Hybrid cross-linked chitosan/protonated-proline:glucose DES membranes with 1 superior pervaporation performance for ethanol dehydration 2 3 Roberto Castro-Muñoz^{1,2*}, Emilia Gontarek³, Jakub Karczewski⁴, René Cabezas⁵, 4 Gastón Merlet⁶, Claudio Araya-Lopez⁷, Grzegorz Boczkaj^{1,8} 5 6 ¹ Faculty of Civil and Environmental Engineering, Department of Sanitary Engineering, Gdansk University 7 8 of Technology, 11/12 Narutowicza St., 80-233, Gdansk, Poland ²Tecnologico de Monterrey, Campus Toluca. Av. Eduardo Monroy Cárdenas 2000 San Antonio 9 Buenavista, 50110, Toluca de Lerdo, Mexico. 10 ³ Faculty of Chemistry, Department of Process Engineering and Chemical Technology, Gdansk University 11 12 of Technology, 11/12 Narutowicza St., 80-233, Gdansk, Poland ⁴ Institute of Nanotechnology and Materials Engineering, Faculty of Applied Physics and Mathematics, 13 Gdansk University of Technology, 11/12 Narutowicza St., 80-233, Gdansk, Poland 14 ⁵Departamento de Química Ambiental, Facultad de Ciencias, Universidad Católica de la Santísima 15 16 Concepción, Concepción, Chile ⁶Departamento de Agroindustrias, Facultad de Ingeniería Agrícola, Universidad de Concepción, Chillán, 17 18 Chile ⁷Laboratory of Membrane Separation Processes (LabProSeM), Department of Chemical Engineering, 19 20 University of Santiago de Chile, Av. Libertador Bernardo O'Higgins 3363, 9170022, Estación Central, 21 Región Metropolitana, Chile ⁸Advanced Materials Center, Gdansk University of Technology, 11/12 Narutowicza St., 80-233, Gdansk, 22 23 Poland *E-mail: food.biotechnology88@gmail.com; castromr@tec.mx (R. Castro-Munoz) 24 25 *Corresponding Author 26

Abstract

This work explores a protonated L-proline:glucose (molar ratio 5:1) deep eutectic solvent (DES) in fabricating biopolymer membranes utilizing chitosan (CS). Initially, the miscibility of CS and DES to prepare homogeneous dense blend membranes has been investigated. Different techniques, such as scanning electron microscopy, contact angle (CA), atomic force microscopy (AFM), Fourier transformed infrared spectroscopy (FTIR) and swelling degree (uptake), were used to characterize the structure of the resulting membranes. Within the pervaporation performance for ethanol dehydration, Arrhenius and mass transfer analysis were analysed in detail. Interestingly, the addition of DESs provided superior performance to crosslinked CS: DES membranes compared with the ones lacking DES. Based on the morphology and properties observed, this new concept of CS-based membranes can be alternatively applied in other solvent separations requiring hydrophilic membranes.

- **Keywords:** Proline:glucose, deep eutectic solvents; chitosan; water-ethanol; hydrophilic

pervaporation.

1. Introduction

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A current trend of research deals with the implementation of deep eutectic solvents (DESs), which are an emerging class of sustainable solvents, into new applications and processes. Some examples of applications are as coatings [1], eco-friendly media in reactors [2], extraction of biomolecules from natural products, [3], separation of heavy metal from complex mixtures, CO₂ capture [4], fuel desulfurization [5], assisting chromatographic techniques [6], biodiesel synthesis aided with enzymes [7], chemical and biochemical reactions [8], among others. The importance of DESs relies on their non-toxic character, low cost in synthesis, easy to use, biodegradable, environmentallysafe and reusable [9]. As for DES synthesis, these eutectic mixtures tend to be typically formed by linking a hydrogen bond acceptor (HBA), usually quaternary ammonium salts, and a hydrogen bond donor (HBD) [1,10]. As for membrane fabrication, DESs are initially utilized as additive agents in DES-supported membranes [11,12], since they can confer a synergistic effect in the separation features of polymer membranes. In fact, such performance improvement showed by DESs in membranes is ascribed to a facilitated diffusion and adsorption of molecules within the functional groups present in DESs [13-15]. Apart from their application as additives, DESs were proposed as poreforming additives in phase inversion process via immersion precipitation for synthesizing membranes with asymmetrical patterns [16]. Very recently, our research group has synthetized and implemented a new DESs, based on an amino acid and an organosulfur solute, for fabricating dense chitosan (CS) membranes, which were subsequently applied into the pervaporation (PV) separation of methanol/MTBE mixtures [17]. CS was proposed in this study since it is the most

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investigated biopolymer for PV membranes with high performance for the removal of water molecules from less polar or non-polar molecules [18,19], e.g., CS has shown interesting permeation rates and salt rejection in seawater desalination via PV [20]. It is worth mentioning that biopolymer-based membranes (including CS, sodium alginate, cellulose acetate, polylactic acid, among others) are a current scope of study according to the need of replacing chemically synthesized polymers [21,22]. Apart from the implementation of DES in biopolymer membranes, the fabrication of mixed matrix membranes (or nanocomposites) through the incorporation of nanomaterials is also a current trend in the field [23-25]. Unfortunately, there is still a big need in finding new DES to assist the fabrication of polymer membranes and improve somehow their physicochemical properties and thus separation performance. It is worth mentioning that the main challenge comprises the rapport of the eutectic mixture and organic phase (i.e., polymer) towards their compelling joining. Here, in this research, an original hydrophilic eutectic mixture, like protonated L-proline:glucose at molar ratio 5:1, was used for the unprecedented tailoring of cross-linked CS membranes to beaten the typical issues associated with the suitable distribution of the eutectic mixtures inside the biopolymer. Strategically, the new DES owning hydrophilic character and soluble in water, such as protonated-L-proline:glucose under optimized molar ratio 5:1, has been eventually elaborated [26]. After this, the DES was blended with CS phase to fabricate a homogeneous dope solution, succeeded by an in-situ cross-linking via glutaraldehyde (GA). These final membranes were fully characterized and evaluated for practical separation application. PV experiments towards the removal of water from azeotropic water-ethanol model mixtures demonstrated the membranes' appropriateness. Finally, a model based on the resistances-in-series model allowed us to characterize the mass transfer phenomenon.

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2. Materials and methods

2.1. Chemicals and supplies

As for L-proline, this has been acquired with a purity of at least 98% from Sigma Aldrich.

Glucose (pure, WarChem), hydrochloric acid (analytical reagent, POCH S.A.) and

glutaraldehyde (at 25 wt.% concentration) were obtained and utilized lacking in

additional treatment. The chitosan polymer has been purchased from Sigma Aldrich.

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2.2. Eutectic mixture preparation

A protonated-L-proline:glucose (5:1) has been preliminarily prepared. Experimentally,

10 g of L-proline and 86.86 mL of HCl 1M were continuously blended at given

conditions (1000 rpm, 70 °C) up to results in a homogenous and translucent mixture.

Next, glucose (3,13 g) was incorporated into the mixture. Residual water has been later

eliminated via evaporation (BUCHI Rotavapor R-300, V-300 vacuum pump).

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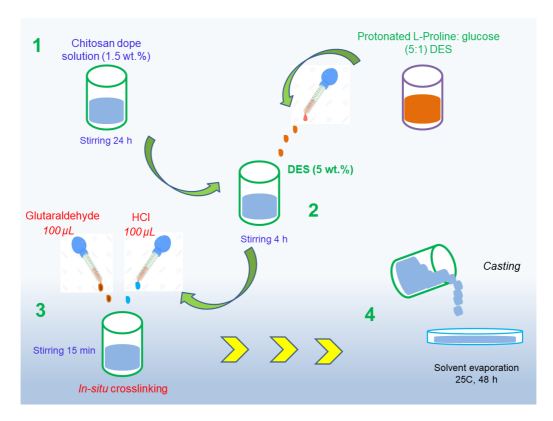
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2.3. Membrane fabrication

Overall protocol for membrane preparation is graphically described in **Figure 1**. All chitosan-based membranes have been fabricated using dense-film casting method. To formulate the dope solutions, the right quantity of CS (1.5 wt.%) was dissolved in 2 wt.% acetic acid in water. The polymeric dope mixtures were kept in agitation during a day.

After that, eutectic mixture L-proline:glucose (5 wt.%) has been then incorporated in dope mixtures. Importantly, 5 wt.% of eutectic mixture corresponds to the best ratio for the membranes, agreeing with previous studies [17,27]. The final solution was mixed during 4 h before the application of the *in-situ* cross-linking with GA, which was initially applied to confer a better chemical and solvent stability of the membranes, while ensuring the DES restraint into the polymer membrane [28]. The in-situ cross-linking has been performed using GA (100 µL), followed by HCI (100 µL). This final mixture was homogenized during 15 min, and then cast. Membrane samples were subjected to drying at 25 °C over 2 days. In general, the membranes' physical aspect has been found as typical homogeneous and continuous film phase with a thickness ca. 25 µm. Finally, the final membranes were named as follows: cross-linked CS & cross-linked CS:L-proline:glucose, labelled as CS:PRO:GLU.



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Figure 1. Preparation strategy of crosslinked chitosan membranes containing L-proline: glucose eutectic mixture.

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2.4. Membrane analysis and characterization

Microscopy techniques: SEM and AFM: Structure-morphological property was studied via FEI Quanta FEG 250 SEM microscope. The secondary electron detector was also utilized for the investigation, recording micrographs (at 5 kV accelerating voltage) in high vacuum mode. Preliminarily, the membrane samples were essentially coated via sputtering (gold layer ca. 10 nm), which was applied to balance the scares surface conductivity. The respective micrographs were obtained at proper magnification. Regarding cross-section, membrane specimens were fractured by immersing in liquid N₂. On the other hand, AFM studies were done using Nanosurf EasyScan 2 by contact mode using silicon tips (AppNano - SICON series) with 10nN constant force [29]. FTIR analysis: Membrane samples have been analysed via Nicolet iS10 spectrometer (from Thermo Fisher Scientific) presenting deuterated triglycine sulfate (DTGS) detector, along with a Golden Gate diamond ATR. Under resolution of 16 cm⁻¹, spectra data was acquired in the range 4000–400 cm⁻¹. Contact angle determination: The surface CA determination has been done with ultrapure water. In this analysis, goniometer OCA15 (Data Physics) was utilized. The data were reported as the average and standard deviation (SD) of at least five assays. 2.4.1. Uptake: As for solvent uptake, which gives the solvent adsorption ability [30], has been investigated for pure ethanol and different water-ethanol mixtures ranging from 0-

50 wt.% water in ethanol. Membrane sample (approximately 1×5 cm) was first weighed;

after this, it was then subjected to immersion in the binary solvent mixtures (at room temperature, 2 days). As documented in other investigations [31], the wet membrane pieces need to be fast cleaned to eliminate the possible residual solvent from surface. Right away, the weight of the samples was again determined (digital balance, Gibertini, Crystal 500, Italy). The uptake calculation was done accordingly [32]:

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$$Uptake \ (\%) = \frac{W_f - W_i}{W_f} \cdot 100$$
 Eq. (1)

where W_f and W_i are the weight values for wet samples after immersion and dry before immersion, respectively.

2.5. Pervaporation separation performance

PV tests have been done in a lab operation unit; the schematic illustration and additional information are observed in previous works [33]. In general, an azeotropic water-ethanol (10–90 wt.%, respectively) mixture was poured into a PV cell. The solvent temperature was controlled (20, 30, 40 and 50 °C) and kept constant utilizing a thermo digital circulating bath. At permeate side, the vacuum pressure was controlled at 1 mbar via CRVpro 4 vacuum pump (Welch Vacuum Products, USA).

Membrane samples with an active area of 5.3 cm² were located on a porous metal support at PV cell. Experimentally, permeating vapour was simultaneously condensed and collected by means of glass trap located inside a condenser containing liquid nitrogen. When steady-state was observed, the permeate samples were obtained from for 4 h running test and right away weighted to estimate the total permeate flux (J):

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$$J = \frac{Q}{A \cdot t} \qquad \text{Eq. (2)}$$

where Q belong to the weight of the permeate (expressed in kg), A is the active membrane area (m²) while t is the testing time (h). The partial flux (J_i) has been evaluated as the product of its respective weight fraction (y_i) accordingly [34]:

$$J_i = Y_i \cdot J \qquad \text{Eq. (3)}$$

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The separation factor (α) was estimated as follows [34]:

$$\alpha = \frac{y_{water}/y_{ethanol}}{x_{water}/x_{ethanol}}$$
 Eq. (4)

where *y* and *x* belong to the weight fraction of each compound present in permeate and feed, respectively. The permeate samples composition was analyzed using Autosystem XL gas chromatograph with flame ionisation detector (FID) and split/splitless injector (Perkin Elmer, USA). Separation was done using a 60,0 m x 0,32 mm ID x 1 um (DB-624) capillary column (Agilent, USA).

The J and α data were given as the average of at least three tests to guarantee the results' accuracy.

188 2.6. Mass transfer analysis

To understand the mass transfer in the pervaporation system, the resistances-in-series theory has been used based on the solution-diffusion mechanism [35]. Considering the solution-diffusion model, a mass transfer flux of a component i can be denoted as a function of an overall mass transfer coefficient and a driving force, as follows:

$$J_i = K_{overall,i}(P_i^{\circ} \gamma_i x_i - P_P y_i) \quad \text{Eq. (5)}$$

Where, $K_{overall,i}$ denotes the total mass transfer coefficient for the component i, P_i° is the

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vapour pressure for the component i, γ_i is the activity coefficient of the component i, P_P is the total pressure at the permeate side, x_i is the molar fraction of the component i at the liquid side, and y_i is the molar fraction of the component i at permeate side. In a pervaporation system, the permeate side is under vacuum pressure so that the total pressure at the permeate side can be neglected, and Eq. (5) can be rewritten and simplified as follows:

$$J_i = K_{overall,i} \cdot (P_i^{\circ} \cdot \gamma_i \cdot x_i) \quad \text{Eq. (6)}$$

- According to García et al. [36] and Arregoitia-Sarabia et. al. [35], overall mass transfer coefficient can be described as the combination for individual resistances
- $\frac{1}{K_{overall\,i}} = \frac{1}{K_{I,i}} + \frac{1}{K_{m\,i}} = \frac{P_i^{sat} \gamma_i}{k_{I,i} \rho} + \frac{\delta}{P_i} \quad \text{Eq. (7)}$
- where $k_{L,i}$ is the mass transfer coefficient for the component i at the liquid side, ρ is the density of the liquid feed, δ is the membrane thickness, and P_i is the permeability of the membrane for the component i.
- The mass transfer coefficient has been calculated using the following correlation based on the work of Johnson et. al. [37]:

$$Sh = \frac{k_{L,i} \cdot d_{pervaporation cell}}{D_{Sol}} = 0,0924 \cdot \left[\frac{\mu}{\rho \cdot D_{AB}}\right]^{0,5} \cdot \left[\frac{R \cdot d^2 \cdot \rho}{\mu}\right]^{0,71} \text{Eq. (8)}$$

- where $d_{pervaporation\ cell}$ is the diameter of pervaporation vessel, D_{AB} represents the diffusion coefficient, μ regards the viscosity of the mixture, d is the diameter of propeller, and R is the revolution per second of the propeller.
- Finally, the membrane selectivity $(S_{i,j})$ and pervaporation separation index (PSI) are determined as follow:

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$$S_{i,j} = \frac{P_i}{P_j}$$
 Eq. (9)

 $PSI = J(S_{i,j} - 1)$ Eq. (10)

3. Results and discussion

3.1. Structure-morphological analysis.

As for prepared membranes either with/without DES, they exhibited a convincing flat and smoothie surface with no apparent defects. It was visibly clear that there is no sign of being plastically deformed, this latter is characteristic of dense polymeric membranes [38]. For pristine CS membrane, cross-section view (Figure 2b) showed clear crater-like structure. This is typically observed when deforming via freeze-fracture. Additionally, this typical structure has been also observed in other pristine biopolymer membranes (like chitosan) [28,39]. Referring to CS:PRO:GLU membrane (Figure 2d), it also showed a homogeneous dense structure lacking in pores or pinholes between the organic phase and eutectic mixture. Essentially, the obtained morphology is considered as convincing proof of exceptional blending/miscibility among the hydrophilic eutectic mixture and CS, this has been reported in precedented studies, e.g., in polymer/polyethylene glycol blend membranes [40,41].

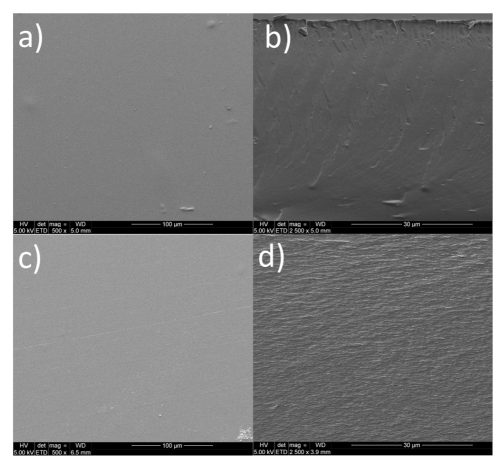


Figure 2. SEM surface and cross-section micrographs of the fabricated CS and L-proline:glucose membranes. (a, b) cross-linked chitosan and (c, d) cross-linked chitosan:L-proline:glucose (CS:PRO:GLU).

Herein, the incorporation of this eutectic mixture has interestingly contributed to obtain a compact but less smooth morphology. As mentioned previously, it has been documented that DESs can act as pore former additive to prepare of porous membranes [16,42]. In this case, our scope is proposed to physically merge and enmesh the chosen hydrophilic solvent in the hybrid membrane for potentialize its separation effect once maintained in the membrane structure. Our hypothesis relies on

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the tremendously success of DESs for the extraction of low molecular weight molecules based on their property to tremendously generate H-bonds by means of dipole-dipole and coulombic forces, among other solute-solvent complexing synergy [43,44]. The eventual success of obtaining homogeneous dense membranes, with convincing evidence of non-DES encapsulation, is due to the synergy of CS at linking some agents thanks to its functional groups. In other words, the lack of functional groups among both phases (i.e., polymer and DES) will consequently lead to not enough interacting properties and consequently minimal miscible features [45]. These latter properties become relevant when dealing with the merging of inorganic/organic dispersing agents into polymers, where potential presence of functional groups or chemically functionalized nanosized materials eventually achieves good synergy and contact at the interface among polymer and filler [46]. CS itself presents a large number of NH₂ and OH terminal domains, which makes the biopolymer an exceptional polymer for polymer blend formation [19]. While specific DESs should also possess functional groups to synergistically interact with CS conferring specific properties to the final membrane. Here, it is quite possible that this hydrophilic DES (PRO: GLU), presenting various functional groups (e.g., amino, hydroxyl, oxygen and carbonyl containing groups), could interconnect with CS. Clearly, the surface micrograph of the blend membranes proves a smooth, continuous and defect-free surfaces with lack of any pinholes. In literature, other DES, such as choline chloride-malonic acid, was able to turn the flat surface of chitosan films to a nonhomogenous morphology while keeping a compact pattern [47]. In this study, we also confirm a smoother surface by adding the DES agent into CS, as observed in **Figure 3**.

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It is worth mentioning that the DES-free crosslinked CS membrane displayed an average root mean square roughness of $S_q=4.0\pm0.5$ nm, where the incorporation of DES slightly lowered such value to $S_q=3.0\pm0.5$ nm. Certainly, upon the nature of the eutectic mixture, the exaggerated load of the DES may promote a rough structure of the final eutectic-chitosan blend membranes [48].

Crosslinked CS membrane 20.0 15.0 10.0

Crosslinked CS PRO:GLU (5:1)

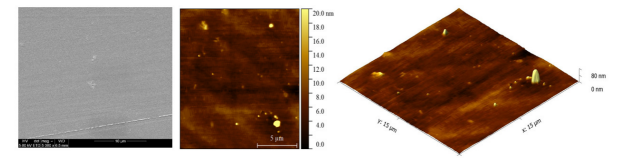


Figure 3. AFM surface and 3D images (15 \times 15 μ m) of pristine CS and CS:PRO:GLU membranes.

It is worth mentioning that the blending of DES into CS membrane may influence the mechanical and thermal properties of the resultant membranes. In a previous work, it has been stated that eutectic mixtures (e.g., based on L-proline:sulfolane) worsened specific mechanical properties, such as Young's modulus and tensile strength, but such an effect was compensate due to the *in situ* cross-linking [17]. DESs tend to decrease the intermolecular interactions in the CS network, causing the co-called plasticization. The incorporation of plasticizers (like DES) in CS, in fact, produces a transition from a rigid to a softer material with elastic properties [49]. Aside from this effect, DES has been demonstrated to affect the thermal stability of the CS membranes in terms of glass transition temperature (T_g), e.g., DES-modified CS films exhibited a lower T_g in comparison with bare CS membranes. Eventually, this can be ascribed to eutectic mixture decomposition. Interestingly, T_g values decreased proportionally as the DES content increased. According to Jakubowska et al. [47], DES concentration increment results in a decrease in temperature at which decomposition starts.

3.2. FTIR and water CA characterization

As seen in **Figure 4**, the FTIR data essentially confirms the effective merging of the hydrophilic PRO:GLU solvent and the biopolymeric phase. Full spectrum data show a firm and vast patterning with asymmetry ranged approximately 3350-3450 cm⁻¹, which a consequence of overlaying the O-H and N-H stretching oscillations of terminal groups connected by H-bonds. Typically, CS spectrum exhibits a various absorption patterns around 1650 cm⁻¹ (which corresponds to C=O stretching of possible CO=NH₂ terminal group), 1520 cm⁻¹ (which corresponds to N-H wagging of non-acetylated 2-aminoglucose) and ca. 1580 cm⁻¹ (which corresponds to N-H wagging of CO=NH₂ terminal), which evidenced by other reports [50]. Apart from that, absorption patterning located ~ 1150 cm⁻¹ refer to a skew-symmetric stretching for C-O-C links, 1060 cm⁻¹ and 1050 cm⁻¹, ascribed to possibly skeletal frequencies comprising a typical C-O

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conjunction. Concerning to the DES, the amino acid (L-proline) owns an extra -NH₂ terminal, identified as an RR'C=NR". Hence, proline is also identified as an imino acid. As proline's three-carbon R-grouping tends to be merged to the alpha-nitrogen terminal, this molecule presents a restrained rigid-ring with rotational behaviour [26]. In the case of glucose, it is a monosaccharide presenting six carbon atoms and an aldehyde group. Considering the typical structure of any DES, both molecules (HBA and HBD) are structurally well attached via electrostatic interaction (i.e., H-bonds) to form a final eutectic system [51], this latter interaction is apparently observable with a firm shifting and oscillation for the peak ranged from 3750-1950 cm⁻¹ [52]. When this PRO:GLU eutectic system is fused in the CS, this results in modest but clear motion on the usual polymer patterning. This latter interaction reveals an exceptional attraction between phases and is translated to affinity. Clear motions, together with the classic behaviour of the patterning, in the range of 3650- 3100 cm⁻¹, must also be noted showing an overlying of the spectra, which are ascribed to O-H, N-H and C-O oscillations in chemical functionalities of the eutectic mixture and polymer. Jakubowska et al. [47] have also reported similar molecular interactions when prepared and characterized chitosan/DES hybrid materials. Importantly, the resulting reticulated flat films containing glutaraldehyde commonly demonstrate clear absorption increment among 1600-1650 cm⁻¹ thanks to N=C bonds [53,54]. The widening at 1550, 1740 and 2850 cm⁻¹ refers to the free aldehyde chain and raised C-H length, respectively. Aliphatic -NH₂ terminals decay directly proportional as the peak 1150 cm⁻¹ decreases. Alternatively, DES have been identified as potential cross-linker agents for CS. Especially, eutectic mixtures containing carboxyl groups are able to interact with the NH₂ groups of CS creating an amide connection [48]. Such an amide connection has been also been observed in CS membranes blended with choline chloride -citric acid [52]. Interestingly, the blending of choline chloride-urea DES into the CS structure was able to create the saccharide ring (C–O–C) between DES and CS, proving the formation of a resilient solid biopolymer membrane without adding cross-linking mediator [55].

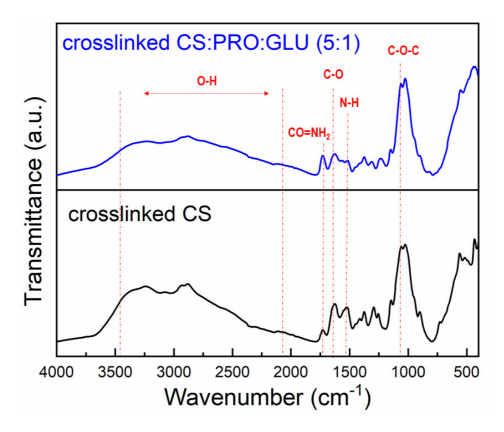


Figure 4. Spectrum data from FTIR measurement for pristine crosslinked chitosan and its eutectic mixture-based blend membrane.

A relevant point to highlight during the addition of this hydrophilic DES in membranes

regards its remarkable response on either hydrophilic or hydrophobic nature of the membrane. The DES-free cross-linked chitosan flat membrane revealed angle values of ca. 70° (see **Figure 5**), which agrees with outcomes published previously, e.g., angle values of chitosan nearly 74-88° [50,56]. In principle, the nature (either hydrophilic or hydrophobic) of the biopolymer indeed stands on its degree of deacetylation, e.g., large deacetylation gives as a result exceptional hydrophilicity in membranes due to many NH₂ terminals tend to be available in the biopolymer [57], this becomes relevant since hydrophilicity is needed for enhanced water adsorption over a membrane interface [58]. Hydrophilic nature of chitosan follows from its polar hydrophilic terminals, including -NH₂ and -OH; unfortunately, this polar nature is compromised when applying cross-linking. However, the addition of the PRO: GLU DES in the CS contributed to an enhanced hydrophilicity translated to lower CA values of ca. 50°. This gives an idea that the polar groups given by the DES have a meaningful influence at improving the hydrophilic nature of the membrane. Importantly, glucose is classified as polar since hydroxyl groups presents high affinity to hydrogen bonds and effective electronegativity.

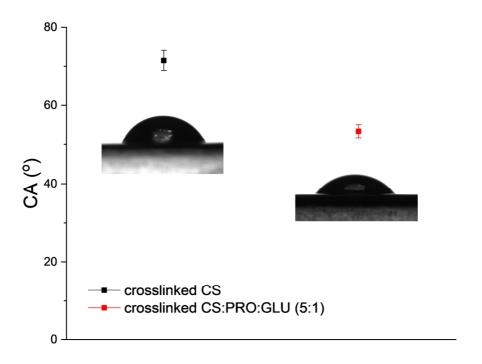


Figure 5. CA values pristine crosslinked CS and CS:PRO: GLU membranes.

3.3. Pervaporation testing

3.3.1. Operating temperature dependence of permeation and separation factor.

PV data for all assayed membranes are reported in **Table S1**. **Figure 6** shows the influence of the feed temperature on the total permeation flux, in which a permeation increase was noted in the range of 20-50 °C in pristine cross-linked and its blend with PRO: GLU DES. It is actually traditional behaviour for polymers as their chains tend to present improved flexibility under high temperature, which consequently contribute to enhanced solvents' sorption. This consequently increases the transport of molecules

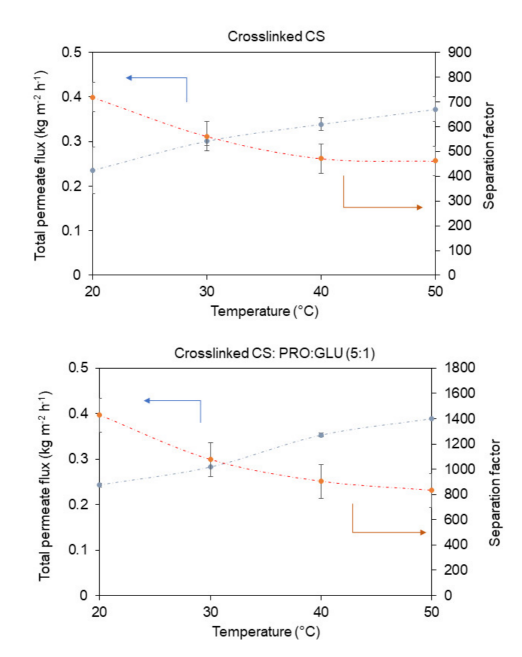


Figure 6. Permeation and separation factor behaviour in respect to operating temperature (10/90 wt.% H₂O/EtOH, pressure: 1 mbar). The semi continuous loops are only guiding.

Essentially, an influence of temperature on overall flux was further analysed by means of the Arrhenius equation, as denoted in Eq. 11.

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$$J = J_o \cdot \exp\left(-\frac{E_{app}}{R \cdot T}\right) \qquad \text{Eq. (11)}$$

In theory, J_{o} refers to the pre-exponential element, while E_{app} corresponds to the apparent activation energy for the transport. The product $R \cdot T$ expresses the typical term of energy. By applying mathematical logarithms in previous Eq. (11), E_{app} is determined directly from the straight line proving a compelling relationship between flux and temperature; it means, an increment in overall flux occurs with temperature increase. As **Table 1** reports, it is noted that water displays lower E_{app} values (ca. 5.07 kJ mol⁻¹) in respect to ethanol (~10.27 kJ mol⁻¹) in crosslinked CS membrane, confirming the water affinity of chitosan. On the other hand, the eutectic solvent blending slightly increased the E_{app} in water molecules approximately 5.57 kJ mol⁻¹ for the hybrid CS:PRO:GLU membrane, however, E_{app} value for ethanol has been more impacted by DES incorporation. Particularly, the E_{app} keeps unchanged towards water in respect to ethanol ranging from 20 to 50 ℃. It is important to point out that the temperature increment impact mainly the water permeation, and it is greatly restricted the permeating ethanol; this is supported by Almeida et al. [60], who documented an improved water solubility in CS films containing DES, in which the water solubility increased when increasing the DES content. In this study, the existence of this hydrophilic PRO: GLU increases the energy demanded for the permeating solvent to be transported over the membrane interface, which is more prominent for ethanol. This latter aspect is in agreement with the DES hydrophilicity, which favours for the selective properties for the more polar compounds (like water) [61].

Table 1. Apparent activation energy values for overall flux, and single water and ethanol across prepared hybrid materials.

Hybrid formulation		E _{app} (kJ mol	1)
_	Overall	Water	Ethanol
Crosslinked chitosan	5.15	5.07	10.27
Crosslinked CS:PRO:GLU	5.63	5.57	11.76

When dealing with selective affinity, α parameter in pure CS membrane decayed when the temperature decreased, as seen in **Figure 6**. Promisingly, separation factor was substantially enhanced via DES incorporation observing data of ca. 1,427 (at 20 °C). Obviously, large α parameters with lower permeating yield were indeed recorded at the minimal testing temperature. It somehow in accordance to the polymer "free-volume" theory, as it declares a thermal movement in chains specifically at amorphous localities boosting an increment in free-volume. It is known that as temperature raises, the frequency and magnitude of the chain swing increases provoking a free volume increment [62]. Although the kinetic diameters of water and ethanol are substantially different (2.6 and 4.3 Armstrong, respectively), the thermal movement in polymer chains

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can indeed facilitate the diffusivity for bigger solvent molecules (like ethanol) over a membrane interface compromising the separation factor. In addition to this, hydrophilic eutectic mixtures have demonstrated to break the intermolecular structure of CS and thus open the network structure, allowing the permeation of solvent molecules [48,60].

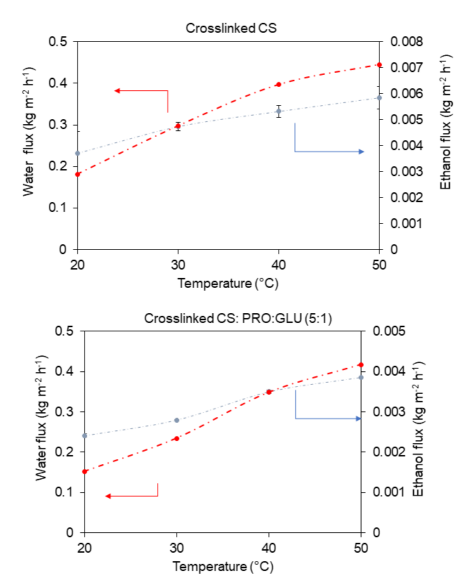


Figure 7. Water and ethanol flux in respect to operating temperature (10/90 wt.% H₂O/EtOH). The semi continuous loops are only guiding.

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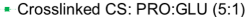
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Very recently, Jakubowska et al. [47] have reported that the DES addition promotes the free volume increment in polymeric matrices. This is ascribed to the enlargement of free-volume fostering the chain motion, which can produce a decrement for the membrane's selective efficiency. However, an application regarding this hydrophilic PRO: GLU DES did not affect the selective properties of CS membrane. Interestingly, both parameters, such as permeation rate and selectivity, were significantly improved. As shown in Figure 7, DES preferentially promoted the passage of water molecules over the polymeric phase while concurrently hindering a possible ethanol permeation. At this point, the polarity of the solvents becomes relevant for their own transport and extraction from complex systems [63]. The analysis for the solvent uptake in the developed membranes is represented in Figure 8. It is clearly observed how both membranes show low solvent uptake when there is a minimal concentration of water in the ethanolic solutions. Experimentally, a water concentration increment of the feeding solvent mixture resulted in a higher membrane swelling in a water concentration from 10 to 50 wt.%. Surprisingly, the incorporation of PRO: GLU DES provoked a decrement in terms of solvent uptake performance compared with the pristine CS. This latter phenomenon supports Jakubowska's findings [47] in which a DES confers stability in chitosan membranes. The uptake is generally expected to be restricted once a crosslinking protocol is implemented as this treatment provides resistance to polymer materials against polar solvents thanks to an enhanced stricture in mobility of chains [64]. Swelling phenomenon is adequately identified as one of the primary bottlenecks of hydrophilicbased polymer materials when separating polar molecules [20]. Therefore, less prone to be swollen membranes are preferentially needed to acquire a stable separation during long-term testing.

Crosslinked CS membrane



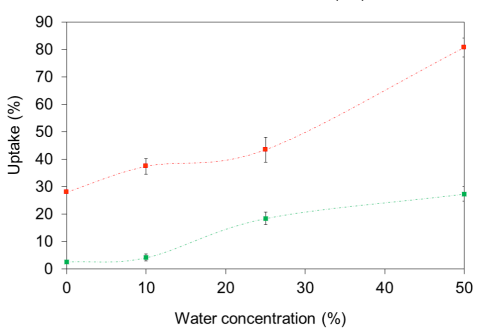


Figure 8. Solvent pptake of CS-eutectic solvent membranes under various water percentage in ethanol (at room temperature). The semi continuous loops are only guiding.

3.3.2. PV data comparability of cross-linked CS: PRO:GLU membranes with other studies

As for PV testing, it is obvious that the PV separation efficiency of both hybrid and polymer materials is primarily dependent on distinct factors, such as membrane properties (e.g., material type, physiochemical and intrinsic properties, structure, etc.), along with operating parameters including feed composition, operating temperature,

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pressure gradient, etc. [34]. Especially, the membrane structure is somehow dictated by the applied membrane preparation strategy [65], while most PV evaluation of membranes has been experimented at different feed concentration and operating parameters. This makes particularly tough to give a fairer comparability among pervaporation results from various works [66]. In our research, we eventually make a comparison of performance outcomes among distinct membrane concepts either unmodified polymer, blends, composites or inorganic tested under close operational parameters, as reported in Table 2. Here, the outstanding selectivity, expressed as separation factor, for crosslinked CS:PRO:GLU membrane was found at 20 °C (approximately 1,427), which represents almost 2-folds bigger α value in comparison to unmodified crosslinked chitosan. In addition to this, the highest permeation rates were acquired at the highest tested temperature (at 50 °C) in both membranes while showing a decrement in selectivity. The membrane containing the DES showed a slight improvement in permeation (see Table S1). When compared with other reports, crosslinked CS:PRO:GLU membranes exhibited better selectivity than other composite membranes, such as crosslinked PVA-filled GO, CS-filled H-ZSM-5, polyimide (PI)-filled ZIF-8, CS-filled titanium dioxide (TiO₂), PI-filled MSS-1, crosslinked PVA-filled ZIF-8-NH₂, among others (see **Table 2**). Depending on the used inorganic fillers filled into the polymer membranes, the aforementioned composites can offer higher permeation rates than our findings. Unfortunately, crosslinked CS:PRO:GLU membranes did not overcome the exceptional selectivity of crosslinked sodium alginate-filled beta zeolite and NaP1 zeolite membranes with unprecedented separation factor values. It is worth pointing out that these membranes (i.e., crosslinked CS:PRO:GLU) are overcoming the

selective-permeable trade-off of the pristine CS membranes.

Table 2. Comparison of crosslinked CS: PRO: GLU membrane performance with some composites and inorganic membranes tested with similar water-ethanol mixtures.

-	Filler brane concept Water % content:				Separation	1
Membrane concept			Testing conditions	J (kg m ⁻² h ⁻¹)	factor	Reference:
Crosslinked CS:PRO:GLU	-	10 wt.%	20 °C, 1 mbar	0.242	1,425	This work
Crosslinked CS:PRO:GLU		10 wt.%	50 °C, 1 mbar	0.389	831.7	This work
Crosslinked PVA- GO	1 wt.%	10 wt.%	40 ℃, 3 mbar	0.137	263	[67]
CS-filled H-ZSM-5	8 wt.%	10 wt.%	80 ℃, 10 mbar	0.230	152	[68]
Crosslinked sodium alginate- filled beta zeolite	10 wt.%	10 wt.%	30 °C, 0.6 mbar	0.130	1,600	[69]
Polyimide-filled ZIF-8	12 wt.%	10 wt.%	42 ℃, 44 mbar	0.260	300	[70]
CS-filled TiO ₂	6 wt.%	10 wt.%	80 ℃, 50 mbar	0.340	196	[71]

Polyimide-filled MSS-1	12 wt.%	10 wt.%	42 ℃, 44 mbar	0.310	190	[70]
Crosslinked CS-filled silica	5 wt.%	10 wt.%	70 ℃, 10 mbar	0.410	919	[72]
Crosslinked PVA-filled ZIF-8-NH ₂	7.5 wt.%	15 wt.%	40 °C, 1 mbar	0.120	200	[73]
NaP1 zeolite membranes	-	10 wt.%	75 °C, 4 mbar	0.45	200 000	[74]
PVA composite membrane	-	10 wt.%	60 ℃, 5 mbar	0.140	170	[75]
PVA composite membrane (PERVAP 2201, Sulzer Chemtech)	-	10 wt.%	60 ℃, 10 mbar	0.100	100	[76]

3.4. Mass transfer performance

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Through the mass transfer model developed for the pervaporation process, and equations 7 and 8, mass transfer resistances can be determined for the different membranes, which are shown in Table 3.

Table 3. Mass transfer resistance distribution

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			Crosslinked CS	3	Cro	osslinked CS:F	PRO:GLU
	Т	$R_{liq}\left(\frac{s}{m}\right)$	$R_{\text{membrane}}\left(\frac{s}{m}\right)$	$R_{\text{overall}}\left(\frac{s}{m}\right)$	R _{liq}	R _{membrane}	$R_{\text{overal}} l\left(\frac{s}{m}\right) \cdot 10^8$
	(℃)	10^{4}	Thembrane $\binom{m}{m}$	10 ⁸	$\left(\frac{s}{m}\right) \cdot 10^4$	$\left(\frac{s}{m}\right)10^8$	Thoverall $\binom{m}{m}$
Water	20	0.799	0.155	0.155	0.799	0.149	0.149
	30	1,37	0.222	0.222	1,39	0.219	0.219
	40	2,26	0.346	0.346	2,32	0.329	0.329
	50	3,60	0.529	0.529	3,72	0.502	0.502
Ethanol	20	1,07	58.3	58.3	1.07	113	113
	30	1,79	64.4	64.4	1.81	132	132
	40	2,90	82.3	82.3	2.96	153	153
	50	4,52	120	120	4.68	209	209

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Table 3 shows how the transport stage in the membrane corresponds to the one that presents the greatest resistance to mass transfer for the two membranes under study, as well as for water and ethanol, reaching approximately 99% of the total resistance



distribution of them. On the other hand, despite the great importance of the resistance value to mass transfer of the membrane, differences in behavior can be observed when comparing the same membrane with water and ethanol, at the same temperature. It is seen that in crosslinked CS membrane, the resistance for ethanol transport is two orders of magnitude bigger compared to water, and as for the membrane modified with DES, a difference of 3 orders of magnitude is obtained, which implies that the modification by including DES improves the separation performance, fulfilling the task of obtaining a membrane with a greater hydrophilic character, offering approximately twice the resistance to ethanol than the crosslinked CS membrane, whereas for water, the resistances are similar.

To compare the performance of membrane modification with DES, it is necessary to compare the selectivity factors, this information is presented in **Table 4**.

Table 4. Water/ethanol selectivity for the different membranes.

			RO:GLU
$S_{W/E}$ PS	$SI\left(\frac{kg}{m^2h}\right)$	$S_{W/E}$	$PSI\left(\frac{kg}{m^2h}\right)$
375	87	756	183
290	87	600	181
238	80	464	163
228	85	416	162
	375 290 238	m^2h (m^2h) 375 87 290 87 238 80	m^2h (m ² h) 375 87 756 290 87 600 238 80 464

Regarding the selectivity factors that both membranes present, it is observed how the incorporation of proline-glucose as DES allows a greater selectivity towards water, mainly because they prevent the passage of ethanol molecules through the membrane, which explains the increase in the resistance of the membrane to ethanol doubling its value respect to the resistance towards ethanol in the Crosslinked CS membrane. The addition of PRO:GLU DES to the membrane allows the PSI of the membrane to improve more than double in comparison with the crosslinked CS membrane, even the membrane performance is greater than that reported in the literature for the water-ethanol system [67].

Finally, the increase in temperature results in selectivity decrement of the Crosslinked

CS:PRO:GLU membrane, being the temperature at 20°C the one that presents the best

performance, which is corroborated by the PSI value presented by the membrane.

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4. Conclusions and future research

This research reveals the fabrication and characterization, for the first time, of dense crosslinked CS-hydrophilic protonated-L-proline: glucose. The membranes present a compelling miscible properties and integration of the original hydrophilic eutectic mixture (PRO:GLU) along the organic biopolymer interface. In general, it was utilized ecofriendly items (including biopolymer, water as primary solvent, "green" eutectic mixture), making the developed hybrid materials as good candidates for fabricating sustainable and eco-friendly dense membranes. For water-ethanol pervaporation separation, these hybrid membranes offer a 2-fold improved pervaporation yield than the DES-free crosslinked chitosan and slight enhancement in permeation. As a perspective, the forthcoming research must be emphasized on enhancing the permeation yield and membranes' selective properties when incorporating inorganic phases like nanomaterials. By smartly selecting the hydrophilic nanostructured materials (such as graphene oxide, MXene, UiO-66 MOF), the resultant mixed matrix membranes based on CS may offer unprecedent permeation rates while improving the selective properties as well [23,67]. Also, these membranes can be assayed in other attractive PV separation applications, such as methanol/MTBE [77], water/isopropanol [78], water/hydrazine hydrate [79], requiring hydrophilic membranes. Due to their interesting capability to form H-bonding intermolecular forces, DESs could foster an outperforming extraction of some other polar solvents present in azeotropic systems [17,43]. Variety of available DESs, as well as high potential for new developments in this field, tries to design new DES-based membranes having tailored selectivity. Lately, visualizing a possible preparation protocol in a more sustainable way, the crosslinker agent (i.e., glutaraldehyde) could be substituted by another less harmful substance. Here, green substances, such as cinnamaldehyde [80], genipin [81], could be an alternative.

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Conflict of Interest

571 The authors declare no conflict of interest.

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