Destabilization and removal of immobilized enzymes adsorbed onto polyethersulfone ultrafiltration membranes by salt solutions

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Abstract

In this work the effectiveness of two saline solutions (NaCl and Na$_2$SO$_4$) to clean a permanently hydrophilic polyethersulfone (PESH) ultrafiltration (UF) membrane with a molecular weight cut-off (MWCO) of 30 kDa previously fouled with enzymatic solutions
was investigated. The influence of protein concentration in the enzymatic solution during the fouling step and the effect of salt type during the cleaning procedure were studied. The protein aggregation was analysed in solution and onto the membrane surface by using several techniques including Dynamic Light Scattering (DLS), Atomic Force Microscopy (AFM) and Infrared Spectroscopy with Attenuated Total Reflectance (ATR-FTIR). In addition, mechanisms that dominate membrane fouling were studied by fitting some mathematical models (Hermia’s models adapted to crossflow filtration, a combined model based on the complete blocking and cake formation equations and a resistance-in-series model) to the experimental data.

Fouling results showed that the complete blocking/adsorption on membrane surface was the predominant fouling mechanism. Regarding the cleaning results, higher cleaning efficiency and low residual protein concentration was obtained with NaCl solutions for all the feed solutions tested due to the favourable interaction between Cl⁻ and proteins.

**Keywords:** membrane cleaning; protein fouling; salt solutions; ultrafiltration; mathematical models

### 1. Introduction

Fouling due to protein-membrane interactions can influence and strongly limit the performance of membranes in several processes such as: proteins separation/fractionation, removal of denatured enzymes in biocatalytic membrane reactors and clarification of food streams.

It is well known that pectinases are enzymes largely used in fruit juice microfiltration (MF) or ultrafiltration (UF) in order to hydrolize pectic substances, i.e. complex glycosidic macromolecules with high molecular weight and negative charge [1] which are responsible
for membrane fouling. In addition, pectinases can be also immobilized on different supports depending on the industrial application including fruit juice and olive mill wastewater processing [2, 3].

Enzyme immobilization techniques and membrane fouling mechanisms have similar characteristics, physical and/or interfacial donor/acceptor interactions being surface controlling factors [4-7]. In this way, Luo et al. [8] stated that enzymatic entrapment on membranes may be considered as a pore blocking mechanism and the adsorption fouling can be related to the enzymatic adsorption on the membrane surface due to hydrophobic and electrostatic interactions. In their work, fouling mechanisms that dominate the enzyme filtration were analyzed by two different mathematical models: a resistance-in-series model considering the intrinsic membrane resistance, the reversible fouling resulting from concentration polarization layer or particle deposit and the irreversible fouling including pore blocking or cake deposit and the classical Hermia’s models applied to dead-end filtration. Other authors also used these models to investigate membrane fouling. Vincent-Vela et al. [9] fitted the Hermia’s models adapted to crossflow filtration to the experimental data obtained during the UF of polyethylene glycol. They concluded that the intermediate and complete blocking mechanisms were the models with the highest fitting accuracy for most of the operating conditions tested. De Barros et al. [10] identified the fouling mechanism as a function of the membrane material used during the crossflow UF of pineapple juice previously treated with enzymes. Results after fitting the Hermia’s models adapted to crossflow showed that complete pore blocking and cake formation were the predominant mechanisms when using the ceramic and polymeric membranes, respectively. Similarly, Cassano et al. [11] established that the fouling mechanism involved in the crossflow UF of blood orange juice with a tubular polyvinylidene fluoride (PVDF) membrane evolved from a partial to a complete pore blocking in dependence of the axial
feed velocity. Choi et al. [12] applied a resistance-in-series model to the MF of bovine serum albumin (BSA) taking into account the membrane resistance, the gel layer resistance and the fouling resistance due to the foulant deposits inside the membrane pores. They obtained good agreement with the experimental data recorded. Machado et al. [13] investigated the effect of an enzymatic treatment with pectin lyase of açai pulp on its crossflow MF. They also studied the predominant fouling mechanisms by a resistance-in-series and the Hermia’s models adapted to crossflow ones. They observed that fouling resistance decreased after the enzymatic treatment as well as the cake formation followed by intermediate and complete pore blocking mechanisms dominated the MF process.

However, as Luo et al. described [8], during the immobilization of enzymes the convective transport of enzymes to the membrane surface makes these particles to be retained on it and thus, enzymes cause an increase in the local concentration. As a result, concentration polarization increases and a fouling layer can be formed on the membrane surface or inside its pores. In order to remove such a fouling layer, membranes have to be cleaned. Conventional cleaning protocols involve chemical cleaning agents such as alkalis, acids, disinfectants, surfactants or combinations of them [14]. However, these conventional cleaning methods may be aggressive for the membranes and their lifetime and selectivity may be reduced. In addition, these conventional cleaning agents cause a negative environmental impact when they are discharged as wastewaters after the cleaning step. For all these reasons, new alternative cleaning techniques, including ultrasounds [15], electromagnetic fields [16] and saline solutions [17] have been developed in the recent years to overcome these problems. Regarding the use of saline solutions, previous studies reported the salting-in and salting-out effect of different cations and anions to increase or decrease, respectively, protein solubility at pH values above and below the isoelectric point of the protein [18, 19]. Lee et al. [17] also investigated the cleaning mechanism of saline
solutions: in a first stage, the difference between the bulk solution concentration and the
gel layer concentration on the membrane surface causes changes on the cross-linked
fouling layer. Then, an ion-exchange reaction takes place between the ions forming the gel
layer on the membrane surface and the salt ions of the cleaning solution that diffuse in the
gel layer. This ion exchange results in a swelling and removal of the gel layer by freeing
the protein molecules and favouring their transport to the bulk solution.

The aim of this work is to investigate the effectiveness of NaCl and Na$_2$SO$_4$ solutions for
cleaning a PESH UF membrane with a molecular weight cut-off (MWCO) of 30 kDa that
was previously fouled with enzymatic solutions of pectinases at different protein
concentration (2, 7.5 and 15 g/L). The influence of protein concentration on membrane
fouling was studied by AFM and ATR-FTIR techniques and the fouling mechanism that
dominates the UF of each feed solution tested was also determined by fitting three
mathematical models to the experimental data: Hermia’s models adapted to crossflow
filtration, a combined model based on the complete blocking and cake formation
mechanisms and a resistance-in-series model. The membrane cleanliness was evaluated by
ATR-FTIR measurements and the residual protein concentration adsorbed on the
membrane surface was also quantified.

2. Materials and methods

2.1. Chemicals

Enzymatic solutions of pectinases (Pectinex Smash XXL, Novozymes) at different protein
concentrations (2, 7.5 and 15 g/L) were used as feed solutions during the fouling step.
According to the manufacturer information, the enzymatic solution is mainly composed of
pectin lyases from *Aspergillus niger*. NaCl and Na$_2$SO$_4$ solutions were the cleaning agents
tested. Protein concentration in the enzymatic solutions was determined by the Bicinchoninic Acid (BCA) assay (Sigma Aldrich).

2.2. Experimental set-up

A permanently hydrophilic polyethersulfone (PESH) ultrafiltration membrane (Microdyn Nadir, Germany) with a MWCO of 30 kDa was tested in the experiments. The main characteristics of the virgin membrane are shown in Table 1.

All the fouling and cleaning experiments were performed in a conventional crossflow UF system. It consisted of a feed tank of 1 L, a variable speed pump, a stainless-steel cell able to accommodate a flat-sheet membrane and a permeate tank. Transmembrane pressure was measured by two manometers allocated before and after the membrane module and regulated by a pressure control valve located on the retentate line. Crossflow velocity was controlled by a digital flowmeter. Temperature was set at 25 ± 1 ºC during the fouling and rinsing steps using a cooling system fed with tap water, while during the cleaning step temperature was set at 50 ± 3ºC using a heater.

2.3. Experimental procedure

Fouling tests were carried out with an enzymatic solution at three different protein concentrations (2, 7.5 and 15 g/L) without pH adjustment (pH value of about 4.3). Experimental conditions of transmembrane pressure, crossflow velocity and temperature were set at 2 bar, 2 m/s and 25 ºC, respectively. The duration of the fouling tests was 2 h. These conditions were selected according to previous studies on protein solutions ultrafiltration [20-22].

After the fouling step, a first rinsing with distilled water was performed during 30 min at a transmembrane pressure of 1 bar and a crossflow velocity of 2.18 m/s. According to the
literature [23], low transmembrane pressures and high crossflow velocities favour the removal of solute molecules deposited on the membrane surface. Then, a cleaning step with saline solutions during 60 min and a second rinsing step with distilled water during 30 min were carried out at the same experimental conditions of transmembrane pressure as the first rinsing step. NaCl and Na$_2$SO$_4$ solutions were tested during the cleaning protocol at a concentration of 5 mM and a temperature of 50 °C. These conditions were the optimal ones in previous studies about saline cleaning of membranes fouled with protein solutions [20, 21].

Permeate flux and hydraulic resistance were monitored during all the steps of the experimental procedure. Once the cleaning procedure was finished, water permeability was measured again in order to evaluate the cleaning efficiency, expressed as:

$$CE = \frac{WP_1}{WP_0} \times 100$$  \hspace{1cm} \text{Eq. 1}

where $WP_1$ and $WP_0$ are the water permeability of the cleaned and virgin membrane, respectively.

2.4. Characterization of the enzymatic solutions

The BCA assay was used to determine the protein concentration in the enzymatic solutions [24, 25]. This assay is a colorimetric method based on the formation of a Cu$^{+2}$-protein complex under alkaline conditions. Then, reduction of Cu$^{+2}$ to Cu$^{+1}$ takes place and the amount of reduction is directly proportional to the amount of protein in the sample. According to the standard assay protocol, 0.1 mL of sample were mixed with 2 mL of the BCA working reagent and incubated at 37 °C during 30 min. Then, samples were cooled at room temperature and absorbance was measured at a wavelength of 562 nm by a UV-visible spectrophotometer. By using a calibration test performed with the bovine serum
albumin (BSA) standard solution, protein concentration in the enzymatic solutions was estimated. Each sample was duplicated for the measurement.

Size measurements of protein particles were carried out by Zetasizer nano (Malvern Instruments). The Zetasizer system determines the particles size by measuring the Brownian motion of the particles in a sample using Dynamic Light Scattering (DLS). Previous studies demonstrated that these techniques were suitable for protein characterization [26, 27]. In order to analyse the particle size in pectinase solution in the conditions in which membrane was fouled with pectinase (at 25 ºC) and during cleaning treatment (at 50 ºC in presence of NaCl), different analysis were carried out by DLS varying protein concentration (2, 7, 15 g/L). In addition protein size measurements at 15 g/L were also carried out in presence of NaCl (5mM). Hypothesis testing considering a confidence interval of 95 % and using the Statgraphics® Centurion XVI software were performed in order to determine if statistically significant differences exist among the particle size distributions at 25 ºC, 50 ºC and the combination of 50 ºC and NaCl [28].

2.5. Membrane characterization

Prior to the fouling experiments, the membrane was compacted with ultrapure water by increasing and decreasing the transmembrane pressure from 1 bar to 4 bar until a constant hydraulic permeability was obtained. Accordingly, the intrinsic membrane resistance ($R_m$) was calculated. Ultrapure water was used in order to prevent additional dirty on the membrane surface.

Infrared spectra in ATR mode were collected onto the membrane surface before and after each fouling and cleaning experiments by using an ATR-FTIR spectrometer (Spectrum One, Perkin Elmer). The adsorption and deposition of proteins on the membrane surface and related changes in the surface roughness were detected by using Atomic Force
Microscopy (AFM), Nanoscope III (Digital Instruments, VEECO Metrology Group). Tapping Mode AFM operated by scanning a tip attached to the end of an oscillating cantilever across 2.5x2.5 µm of sample surface.

The amount of residual material adsorbed onto the membrane surface was quantified according to the following equation [29]:

$$\frac{H_{1539}}{H_{1240}} = 0.0034C_{protein} + 0.0165$$  \hspace{1cm} \text{Eq.2}

where $H_{1539}$ is the height of the absorption band at 1539 cm$^{-1}$ (amide 2), $H_{1240}$ is the height of the absorption band at 1240 cm$^{-1}$ (PES membrane) and $C_{protein}$ is the residual protein concentration deposited on membrane surface. This equation is valid in the protein concentration range of 0.5-350 µg/cm$^2$ with a maximum deviation of 1 µg/cm$^2$, according to [30].

2.6. Mathematical modelling

Different mathematical models were fitted to the experimental data obtained during the UF of pectinase solutions using the MathCad® Genfit algorithm, which minimizes the overall difference between experimental and predicted results by the Levenberg-Marquadt method. The success of fitting was evaluated in terms of regression coefficient $R^2$ and standard deviation SD.

2.6.1. Hermia’s models

Hermia [31] developed four models based on classical constant pressure dead-end filtration equations in order to describe four main types of membrane fouling: complete blocking, intermediate blocking, standard blocking and cake layer formation. Several authors [9-11, 32] adapted these models to crossflow configuration by incorporating the flux associated
with the back-transport mass transfer, which is evaluated at the steady-state [33]. Eq. 3 shows the general equation for Hermia’s models adapted to crossflow UF:

\[
-\frac{dJ}{dt} = K(J - J_{ss})J^{2-n}
\]

Eq. 3

where \(J\) is the permeate flux, \(K\) is a model constant, \(n\) is the model parameter and \(J_{ss}\) is the permeate flux when the steady-state is achieved.

Four different models corresponding to the four fouling mechanisms above mentioned are considered depending on the value of the parameter \(n\): complete blocking (\(n = 2\)), intermediate blocking (\(n = 1\)), standard blocking (\(n = 1.5\)) and gel layer formation (\(n = 0\)).

Complete blocking model takes into account that a solute molecule that reaches to membrane surface blocks a pore entrance completely without penetrating inside the pores. This model assumes that a monomolecular layer is formed on the membrane surface.

The intermediate blocking model considers that a solute molecule can deposit on previously settled ones. However, as in the complete blocking model, fouling takes place only on the membrane surface.

When solute molecules are smaller than membrane pore size, these molecules can penetrate inside the pores. This is the main hypothesis of the standard blocking model.

Cake formation model is based on the assumption that solute molecules accumulate on membrane surface because they have a larger size than membrane pores. Therefore, a permeable layer is formed on the membrane surface.

2.6.2. Combined model

Previous studies [32-35] reported that the typical variation of permeate flux with time includes two fouling mechanisms: a rapid flux decline during the first minutes of operation due to pore blocking and an accumulation of foulants on the membrane surface that causes a long term flux decline due to a cake formation. In this work, a combined model
considering the Hermia’s equations for crossflow UF corresponding to complete blocking and cake formation was used to predict the permeate flux decline. Therefore, the general equation of the combined model is Eq. 4, where $\alpha$ is the fraction of membrane pores that are completely blocked:

$$J_{\text{combined model}} = \alpha J_{\text{complete blocking model}} + (1 - \alpha) J_{\text{cake formation model}}$$  \hspace{1cm} \text{Eq. 4}

Eq. 4 involves two different constants depending on the fouling mechanism: $K_c$ for the complete blocking model and $K_{cf}$ for the cake formation model. The parameter $K_c$ corresponds to the blocked membrane surface per unit of total permeate volume and unit of membrane surface porosity. On the other hand, $K_{cf}$ represents the ratio between the cake characteristics (specific cake resistance and cake mass per unit of total permeate volume) and the original membrane ones [9].

2.6.3. Resistance-in-series model

According to the Darcy’s law (Eq. 5), permeate flux is related to the transmembrane pressure and the total hydraulic resistance:

$$J = \frac{\Delta P}{\mu R}$$  \hspace{1cm} \text{Eq. 5}

where $\Delta P$ is the transmembrane pressure, $\mu$ is the feed solution viscosity and $R$ is the total hydraulic resistance.

In the resistance-in-series model, the total hydraulic resistance is the sum of different resistances that contribute to the permeate flux decline during the UF process. In this work, the membrane resistance, the cake resistance and the adsorption and concentration polarization resistances were considered (Eq. 6):

$$J = \frac{\Delta P}{\mu(R_m + R_{ad} + R_c)}$$  \hspace{1cm} \text{Eq. 6}
where $R_m$ is the virgin membrane resistance, $R_c$ is the cake resistance and $R_{ad}$ is the resistance due to adsorption on/in membrane surface and pores and concentration polarization. In addition, $R_{ad}$ can be expressed as an exponential function of the steady-state adsorption and concentration polarization resistance and the rate at which foulant molecules are deposited on the membrane [12, 36]. Therefore, Eq. 7 shows the general equation for the resistance-in-series model:

$$J = \frac{\Delta P}{\mu \left(R_m + R'_{ad} \left(1 - \exp(-bt)\right) + R_c\right)}$$  

where $R'_{ad}$ is the steady-state adsorption and concentration polarization resistance and $b$ is the fouling rate due to adsorption.

3. Results and discussion

3.1. Fouling results

3.1.1. Determination of protein fouling

DLS measurements on protein solutions (Table 2) showed a bimodal distribution for all the protein concentrations tested at 25 °C. As it is possible to observe, the particle size belonging to distribution 2 increased as a function of concentration, reaching a value of 410 ± 221 nm at 15 g/L. The increase in size, as well as the high PDI, suggested a severe protein aggregation which is expected to lead to a higher fouling during the UF process when the protein concentration is increased.

AFM and ATR-FTIR measurements also demonstrated that protein fouling increased with the protein concentration in the enzymatic solution, causing a general increase in the surface roughness as well. Fig. 1 shows the 2D AFM images of the virgin and fouled membranes at different protein concentration (2, 7.5 and 15 g/L). When protein concentration was 2 g/L big protein aggregates can be well distinguished from membrane surface while the membrane surface continues to exhibit the same surface morphology of
the initial one, causing a dramatic increase in the surface roughness with value of $R_a$ of 28±8 nm (Fig. 1b). At 7.5 g/L the layer is partially formed on the membrane surface, revealing heterogeneous regions where initial membrane structure and isolated protein aggregates are well distinguished (Fig. 1c). In comparison with previous situation (2 g/L) the protein clusters decrease in size but increase in number. At the highest concentration tested, a gel layer that completely covers the original membrane surface can be observed, producing an increase in surface roughness (Fig. 1d) as compared with the virgin membrane. On this basis it can be assumed that protein-membrane interaction, as well as protein cluster disassembling, is favoured when protein concentration is raised. On the other hand, it is relevant to underline that the convective transport to the membrane enhances protein overlapping with gel formation. These results are in agreement with those obtained by Ohnishi et al. [37], who demonstrated protein layer was denser at the highest concentration tested, as it occurs in our work.

ATR-FTIR spectra were collected onto virgin and fouled membranes. An intense absorption band can be detected at about 1650 cm$^{-1}$ due to the stretching vibration of C=Oamide typical of the amide I group (Fig. 2a). This band becomes more intense for membranes fouled with solutions at 2 g/L. The broadness of this band along with the appearance of a strong IR mode at 1504 cm$^{-1}$, which is typical of a C-H$_{arom}$ bond, suggests a strong contribution of the aromatic component to the spectrum due to different rearrangement of the protein aggregates when binding the membrane surface. Also, at 1168 cm$^{-1}$ an IR mode associated to C-O-C stretching vibrations appears to be more intense confirming a different local chemical environment due to a varied rearrangement of the protein assembling. It is noteworthy that stretching located at 1620 towards higher frequency suggests intermolecular β-sheets bonds confirming a protein aggregation state [38, 39].
Another diagnostic weak band is observed at around 1730 cm\(^{-1}\), which corresponds to C=O ester bond. This infrared mode was detected in the spectra for all the fouled membranes with the same band intensity, whereas it was never observed in the new membrane spectrum. It is relevant to underline that the strong IR absorption bands, which characterize the spectrum of fouled membranes with 2.0 g/L, are due to local adsorption of protein aggregates completely spanned at higher concentration as confirmed by AFM images (Figs. 1b,d).

3.1.2. Mathematical modelling

The mean value of the intrinsic membrane resistance (R\(_m\)) was \(3.8 \times 10^{12}\) m\(^{-1}\). This value was taken as a constant in the resistance-in-series model.

Fig. 3 shows the time evolution of the experimental permeate flux obtained for all the feed solutions tested. As expected, a lower permeate flux was measured by increasing the protein concentration due to a more severe membrane fouling. This is in accordance with the AFM image (Fig. 1d) where all the membrane pores appear to be fully covered by a protein layer. Fig. 3 also shows the permeate flux predictions by the two models with the highest fitting accuracy (highest regression coefficient, R\(^2\), and lowest standard deviation, SD). In all cases, resistance-in-series and the combined model were the models with the best fitting accuracy for all the feed solutions tested, with values of R\(^2\) ranging from 0.91 to 0.99 and SD values from 0.01 to 0.04, while the accuracy of Hermia’s models varied from 0.85 to 0.96 for R\(^2\) values and from 0.03 to 0.06 for SD. Both resistance-in-series and combined models considered that complete blocking/adsorption and cake formation were the main mechanisms contributing to membrane fouling. Therefore, both mechanisms should be considered to explain the membrane fouling with enzymatic solutions at the experimental conditions tested (2 bar, 2 m/s and 25 °C).
Table 3 shows the values of model parameters for the best fitting models. The values of the parameters $R_{\text{ad}}'$, $R_c$, $K_c$, $K_{cf}$ and $\alpha$ increased when protein concentration increased. This is due to the fact that more severe membrane fouling occurs with increasing protein concentration in the feed solutions. A similar behaviour was observed in the MF process of BSA solutions [35].

Regarding the value of the parameter $\alpha$, which indicates the fraction of completely blocked pores (Eq. 4), the predominant fouling mechanism was complete blocking for all the feed solutions tested. This is in agreement with the particle size measured: the largest size of pectinase molecules (50 kDa) compared to the membrane MWCO (30 kDa) allows these particles to be deposited on the membrane surface blocking the pore entrance. On the other hand, concentration polarization increased when protein concentration increased [33] and thus, the value of the resistance $R_{\text{ad}}'$ that considers both adsorption and concentration polarization phenomena at steady-state, was the highest at 15 g/L. In addition, as displayed in AFM images (Fig. 1d), at the highest concentration tested, the cake formed on the membrane surface was denser. This resulted in the increased parameters related to the cake formation, $R_c$ and $K_{cf}$. Regarding the intermediate pectinase concentration (7.5 g/L), the values of all these parameters also increased from those obtained at 2 g/L, due to the partial formation of the protein layer on the membrane surface (Fig. 1c). However, the values of the model parameters were much lower than those obtained at 15 g/L, as there were initial membrane regions uncovered at the intermediate protein concentration. However, the same pattern was not observed for the parameter $b$, which represents the rate at which foulant molecules are adsorbed on membrane surface. In this case, when protein concentration increased, the value of $b$ decreased. As the ATR-FTIR spectra of the fouled membranes indicated, at the lowest concentration used (2 g/L), the protein-membrane interaction became greater due to the low amount of proteins compared to that when working at 15
g/L. As a consequence, proteins were deposited on membrane surface at a higher rate than in the case of high protein concentration.

3.2. Membrane cleaning

Fig. 2b shows the spectra of the virgin and the cleaned membranes that were fouled with enzymatic solutions at protein concentrations of 2, 7.5 and 15 g/L. As it can be observed, an almost total overlapping among all the spectra was obtained, which is an indicator of the removal of protein deposits on the membrane surface using NaCl as cleaning agent at a concentration of 5 mM and 50 °C. The only exceptions are the appearance of a band at a frequency of 1750 cm$^{-1}$, which can be attributed to ester carbonyl and a band at a frequency of 1040 cm$^{-1}$, which can be related to the C-N stretching vibrations typical of proteins [40]. The band intensity is somewhat significant on fouled membranes (Fig. 2a) whereas it remained becomes negligible after the cleaning procedure (Fig. 2b). This provides indications about the efficiency of the cleaning procedures, the presence of some residual contaminants being onto the membrane surface.

Table 4 shows the results of residual protein concentration adsorbed on membranes and of the cleaning efficiency. Regarding the values of protein concentration after NaCl cleaning, it can be observed that all membranes showed a low amount of proteins on their surface, even if a gradual increase in the residual adsorption was detected for membranes fouled at higher protein concentration. Indeed the deposition of larger amount of protein at higher concentration was also revealed by AFM analyses. These results are in good agreement with those obtained by Rabiller-Baudry et al. [30] which studied the efficiency of several cleaning agents for PES UF membranes fouled with skim milk. In this case the best cleaning agents led to a residual protein concentration from 3 to 11 µg/cm$^2$ at a transmembrane pressure of 1 bar. On the other hand, according to the cleaning efficiency
evaluation (Table 4), it can be concluded that the low protein amount deposited on the membrane surface for the different protein concentrations tested did not greatly affect the membrane permeation properties.

The effect of salt type on the cleaning efficiency was also studied by means of ATR-FTIR measurements. Fig. 2c shows the spectra of the virgin and cleaned membranes, previously fouled with a 15 g/L protein concentration, by using NaCl and Na\textsubscript{2}SO\textsubscript{4} solutions. It is relevant to observe the appearance of two bands at 1504 cm\textsuperscript{-1} and 1168 cm\textsuperscript{-1} in the spectrum associated to membranes cleaned with Na\textsubscript{2}SO\textsubscript{4} (Fig. 2c). This suggests the presence of residual proteins not completely removed by the cleaning procedure. As previously described, these two bands indicate a different protein rearrangement on the membrane surface at lower protein concentration. Indeed, the spectrum is comparable to that recorded onto membranes fouled with 2 g/L pectinase (Fig. 2a). In addition, the shift of the infrared mode located around 1620 cm\textsuperscript{-1} towards higher frequency suggests a contribution of the band usually assigned to intermolecular \(\beta\)-sheets bonds, well known as “aggregation band” [38, 39]. This shift is also observable for membranes coming in contact with 2 g/L pectinase solution (Fig. 2a).

The residual protein concentration quantified from the ATR-FTIR spectrum was calculated for the Na\textsubscript{2}SO\textsubscript{4} cleaning protocol (Table 4). In this table, the greatest amount of protein deposited on the membrane surface after cleaning procedure was obtained with Na\textsubscript{2}SO\textsubscript{4} (24.68 µg/cm\textsuperscript{2}) and the cleaning efficiency was slightly lower than those obtained when NaCl was used as cleaning agent.

The different cleaning efficiency obtained with the two salt solutions can be ascribed to salting-in and salting-out mechanisms. Na\textsubscript{2}SO\textsubscript{4} showed a strong salting-out effect and enhanced protein aggregation and adsorption according to Tsumoto et al. [41]. In addition, SO\textsubscript{4}\textsuperscript{2-} exhibits a salting-out effect higher than Cl\textsuperscript{-}, causing a protein precipitation. This is
due to the different ability of anions to interact with water molecules [18]. According to the law of matching water affinities, monovalent anions (as Cl⁻) are weakly hydrated because they have a larger size than monovalent cations and they preferably interact with the positive-charged side chains of proteins and with the non-polar functional groups that are weakly hydrated as well [19]. This behaviour was also observed in a previous study about salt cleaning of membranes fouled with whey model solutions [21], in which similar efficiencies to those reported in Table 4 after the cleaning procedure were achieved.

On the basis of these experimental findings, DLS analyses were also performed in order to evaluate the combined effect of NaCl and temperature on protein aggregates destabilization/solubilisation. As it can be observed in Table 2, membrane cleaning with NaCl at 50 °C led to an additional protein aggregates destabilization that enhances salting-in effects. In particular, when the temperature was increased from 25 °C to 50 °C, with and without NaCl, the changes in size observed for the particles belonging to both the first and second distribution were statistically significant (p-values lower than 0.05), while a big effect of particle decrease is observed for the distribution 2. This effect demonstrated that the distribution 2 is characterized by big aggregates that, thanks to a temperature effect, are destabilized. In addition, for what concerns particles belonging to distribution 1, any substantial change is observed both increasing temperature and in presence of NaCl (5mM), but a greater change in size is observed for particles belonging to distribution 2. In this last case, the combination of NaCl and temperature (50 °C) resulted in a decrease in particle size from 288 to 240 nm. At higher temperature protein hydration is favoured. This means that protein-water bonds are predominant, causing solubilisation of protein aggregates. In the presence of NaCl, protein-protein interactions are further broken due to salting-in effects, thus promoting higher hydration shell [41,42].
4. Conclusions

Fouling experiments performed with enzymatic solutions of pectinases at different protein concentrations (2, 7.5 and 15 g/L) demonstrated that a higher protein concentration caused a more severe UF membrane fouling. AFM images as well as ATR measurements carried out on the virgin and fouled membranes confirmed that a continuous protein layer was formed on the membrane surface when a concentration of 15 g/L was used.

The mathematical models studied in this work predicted with high accuracy the experimental permeate flux decline corresponding to the UF of enzymatic solutions of pectinases at different protein concentration (2, 7.5 and 15 g/L), 2 bar and 2 m/s. The Hermia’s standard blocking model did not fit well the experimental data since the solute molecules size was larger than the membrane pore size and thus, these molecules cannot penetrate inside the membrane pores. Adversely, the models that showed the best fitting accuracy were the combined model and the resistance-in-series, one of which suggested a predominant pore blocking/adsorption mechanism, followed by a cake formation.

Among the salts tested, NaCl showed better results than Na$_2$SO$_4$ due to a better ability to dissolve proteins from membrane surface for salting-in effect. The cleaning efficiency of NaCl in combination with temperature was also investigated by DLS experiments, thus confirming a cooperation of the two parameters in the protein aggregates destabilization.

The cleaning efficiency was investigated through water permeability measurements. Although some residual contaminants were still present after cleaning with NaCl solution, the initial permeability value was completely restored. The suitability and reliability of the cleaning procedure with NaCl was confirmed with other protein solutions (whey model solutions), achieving similar cleaning efficiencies.
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References


**Table 1.** Main properties of the membrane used.

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<td>Commercial code</td>
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</tr>
<tr>
<td>Manufacturer</td>
<td>Microdyn Nadir</td>
</tr>
<tr>
<td>Type</td>
<td>Flat-sheet</td>
</tr>
<tr>
<td>Nominal MWCO (kDa)</td>
<td>30</td>
</tr>
<tr>
<td>Active layer</td>
<td>PESH</td>
</tr>
<tr>
<td>Effective area (cm²)</td>
<td>35.25</td>
</tr>
<tr>
<td>Water permeability at 25°C (L/m²·h·bar)</td>
<td>106.00</td>
</tr>
<tr>
<td>Maximum operating temperature (°C)</td>
<td>95</td>
</tr>
<tr>
<td>pH range</td>
<td>0-14</td>
</tr>
</tbody>
</table>

**Table 2.** Particle size measurements of pectinase solutions by DLS at various concentration and temperature.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Protein concentration (g/l)</th>
<th>Particle size distribution 1 (nm)</th>
<th>Particle size distribution 2 (nm)</th>
<th>PDI₁ (%)</th>
<th>PDI₂ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25 °C</td>
<td>15</td>
<td>15±4</td>
<td>410±221</td>
<td>27</td>
<td>54</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>6±1</td>
<td>91±38</td>
<td>17</td>
<td>42</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>6±0.6</td>
<td>61±18</td>
<td>10</td>
<td>29</td>
</tr>
<tr>
<td>50 °C</td>
<td>15</td>
<td>12±3</td>
<td>288±105</td>
<td>25</td>
<td>36</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>6±1</td>
<td>17±4</td>
<td>17</td>
<td>23</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>6±0.8</td>
<td>19±4</td>
<td>13</td>
<td>21</td>
</tr>
</tbody>
</table>

PDI: polydispersity index percentage = standard deviation/ intensity weighted Z average

**Table 3.** Values of model parameters for the best fitting models.

<table>
<thead>
<tr>
<th>Protein concentration (g/L)</th>
<th>Resistance-in-series model</th>
<th>Combined model</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R’ad (m⁻¹)</td>
<td>b (s⁻¹)</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>------------</td>
<td>---------</td>
</tr>
<tr>
<td>2</td>
<td>1.542·10¹³</td>
<td>3.637·10⁻⁴</td>
</tr>
<tr>
<td>7.5</td>
<td>8.865·10¹⁸</td>
<td>3.388·10⁻⁵</td>
</tr>
<tr>
<td>15</td>
<td>3.717·10¹⁷</td>
<td>8.994·10⁻⁹</td>
</tr>
</tbody>
</table>

**Table 4.** Residual protein concentration after cleaning procedure and water permeability recovery.

<table>
<thead>
<tr>
<th>Membrane</th>
<th>Protein concentration during fouling (g/L)</th>
<th>Residual protein concentration (µg/cm²)</th>
<th>Cleaning efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cleaned with NaCl</td>
<td>2</td>
<td>4±1</td>
<td>100±3</td>
</tr>
<tr>
<td></td>
<td>7.5</td>
<td>9±1</td>
<td>100±3</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>14±2</td>
<td>100±1</td>
</tr>
<tr>
<td>Cleaned with Na₂SO₄</td>
<td>15</td>
<td>25±1</td>
<td>97±1</td>
</tr>
</tbody>
</table>
Fig. 1. AFM images of new membrane (a) and membranes fouled with enzymatic solutions at 2 g/L (b), 7.5 g/L (c) and 15 g/L (d).
Fig. 2. ATR-FTIR spectra of new membrane and (a) membranes fouled with enzymatic solutions at 2, 7.5 and 15 g/L; (b) membranes cleaned with NaCl (enzymatic solution
concentration during fouling: 2, 7.5 and 15 g/L); (c) membranes cleaned with NaCl and Na₂SO₄ (enzymatic solution concentration during fouling: 15 g/L).
Fig. 3. Permeate flux predictions for the best fitting models using enzymatic solutions (lines: estimated results; symbols: experimental data).