of transport can be obtained through nonporous membranes by means of a solution and diffusion mechanism. Pervaporation is a special case of this separation where the feed is in the liquid phase while the permeate, typically drawn under subatmospheric conditions, is in the vapor phase.

6. In facilitated transport, separation is achieved by reversible chemical reaction in the membrane. High selectivity and permeation rate may be obtained because of the reaction scheme. Liquid membranes are used for this type of separation.

Ultrafiltration

Suspended materials and macromolecules can be separated from a waste stream using a membrane and pressure differential, called Ultrafiltration. This method uses a lower pressure differential than reverse osmosis and doesn't rely on overcoming osmotic effects. It is useful for dilute solutions of large polymerized macromolecules where the separation is roughly proportional to the pore size in the membrane selected. Ultrafiltration membranes are commercially fabricated in sheet, capillary and tubular forms. The liquid to be filtered is forced into the assemblage and dilute permeate passes perpendicularly
through the membrane while concentrate passes out the end of the media. This may prove useful for the recovery and recycle of suspended solids and macromolecules.

Ultrafiltration (UF) designates a membrane separation process, driven by a pressure gradient, in which the membrane fractionates components of a liquid as a function of their solvated size and structure. The membrane configuration is usually cross-flow. In UF, the membrane pore size is larger allowing some components to pass through the pores with the water. It is a separation/fractionation process using a 10,000 MW cutoff, 40 psig, and temperatures of 50-60°C with polysulfone membranes. In UF milk, lactose and minerals pass in a 50% separation ratio; for example, in the retentate would be 100% of fat, 100% of protein, 50% of lactose, and 50% of free minerals.

Excellent results have been achieved in textile finishing applications and other situations where neither entrained solids that could clog the filter nor dissolved ions that would pass through are present. Membrane life can also be affected by temperature, pH, and fouling.

**Microfiltration**

Microfiltration (MF) designates a membrane separation process similar to UF but with even larger membrane pore size allowing particles in the range of 0.2 to 2 micrometers to pass through. The pressure used is generally lower than that of UF process. The membrane configuration is usually cross-flow. MF is used in the dairy industry for making low-heat sterile milk as proteins may pass through but bacteria do not. Please click above link for a schematic diagram of these membrane processes.

**Uses of membrane separation**

The use of thin barriers (membranes) between miscible fluids for separating a mixture; a suitable driving force across the membrane, for example concentration or pressure differential, leads to preferential transport of one or more feed components.

Separations of liquids and gases are commonly accomplished using membrane separation methods which include dialysis, reverse osmosis, and ultrafiltration Hybrid and more exotic membrane methods that have also proven effective are electrodialysis, helium separation through glass, hydrogen separation through Palladium and alloy membranes, immobilized solvent and liquid-surfactant membranes. Permeation of liquids and gases through polymeric membranes occurs where a constituent passes through the
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