

**Communication**

Reverse Osmosis Process Membranes Modeling – A Historical Overview

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Abstract: This paper presents an overview to the state-of-the-art area of mathematical modeling for reverse osmosis (RO) process. As a liquid-solid process, RO constitutes a valuable and vital physical solution for seawater desalination, wastewater depollution, and water treatment comparatively to the largely controversial and polluting chemical processes such as chlorination and coagulation/flocculation. Great works are required in modeling of RO technique in order to obtain a complete and exhaustive model. Complicated and varying raw water qualities and quantities parameters through time and space are rendering RO modeling hard. Be facing the increasing pollution levels and trying to satisfy the drinking water guidelines, RO membrane modification is often required. This situation made modeling this highly dynamic process more difficult to accomplish.

Keywords: Reverse Osmosis (RO), Thin-Film Composite (TFC), Desalination, Water/Wastewater Treatment, Seawater, Brackish Water, Concentration Polarization (CP)

1. Introduction

Invented 60 years ago, reverse osmosis (RO) became a frequent unit operation in chemical and environmental engineering plants [1-4]. Nowadays, this membrane process is occurring in a larger interval of industrial applications such as desalination of seawater and brackish water, treatment of municipal and industrial wastes, concentration of food products, production of ultra-pure water for several applications comprising the semi-conductor industry, and recovery of important materials in chemical and petrochemical industries [5-12].

RO is an interesting process since it is simple, it can be used for a wide interval of issues in diverse fields, it is economically competitive, and it needs no phase variation [13-15]. The last characteristic is especially crucial for solutions which are responsive to heating, such as food products and pharmaceutical materials [5, 16-18].

The new idea, which conducted to RO, begun in the early 1950's at the University of California at Los Angeles (UCLA), where Samuel T. Yuster imagined the concept of utilizing the Gibb's adsorption equation as a road map to discover methods

for making fresh water from brackish water and seawater [7, 19, 20]. Following Gibb's equation, a comparatively pure water film should occur at the interface of brine and air (or any other hydrophobic surface) which may be skimmed off and, consequently, fresh water may be made. Initially, the implementation of the concept failed until solid-liquid (rather than liquid-gas) interface was studied [21]. The initial idea was to allow pure water create at the solid-liquid interface and then skim the water off by pushing the interfacial water to flow, under an elevated pressure, through the pores of the solid. At this period, the first victorious experiment was realized utilizing a flat plastic film, supported by a porous plate, in 1958 at UCLA [5, 19, 22, 23]. On one's own, approximately the same period, Breton and Reid performed the first effective RO experiments utilizing cellulose acetate membranes and seawater, at the University of Florida [24, 25]. Nevertheless, at that period, the permeation flux of water was very low (approximately a few drops per week) and, consequently, the process was not convenient for industrial application. The advance, which pushed RO to be commercially viable, arrived

in 1958 when Loeb and Sourirajan produced the first asymmetric cellulose acetate membrane [26]; the membrane was not a uniform membrane as previously but was produced of a thin skin film, which was as dense as the precedent homogeneous membranes, and a porous support film. In the Loeb-Sourirajan membrane, important fluxes may be achieved because the flow resistance, in the asymmetric membranes, was much lower than in the uniform membranes [5, 27-30].

To report conveniently the efficiency of a RO membrane, mathematical models are required [31, 32]. These mathematical equations may be subsequently employed for convenient design of RO units [33]. This need has conducted to the expansion of some transport models [34]. The main goal of a transport model is to describe the membrane efficiency, habitually shown as permeation flux and separation (percentage of solute removal from feed solution), to the conditions in operation (such as pressure or feed concentration) or the driving forces (frequently pressure and concentration gradients) through some coefficients (known as phenomenological transport coefficients) which comprise the model parameters [35, 36]. The coefficients (or the parameters) have to be established from experimental information. The triumph of a model can be quantified in matter of the capacity of the model to show mathematically the information with coefficients (or parameters) that are rationally constant through the interval of working conditions. Finally, the model with the established transport coefficients can illustrate the efficiency of a membrane through a large interval of working conditions. This capacity to anticipate the efficiency is the real potential of a transport model. This can be employed, partially, to avoid the elevated costs of experimentation. Integrated with a research program in membrane manufacturing, this may conduct to better conception standards for customizing producing membranes, and joined with a process design program may conduct to a more rational scale-up for RO systems [5, 37, 38].

This review presents a global view on the state-of-the-art area of mathematical modeling for RO process.

2. First Principles of Membrane Process

This section covers the primary definitions and concepts employed usually in RO writings [5].

2.1. Osmosis, Osmotic Pressure and RO

While a semi-permeable membrane (permeable to solvent but not to solute) is put in the midst of two compartments, one containing pure solvent and the other containing a solution (the solvent plus a solute), the solvent penetrates over the membrane to the solution part. This process, which is named “osmosis” (Figure 1(a)), occurs since the system looks for thermodynamic equilibrium. The chemical potential of the pure solvent is bigger than that of the solvent in the solution part thus solvent flows to the solution part to tend to recover equilibrium [5, 39, 40].

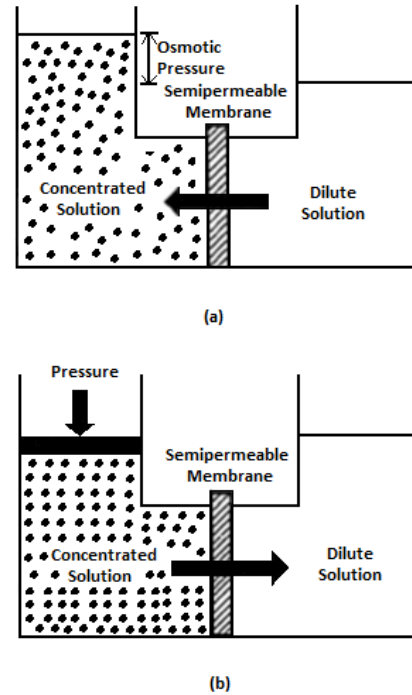


Figure 1. (a) Osmosis, osmotic pressure, and (b) RO [6].

In a device, such as shown in Figure 1, the equilibrium is obtained when there is no net solvent flow over the membrane. The pressure head produced on the solution part is called the “osmotic pressure” (Figure 1(a)), which is a thermodynamic characteristic of the solution and independent of the membrane. When a pressure bigger than the osmotic pressure is imposed to the solution part, augmenting the chemical potential, the solvent flow moves back from the solution part to the pure solvent part; the process is called the “reverse osmosis” or RO as illustrated in Figure 1(b) [6]. For an actual membrane, some solute may be displaced over the membrane and, consequently, the osmotic pressures of the solutions on both parts of the membrane have to be taken into account in modeling. An “efficient pressure driving force” from one side to the other of the membrane can be defined as the applied pressure difference, ΔP , minus the osmotic pressure difference, $\Delta\pi$, from one side to the other of the membrane. For most models, the solvent flux, N_B , is looked to be proportional to the effective pressure driving force, $(\Delta p - \Delta\pi)$ [5, 41, 42].

The osmotic pressure of a solution, π , is linked to the mole fraction and the partial molar volume of the solvent as [43]:

$$\pi = -\frac{RT}{v_B} \ln X_B \quad (1)$$

For dilute solutions, Eq. (1) is reduced to van't Hoff equation [7, 43]:

$$\pi = C_A RT \text{ for nondissociable solutes} \quad (2)$$

$$\pi = (v^+ + v^-) C_A RT \text{ for dissociable solutes} \quad (3)$$

It is more suitable to utilize experimental values of osmotic pressures, which can be obtained for diverse solutions, from different references [5, 7, 44, 45].

2.2. Driving Forces for RO

In RO, the important driving forces are pressure and concentration gradients which conduct to permeation flux of solvent and solute, respectively. The cross effect of solute concentration driving force on solvent flux is symbolized by the osmotic pressure term in the solvent flux equation (see the “effective pressure driving force” in Section 2.1). The cross effect of pressure driving force on solute flux is usually low, for high separation membranes, and is frequently ignored; at what time taken into account this influence is termed by the Staverman (or reflection) coefficient, σ [46]. At what time the Staverman coefficient is considered, the equation of solvent flux is formulated, in most models [47, 48], as:

$$N_B = A[\Delta P - \sigma(\pi_2 - \pi_3)] \quad (4)$$

where A is the solvent permeability coefficient [5].

The Staverman coefficient, σ , is a separation property of the membrane which may take values in the interval $0 \leq \sigma \leq 1$. The closer the σ is to 1, the greater the membrane separation; consequently, $\sigma = 0$ identifies a membrane with very small solute refusal, and $\sigma = 1$ identifies a membrane with total solute refusal [5].

2.3. Membrane Efficiency and Concentration Polarization (CP)

Membrane efficiency is frequently described in matter of “permeation flux” and “separation” [49]. The flux is the rate of material conveyed per unit membrane area, and separation is the fractional concentration decrease of solute from feed stream (i.e., the high-pressure side solution) as juxtaposed with permeate stream (i.e., the low-pressure side solution) [5].

Separation, f (which is occasionally named “rejection” or “retention” in some references), is described in matter of the feed bulk molality, m_{A1} , and the permeate molality, m_{A3} [7]:

$$f = \frac{m_{A1} - m_{A3}}{m_{A1}} \quad (5)$$

For fairly dilute solutions, the molal concentration, m_{Ai} , may be estimated by molar concentration, C_{Ai} , and Eq. (5) be formulated as [5]:

$$f = \frac{C_{A1} - C_{A3}}{C_{A1}} \quad (6)$$

On the other hand, separation may be described in matter of the concentration of the boundary-film solution just outside the membrane on the high-pressure part, C_{A2} . This separation, \hat{f} , named the “theoretical separation”, is formulated, for quite dilute solutions, as [5]:

$$\hat{f} = \frac{C_{A1} - C_{A3}}{C_{A2}} \quad (7)$$

The boundary-layer concentration, C_{A2} , can vary from the feed bulk concentration, C_{A1} , because of a process named “concentration polarization” (CP) [50]. The polarization influence takes place since as solute is rejected by the membrane the solute concentration augments close the

membrane; this situation, which is known as the “solvent-membrane affinity” situation, has been represented in Figure 2. Compared to the solvent-membrane affinity situation is the “solute-membrane affinity” situation in which the solute molecules are caught, rather than refused, by the membrane; this situation is illustrated in Figures 3 and 4 for positive and negative separations, respectively. At steady state, C_{A2} does not change as a function with time and the “film theory” is used to illustrate the CP influence [51-53]. A mass balance for solute through the membrane gives a form of Fick’s first law which by integration conducts to [5]:

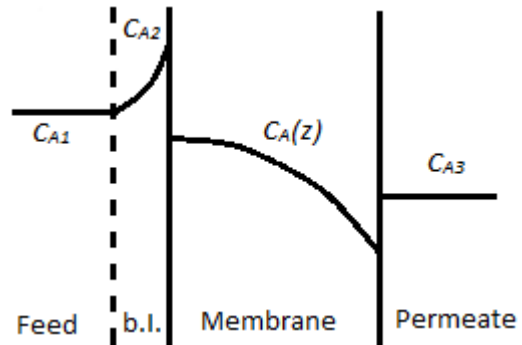


Figure 2. Partitioning influence in a solvent-membrane affinity system (e.g., NaCl-water-cellulose acetate membrane) [5].

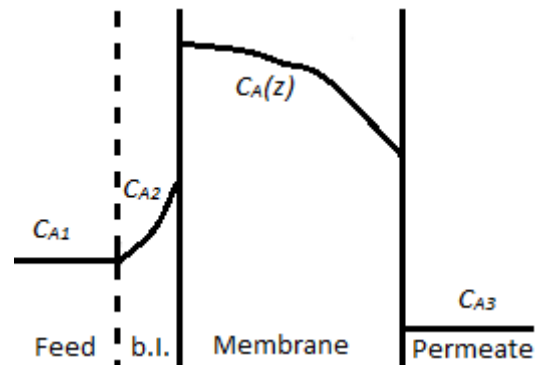


Figure 3. Partitioning influence in a solute-membrane affinity system with positive separation (e.g., toluen water-cellulose acetate membrane) [5].

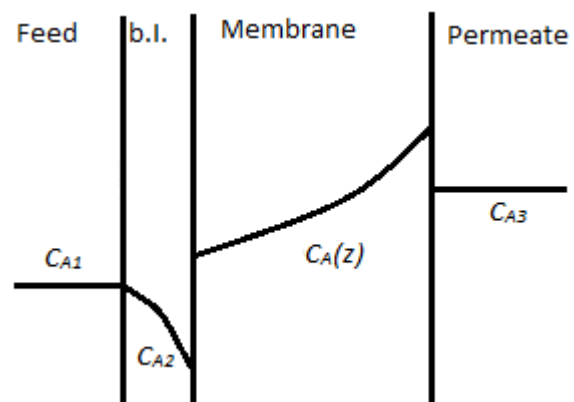


Figure 4. Partitioning influence in a strong solute-membrane affinity system with negative separation (e.g., p-chlorophenol-water-cellulose acetate membrane) [5].

$$C_{A2} = C_{A3} + (C_{A1} - C_{A3})e^{(J_v/k)} \quad (8)$$

where k is the mass transfer coefficient, a characteristic of the boundary layer [7]. As mixing on the elevated-pressure part of the membrane is augmented, Eq. (8) tells in advance that the mass transfer coefficient augments decreasing the polarization influence (i.e., C_{A2} moves toward C_{A1}). When mixing is infinite (i.e., $k \rightarrow \infty$) \hat{f} moves toward f ; consequently, \hat{f} is the theoretical separation that would be metered with ideal mixing on the elevated-pressure part of the membrane [7]. CP is an unwelcome influence [54]. As the polarization effect augments larger the boundary-layer concentration elevates, the osmotic pressure elevates, and, consequently, the effective pressure driving force ($\Delta p - \Delta \pi$) reduces; therefore the solvent flux reduces (see Eq. (4)) [5].

The mass transfer coefficient, k , depends on feed flow rate, cell geometry, and solute system. Generalized correlations of mass transfer coefficient propose that the Sherwood number, Sh , is a function of Reynolds, Re , and Schmidt, Sc , numbers [7, 55]:

$$Sh = \hat{a} Re^{\hat{b}} Sc^{1/3} \quad (9)$$

where \hat{a} and \hat{b} are parameters to be evaluated by experiments [5].

For a constant flow rate and cell configuration, Eq. (9) suggests that k changes as a function of solute diffusivity to the $2/3$ power, because the Reynolds number wipes out and the Schmidt number is inversely proportional to the solute diffusivity. Reformulating Eq. (9) comparatively to a reference solute at the identical working states produces [7]:

$$k = k_{ref} \left(\frac{D_{AB}}{D_{AB,ref}} \right)^{2/3} \quad (10)$$

As a result, if k_{ref} is known for a reference solute, therefore k for any other solute can be evaluated using Eq. (10) if the working states are identical [5].

A significative correlation in RO transport principles, which is employed in several transport models, is the correlation between the solute and solvent fluxes. These two fluxes are linked to the permeate concentration by a global mass balance as [5]:

$$C_{A3} = C \frac{N_A}{N_A + N_B} \quad (11)$$

3. RO Membranes and Membrane Modules

First RO membranes were uniform (i.e., with an isotropically dense film); for this reason, water permeation flux was exceptionally low because of the big flow resistance given by the homogeneous thickness of the membrane [5, 25, 56].

The triumph of the RO technique is attributed to a big range to the expansion of an “asymmetric membrane”, initially advanced by Loeb and Sourirajan [26]. An asymmetric membrane is manufactured to produce a relatively ultrathin, dense surface film supported underneath by a much thicker,

porous film. The asymmetric structure is a direct result of the casting technique employed. While a polymer solution is cast on a flat surface, the evaporation of the solution gives a surface skin. Ulterior gelation in cold water fixes the structure; the porous film is constituted by the substitution of the solvent by the nonsolvent water [7]. Scanning electron microscopy specifies that three films occur in an asymmetric membrane: a relatively dense surface skin, a transition film, and a porous support film [57]. The transition film is in-between in both density and position comparatively to the other two films. The great part of the resistance to mass transfer over the membrane occurs in the surface skin. Consequently, it can be supposed that the membrane efficiency is a function mostly of the chemical nature and physical structure of the surface skin [5, 58, 59].

To a certain degree brevity and completeness, new generation of membranes, which is a member of the family of asymmetric membranes, are the “thin-film composite” (TFC) membranes [60-62]. These membranes possess an alike composition to asymmetric membranes except that the films are manufactured one by one: the porous support is manufactured before anything else, from one polymer material, and a thin layer of a different polymer is covered (usually by interfacial polymerisation) on the porous substrate [5, 63, 64].

Concerning the skin film, the fundamental interrogation is if it is porous. Scanning electron microscopy has shown that ultrafiltration membranes are obviously porous; however, while it concerns the extremely fine structure of RO membranes, no pores have been identified. The presence or absence of pores in RO membranes may not be shown by nowadays knowledge [5]. This is the heart of the discussion in which certain transport models suppose the membrane to be porous and some suppose the membrane as non-porous; certain models, which are frequently based on irreversible thermodynamics (IT), suppose no specific membrane composition or transport phenomena [5, 65].

One tradely triumphant type of TFC membrane is the aromatic polyamide FT30 membranes produced by FilmTec Corporation [63, 66-69]. The FT30 membrane comprises three portions: an ultrafin, skin film of $\sim 0.25 \mu\text{m}$ thick; a microporous support film of polysulfone ($\sim 50 \mu\text{m}$ thick); and a polyester carrier web with extremely big pores and a thickness of $\sim 125 \mu\text{m}$. The chemical material of the skin film is a crosslinked aromatic polyamide with some anionic functionality (i.e., negative charges). The skin film in the FT30 membrane possesses a thickness some times that in other TFC membranes [66]; this forces the membrane to be much more resistant to mechanical and oxidizing damage. The membrane may be employed in a large interval of pH: ~ 3 -11 for continuous runnings and 1-13 for short-term runnings. This lets the membrane to be washed with strong acid and base solutions at temperatures reaching 50°C . One more crucial property of FT30 membranes is the durability at elevated temperatures; experiments have been performed up to 85°C [66]. The FT30 membranes have illustrated reasonable resistance to chlorine aggression, an experiment missed by most of the other noncellulosic RO membranes. The minimum

rate of oxidation aggression has been detected in the pH interval of 5-6 for small durations of time [69]. Fixed exposure or elevated chlorine concentrations (> 5 ppm) will demolish FT30 membranes quickly. Seawater experiments (35 000 ppm total dissolved salts (TDS)) have demonstrated salt refusals of ~ 99.0 to 99.2 percent for spiral-wound elements of FT30 membranes at 5.5×10^5 Pa pressure and 25°C temperature; the corresponding permeation fluxes of $\sim 1.04 \times 10^{-5} \text{ m}^3/\text{m}^2 \text{ s}$ have been mentioned at these situations [68]. For brackish water solutions (2000 ppm TDS), fluxes of $\sim 0.75 \times 10^{-5} \text{ m}^3/\text{m}^2 \text{ s}$ and salt rejections of ~ 96 percent have been mentioned for 1.38×10^5 Pa pressure and 25°C temperature [5, 68, 70-72].

Two tradely crucial members of the family of FT30 membranes are the so-called SW30HR (Sea Water-High Rejection) and BW30 (Brackish Water) membranes. SW30HR is an elevated refusal membrane employed for seawater desalination, and BW30 is utilized for brackish water desalination. These two membranes are chemically identical but possess distinguished efficiency; the correct difference between these membranes is proprietary. Despite the elevated flux and elevated separation properties of TFC membranes, comprising the FT30 membranes, small effort has been performed to know the transport properties of these membranes [5].

Some technologies have been advanced to put a big membrane area into a relatively little volume, for industrial usage. The most well-known of these conceptions are: spiral wound, hollow fibre, and tubular. The proportional qualities of each of these conceptions is function of the specific use [5, 6, 73-77].

4. RO Transport Mechanisms and Models

In the last few decades, a big number of models and theories have been suggested to illustrate the transport mechanism of solute and solvent over RO membranes [5]. Some reviews may be found in the literature [8, 34, 78]. The models have been frequently suggested in order to illustrate the transport mechanisms in RO processes (for those models which suppose several types of transport mechanism in the membrane). However, large approval subsists concerning the mechanism. These models have been obtained from two distinct large approaches. The first group of models, named phenomenological, are frequently founded on IT where the membrane is considered as a black box in which comparatively steady processes appear near equilibrium [79, 80]. In these models, fluxes are linear combinations of forces in the system [5].

In the second group of models (which may be named the "mechanistic models"), a mechanism of transport is supposed and, appropriately, fluxes are linked to the driving forces that occur in the system. The general models can be partaged into general and electrokinetic models [81]. The general models can be put in an application for any type of solute; but, the electrokinetic modes apply only to electrolytic solution feeds.

From another viewpoint, the mechanistic models can be partaged into porous and non-porous membrane models. The transport models in the porous membrane group suppose that the membrane is porous, and those in the non-porous membrane group assumes the membrane is dense and not porous [5].

4.1. Transport Mechanisms

The mechanism of membrane transport in RO remains a matter of debate [5]. Some mechanisms have been proposed; these mechanisms are discussed shortly here.

4.1.1. Sieve Mechanism

This extremely easy idea of sieve-filtration affirms that membrane separation happens because of the difference between the molecular sizes of solute and solvent; the membrane pore size must be in the midst of the two molecular sizes [82]. The mechanism is excluded in RO because it may not illustrate systems such as sodium chloride-water in which the molecular sizes of the solute and water are approximately identical [5].

4.1.2. Wetted Surface Mechanism

Reid and Breton recognized that, because of wettability of membrane materials, water is sorbed into the membrane by hydrogen bonding [25]. The long chained water structure, blocking up the membranes pores, subsequently prohibits the solute transfer across the membrane [5]. The water thereafter permeates by diffusing from one wetted part to another.

4.1.3. Solution-Diffusion (SD) Mechanism

Following this mechanism, proposed by Lonsdale et al., the solute and solvent primarily dissolve in the dense skin film of a membrane and after that diffuse across the membrane; consequently, the solubilities and diffusivities of the solution component are of great significance [83]. Pore flow (i.e., convection [84]) is avoided in this mechanism. After that, Sherwood et al. enlarged the SD model by supposing some imperfections, or holes, on the membrane surface film which participate to the pore flow of solute and solvent, which is famous as the Solution-Diffusion-Imperfection model [5, 85].

4.1.4. Preferential Sorption-Capillary Flow (PS-CF) Mechanism

At the opposite side to SD mechanism, previously, the PS-CF mechanism, primarily proposed by Sourirajan, supposes that the skin film of the membrane is porous [19]. Following this mechanism, the membrane transport is to a certain degree directed by surface phenomena, at the solution-skin film interface, and partially by fluid transport over the micro-capillaries. The physicochemical characteristic of the solute-solvent-membrane surface system establishes which compound of solution is discriminatory sorbed by the membrane. As an example, for the system NaCl-water-cellulose acetate, in which the membrane possesses a low dielectric constant, the ions are repulsed by the membrane and a solute-free film of water is sorbed on the membrane. The pure water film is subsequently pushed to

flow across the capillaries under the system pressure. Once upon a time, Glueckauf had evaluated, using surface tension information, that the thickness of the pure water film is ~0.35 nm [5, 86, 87].

4.2. Transport Models

This section shortly presents certain significant transport models of RO. More information may be discovered in the literature [34, 78].

A small number of models have considered the membrane as a “black box” in which no mechanism has been supposed. Subsequently, the principles of IT have been put in an application to obtain equations for the membrane separation and flux. One of the concepts of IT is that the system may be partaged into a small subsystems in which local equilibrium may occur and consequently thermodynamic quantities may formulated for these subsystems. For systems that are not too distant from equilibrium, IT proposes logical formulas between forces and fluxes [5].

Onsager proposed that the fluxes and forces may be formulated by the following linear equations [88]:

$$J_i = L_{ii}F_i + \sum_{i \neq j} L_{ij}F_j ; i = 1, n \quad (12)$$

where the fluxes, J_i , are linked to the forces, F_j , by the phenomenological coefficients, L_{ij} . For membrane systems, the driving forces may be linked to the pressure and concentration differences through the membranes, and the fluxes are solvent and solute permeate fluxes. Eq. (12) may be made easier to understand by supposing that cross coefficients are the same [88]:

$$L_{ij} = L_{ji} \text{ for } i \neq j \quad (13)$$

The above Onsager reciprocal relationship (ORR), Eq. (12), is acceptable if the system is near equilibrium, the linear laws (i.e., Eq. (12)) are acceptable, and the exact selection of fluxes and forces has been performed. For systems that are distant from equilibrium, as is frequently the situation in RO, Eq. (13) may not be exact. The veracity of the ORR has been treated by Soltanieh and Gill [34].

Two most important phenomenological transport models based on IT are discussed by [5].

5. Computational Fluid Dynamics (CFD) Modeling for Membrane Filtration

Noticing and measuring the flow and solute transfer inside the membrane system, particularly the concentration close to the membrane surface, is hard using experiments; therefore, analytical and numerical CFD solutions have been utilized to examine the hydrodynamics [89, 90]. Primarily analytical and numerical solutions were not able to adress spacerfilled membrane channels since it was difficult to construct such complex geometry in the model and they required to use facilitated or predefined velocity profiles [91, 92]. Several following researches have mentioned that spacers have a

crucial action in dictating velocity and concentration distribution within membrane channels; therefore, the primary models without spacers were restricted in their usage [93-95].

An additional facilitation in primary CFD modeling was that the permeate flow across the membrane (flux) was frequently neglected or placed as a constant number. By performing so, the reciprocal action enter the fluid flow and solute transport was neglected. This signified the models were incapable to assess a crucial property of membrane filtration and CP [90, 96-103].

6. Development of the Stagnant Film Model

As seen in Section 2.3, precise models for CP, the accumulation of solutes at the membrane–solution interface in RO tubes, are crucial for anticipating system efficiency [12]. Regardless of its empirical good results, several modeling approaches used in the deduction of the frequently-utilized stagnant film model appear to restrict the model's usage to actual situations. Moreover, several subsisting models for CP utilize an average mass transfer coefficient with a local mass transfer driving force, which conducts to inaccurate anticipations for the osmotic pressure at the membrane–channel interface. Rohlfs et al. decreased the Zydney-transformed governing equations for solute mass transfer to an analogous convective heat transfer problem [12]. They then applied the principle of superposition to fit solutions from the heat transfer problem to the RO tube boundary conditions, producing a solution that exactly takes into account a local transport coefficient with a local mass transfer driving force. The finished relationships for RO element sizing and rating illustrates good accord with experimental information and gives a theoretical construction for CP modeling that considers the characteristic formation of the mass transfer boundary film not taken into account for by several existing, more empirical models. The model has crucial results for the conception of RO systems with high permeability membranes, as the reduction in membrane resistance in these systems conducts to a relative augmentation in the significance of CP in system efficiency [12].

7. Conclusion

The main points drawn from this study may be listed as below:

From the idea, which conducted to RO, where Samuel T. Yuster imagined the concept of utilizing the Gibb’s adsorption equation as a road map to discover methods for making fresh water from brackish water and seawater, started a long history for a new revolutionary technology for desalination and water treatment.

Modeling RO technique still has great tasks to perform until a complete and exhaustive model will be obtained. Complicated and varying raw water qualities and quantities parameters through time and space are making hard to modelise perfectly this highly dynamic process. Facing to the increasing pollution

levels and trying to satisfy the drinking water guidelines, RO membrane modification is often required which made modeling RO process harder to accomplish.

In spite of its large success, RO remains a limited physical process which need more research in terms of design and construction to increase its performance.

List of Abbreviations

A	Solvent permeability coefficient
\acute{a}	Parameter in Eq. (9), dimensionless
\acute{b}	Parameter in Eq. (9), dimensionless
C_A	Solute concentration (mol m^{-3})
C_{Ai}	Molar concentration
C_{A1}	Feed bulk concentration
C_{A2}	Concentration of the boundary-film solution just outside the membrane on the high-pressure part
C_{A3}	Permeate concentration
CFD	Computational fluid dynamics
CP	Concentration polarization
D_{AB}	Solute diffusivity ($\text{m}^2 \text{s}^{-1}$)
$D_{AB, ref}$	Reference solute diffusivity ($\text{m}^2 \text{s}^{-1}$)
f	Separation (rejection or retention)
\acute{f}	Theoretical separation
F_j	Force
IT	Irreversible thermodynamics
J_i	Flux of component i through membrane, in Eq. (11)
k	Mass transfer coefficient for any solute (m/s)
k_{ref}	Mass transfer coefficient for a reference solute (m/s)
L_{ij}	Phenomenological coefficient
m_{A1}	Feed bulk molality (molal concentration)
m_{A3}	Permeate molality
N_A	Solute flux ($\text{kg m}^{-2} \text{s}^{-1}$)
N_B	Solvent flux ($\text{kg m}^{-2} \text{s}^{-1}$)
ORR	Onsager reciprocal relationship
PS-CF	Preferential sorption-capillary flow
R	Ideal gas constant ($8.314 \text{ kg m}^2 \text{ s}^{-2} \text{ K}^{-1} \text{ mol}^{-1}$)
Re	Reynolds number, dimensionless
RO	Reverse osmosis
Sc	Shmidt number, dimensionless
Sh	Sherwood number, dimensionless
SD	Solution-diffusion (SD)
T	Temperature (K)
TDS	Total dissolved salts
TFC	Thin-film composite
UCLA	University of California at Los Angeles
v_B	Partial molar volume of the solvent ($\text{m}^3 \text{ mol}^{-1}$)
v_i	Partial molar volume of component i ($\text{m}^3 \text{ mol}^{-1}$)
v^+	Number of kmol of cations from dissociation of 1 kmol salt, dimensionless
v^-	Number of kmol of anions from dissociation of 1 kmol salt, dimensionless
X_B	Mole fraction of the solvent

Greek Symbols

ΔP	Applied pressure difference, from one side to the other of the membrane (Pa)
$\Delta P - \Delta \pi$	Efficient pressure driving force (Pa)
$\Delta \pi$	Osmotic pressure difference, from one side to the other of the membrane (Pa)
π	Osmotic pressure of a solution (Pa)
π_2	Osmotic pressure of a solution at one side of the membrane (Pa)
π_3	Osmotic pressure of a solution at the other side of the membrane (Pa)
σ	Staverman (or reflection) coefficient

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