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Electrically Driven Membrane Processes

Downstream Processing Søren Prip Beier



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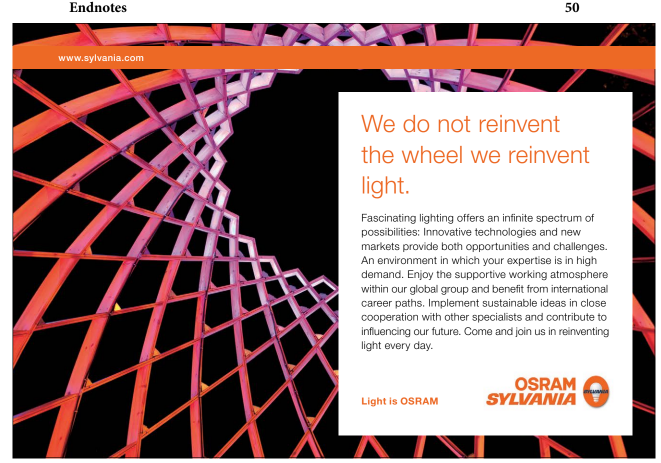
Electrically Driven Membrane Processes

Downstream Processing

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Contents

	List of examples	5
	Nomenclature	7
1	Introduction	9
2	Electrodialysis	18
2.1	Basic concept	19
2.2	Critical current and critical current density	21
2.3	Desalination degree	38
2.4	Current efficiency	40
2.5	Energy requirements	42
2.6	Anti-fouling mechanism	47
3	Summary	48



List of examples

Example A: Transport numbers	17
Example B: Experimental determination of critical current	31
Example C: Determination of critical current from literature correlations	33
Example D: Influence of hydrodynamic conditions on the critical current	37
Example E: Desalination degree	39
Example F: Current efficiency	41
Example G: Thermodynamic efficiency	46

Electrically Driven Membrane Processes: Downstream Processing

The world is developing rapidly. New products are constantly being developed, new technologies and concepts emerge. This calls for constant development of new production processes and education of skilled scientists and engineers.

This book is written to you who have an interest in natural science and especially in downstream production processes in which a separation process is required. The book is written to all chemical engineering students who are participating in courses about downstream processing, membrane processes and/or membrane technology. And it is written to scientists, chemist and/or engineers working with downstream processing and especially electrically driven membrane processes, as well. Membrane processes find applications in almost all kinds of industries as one or more downstream purification/ separation processes:

- Chemical industry
- Pharmaceutical industry
- Food industry
- Dairy industry
- Wastewater treatment industry
- Etc...

For reading and understanding this book you are supposed to have basic skills in mathematics, engineering and chemistry in general. After introduction of certain equations, the SI-units will be thoroughly explained in order to give the reader an overview of the different terms and parameters in that particular equation. I have chosen to do so as such approach helped me when I was studying. Relevant examples will be included in order to show how the described theory can be applied in practice.

I alone am responsible for any misprints or errors in the book and I will be grateful to receive any critics and suggestions for improvement. The book will give you an introduction to basic principles behind electrically driven membrane processes. Relevant theory and models will be presented together with key terms widely within the area of membrane technology.

October 2015 Søren Prip Beier

Nomenclature

a	Activity			
A	Anion exchange membrane			
$A_{membrane}$	Area of membrane			
BC	Boundary condition			
C	Cation exchange membrane			
c	Concentration			
C_1, C_2, C_3	Constants			
c_c Concentrate bulk concentration				
C' _c Concentrate concentration at membrane surface				
c_d	Diluate bulk concentration			
C'_{d} Diluate concentration at membrane surface				
D Diffusion coefficient				
d_h	Hydraulic diameter			
$\mid E \mid$	Voltage			
E_{don}	Donnan potential			
E	Energy consumption			
eq.	quivalent			
F	Faraday number (96485 C/eq.)			
i	Current density			
I	Current			
$\int J$	Flux			
$\mid k \mid$	Mass transfer coefficient			
ig L	Length of flow channel			
$\mid m \mid$	Mobility			
n	Number of cell pairs			
Q	Volume flow rate			
Q_c	Concentrate volume flow rate			
Q_d	Diluate volume flow rate			
Q_e	Electrode solution volume flow rate			
R	Electrical resistance			

Re	Reynolds number
R_{gas}	Gas constant
S	Desalination degree
Sc	Schmidt number
Sh	Sherwood number
t_{+}	Transport number of cation in solution
t_{-}	Transport number of anion in solution
$-\frac{1}{t_+}$	Transport number of cation in membrane
\overline{t}	Transport number of anion in membrane
T	Absolute temperature
u	Flow velocity
x	Direction
Z	Valence of ion
Z_{+}	Valence of cation
Z_{-}	Valence of anion
Greek letters	
δ	Boundary layer thickness
Δp	Pressure drop
η	Current efficiency
η_x , $x = c, d, e, pump$	Pump efficiency
μ	Chemical potential
μ^0	Standard chemical potential
ν	Kinematic viscosity
σ	Electrical conductance
$ \psi $	Electrical potential
ξ	Current utilization

1 Introduction

A membrane process is capable of performing a certain separation by use of a membrane. The core in a membrane process is a membrane that allows certain components to pass while retaining others. Initially some key terms used in membrane technology are shown in Figure 1.

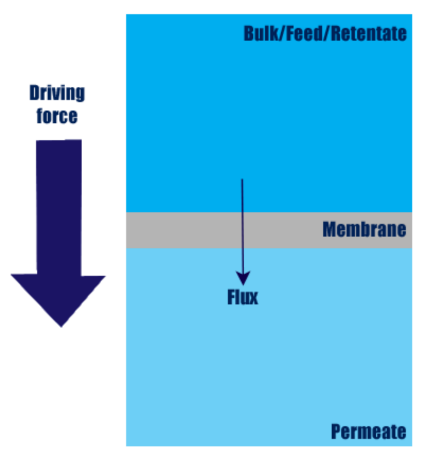


Figure 1: Membrane process

Sketch of a membrane process. The core a membrane through which a driving force induces a flux from the bulk to the permeate side.

The feed side is often called the *bulk solution*. The components in the bulk solution that are retained can also be referred to at the *retentate*. When a driving force is established across the membrane a *flux* will go through the membrane from the bulk solution to the permeate side. The flux is referred to with the letter "J".

A particular separation is accomplished by use of a membrane with the ability of transporting one component more readily than another. In other words, the membrane is more permeable to certain components than other components because of differences in physical or chemical properties between the membrane and the components that are transported through the membrane. As seen in Figure 1, a driving force across the membrane induces a flux from the bulk solution to the permeate side. Different driving forces are used in different membrane processes (listed in Table 1).

Driving force	Membrane process
Pressure gradient Micro-, ultra-, nanofiltration and reverse osmosis	
Concentration gradient Gas separation, pervaporation, dialysis	
Temperature gradient	Membrane distillation, thermo osmosis
Electrical voltage gradient	Electrodialysis, membrane electrolysis

 Table 1: Different membrane processes

Different driving forces, different membrane processes.

In this book we will focus on membrane processes in which the driving force is an electrical voltage difference. Electrically driven membrane processes are widely used to remove charged components from a feed solution or suspension. In contrast to pressure driven membrane processes where you have a volume flux through the membrane, you have a flux of ions through the membrane in electrically driven membrane processes. In order to establish an electrical driving force you need an electrical field. Therefore two electrodes are required; a cathode and an anode. The positive ions (cations) in a solution will migrate to the negative electrode (cathode), the negative ions (anions) will migrate to the positive electrode (anode) and the uncharged molecules will not be affected by the electrical field. One of the greatest applications of electrically driven membrane processes is the desalination of saline water in the production of potable water. The membranes used for this purpose are ion exchange membranes which only allow transport of certain ions.

- *Cation exchange membranes*: Cation exchange membranes are incorporated with negatively charged groups (for example sulfonic or carboxylic acid groups) which will repel anions and only allow transport of cations.
- *Anion exchange membranes:* Anion exchange membranes are incorporated with positively charged groups (for example those derived from quartenary ammonium salts) which will repel cations and only allow transport of anions.

Various types of ion exchange membranes can be distinguished. You can either have heterogeneous or homogeneous ion exchange membranes:

- Heterogeneous ion exchange membranes: Heterogeneous ion exchange membranes are prepared from ion exchange resins and a film-forming polymer. These materials are combined and made into a film by dry-molding for example. The mechanical strength is relatively poor especially at high swelling degrees and the electrical resistance is relatively high which of course is unwanted.
- *Homogeneous ion exchange membranes*: In homogeneous ion exchange membranes the charged groups are attached directly to the polymer chains. The charge is thus distributed uniformly within the whole membrane structure. The swelling of these membranes can be reduced by crosslinking the polymers.

In order to have a good ion exchange membrane, the membrane has to fulfill certain criteria:

- High selectivity
- High electrical conductivity
- High mechanical strength
- High chemical strength
- High ion permeability
- Low electrical resistance





The separation principle when using ion exchange membranes is based on Donnan exclusion which is sketched in Figure 2. The figure shows the case with anions being excluded by cations at the surface of a cation exchange membrane.

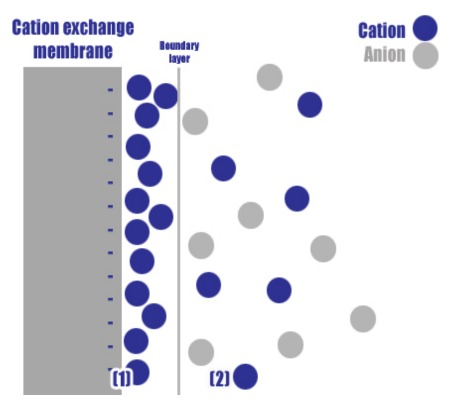


Figure 2: Donnan exclusion at membrane surface

The separation principle associated with ion exchange membranes is based on Donnan exclusion. The cation exchange membrane is incorporated with negative charges and thus a layer of oppositely charges cations occupy the region close the membrane surface in the boundary layer (1). Beyond the boundary layer the concentration of cations and anions is equal (2).

Donnan exclusion is named after the British chemist Frederick George Donnan, and as sketched in Figure 2 ions which the same charge as the membrane are excluded because a layer of oppositely charged ions are located closest to the membrane surface in the boundary layer. The chemical potential of the cations in the membrane (phase 1, Figure 2) and outside the boundary layer (phase 2, Figure 2) can be expressed as follows:

$$\mu_{+}^{1} = (\mu_{+}^{1})^{0} + R_{gas} \cdot T \cdot \ln(a_{+}^{1}) + z_{+} \cdot F \cdot \Psi^{1}, \text{ chemical potential in the membrane}$$

$$\mu_{+}^{2} = (\mu_{+}^{2})^{0} + R_{gas} \cdot T \cdot \ln(a_{+}^{2}) + z_{+} \cdot F \cdot \Psi^{2}, \text{ chemical potential in the bulk}$$
(1)

At equilibrium the chemical potential of the cations in the membrane and in the bulk solution must equal according to thermodynamic considerations. The Donnan potential E_{don} is defined at the difference between the potential in the membrane \Box^1 and in the bulk solution \Box^2 . If the chemical potential at reference state μ^0 is assumed to be equal the following expression for the Donnan potential can be derived from equation (1):

$$E_{don} = \psi^1 - \psi^2 = -\frac{R_{gas} \cdot T}{z_+ \cdot F} \cdot \ln \left(\frac{a_+^1}{a_+^2} \right)$$
 (2)

The Donnan potential is thus determined from the activities of the cations. A similar expression can be written for the anions. It is seen from equation (2) that the Donnan potential is proportional to the natural logarithmic ratio between the activity of the ions in the membrane (phase 1) and the activity of the ions in the bulk solution (phase 2). Thus it is the higher concentration of one of the ions inside the membrane that induces the Donnan potential. The Donnan exclusion can also be depicted in another way that might explain the situation better. In Figure 3 a cross sectional cut of a cation exchange membrane is sketched. You can see a pore through the membrane and the walls are incorporated with negatively charges just as the membrane surface sketched in Figure 2.

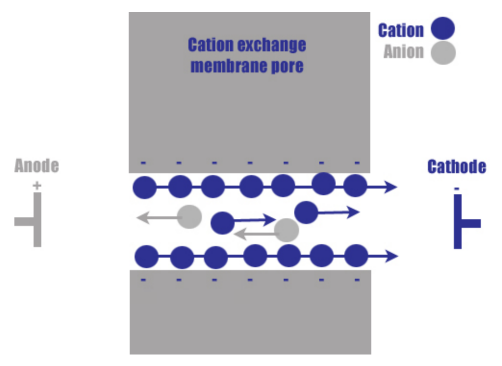


Figure 3: Donnan exclusion inside membrane pore

The walls of a cation exchange membrane pore are covered with negative charges. Thus cations will cover the walls because of electrostatic interactions. In the rest of the pore volume both cations and anions can in principle be found. When a voltage difference is applied the anions will migrate towards the anode and the cation toward the cathode.

In Figure 3 it is seen that because of the negatively charges incorporated in the cation exchange membrane there are much more cations present inside the membrane than anions. When an electrical voltage is applied across the membrane the cations will migrate towards the cathode and the anions towards the anode. Because there are much more cations present inside the membrane than anions, much more cations will be transported through the membrane than anions. The same is the case in anion exchange membranes where much more anions are transported through because much more anions are present inside anion exchange membranes. This is the principle behind the separation of differently charged ions in ion exchange membranes. Some anions are able to pass through a cation exchange membrane but compared to the number of cations that pass through the cation exchange membrane this amount of transported anions is very low. This gives the ion selectivity. Ions are transported through ion exchange membranes as sketched in Figure 3 but also water molecules can convectively be dragged in the same direction as the ions. This is called electroosmotic water transport. In the situation sketched in Figure 3 we will thus have an electroosmotic transport of water molecules toward the cathode (water molecules are convectively dragged along with cations) that is very much larger that the small electroosmotic water transport towards the anode (water molecules are convectively dragged along with the anions). Thus the overall electroosmotic water transport will be in the direction of the cathode in the situation sketched in Figure 3. In anion exchange membranes the overall electroosmotic water transport will of course be in the direction of the anode.



If the concentration of negatively incorporated charges in the cation exchange membrane is known (or concentration of positively incorporated charges in the anion exchange membrane) the concentration of co-ions inside the membrane can be calculated. We are looking at the case with a cation exchange membrane as sketched in Figure 2 and Figure 3; when the Donnan equilibrium is established there is a connection between the concentrations of negative charges in the bulk solution, in the membrane pores and the negative charges incorporated into the membrane. If we are looking at an example with sodium chloride in solution in equilibrium with sodium chloride in a cation exchange membrane an expression for the Donnan equilibrium can be derived since there has to be electrical neutrality overall:

$$\frac{c_{CI^-}}{\overline{c}_{CI^-}} = \sqrt{\frac{\overline{c}_{R^-}}{\overline{c}_{CI^-}}} + 1 \tag{3}$$

Here the concentration of chloride in the bulk solution is denoted $c_{_{C\!\Gamma}}$ and the concentration of chloride in the cation exchange membrane is denoted C_{Cl} . The concentration of fixed negative charges inside the cation exchange membrane is denoted c_{R^-} . This expression can be used to determine the concentration of anions inside a cation exchange membrane. This can be useful in the calculation of so-called *transport* numbers which we will see later.

Since we are dealing with an electrical field and the flow of current we first have to introduce the difference between two important terms:

-
$$I$$
, Current (flow of charges) $SI - units$: $I = \left[I = \frac{C}{s}\right]$

-
$$I$$
, Current (flow of charges) $SI - units$: $I = \left[I = \frac{C}{s}\right]$
- i , Current density (flux of charges) $SI - units$: $i = \left[i = \frac{C}{m^2 \cdot s}\right]$

As one can see the current is the total flow of charges in a given cross sectional area (wire, membrane area etc.) and can be obtained by multiplying the current density with the cross sectional area of the flow of charges. We denote the current with a capital "I" and the current density with a small "i".

If we are looking at cations, the transport through a bulk solution and through a cation exchange membrane can be written based on phenomenological equations:

$$J = t_{+} \cdot \frac{i}{z_{+} \cdot F} \quad \text{(flux in bulk solution)}$$

$$J = \overline{t_{+}} \cdot \frac{i}{z_{+} \cdot F} \quad \text{(flux in membrane)}$$

$$SI - units : J = \left[\frac{C}{s \cdot m^{2}} \right] = \frac{mole}{s \cdot m^{2}}$$

$$\frac{eq}{mole} \cdot \frac{C}{eq} = \frac{mole}{s \cdot m^{2}}$$
(4)

The flux is proportional to the current density i and z_+ is the valence of the cations (eq. pr. mole) and F is the Faraday constant (96485 C/eq.). The flux is a flux of moles pr. time pr. area (moles of charges). This is in contrast to pressure driven membrane processes where we have a volume flux (volume pr. time pr. area). The proportionally constants in equation (4) are *transport numbers*.

$$t_{+}$$
 (transportnumber of cations in solution)
$$\bar{t}_{+}$$
 (transportnumber of cations in membrane) (5)

Transport numbers can be defined for specific ions in specific phases based on their concentrations and mobility in that respective phase. Here the transport numbers of cations are defined in solutions and in the membrane phase:

$$t_{+} = \frac{m_{+} \cdot c_{+}}{m_{+} \cdot c_{+} + m_{-} \cdot c_{-}}, \quad \text{transportnumber of cations in solution}$$

$$\bar{t}_{+} = \frac{m_{+} \cdot c_{+}}{m_{+} \cdot c_{+} + m_{-} \cdot c_{-}}, \quad \text{transportnumber of cations in membrane}$$
(6)

Here m is the mobility of the ions. The mobility of cations and anions respectively is almost the same in the membrane and in the solutions which is also visualized in Figure 3 where anions and cations are equally mobile. Thus it is the concentrations that roughly determine the size of the transport number. Sodium chloride (NaCl) can be used as an example. In a solution sodium ions and chloride ions are transported almost equally since they are equal in concentration and in mobility. Thus t_{\perp} and t_{\parallel} are almost the same according to equation (6), but in a cation exchange membrane almost only sodium ions are present (see again Figure 3). Thus the cation concentration in the membrane is much larger than the anion concentration which according to equation (6) gives a transport number of almost 1. The sum of transport numbers of the cations and anions in solutions and in membrane respectively equals 1 (can be derived from equation (6)), and thus the following can be stated for solutions containing sodium chloride:

$$t_{+} + t_{-} = 1 \Rightarrow t_{+} \approx t_{-} \approx \frac{1}{2}$$
 (transport numbers in solutions)
$$\bar{t}_{+} + \bar{t}_{-} = 1 \Rightarrow \bar{t}_{+} \approx 1 \quad (\bar{t}_{+} >> \bar{t}_{-}, \text{ transport numbers in cation exchange membrane})$$
(7)
$$\bar{t}_{+} + \bar{t}_{-} = 1 \Rightarrow \bar{t}_{-} \approx 1 \quad (\bar{t}_{-} >> \bar{t}_{+}, \text{ transport numbers in anion exchange membrane})$$

Thus in a solution both sodium and chloride ions have transport numbers of $\sim \frac{1}{2}$. In cation exchange membranes the transport number of anions almost equals zero and thus the transport number of cations in a cation exchange membrane almost equal 1. In anion exchange membrane on the other hand the transport number of cations almost equals zero leading to a transport number of anions close to 1. As seen in equation (5), (6) and (7) a "bar" is placed on top of the "t" in order to show that the corresponding transport number refers to the transport number in the membrane.

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We have now seen in equation (7) that for sodium chloride the transport numbers almost equal ½ and 1 in solution and in the membrane respectively. This is also true when we are dealing with dilute solutions when the concentration of co-ions inside the membrane is very low. Of course this co-ion concentration will increase if the bulk concentration increases and thus according to equation (6) the co-ion concentration can no longer be neglected when the transport numbers inside the membrane is to be calculated. We will look more into that scenario in the following example:

Example A: Transport numbers

The transport number of sodium ions inside a cation exchange membrane is to be calculated when the membrane is in equilibrium with a solution of 0.1 M and 1.0 M respectively. The producer of the membrane informs that the fixed negative charge concentration inside the cation exchange membrane is $c_{R^-} = 1.54$ M. The transport number of cations inside the cation exchange membrane can be calculated according to equation (6). We assume that the mobility of cations and anions respectively are equal:

$$\bar{t}_{+} = \frac{\bar{m}_{Na^{+}} \cdot \bar{c}_{Na^{+}}}{\bar{m}_{Na^{+}} \cdot \bar{c}_{Na^{+}} + \bar{m}_{C\Gamma} \cdot \bar{c}_{C\Gamma}} = \frac{1}{1 + \frac{\bar{c}_{C\Gamma}}{\bar{c}_{Na^{+}}}}$$

The concentration of sodium ions inside the membrane must equal the sum of chloride ions and fixed negative charges because of electrical neutrality. This means the transport number can be calculated as follows:

$$\bar{t}_{+} = \frac{1}{1 + \frac{c_{C\Gamma}}{c_{Na^{+}}}} = \frac{1}{1 + \frac{c_{C\Gamma}}{c_{C\Gamma}}} + \frac{1}{c_{C\Gamma}}$$

The concentration of chloride inside the membrane can be calculated from the Donnan equilibrium expression given in equation (3). For the two bulk concentrations of 0.1 M and 1.0 M this give the following chloride concentrations inside the membrane:

$$C_{\text{NaCI}} = 0.1 \text{ M}:$$
 $C_{\text{NaCI}} = 0.1 \text{ M}:$ $\frac{c_{Cl^-}}{\overline{c}_{Cl^-}} = \sqrt{\frac{\overline{c}_{R^-}}{\overline{c}_{Cl^-}}} + 1 \Leftrightarrow \frac{0.1M}{\overline{c}_{Cl^-}} = \sqrt{\frac{1.54M}{\overline{c}_{Cl^-}}} + 1 \Rightarrow \overline{c}_{Cl^-} = 0.0065M$

$$\mathsf{C}_{\mathsf{NaCl}} = \mathsf{1.0 \, M:} \qquad \frac{c_{\mathit{Cl}^-}}{\overline{c}_{\mathit{Cl}^-}} = \sqrt{\frac{\overline{c}_{\mathit{R}^-}}{\overline{c}_{\mathit{Cl}^-}}} + 1 \Leftrightarrow \frac{1.0M}{\overline{c}_{\mathit{Cl}^-}} = \sqrt{\frac{1.54M}{\overline{c}_{\mathit{Cl}^-}}} + 1 \Rightarrow \overline{c}_{\mathit{Cl}^-} = 0.492M$$

These two concentration of chloride inside the membrane are inserted into the expression for the transport numbers which are then calculated:

$$C_{\text{NaCl}} = 0.1 \text{ M:} \qquad \bar{t}_{+} = \frac{1}{1 + \frac{\bar{c}_{Cl^{-}}}{\bar{c}_{Cl^{-}} + \bar{c}_{R^{-}}}} = \frac{1}{1 + \frac{0.0065M}{0.0065M + 1.54M}} = 0.996$$

$$C_{\text{NaCl}} = 1.0 \text{ M:} \qquad \bar{t}_{+} = \frac{1}{1 + \frac{\bar{c}_{Cl^{-}}}{\bar{c}_{Cl^{-}} + \bar{c}_{R^{-}}}} = \frac{1}{1 + \frac{0.492M}{0.492M + 1.54M}} = 0.805$$

It is thus seen that the transport numbers inside the ion exchange membranes are relatively dependent on the bulk concentrations. At low concentrations the transport number of cations is close to 1 which is also stated in equation (7) but when the bulk concentration is increased the transport number decreases.

The concepts mentioned and explained in this introduction section are all important terms concerning electrically driven membrane processes. In the following section the heavyweight of all electrically driven membrane processes, electrodialysis, will be introduced and explained in details.

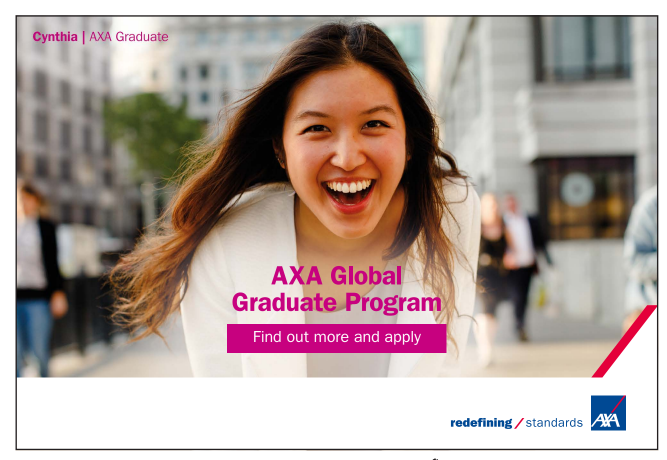
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2 Electrodialysis

Electrodialysis is an electrically driven membrane process in which electrically charged membranes (ion exchange membranes) are used to remove ions from aqueous solutions by use of an electrical field. Electrodialysis finds applications such as:

- Production of potable water by desalination
- Production of salt from seawater
- Removal of salts and acids from pharmaceutical solutions
- Removal of salts and acids in food processing
- Recovery of water and valuable metal ions from industrial effluents

In this chapter the basic concepts of electrodialysis will be presented. Examples from experiments done on a smaller scale electrodialysis system will be included in order to show how the described theory can be applied in practice. The energy consumption will also be described.





2.1 Basic concept

The basic principle in electrodialysis is that two electrodes are separated by cation exchange membranes and anion exchange membrane placed in an alternating way. A sketch of an electrodialysis system is shown in Figure 4.

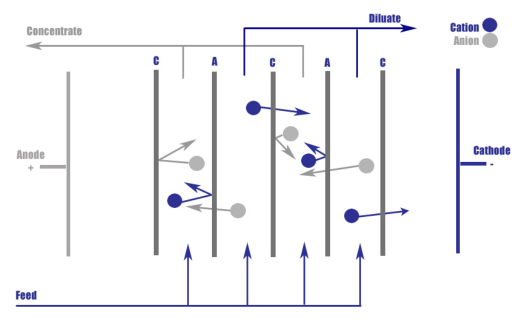


Figure 4: Electrodialysis system

Schematic representation of the principle behind electrodialysis. A: Anion exchange membrane, C: Cation exchange membrane. Two electrodes (anode and cathode) are separated between cation exchange membranes and anion exchange membranes placed in an alternating way. In the electrical field, anions will migrate towards the anode and cations towards the cathode.

The feed solution (saline water for example) is pumped into the chambers between the ion exchange membranes. When a voltage difference is established between the anode and the cathode, the anions will start to migrate towards the anode and the cations will start to migrate towards the cathode. The anions are only able to pass anion exchange membranes and cations are only able to pass cation exchange membranes. Thus an anion is only able to pass one anion exchange membrane whereas it is rejected by cation exchange membranes. On the other hand cations are only able to pass cation exchange membranes and are rejected by anion exchange membranes. This means that when you are looking at Figure 4 every second chamber will increase in concentration (concentrate) and every second chamber will decrease in concentration (diluate). The two chambers closest to the electrodes are called electrode chambers. In these electrode chambers the following electrode reactions take place when sodium chloride solutions are used as feed solution:

$$\frac{2Cl^{-} \rightarrow Cl_{2} + 2e^{-}}{H_{2}O \rightarrow \frac{1}{2}O_{2} + 2H^{+}}$$
Anode reaction

$$2H_2O + 2e^- \rightarrow H_2 + 2OH^-$$
 Cathode reaction

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It is thus seen that when electrodialysis is done on sodium chloride solutions, chlorine and oxygen gas is produced at the anode and hydrogen gas is produced at the cathode. This is not wanted and this is the reason (among other aspects) that electrodialysis systems often consists of up to several hundreds of cell pair placed in parallel in order to minimize the irreversible energy consumption associated with producing gasses at the electrodes. By definition one cell pair consists of the following:

One cation exchange membrane
 One anion exchange membrane
 One cell pair
 One dilution chamber

In commercial electrodialysis systems up to 500 cell pairs (n = 500) are placed in parallel in a so-called membrane stack. The maximum number of cell pairs is determined by practical considerations such as maximum allowed voltage over the whole stack, stability of the stack and the ability of the system to supply the individual chamber equally with feed solution. By using such a membrane stack the applied driving force is utilized very efficiently. The net result of an electrodialysis process is that at the outlet from the membrane stack diluted and concentrated solutions can be collected from the alternating chambers. The ion exchange membranes are separated by spacers. The flow channels in these spacers can have different geometries that can be optimized in order to minimize the concentration polarization in the boundary layer and thus increase the mass transfer coefficient. We will go more into details about that later. Normally two types of spacers can be distinguished: Sheet-flow spacers and tortuous-path spacers. In the sheet flow spacers the velocity is relative low and the residence time is low as well, whereas the residence time in tortuous-path spacers is much larger. The velocity in the tortuous-path spacers is also much larger (which decreases the concentration polarization problems) since the flow channel constitutes a long flow path with many turns. On the other hand the pressure drop is generally larger in a tortuouspath spacer than in a sheet-flow spacer. This gives a larger pump energy consumption (this will be explained in section 2.5.1 Pump energy). In electrodialysis systems three pumps are normally required:

- Pump for electrode solution
- Pump for concentrate solution
- Pump for diluate solution

Thus in energy calculation those three pumps normally have to be taken into consideration when the total energy consumption is to be calculated and evaluated. This will be described in more details in section 2.5 Energy requirement.

2.2 Critical current and critical current density

When a voltage different is applied between the two electrodes, current flows between the two electrodes in the form of ions. Below a certain *critical current* the voltage difference is proportional to the current according to Ohm's law which can be written as follows:

$$I = \frac{1}{R} \cdot E \quad \text{(Current)}, \quad SI - units : \quad I = \left[\frac{C}{s} = \frac{1}{\left(\frac{J \cdot s}{C^2} \right)} \cdot \frac{J}{C} = \frac{C}{s} \right]$$

$$i = -\sigma \cdot \frac{dE}{dx} \quad \text{(Current density)}, \quad SI - units : \quad i = \left[\frac{C}{s \cdot m^2} = \frac{C^2}{J \cdot s \cdot m} \cdot \frac{\left(\frac{J}{C} \right)}{m} = \frac{C}{s \cdot m^2} \right]$$
(8)

In the general form of Ohm's law we see proportionality between the current (C/s) and the voltage (J/C). The term "R" is the electrical resistance (J·s/C² = \square , Ohm). Ohm's law can also be written in a "flux form" by dividing the current with the area (of the membrane). By doing this you get the current density (Coulomb/sec/area = C/(s·m²)) i and then the term " σ " is the electrical conductance (C²/(J·s·m)). The current density is then proportional to the voltage gradient (J/(C·m)).



If an ion exchange membrane is completely selective, one equivalent of ions will be transported through the ion exchange membrane pr. Faraday used electrical current. The Faraday number is 96485 Coulomb pr. equivalent. This means that one mole of salt (NaCl) is removed pr. Faraday electrical current because sodium and chloride ions both have valences z of 1 eq./mole. If the electrodialysis membrane stack then consists of n cell pairs, n moles of NaCl will be removed pr. Faraday electrical current. In general, n equivalents of salt are removed pr. Faraday electrical current.

Ohm's law is only valid below a certain *critical current*. A critical current exists because of concentration polarization phenomena in the laminar boundary layer at the membrane surfaces. The polarization phenomenon is explained in Figure 5. The description here concerns the polarization phenomenon at cation exchange membranes.

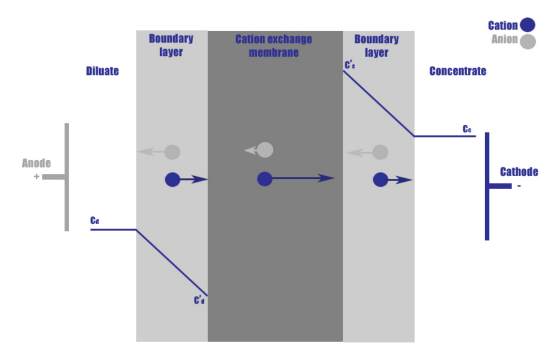


Figure 5: Concentration polarization

Concentration polarization of cations in electrodialysis in the boundary layers at both sides of a cation exchange membrane at steady state. This phenomenon results in the existence of a critical current.

In Figure 5 a cation exchange membrane is sketched and the concentration levels shown concerns the concentration of cations. The flow of anions is also sketched in Figure 5 in order to show the full picture. It is seen that the flow of anions inside the cation exchange membrane is almost zero which is explained by the fact that the transport number of anions inside the cation exchange membrane is almost zero according to equation (7). The cations will migrate through the cation exchange membrane toward the cathode. On the right side of the membrane we have a concentration chamber and on the left side we have a dilution chamber. In the solutions both cations and anions (Na+ and Cl-) have almost equal transport numbers (value ~ ½) according to equation (7). This means that they are transported in almost the same amount at constant applied voltage. Cations and anions are thus transported in the bulk solution and in the boundary layer with a transport number of ~1/2. In the membrane cations are transported in a much larger amount than anions (see also Figure 3) because of the negative incorporated charges which means that they are transported with transport number of almost 1. The ions are transported by diffusion and by the applied electrical driving force. This means that in the laminar boundary layer at both sides of the membrane at steady state there will be a linear concentration gradient as sketched in Figure 5. This is because the cations are transported in a much larger amount than anion inside the membrane than in the boundary layer. If we look at the left side of the membrane the ions are removed away from the left surface through the membrane faster than they are supplied by diffusion in the boundary layer. Thus a decreasing cation concentration is established in the left side boundary layer. At the right side of the membrane an accumulation of cations will initially take place because the cations are transported in a much less amount in solution that in the membrane and thus they will be a concentration increase when leaving the membrane. When steady state is reached the cation concentration profile sketched in Figure 5 will be established.

Why is the concentration polarization a problem? In order the run an economically rentable electrodialysis process the overall electrical resistance of the membrane stack must not be too high. The electrical resistance of a single cell pair can be divided into four sub resistances which are all sketched in Figure 6.

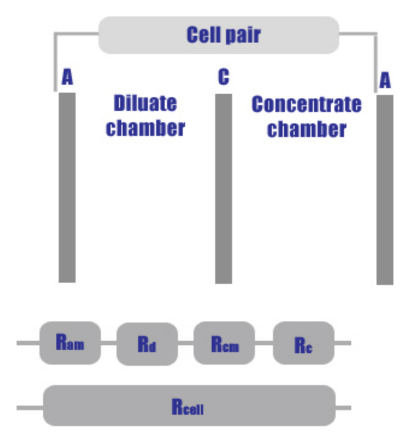


Figure 6: Electrical sub resistances in a cell pair

The overall electrical resistance of a cell pair can be divided into four sub resistances. A: Anion exchange membrane, C: Cation exchange membrane, R_{am} : Resistance of anion exchange membrane, R_{cm} : Resistance of cation exchange membrane, R_{d} : Resistance of diluate solution, R_{c} : Resistance of concentrate solution, R_{cell} : Total resistance of cell pair.

From Figure 6 it is seen that the electrical resistance of a single cell pair can be divided into the four following sub resistances:

- Resistance of anion exchange membrane, R_{am}
- Resistance of diluate chamber, R_d
- Resistance of cation exchange membrane, R_{cm}
- Resistance of concentrate chamber, R_c

The overall resist of a single cell pair (R_{cell}) is the sum of the four sub-resistances. The resistance of the whole membrane stack consisting of n cell pair is thus given in equation (9)

$$R_{membrane\ stack} = n \cdot R_{cell} = n \cdot \left(R_{am} + R_d + R_{cm} + R_c\right), \quad SI - units : \quad R = \left\lfloor \frac{J \cdot s}{C^2} \right\rfloor$$
(9)

The electrical resistance is very dependent on the ease of which the ions are transported and on the presence of ions since the ions constitute the electrical current. The electrical resistance of the ion exchange membranes is relatively low whereas the resistance in the diluate can become quite high because of low ion concentration. A large polarization on the left side of the membrane (Figure 5) gives a very low ion concentration resulting in large electrical resistance. The polarization on right side of the membrane (Figure 5) is not of great importance since it results in higher ion concentration which will not result in an enhanced electrical resistance. Thus it is often the resistance in the diluate chambers that determines the overall resistance of the membrane stack because of low ion concentration at the membrane surface (low c'_{d}) due to concentration polarization. Therefore the concentration polarization is a problem that has to be minimized in order to keep the overall electrical resistance and the energy requirement low.

When the current reaches a certain level the concentration polarization on the diluate side of the membrane (see Figure 5) reaches a level where the cation concentration at the membrane surface $c'_{\rm d}$ reaches zero². At that point the *critical current* is reached and Ohm's law (equation (8)) can no longer describe the association between current and voltage). When the concentration of cations is zero, water splitting at the membrane surface will occur:

Water splitting: $H_2O \rightarrow OH^- + H^+$



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The water splitting occurs in order to generate ions that can constitute an electrical current. When water starts to split, the OH⁻ ions can pass through the anion exchange membranes and the H⁺ ions can pass through the cation exchange membranes. This will affect the pH in the different chambers. When the current through a membrane stack is measured at increasing values of the voltage, the Current-Voltage plot will typically look as sketched in Figure 7.

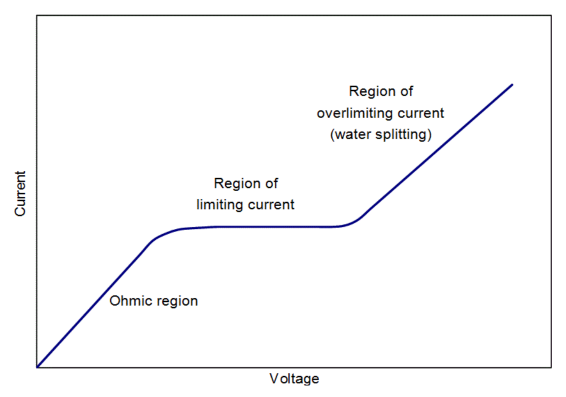


Figure 7: Typically Current-Voltage profile plot

The current at increasing values of voltage. Three regions can normally be observed. I) Ohmic region, II) Region of limiting current and III) Region of overlimiting current (water splitting).

The initial region is described by Ohm's law and is thus called the Ohmic region. At a certain voltage there are not enough ions to transfer the charges which mean that the critical current is reached and the cation concentration at the cation exchange membrane surface on the diluate side is close to zero. Thus the current can not increase anymore even though the voltage is increased. This is the region of limiting current. When the voltage is further increased water molecules start to split at the membrane surface and thus "new" ions are generated (H⁺ and OH⁻) that are able to increase the current. Thus at this point the current increases again at increasing voltage. This is the region of overlimiting current.

By setting up mathematical models for the cation transport through the cation exchange membrane sketched in Figure 5, an expression for calculation of the critical current can be derived. First we can look at Figure 8 in which the cation concentration profile between the dilute stream and the cation exchange membrane is sketched. It is in this area the concentration decrease towards zero leads to the existence of a critical current.

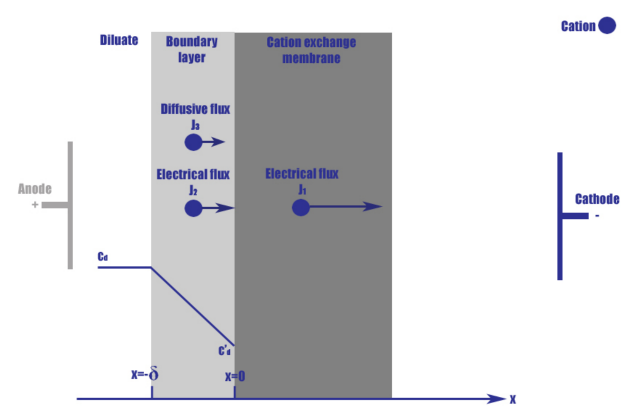


Figure 8: Cation concentration profile

The concentration profile of cations in the laminar boundary layer between the diluate stream and the cation exchange membrane. Two types of cation transport is sketched: Electrical transport due to the electrical field (E-field) and diffusive transport due to the concentration gradient.

In Figure 8, electrical and diffusive transport of salt ions (only cations in this case) is shown³. We are looking at the situation in steady state where a linear concentration decrease in the laminar boundary layer exists. When the current density (flux of charges) through the system is denoted i, the flux of cations in the membrane and in the boundary layer can be expressed as follows:

$$J_{1} = \overline{t}_{+} \cdot \frac{i}{z_{+} \cdot F}, \text{(Flux in membrane due to E - field)}$$

$$J_{2} = t_{+} \cdot \frac{i}{z_{+} \cdot F}, \text{(Flux in boundary lay er due to E - field)}$$

$$SI - units : J = \left[\frac{C}{\frac{m^{2} \cdot s}{m^{2} \cdot s}} - \frac{mole}{m^{2} \cdot s}\right]$$

$$J_{3} = -D \cdot \frac{dc}{dx} \quad \text{(Diffusive flux in boundary lay er)}$$

$$SI - units : J = \left[\frac{m^{2}}{s} \cdot \left(\frac{mole}{m^{3}}\right) - \frac{mole}{m^{2} \cdot s}\right]$$

The fluxes J_1 and J_2 are caused by the electrical field (E-field) according to equation (4). The flux J_3 is caused by the concentration gradient of cations in the boundary layer according to Darcy's law. At steady state a mole balance for the cations can be set up at the membrane surface which gives the following first order differential equation with the following boundary conditions (BC). The direction of x is defined to go from the left to the right according to Figure 8.

$$J_{1} = J_{2} + J_{3} \Leftrightarrow \overline{t}_{+} \cdot \frac{i}{z_{+} \cdot F} = t_{+} \cdot \frac{i}{z_{+} \cdot F} - D \cdot \frac{dc}{dx} \Leftrightarrow$$

$$\frac{i}{z_{+} \cdot F} \cdot (\overline{t}_{+} - t_{+}) = -D \cdot \frac{dc}{dx}$$

$$BC1: \quad x = -\delta \to c = c_{d}$$

$$BC2: \quad x = 0 \to c = c'_{d}$$
(10)



28

Equation (10) can be integrated by use of the boundary conditions to give an expression for the dependency between the current density and the bulk concentration (of the dilute stream) and the boundary layer thickness:

$$\frac{i}{z_{+} \cdot F} \cdot (\bar{t}_{+} - t_{+}) = -D \cdot \frac{dc}{dx} \Leftrightarrow \int_{-\delta}^{0} \frac{i}{z_{+} \cdot F} \cdot (\bar{t}_{+} - t_{+}) dx = -\int_{c_{d}}^{c_{d}} D \cdot dc \Leftrightarrow \frac{i}{z_{+} \cdot F} \cdot (\bar{t}_{+} - t_{+}) \cdot \delta = -D \cdot (c'_{d} - c_{d}) \Leftrightarrow i = \frac{D \cdot (c_{d} - c'_{d}) \cdot z_{+} \cdot F}{\delta \cdot (\bar{t}_{+} - t_{+})}$$

As mentioned earlier the critical current density is reached when the concentration of cations at the membrane surface (c'_{d}) at the diluate side reaches zero. At this point water molecules start to split in order to generate ions to conduct the current. An expression for the calculation the critical current density (i_{crit}) can thus be derived by letting c'_{d} go to zero:

$$\begin{split} i_{crit} &= \lim_{\mathbf{c_m} \to 0} \left(\frac{D \cdot \left(c_d - c_d^{\prime} \right) \cdot z_+ \cdot F}{\delta \cdot \left(\overline{t}_+ - t_+ \right)} \right) = \frac{D \cdot c_d \cdot z_+ \cdot F}{\delta \cdot \left(\overline{t}_+ - t_+ \right)} = \frac{k \cdot c_d \cdot z_+ \cdot F}{\left(\overline{t}_+ - t_+ \right)} \quad \text{(Critical current)} \\ I_{crit} &= A_{membrane} \cdot \frac{k \cdot c_d \cdot z_+ \cdot F}{\left(\overline{t}_+ - t_+ \right)} \quad \text{(Critical current)} \\ SI - units : \quad I_{crit} &= \left[m^2 \cdot \frac{m}{s} \cdot \frac{mole}{m^3} \cdot \frac{eq.}{mole} \cdot \frac{C}{eq.} = \frac{C}{s} \right] \end{split}$$

Again it is seen that the current is just the current density multiplied by the cross sectional area, which in this case is the area of one of the ion exchange membranes. The ratio between the diffusion coefficient and the boundary layer thickness can be replaced by k which is the mass transfer coefficient ($k = D/\delta$). It is thus seen that the critical current density is a function of the concentration of cations in the bulk (diluate stream) and the boundary layer thickness (mass transfer coefficient). Therefore the critical current density is highly dependent of the hydrodynamic conditions and can be increased by decreasing the boundary layer thickness (increasing the mass transfer coefficient) for example by increasing the flow rate along the membrane surface in the diluate chambers. The critical current density will also increase with increased ion concentration in the diluate since more ions will then be present to constitute the flow of charges (the current)⁴. Equation (11) can be used to calculate the critical current for a given electrodialysis system but often you don't know the mass transfer coefficient. Therefore you can do two things to determine the critical current for a given electrodialysis system:

- Experimental determination of critical current
- Determination of critical current from literature correlations

In the experimental determination you don't need to know the mass transfer coefficient. On the other hand, by estimation of the critical current from literature correlation you actually calculate the mass transfer coefficient (diffusion coefficient divided with the boundary layer thickness) and insert it into equation (11). It can often be a good idea to do both things in order to get a more accurate picture of the level of the critical current and to check whether your flow and mass transfer in the system can be described by the well known correlation from the literature. In the following two sections we shall see how the critical current can be determined from experiments and from literature correlations.

2.2.1 Experimental determination of critical current

In an electrodialysis system the critical current can be determined by measuring the current (with an ampere meter) through the membrane stack as a function to the voltage. Then you can plot the current vs. the voltage and determine the critical current as the point where the curve flattens as in Figure 7. This can however often be quite uncertain because the curve often does not flatten so much. Another approach can sometimes be more accurate. The resistance can be plotted vs. the reciprocal value of the current. By doing this, the level of the current (reciprocal value of the current) where the resistance is constant (the Ohm's law region) and the level of the current where the resistance starts to increase (the region of limiting current) can be determined more accurate. The resistance is calculated as E/I according to Ohm's law (equation (8)). In the following example the critical current will be determined from actual smaller scale electrodialysis data according to these two approaches.



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Example B: Experimental determination of critical current

For an electrodialysis system consisting of 20 cell pairs the current is measured at different values of the voltage between 0 and 50 volts. The data are plotted in an "I vs. E plot" and an "E/I vs. 1/I plot" which is shown in Figure 9. Sodium chloride solutions are used as feed solutions.

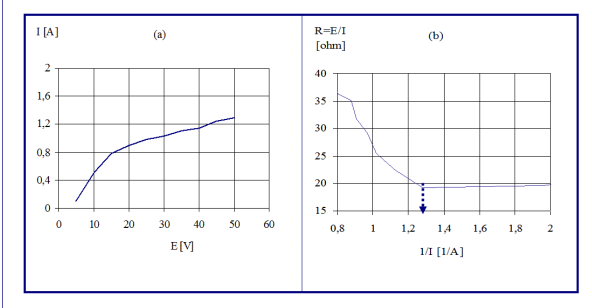


Figure 9: Experimental determination of critical current (a)The current through the membrane stack as a function of the voltage. (b) The resistance (E/I) as a function of 1/I. The critical current in this case is most easily identified on the right plot indicated by the blue arrow. Constant flow rate.

It is seen that the curve (Figure 9a) has the same sequence as the initial sequence of the typical current-voltage plot shown in Figure 7 page 23. Up to around 15 volt linear relationship between the current and voltage exists according to Ohm's law. Above ~ 15 volt the curve is flattened but identification of an actual critical current is from this plot difficult. By plotting the resistance of the membrane stack (voltage divided by the current) as a function of the reciprocal value of the current, the critical current can be identified as the point where the resistance starts to increase when going from the right to the left on the curve in Figure 9b. Ohm's law assumes a constant resistance which is seen on the right part of the curve. At high values of the current (low values of 1/l) the resistance starts to increase indicating that the Ohmic region is exceeded and the region of limiting current is entered. The point where the curve breaks is identified as the critical current, which in this case can be estimated to:

$$I_{crit} = \frac{1}{1.28A^{-1}} = 0.78A$$

This value of 0.78 A would have be more uncertain determined if only Figure 9a have been used.

The critical current density can then be calculated by dividing the critical current with the area of one of the ion exchange membranes in the membrane stack.

2.2.2 Determination of critical current from literature correlations

As mentioned earlier the critical current can experimentally be determined by plotting data of the current as a function of the voltage. On the other hand in order to calculate the critical current by use of equation (11) you need to have a value for the mass transfer coefficient.

One way to get a value for the mass transfer coefficient is to use different flow correlations from the literature that combines different dimensionless numbers concerning flow and mass transfer. These dimensionless numbers are:

Rey nolds number: Re =
$$\frac{d_h \cdot u}{v}$$
, $SI - units$: Re = $\left[\frac{m \cdot \left(\frac{m}{s}\right)}{\left(\frac{m^2}{s}\right)}\right] = [-]$

Schmidt number: Sc = $\frac{v}{D}$, $SI - units$: Sc = $\left[\frac{\left(\frac{m^2}{s}\right)}{\left(\frac{m^2}{s}\right)}\right] = [-]$

Sherwood number: Sh = $\frac{k \cdot d_h}{D}$, $SI - units$: Sh = $\left[\frac{\left(\frac{m}{s}\right) \cdot m}{\left(\frac{m^2}{s}\right)}\right] = [-]$

The hydraulic diameter of the flow channel is denoted d_h . The Reynolds number includes the flow velocity u and thus tells whether the flow is in the laminar or turbulent region. The Schmidt number is the ratio between the kinematic viscosity v and the diffusion coefficient D and thus the Schmidt number tells how fast velocity is propagated through the fluid compared to how fast mass propagates (diffuses) through the fluid. The Sherwood number includes the mass transfer coefficient k and by using correlations combining these three dimensionless number given in equation (12) you can estimate the mass transfer coefficient. Such correlations are shown in Table 2.

	Laminar flow	Turbulent flow
Tube geometry	$Sh = 1.62 \cdot (Re \cdot Sc \cdot d_h / L)^{0.33}$	GL 0.04 D 0.75 G 0.33
Channel geometry	$Sh = 1.85 \cdot \left(\text{Re} \cdot \text{Sc} \cdot d_h / L \right)^{0.33}$	$Sh = 0.04 \cdot Re^{0.75} \cdot Sc^{0.33}$

Table 2: Flow correlations⁵

Different flow correlations for different flow regimes and flow geometries.

How can these correlations be used to find the mass transfer coefficient and how can we then afterward calculate the critical current? These questions are answered through the following example:

Example C: Determination of critical current from literature correlations

We are looking at a lab scale electrodialysis system at which an experiment with NaCl solution is conducted. From current and voltage data a critical current of 5.55 A has been determined by the method sketched in Example B. Relevant experimental parameters are given in Table 3.

Table 3: Experimental parameters

Relevant experimental parameters for an electrodialysis experiment done with a lab scale electrodialysis system.

Cell	Area of one	Mean log. diluate conc., c _d	Total diluate flow, Q _d	Spacer flow channel		
pairs, n	membrane, A _{membrane}			height	Width	Length, L
20	218 cm ²	7.44×10 ⁻⁵ mole/cm ³	11.72 g/s	0.10 cm	0.61 cm	~ 18 cm



First of all we have to know whether we are in the turbulent or laminar region. For this purpose we have to calculate the Reynolds number according to equation (12). We have to know the kinematic viscosity, the diluate flow velocity and the hydraulic diameter of the diluate flow spacer channel. The kinematic viscosity of water at 20°C is used (0.01 cm²/s) since the solution is very dilute. The total flow in the 20 diluate channels Q_d (we have 20 diluate channels since there are a total of 20 cell pairs) is 11.72 g/s. Since the solution is quite diluted we use the density of water. The flow qd in each of the 20 channels is there

$$q_d = \frac{Q_d}{n} = \frac{11.72 \frac{g}{s}}{20} \approx \frac{11.72 \frac{ml}{s}}{20} = 0.59 \frac{ml}{s} = 0.59 \frac{cm^3}{s}$$

The flow velocity u can be calculated by dividing the flow qd with the cross sectional area of the flow channel. The cross sectional area is product of the channel height and width in the spacer (0.10 cm × 0.61 cm = 0.061 cm²):

$$u = \frac{q_d}{A_{channel}} = \frac{0.59 \frac{cm^3}{s}}{0.061 \ cm^2} = 9.61 \frac{cm}{s}$$

The hydraulic diameter of the spacer channel is also calculated:

$$d_h = 4 \cdot \frac{A_{channel}}{L_{wet nerinhery}} = 4 \cdot \frac{0.061 \text{ cm}^2}{2 \cdot (0.61 \text{cm} + 0.10 \text{cm})} = 0.17 \text{ cm}$$

Now we can calculate the Reynolds number according to equation (12):

Re =
$$\frac{d_h \cdot u}{v} = \frac{0.17 \ cm \cdot 9.61 \frac{cm}{s}}{0.01 \frac{cm^2}{s}} = 165$$

Since the Reynolds number is relatively low we assume that we are in the laminar region. In order to calculate the Sherwood number according to the laminar "channel flow" correlation in Table 2, we first have to calculate the Schmidt number according to equation (12). The diffusion coefficient of NaCl in water is roughly 1.5×10^{-5} cm²/s:

Sc =
$$\frac{v}{D} = \frac{0.01 \frac{cm^2}{s}}{1.5 \cdot 10^{-5} \frac{cm^2}{s}} = 666.7$$

The correlation from Table 2 is used to calculate the Sherwood number. The spacer channel length is approximately 18 cm.

Sh = 1.85 ·
$$\left(\text{Re-Sc} \cdot d_h / L\right)^{0.33} = 1.85 \cdot \left(165 \cdot 666.7 \cdot \frac{0.17 \ cm}{18 \ cm}\right)^{0.33} = 18.31$$

The mass transfer coefficient can be calculated from the Sherwood number according to equation (12):

Sh =
$$\frac{k \cdot d_h}{D} \iff k = \frac{Sh \cdot D}{d_h} = \frac{18.31 \cdot 1.5 \cdot 10^{-5} \frac{cm^2}{s}}{0.17 \text{ cm}} = 1.6 \cdot 10^{-3} \frac{cm}{s}$$

With this value of the mass transfer coefficient the critical current can be calculated according to equation (11) with a mean logarithmic diluate concentration of 7.44×10^{-5} mole/cm³. We use a mean logarithmic concentration in the diluate chambers since the concentratio

$$I_{crit} = A_{membrane} \cdot \frac{k \cdot c_b \cdot z_+ \cdot F}{\left(\bar{t}_+ - t_+\right)} = 218 \ cm^2 \cdot \frac{1.6 \cdot 10^{-3} \frac{cm}{s} \cdot 7.44 \cdot 10^{-5} \frac{mole}{cm^3} \cdot 1 \frac{eq.}{mole} \cdot 96485 \frac{C}{eq.}}{\left(1 - \frac{1}{2}\right)} = 5.0 \ A$$

It is seen that this value is close to the experimentally determined critical current of 5.55 ampere, and thus the flow correlations in this case apparently gives a good approximation for the mass transfer coefficient and thus for the calculation of the critical current. Of course there are uncertainties. For example the diffusion coefficient is concentration dependent and the linear length of the flow channel is also not so easy to define since the channels are equipped with insert and it is a question if you can talk about a linear flow channel length at all. But anyway the use of literature correlations can lead you to get an idea of level of the mass transfer coefficient and thus the critical current.

2.2.3 Influence of hydrodynamic conditions on the critical current

As mentioned earlier and according to equation (11) the critical current is proportional to the mass transfer coefficient. Since the mass transfer coefficient is very dependent on the hydrodynamic conditions in the membrane stack the critical current can be varied by changing for example the flow velocity in the diluate chambers. Experimental values of the critical current at different flow velocities in the diluate chambers can be compared with the literature flow correlations. This can be useful if you want to see whether your flow and mass transfer in your electrodialysis system follow the well-known literature correlations. The mass transfer coefficient can be expressed by the flow correlation for channel geometry given in Table 2:

$$Sh = 1.85 \cdot \left(\operatorname{Re} \cdot \operatorname{Sc} \cdot \operatorname{d}_{h} / L \right)^{0.33} \Leftrightarrow k = 1.85 \cdot \frac{D}{d_{h}} \cdot \left(\frac{u \cdot d_{h}}{v} \cdot \frac{v}{D} \cdot \frac{d_{h}}{L} \right)^{0.33} = u^{0.33} \cdot (C_{1})$$
 correlation for laminar flow
$$C_{1} = 1.85 \cdot \frac{D}{d_{h}} \cdot \left(\frac{d_{h}^{2}}{D \cdot L} \right)^{0.33} = \operatorname{constant}$$
 (13)
$$Sh = 0.04 \cdot \operatorname{Re}^{0.75} \cdot \operatorname{Sc}^{0.33} \Leftrightarrow k = 0.04 \cdot \frac{D}{d_{h}} \cdot \left(\frac{u \cdot d_{h}}{v} \right)^{0.75} \cdot \left(\frac{v}{D} \right)^{0.33} = u^{0.75} \cdot (C_{2})$$
 correlation for turbulent flow
$$C_{2} = 0.04 \cdot \frac{D}{d_{h}} \cdot \left(\frac{d_{h}}{v} \right)^{0.75} \cdot \left(\frac{v}{D} \right)^{0.33} = \operatorname{constant}$$

These expressions for the mass transfer coefficient can be inserted into the expression for the critical current, equation (11):

$$\frac{I_{crit}}{c_d} = k \cdot \frac{A_{membrane} \cdot z_+ \cdot F}{\left(\overline{t}_+ - t_+\right)} = k \cdot C_3 \Rightarrow \begin{cases}
\frac{I_{crit}}{c_d} = u^{0.33} \cdot (C_1 \cdot C_3) & \text{(correlation for laminar flow)} \\
\frac{I_{crit}}{c_d} = u^{0.75} \cdot (C_2 \cdot C_3) & \text{(correlation for turbulent flow)}
\end{cases} (14)$$

$$C_{1} = 1.85 \cdot \frac{D}{d_{h}} \cdot \left(\frac{d_{h}^{2}}{D \cdot L}\right)^{0.33}, \quad C_{2} = 0.04 \cdot \frac{D}{d_{h}} \cdot \left(\frac{d_{h}}{v}\right)^{0.75} \cdot \left(\frac{v}{D}\right)^{0.33}, \quad C_{3} = \frac{A_{membrane} \cdot z_{+} \cdot F}{\left(\bar{t}_{+} - t_{+}\right)}$$

The expressions in equation (14) give the dependency between critical current and hydrodynamic conditions (flow velocity). These expressions can be compared with experimental values of the critical current and diluate concentrations at different flow velocities. We are going to do such a comparison in the following example:



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Example D: Influence of hydrodynamic conditions on the critical current

We are going to investigate the influence of the flow velocity on the critical current. Data from an electrodialysis experiment with NaCl solutions at three different flow velocities in the diluate chamber are given in Table 4. Again an electrodialysis system consisting of 20 cell pairs is used.

Flow velocity in diluate chamber, u [m/s]	Mean log. Conc. in diluate chamber, cd [mole/m3]	Critical current*, Icrit [A=C/s]
6.50×10 ⁻²	77.58	4,35
7.37×10 ⁻²	74.83	4,65
9.61×10 ⁻²	74.45	5,55

Table 4: Critical current at different flow velocities

Experimental values for electrodialysis experiments with NaCl solutions at different flow velocities.

A mean logarithmic value for the diluate concentration is used since the concentration changes from the inlet to the outlet of the membrane stack. The data in Table 4 can be plotted together with the expressions given in (14), but first the constants C_1 , C_2 and C_3 are to be calculated (SI-Units are used as in Table 4):

$$\begin{split} C_1 &= 1.85 \cdot \frac{D}{d_h} \cdot \left(\frac{d_h^{\ 2}}{D \cdot L}\right)^{0.33} = \\ 1.85 \cdot \frac{1.5 \cdot 10^{-9} \frac{m^2}{s}}{0.17 \cdot 10^{-2} m} \cdot \left(\frac{\left(0.17 \cdot 10^{-2} m\right)^2}{1.5 \cdot 10^{-9} \frac{m^2}{s} \cdot 0.18 m}\right)^{0.33} \Leftrightarrow C_1 = 3.49 \cdot 10^{-5} \left(\frac{m}{s}\right)^{0.67} \\ C_2 &= 0.04 \cdot \frac{D}{d_h} \cdot \left(\frac{d_h}{v}\right)^{0.75} \cdot \left(\frac{v}{D}\right)^{0.33} = \\ 0.04 \cdot \frac{1.5 \cdot 10^{-9} \frac{m^2}{s}}{0.17 \cdot 10^{-2} m} \cdot \left(\frac{0.17 \cdot 10^{-2} m}{10^{-6} \frac{m^2}{s}}\right)^{0.75} \cdot \left(\frac{10^{-6} \frac{m^2}{s}}{1.5 \cdot 10^{-9} \frac{m^2}{s}}\right)^{0.33} \Leftrightarrow C_2 = 7.99 \cdot 10^{-5} \left(\frac{m}{s}\right)^{0.25} \\ C_3 &= \frac{A_{membrane} \cdot z_+ \cdot F}{\left(t_+ - t_+\right)} = \frac{0.0218 m^2 \cdot 1 \frac{eq_-}{mole} \cdot 96485 \frac{C}{eq_-}}{\left(1 - \frac{1}{2}\right)} \Leftrightarrow C_3 = 4206.8 \frac{m^2 \cdot C}{mol} \end{split}$$

These constants can now be inserted into the expressions in equation (14):

$$\frac{I_{crit}}{c_b} = u^{0.33} \cdot (C_1 \cdot C_3) = u^{0.33} \cdot 0.15 \left(\frac{m}{s}\right)^{0.67} \cdot \frac{m^2 \cdot C}{mole} \quad \text{(laminar correlation)}$$

$$\frac{I_{crit}}{c_b} = u^{0.75} \cdot (C_2 \cdot C_3) = u^{0.75} \cdot 0.34 \left(\frac{m}{s}\right)^{0.25} \cdot \frac{m^2 \cdot C}{mol} \quad \text{(turbulent correlation)}$$

^{*} Experimentally determined as sketched in Example B

The natural logarithm of these expressions is plotted together with the data from Table 4 in Figure 10 in order to give straight lines.

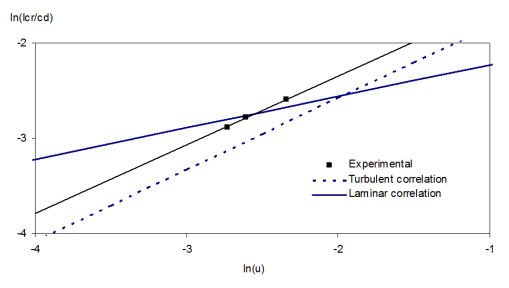


Figure 10: Hydrodynamic influence on the critical current Influence of hydrodynamic conditions (flow velocity) on the critical current visualized by plotting $ln(l_{cr}/c_{d})$ vs. ln(u). Icr: Critical current, cd: diluate concentration (mean logarithmic value), u: flow velocity. Experimental data are plotted together with expressions from literature correlations for mass transfer at turbulent or laminar flow.

In the plot one sees that the slope of the experimental values is close to the slope predicted by the turbulent correlation. This can be explained by the presence of turbulence promoters in the flow channels even though the Reynolds number is low. Overall the hydrodynamic influence (flow velocity) on the critical current seems to be well predicted by the literature correlations. One has to remember that many uncertainties are associated with using the flow correlations. The length of the flow channel for example is difficult to measure / estimate since many spacers have flow channels with many turns (tortuous-path spacers). Whether the flow is to be considered laminar or turbulent is also a question since many spacers are equipped with different insert in order to reduce the laminar boundary layer thickness and thus enhance the mass transfer coefficient. As mentioned earlier the diffusion coefficient is also concentration dependent and not totally constant. Finally the used flow equation from Table 2 arejust a few of many different flow equations for many different flow geometries so there might be correlation in the literature that fits this particular electrodialysis system more accurate.

But overall the experimental values of critical current, diluate concentrations and flow velocities seem to fit the literature correlations concerning flow and mass transfer quite well in this case.

2.3 Desalination degree

An import factor concerning the performance of an electrodialysis system is the desalination degree. The desalination degree indicates how large an amount of the salt in the diluate stream that is removed during one passage through the membrane stack. The desalination degree *S* can be calculated from the following equation:

$$S = \frac{c_{d,in} - c_{d,out}}{c_{d,in}}, \quad SI - units : \quad S = \left| \frac{\left(\frac{mole}{m^3}\right)}{\left(\frac{mole}{m^3}\right)} \right| = [-]$$

$$(15)$$

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The concentration in the diluate is denoted cd and from equation (15) is can be seen that the desalination degree can be determined from the diluate concentration that enters the dilution chambers and the diluate concentration that exits the dilution chambers. From the equation it is seen that a total desalination corresponds to a desalination degree of 1 (100%). The desalination degree is (among different factors) influenced by the flow rate of the feed stream. This will be shown in the following example:

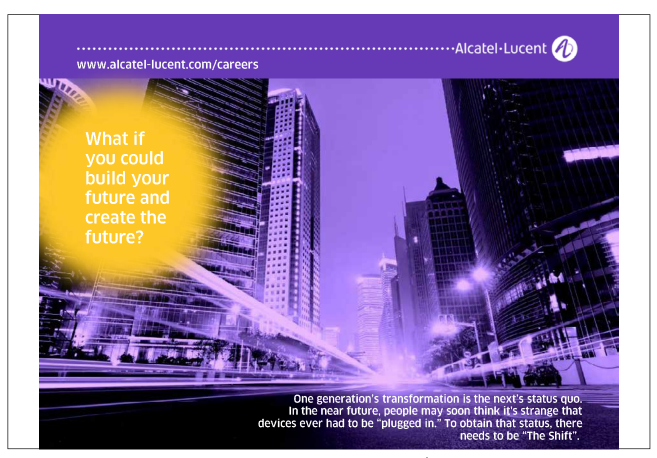
Example E: Desalination degree

In an electrodialysis system consisting of 20 cell pairs, the outlet concentrations from the diluate chambers are measured for experiments conducted at different flow rates at constant inlet concentration of 0.2 M. The concentrations are measured at a voltage difference over the membrane stack of 10 volts and the feed consists of sodium chloride solutions. Data from the experiments are shown in Table 5.

Flow rate [g/s]	Outlet concentration, c _{d,out} [mole/l]	Desalination degree, S
5.74	0.150	0.25
7.52	0.160	0.20
9.22	0.165	0.18
11.29	0.175	0.13

Table 5: Desalination degree

Feed concentration = 0.2 M, outlet concentration and desalination degree for experiments conducted at 10 volts at different flow velocities. Feed solutions consist of diluted sodium chloride.





It is seen that when the feed flow rate is increased the desalination degree decreases. The desalination degree vs. the total feed flow rate is plotted in Figure 11.

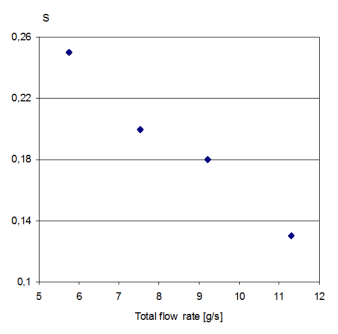


Figure 11: Desalination degree vs. feed flow rate Desalination degree (S) at different feed flow rates at a constant feed concentration (0.2 M). Voltage = 10 V.

The decreasing effect on the desalination degree at increasing feed flow rate is clearly seen in Figure 11. An increased feed flow rate decrease the time the salt "spend" in the diluate chamber in one passage and thus the "possibility" of desalination decreases.

2.4 Current efficiency

The current efficiency is a relevant term to consider. The current efficiency tells how many equivalents of salt that has been removed from the diluate to the concentrate pr. transferred current equivalents. In the ideal case the current efficiency equals 1 which corresponds to the case where one current equivalent is able to remove one salt equivalent. The following equation can be used to calculate the current efficiency η :

$$current \ efficiency = \frac{salt \ equivalents \ removed}{used \ current \ equivalents} \Rightarrow$$

$$\eta = \frac{Q_d \left(c_{d,in} - c_{d,out}\right) \cdot z}{n \cdot \left(\frac{I}{F}\right)}, \quad SI - units: \quad \eta = \left[\frac{\frac{m^3}{s} \cdot \left(\frac{mole}{m^3}\right) \cdot \frac{eq}{mole}}{\left(\frac{C}{s}\right)} - \frac{\left(\frac{eq}{s}\right)}{\left(\frac{eq}{eq}\right)}\right] = [-]$$
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The total flow rate in the dilution chambers is denoted Q_d , the number of cell pairs n, I is the current, z is the valence of the salt ions (eq./mole, z=1 if we are dealing with NaCl solutions) and F is the Faraday number (96485 C/eq.). The number of salt equivalents removed from the diluate to the concentrate can thus be calculated as the difference between "in-" and "out-concentration" multiplied by the flow rate and the valence of the ions. The number of used current equivalents is calculated as the current divided by the Faraday number. The number of cell pairs also has to be multiplied since every cell pair each transfers the same amount of current equivalents. From the SI-Units in equation (16) one sees that the current efficiency is a dimensionless term (just as the desalination degree described in section 2.3 Desalination degree). In the following example we will see an application of equation (16) for calculating the current efficiency from real electrodialysis data.

Example F: Current efficiency

An electrodialysis experiment is conducted on a small scale apparatus consisting of 20 cell pairs. A sodium chloride solution is used as feed stream and the in- and outlet concentration and the current is measured for an experiment conducted with a total diluate flow rate of 11.72 g/s and a voltage difference of 50 volts. Relevant parameters and experimental data are shown in Table 6.

Diluate flow rate, Q _d [m³/s]	11.72×10 ⁻⁶
Diluate inlet concentration, c _{d,in} [mole/m³]	160
Diluate outlet concentration, cd.out [mole/m³]	30
Number of cell pairs, n	20
Current, I [A = C/s]	8.09
Valence of salt ions, z [eq./mole]	1
Faraday number, F [C/eq.]	96485

Table 6: Experimental parameters

Parameters and experimental data for an electrodialysis experiment conducted with a sodium chloride feed solution.

The numbers given in Table 6 are directly inserted into equation (16) in order to calculate the current efficiency:

$$\eta = \frac{Q_d \left(c_{d,in} - c_{d,out}\right) \cdot z}{n \cdot \left(\frac{I}{F}\right)} = \frac{11.72 \cdot 10^{-6} \frac{m^3}{s} \cdot \left(0.16 \cdot 10^3 - 0.03 \cdot 10^3\right) \frac{mole}{m^3} \cdot 1 \frac{eq.}{mole}}{20 \cdot \left(\frac{8.09 \frac{C}{s}}{96485 \frac{C}{eq}}\right)} = 0.91$$

It is thus seen that the number of transferred salt equivalents are calculated from diluate inlet and outlet concentrations and the total diluate volume flow rate whereas the total number of current equivalents are determined from the current and the number of cell pairs. In this case it is thus seen that 0.91 equivalents of salt is removed pr. transferred equivalent of current.

2.5 Energy requirements

Evaluation of the energy consumption when running an electrodialysis system is of very great importance. Generally the energy consumption in electrodialysis can be divided into two parts:

- Pump energy: Energy to pump the solutions through the electrodialysis system
- Electrical energy: Electrical energy to transfer ions through the ion exchange membranes

The overall energy requirement is often dominated by one of these two terms depending on different process parameters and operation conditions.

2.5.1 Pump energy

In chemical engineering it is well know that the energy consumption associated with pumping liquids through a given system is dependent on the volume flow rate Q, the pressure drop in the system $\Box p$ and the efficiency of the pump η_{pump} . The pump energy can thus be calculated by use of the following equation:

Pump energy:
$$E = \frac{Q \cdot \Delta p}{\eta_{pump}}$$

$$SI - units: E = \left[\frac{m^3}{s} \cdot Pa = \frac{m^3}{s} \cdot \frac{N}{m^2} = \frac{m \cdot N}{s} = \frac{J}{s}\right]$$
(17)



Normally in an electrodialysis system three pumps are required as mentioned earlier:

- One pump to pump the solutions through the dilution chambers
- One pump to pump the solutions through the concentration chambers
- One pump to pump electrode solutions through the two electrode chambers

Thus the energy consumption to pump the solutions through an electrodialysis system can be calculated from knowledge of the pressure drops in the dilution-, concentration and electrode chambers, efficiencies of dilution-, concentration and electrode pumps and the volume flow rates in the three different types of chambers:

$$E_{pump,total} = \frac{Q_d \cdot \Delta p_d}{\eta_d} + \frac{Q_c \cdot \Delta p_c}{\eta_c} + \frac{Q_e \cdot \Delta p_e}{\eta_e}$$
(18)

The sup-scripts *d*, *c* and *e* refer to the diluate, concentrate and electrode solutions, respectively. The volume flow rates are normally easy to measure but determining the actual pressure drop in the three circuits can be more uncertain and difficult to measure.

2.5.2 Electrical energy

The energy consumption associated with removing ions from the diluate to the concentrate through the ion exchange membranes is proportional to the current I flowing through the membrane stack and the total voltage drop E between the two electrodes over the whole membrane stack. This energy consumption can be expressed as follows:

Electrical energy:
$$E_{electrical} = I \cdot E$$
, $SI - units$: $E = \left[\frac{C}{s} \cdot \frac{J}{C} = \frac{J}{s} \right]$ (19)

As for the pump energy in equation (17) the unit of the electrical energy in equation(19) is given in "energy pr. time" (Joules pr. second). If one knows the resistance of one single cell pair R_{cell} the total resistance of the membrane stack consisting of n cell pairs can be calculated by multiplying the number of cell pairs with the resistance of one single cell pair (according to equation (9)). Thus equation (19) can be rewritten when the voltage drop over the whole membrane stack E is replaced by product of the total resistance of the membrane stack and the current flowing through the membrane stack according to Ohm's law:

$$E_{electrical} = I^2 \cdot R_{cell} \cdot n \tag{20}$$

The amount of salt that is removed from the diluate to the concentrate is directly proportional to the amount of current flowing through the membrane stack (in the Ohmic region). The amount of current flowing through the membrane stack, the diluate volume flow rate Q_d and the current utilization \square determines the difference between the inlet and outlet concentrations of the diluate $\square c_d$ (the level of desalination). The current utilization is the product of the current efficiency η (from equation (16)) and the number of cell pairs n ($\square = \eta \times n$). This can be gathered in the following equation:

$$I = \frac{\Delta c_d \cdot Q_d \cdot z \cdot F}{\xi}, \quad SI - units: \quad I = \left[\frac{mole}{m^3} \cdot \frac{m^3}{s} \cdot \frac{eq}{mole} \cdot \frac{C}{eq} = \frac{C}{s} \right]$$
 (21)

Again the term z is the valence of the ions and F is the Faraday number. The different equations presented in this sub section can be used to calculate the energy requirements needed to remove a certain amount of ions from the diluate to the concentrate. Thus the electrical energy together with the energy required to pump the solutions through the system determines the overall energy consumption.

2.5.3 Thermodynamic least amount of energy

The energy consumption given in the previous section (2.5.2 Electrical energy) gives the actual energy consumption associated with ion transfer for a given electrodialysis system and this amount of energy is often much larger that the least amount of energy required just by looking at the "thermodynamic least amount of energy required". The thermodynamic least amount of energy can be considered as the numerical value of the amount of energy that is released when a given amount of diluate and concentrate are mixed together. The thermodynamic least amount of energy required can be calculated by use of the following equation:

$$E_{\text{ thermodynamic least amount}} = 2 \cdot R_{gas} \cdot Q_d \cdot T \cdot \left(c_{d,in} - c_{d,out} \right) \cdot \left[\frac{\ln \left(\frac{c_{d,in}}{c_{d,out}} \right)}{1 - \left(\frac{c_{d,in}}{c_{d,out}} \right)} - \frac{\ln \left(\frac{c_{d,in}}{c_{c,out}} \right)}{1 - \left(\frac{c_{d,in}}{c_{d,out}} \right)} \right]$$

$$SI - units: \quad \stackrel{\cdot}{E}_{\text{thermodynamic least amount}} = \left[\frac{J}{mole \cdot K} \cdot \frac{m^3}{s} \cdot K \cdot \frac{mole}{m^3} = \frac{J}{s} \right]$$
 (22)

In this equation it should be mentioned that " $R_{\rm gas}$ " is the gas constant, $Q_{\rm d}$ is the total volume flow rate of the diluate and T is the absolute temperature. As mentioned above, this thermodynamic least amount of energy is often much less that the actual electrical energy requirement for transferring ion through the membranes (equation (19) for example). This is due to the influence of different factors such as:

- The current efficiency η (equation (16)) is less than 1.
- The Ohmic resistance in the concentrate, diluate and electrode solutions and in the anion and cation exchange membranes causes generation of heat and thus loss of energy.

The thermodynamic efficiency can be evaluated by comparing the thermodynamic least amount of energy to remove ions from the diluate to the concentrate with the actual amount of energy used to remove ions from the diluate to the concentrate.

Thermodynamic efficiency =
$$\frac{E_{\text{thermodynamic least amount}}}{E_{\text{electrical}}} = \left(\frac{\text{equation } 22}{\text{equation } 19}\right)$$

SI-units: Thermodynamic efficiency = $\left[\frac{J}{s}\right] = [-]$ (23)

The thermodynamic efficiency is often very low (often below 5–10%) which for an electrodialysis experiment will be sketched in the following example.



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Example G: Thermodynamic efficiency

For an electrodialysis system consisting of 20 cell pairs the in- and outlet concentrations from the concentration and diluate chambers and the current were measured at each step of the voltage between 10 and 50 volts. The diluate flow rate was kept constant (6.6 g/s) and the feed solution consisted of sodium chloride. The temperature was also kept constant. At each voltage-step the electrical energy consumption was calculated by using equation (19):

$$E_{electrical} = I \cdot E$$

The thermodynamic least amount of energy was also calculated at each voltage-step from the volume flow rate of the diluate and the different concentrations by use of equation (22):

$$\dot{E}_{\text{ thermodynamic least amount}} = 2 \cdot R_{gas} \cdot Q_d \cdot T \cdot \left(c_{d,in} - c_{d,out}\right) \cdot \left| \frac{\ln \left(\frac{c_{d,in}}{c_{d,out}}\right)}{1 - \ln \left(\frac{c_{d,in}}{c_{d,out}}\right)} - \frac{\ln \left(\frac{c_{d,in}}{c_{c,out}}\right)}{1 - \ln \left(\frac{c_{d,in}}{c_{c,out}}\right)} \right|$$

The thermodynamic efficiency is the defined as the ratio between the thermodynamic least amount of energy and the actual electrical energy requirement according to equation (23):

Thermodynamic efficiency =
$$\frac{\dot{E}_{\text{thermodynamic least amount}}}{\dot{E}_{\text{electrical}}} = \left(\frac{equation \ 22}{equation \ 19}\right)$$

The thermodynamic efficiency is plotted as a function of the voltage for the experiment in Figure 12.

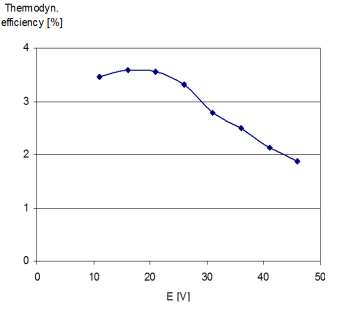


Figure 12: Thermodynamic efficiency vs. voltage

The thermodynamic efficiency vs. the voltage for an electrodialysis experiments conducted at a constant diluate flow rate of 6.6 g/s and a sodium chloride feed solution of 0.16 mole/l.

It is seen in Figure 12 that the thermodynamic efficiency throughout the whole experiment is very low and in all cases below 4%. It is also seen that at low voltage the thermodynamic is higher than at large voltage. An explanation could be that at these conditions the irreversible processes at the electrodes do not play as large a role at when the voltage is increased. Therefore the efficiency at low voltage is "closer" the thermodynamic least amount of required energy than at larger voltage.

2.6 Anti-fouling mechanism

When electrodialysis is applied commercially to desalt seawater, fouling of the ion exchange membranes can be a severe problem as in every other membrane separation process. Particle material, colloids, macromolecules and other charged or uncharged components can block the membrane pores or membrane surface and thus increase the resistance of the membrane. This will increase the operation costs. In electrodialysis in which a current flows in one direction, especially larger charged components can be transferred to the membrane surface but because of their size they are not able to pass through the membrane. Scaling on the membrane surface at high concentrations can also create problems. Because most of the colloids and macromolecules that appear in the water are negatively charged, it is mostly the anion exchange membranes that are affected by the fouling problems. In some commercial electrodialysis system this is avoided by changing the direction of the current every 30 minutes for example. Of course this leads to changes of the chambers every 30 minutes:

- Diluate chambers → Concentrate chambers
- Concentrate chambers → Diluate chambers

This means that the outlet from the chambers in the first short period after changing the direction of the current can not be used as concentrate and diluate. But after this period the concentrate and diluate can again be collected from the outlet of the membrane stack. By doing this changing of current direction the charged larger particles and components that eventually will stick to the membrane surfaces will every 30 minutes be removed and moved to the other side the chamber and thus a build up of a large fouling layer over a longer period is partly avoided. Thus the overall resistance of the membrane stack will by average be kept constant and the running costs will be kept constant.

3 Summary

In this book we have learned about basic terms in electrically driven membrane processes. The core in electrically driven membrane processes are membranes incorporated with fixed charges:

- Anion exchange membranes and
- Cation exchange membranes.

We have been looking at the fluxes of ions through these membranes and we have seen that these fluxes depend on the transport numbers of the ions. The membranes achieve their selectivity because these transport numbers for the oppositely charged ions are very different inside the membrane whereas the transport numbers in the solution for the oppositely charged ions are almost equal for equal valences. It is also explained that the separation principle in electrical driven membrane processes is based on Donnan exclusion.



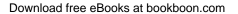
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One major part of the electrical driven membrane processes is electrodialysis. Electrodialysis has many applications and one of the greatest is the desalination of saline water into potable water. The basic concepts of electrodialysis are explained as well as the existence of a critical current. Examples are included in order to show ways to determine this critical current experimentally and from literature correlations based on dimensionless numbers concerning flow and mass transfer. The influence of the hydrodynamic conditions on the critical current is also explained and discussed. Other relevant terms used in the evaluation of electrodialysis systems such as the desalination degree and the current efficiency are also described and discussed through relevant examples. The energy required to run electrodialysis systems can be divided into the energy required to pump the solutions through the system and the energy required to transfer ions through the ion exchange membranes. Relevant equations are given in order to calculate these energy requirements and the thermodynamic efficiency is also defined. Finally some fouling problems are briefly mentioned together with an approach for avoiding these fouling problems.

Other electrically driven membrane processes such at the REED process, Donnan dialysis, Membrane electrolysis, the use of bipolar membranes and finally fuel cells can also be mentioned. Some of these processes can have the potential to grow in the future.

Endnotes

- 1. It shall however be mentioned that the enhanced concentration in the boundary layer on the concentrate side can lead to scaling of ionic solids at the membrane surface on the concentrate side for example if the concentrate contains calcium ions.
- 2. This happens at the cation exchange membrane before it happens at the anion exchange membrane. This is due to fact that the electrical mobility of the cations is generally a bit lower than for the anions. In the case of sodium chloride the transport number of sodium ions is approximately 0.4 and the transport number for chloride ions is approximately 0.6 (and actually not equal which is stated in equation (7). This is because the friction between the water molecules and the ions is largest for cations. This means that at a given current the concentration at the membrane surface on the diluate side (c'_d) will reach zero at the cation exchange membrane before is reaches zero at the anion exchange membrane. Thus the limiting current will be attained faster at the cation exchange membrane than at the anion exchange membrane.
- 3. Other kinds of transport concerning the transport of water molecules through the ion exchange membranes are also observed in electrodialysis. Osmotic transport of water from the diluate to the concentrate can be observed because of the concentration difference. Electroosmotic water transport can also be observed as sketched in Figure 3. where water molecules are dragged along with the electrically transported ions.
- 4. When, however, the feed solution ion concentration is increased and reaches a level of concentration as the fixed charges in the ion exchange membrane, some co-ions may also enter the membrane which of course is unwanted since the membrane then looses it selectivity. That way the current efficiency becomes very low and the electrical energy consumption increases.
- 5. From M. Mulder, "Basic principles of membrane technology", 2nd edition, Kluwer Academic Publishers, 1996.