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Fabrication of novel porous membrane from biobased water-soluble polymer (hydroxypropylcellulose)

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Abstract

Herein, a novel polymeric porous membrane was developed, without the use of any organic solvent in the initial dope solution and using a biobased polymer derived from cellulose: Hydroxypropylcellulose (HPC). HPC was dissolved in water (20 wt\%) and the phase separation was induced by increasing the temperature above the Lower Critical Solution Temperature (LCST) of the polymer solution, around 40 °C in the concentration range concerned in this study. To fix the membrane morphology and to prevent any resolubilization in water during filtration tests, a chemical crosslinking was performed using Glutaraldehyde. The phase diagram of HPC/water system was first studied and not only the cloud point but also the spinodal curves were determined using optical transmission techniques. It was exhibited that HPC phase diagram is very weakly dependent on concentration up to large concentrations and that the metastable region is very small, i.e. the cloud point and the spinodal curves are very close in a large range of concentration. The membrane stability was tested in water and some organic solvents, thus demonstrating the efficiency of the chemical crosslinking during membrane formation. The swelling and mechanical properties of HPC membranes were also investigated depending on the operating condition during membrane formation, showing that the temperature ramp during the membrane formation, from initial to final
temperature, have a significant effect on the crosslinking efficiency and hence on the swelling properties. Finally, adding a porogen (PEG200) into the collodion, filtration tests were performed and exhibited that the membrane filtration properties depend on the temperature ramp as well, showing higher water flux for the highest temperature ramp tested.

**Keywords**

Membranes, green chemistry, biobased polymer, water-soluble polymer

1. **Introduction**

Membrane separation technologies are nowadays extensively used in many engineering applications including water treatment and purification. Polymeric membrane materials account for a large majority (ca 85%) of the membrane material demand. Within the polymeric membrane segment, synthetic polymers including polysulfone, polyimide, nylon, polyacrylonitrile, PVDF and others dominate the market. Polymer phase separation, which consists of lowering the solvency of the solvent to the polymer, is mostly used in manufacturing of such porous materials: a dope solution, i.e. a polymer dissolved in a solvent, is cast as a film on a support or extruded as a hollow fiber. Subsequently, a change in the thermodynamic state disturbs the solution equilibrium. Thermally Induced Phase Separation (TIPS) and Non-solvent Induced Phase Separation (NIPS) are the most convenient and common methods for manufacturing porous polymeric membranes. Many membrane structures can be formed depending on operating conditions (process parameters) and system properties (formulation) [1-3]. NIPS process (including immersion precipitation and Vapour Induced Phase Separation, VIPS) consists of adding a non-solvent into a binary polymer/solvent solution. TIPS process is classically conducted by cooling a hot polymer/solvent solution, inducing thermodynamic instability and thereby phase separation [4-11]. Porous membranes are thus produced by phase separation from polymer solutions and from economic point of view, water is generally used as the non-solvent. Since the polymer concentration is usually in the range 15-20%
and since coagulation and washing baths require being often renewed, large amounts of aqueous solutions must be treated. For instance, 10 m$^2$ of ultrafiltration membrane need about 1 to 1.5 kg of solvent. The major fraction can theoretically be recycled after an expensive distillation stage (ca 90% for slightly volatile solvents), a fraction is lost by evaporation in air (not negligible for highly volatile solvents and high temperature process) and the last fraction is sent for wastewater treatment. Moreover, the use of organic solvent in the membrane formation has consequences on industrial fabrication units, which must be adapted to protect workers from solvent toxicity. Therefore, the use of organic solvents has a significant cost input in the process of membrane manufacturing apart the environmental and public health impact. Facing the challenges of decreasing these impacts, shift of paradigm is needed from conventional solvents towards greener ones including bio-based solvents, ionic liquids and supercritical carbon dioxide [12-13]. Among them, water is obviously the safer and the less demanding on energy. Nevertheless, to the best of our knowledge, very few papers have been devoted to the membrane preparation in water [14-16].

In this respect, the present work aims at preparing polymeric membranes using water as the solvent. This objective clearly addresses the issue of designing greener processes avoiding the use of organic solvents, one of the cornerstones of the green chemistry area. The other objective is to design membrane from bio-sourced materials. The economy of atoms and reduction of energy in the manufacturing of membranes is assumed to be achieved in a friendly environmentally approach using (semi)natural and water-soluble polymers. Cellulose ethers fulfill these criteria. These derivatives are characterized by a lower critical solution temperature (LCST) that can be used to induce the phase separation. The polymer phase separates through a temperature increase above the LCST [17], calling the process LCST-TIPS in contrast to the commonly used upper critical solution temperature (UCST)-TIPS for membrane preparation with synthetic polymers. Phase separation with LCST systems can usually be tuned at moderate, close to ambient temperatures, limiting energy consumption. The available literature on the LCST-TIPS process is very limited. Qin and coworkers prepared membrane from polyethersulfone/poly(vinyl pyrrolidone) K90/N, N-dimethylacetamide
(DMA)/1,2-propanediol (1,2-PD) system [18-19] and cellulose acetate/poly (vinyl pyrrolidone) 360K/N-methyl-2-pyrrolidone (NMP)/1,2-propanediol system [20]. Recently, free organic solvent membranes have been prepared from PVA/water system [16].

Hydroxypropylcellulose (HPC), non-ionic ether of native cellulose, is a polymer with temperature dependent water solubility. Thanks to its renewable origin, ease of production, non-toxicity and optical properties, HPC is widely used (i) for the preparation of dosage drugs forms and coatings, (ii) as viscosity modifier in many product formulations [21]. Furthermore HPC is studied in polymer physics for its lyotropic liquid crystalline phase behavior [21]. HPC has also been largely studied to develop hydrogels for controlled release of active substances above its LCST [22-24].

In this study, porous membranes prepared by LCST-TIPS process from HPC will be demonstrated. The dynamics of membrane formation, including the coupling between the phase inversion dynamics, the crosslinking and the mass transfer by evaporation will be discussed. Since this polymer was poorly studied in the past, the phase diagram was investigated, including both the determination of the cloud point curve and the spinodal curve using light scattering techniques. The membrane morphology of HPC membranes was reported and discussed as a function of the operating parameters including the initial formulation and the temperature ramp. The stability and the mechanical properties of the membranes were then investigated. Finally, preliminary filtration tests in water will be presented.

2. Experimental

2.1. Materials

Hydroxypropylcellulose (HPC) was obtained from Sigma Aldrich (Fig. 1). Its average molecular weight, determined by size-exclusion chromatography, was $M_w$ 80,000 g/mol. Distilled water was used as solvent in polymer/solvent system. The washing bath was exclusively composed of distilled water. Glutaraldehyde (GA solution 25 wt%) was chosen as the crosslinking agent, hydrochloric acid (HCl
solution 37 wt%) as the crosslinking reaction catalyst and polyethylene glycol Mw 200 g/mol as porogen (PEG200). All of these products were purchased from Sigma Aldrich and all the reagents were used without further purification.

2.2. Methods

2.2.1. Preparation of polymer solutions

Hydroxypropylcellulose (HPC) aqueous solution was prepared by dispersing HPC powder in distilled water preheated at 60 °C under magnetic stirring. The initial HPC concentration in water was 20 wt%. The dispersion was then cooled to ambient temperature to promoting polymer dissolution and obtaining homogeneous polymer solution. A given amount of additives was then blended into the solution under magnetic stirring. The solution prepared was degassed under static conditions at 4 °C overnight. The polymer concentration (20 wt%) was chosen in order to reach a convenient viscosity for casting the solution onto the glass plate (the viscosity of HPC solutions was equal to 18.98 Pa.s during casting) and to have a moderate cloud point temperature (close to 40 °C). The chemical crosslinker, Glutaraldehyde, was added in the collodion, at a concentration of 0.5 wt% (versus the total solution weight). The catalyzer, HCl, was added just before casting the polymer solution onto the heating plate. HCl concentration in the collodion was equal to 1 wt%. The composition of the polymer solution (see Table 1) was fixed from previous study of Suto et al. [32] to optimize the crosslinking, i.e. to obtain an isotropic membrane with well distributed chemical bonds in the whole membrane: Glutaraldehyde weight ratio ¼ (GA/HPC) was shown to be the most efficient ratio for obtaining a stable polymer matrix. The concentration of each component in the collodion was summarized in Table 1.

Table 1. Concentration of polymer and additives in the collodion.

<table>
<thead>
<tr>
<th>HPC</th>
<th>GA</th>
<th>HCl</th>
<th>PEG200</th>
</tr>
</thead>
<tbody>
<tr>
<td>20 wt%</td>
<td>0.5 wt%</td>
<td>1 wt%</td>
<td>5 wt%</td>
</tr>
</tbody>
</table>
2.2.2. Cloud point measurements

Polymer solutions were prepared by weighing at different concentrations and put after thorough mixing and heating/cooling steps in quartz cuvettes. These cuvettes were immersed in a water thermostated bath at a temperature in the one-phase region. The temperature was then progressively increased by steps of 0.5 °C per 30 minutes. A laser beam is sent through the cuvette and the transmission of the beam is measured as a function of temperature by means of two photodiodes, one as a reference of the incident beam and the second on the transmitted path.

2.2.3. Membranes preparation

The blend solution was cast onto a glass plate at room temperature with an automatic microcoater (Erichsen Instruments) using two speed casting: 3.4 ± 0.2 cm/s and 15.0 ± 0.5 cm/s. The thickness of the solution was fixed at 250 µm for the whole experiments. The sample was then placed on a Linkam hotstage LTS 420, which was chosen for its accurate thermal control and its built-in stage manipulator; the software Linksys 32 allowed to perfectly control the temperature ramp (from 0.01 to 120 °C/min) and the target temperature (60 °C); two temperature ramps were performed in this study to increase temperature from ambient to target temperature: 2 °C/min and 120 °C/min. The membrane formation was conducted at ambient atmosphere, i.e. under RH close to 50%. HPC concentration was chosen in such a way that the solution viscosity of HPC/water system (close to 7 Pa.s without additives) allowed (i) easy handling of the solution and additives admixture with a fluent homogenization, (ii) casting the polymer solution onto a glass plate for flat membranes. The whole operating conditions were reported in Table 2.

The membrane was then removed from the glass plate and placed in a washing bath containing distilled water in order to remove excess of additives agents. The membrane was stored in distilled water prior to further water flux measurements, or oven dried at 30 °C for further morphological and mechanical characterizations.

**Table 2.** Operating conditions for membrane preparation.
### Characterization of the membranes

#### 2.2.4.1. Scanning electron microscopy

The SEM micrographs were produced using HITACHI S-4800 equipped with a Sputter Coater SC 7620 for sample metallization. The HPC films were fractured in liquid nitrogen; the samples surface and cross section were analyzed by SEM.

#### 2.2.4.2. Viscosity

The viscosity of HPC solutions (with and without additives) has been measured using a Rheometric Scientific (Physica MCR 301, Anton Paar) equipped with a parallel-plate geometry (air gap between 0.01 and 0.1 mm). All the experiments have been performed at 25 °C.

#### 2.2.4.3. Crosslinking

The crosslinking was followed using the same Rheometric Scientific (Physica MCR 301, Anton Paar) equipped with a parallel-plate geometry; variable oscillation amplitude and variable oscillation frequency were tested to determine the best operating conditions (linear domain) and to fix the operating conditions, i.e. shear stress and oscillatory frequency. The variation of the elastic (G') and viscous (G'') modulus were followed under small amplitude oscillatory shear stress (1%) and 1 Hz frequency. The rheological measurements were conducted in closed system (the polymer solution was put between two glass plates and isolated from the external environment) to prevent solvent evaporation during experiments.

#### 2.2.4.4. Mechanical properties

The measurements of mechanical properties were carried out under dry conditions on the Rheometric Scientific (Physica MCR 301, Anton Paar) testing machine at 25 °C with a frequency of

### Table

<table>
<thead>
<tr>
<th>Initial film thickness</th>
<th>Casting speed</th>
<th>Temperature ramp</th>
<th>RH</th>
<th>Water bath temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>250 µm</td>
<td>15.0 ± 0.5 cm/s</td>
<td>3.4 ± 0.2 cm/s</td>
<td>2 °C/min</td>
<td>50 %</td>
</tr>
</tbody>
</table>
1 Hz. A film sample was cut into pieces with 4 x 1 cm². The sample thickness was determined with a digital micrometer (MITUTOYO with an accuracy of 0.001 mm). Five measurements were performed on each membrane at different locations. The sample was attached to the tensile grips and the test was finished once the sample broke. Three successful measurements were used to obtain average values for mechanical properties. The stress \( \sigma \) was determined in terms of the original cross-sectional area, and the strain defined as \( \varepsilon = (l - l_0)/l_0 \), where \( l \) and \( l_0 \) are the length at the time of data collection and original length, respectively.

2.2.4.5. Solubility and water sorption

A sample piece (2 x 1 cm²) was cut from the HPC film oven dried at 30 °C for 24 h and immersed in distilled water for 40 days. The swelling ratio was determined at ambient temperature, based on the following equation: \( \text{SR (\%)} = (W-W_0)/W_0 \times 100 \), where \( W_0 \) is the initial weight of dried sample piece and \( W \) is the weight of swollen sample piece (the surfaces of the film were wiped with soft paper tissue before measurements).

2.2.4.6. Water flux measurements

Ultra-pure water was used to characterize the pure water flux through porous HPC membranes at temperature 25 ± 0.5 °C in a stirred ultrafiltration cell (Millipore). The effective membrane area was 4.1 cm² and the membrane was supported by a non-woven material (Polyethylene Terephthalate) to enhance the mechanical properties of HPC films. The pressure was gradually increased from 0 to 3.5 bars with pressure steps of 0.2 bar during 15 min. Each membrane was conditioned prior to the filtration tests, i.e. the pressure was gradually increased from 0.1 to 3.5 bars (pressure step of 0.2 bar during 30 min).
3. Results and Discussion

3.1. Phase diagram

![Chemical structure of HPC](image)

**Figure 1.** Chemical structure of HPC (R = -H/-CH₂CH-(OR’)CH₃ with R’= -H/-CH₂CH(OR)CH₃)

The usual method found in literature for determining the cloud point temperature (Tₜₚ) of water soluble polymers solutions amounts to measuring the optical transmission through a sample. As the transition is usually spanning a broad interval of a few degrees a (rather arbitrary) method has been adopted for measuring the Tₜₚ temperature. The latter is determined as the abscissa of the intersection of two limiting slopes of the transmission curve, one horizontal slope drawn at temperatures lower than the transition and one inclined slope representing the transmission decrease through the temperature interval (the principle is depicted in T. Kyu, P. Zhuang, P. Mukherjee [25] for the scattering intensity and by E. Marsano and G. Fossati [26] for the transmission). Our measurement is reported in Fig.2 [27-28].
The spinodal temperature, defined as the temperature at which the scattered light intensity extrapolated at zero angle (meaning zero wavevector of transfer) diverges, was also measured (Fig. 2). The values of $T_{cp}$ are in good agreement with the previously reported ones for dilute samples by F. Winnik [28] and for concentrated samples by T. Kyu and P. Mukherjee [29], and E. Marsano and G. Fossati [26]. The spinodal temperature is, to our knowledge, reported here for the first time [30-32]. The place where both temperatures coincide should indicate the location of the critical point (between 5 and 10 wt%) which is shifted to higher concentrations with respect to a monodisperse sample of same $M_w$ [33]. It must be noted, that contrary to pure polydisperse phase diagrams originating from a Flory-Huggins-like modeling, HPC phase diagram is very weakly dependent on concentration up to large concentrations like is PNIPAm phase diagram [34].

**Figure 2.** Cloud point temperatures and spinodal temperatures measured as described in text
3.2. Fabrication of HPC membranes and morphological structure characterization

3.2.1. Simultaneous phenomena involved in the membrane formation mechanisms

The membrane preparation is represented in Fig. 3: once the polymer was dissolved in water at ambient temperature, the polymer solution was cast onto a glass plate at two casting speeds and then placed onto a heating plate. Both the target temperature and the temperature ramp could be modified during the heating of the polymer solution for inducing the phase inversion. At a temperature higher than the cloud point temperature of the HPC/water system, three phenomena occurred concomitantly: phase separation, crosslinking and water evaporation. At the end of this stage, the temperature was reduced until ambient temperature and the membrane was washed in a water bath before storage.

Figure 3. Schematic of membrane preparation

Polymeric membranes made from HPC/water system were prepared using LCST-TIPS method and a chemical crosslinking was performed using Glutaraldehyde (GA). The chemical crosslinking aimed at (i) preventing the membrane dissolution in water solution in filtration operation and (ii) fixing the membrane morphology to obtain a porous structure. As aforementioned, a PEG200 was added to the
formulation as porogen agent. The chemical crosslinking was performed concomitantly with the phase separation, i.e. the crosslinking agent was added in the initial formulation just before casting the solution onto the glass plate. The temperature ramp was carefully controlled during membrane formation and solvent evaporation occurred since the system was open to air; three phenomena were involved in the membrane formation mechanisms: phase inversion, crosslinking and solvent evaporation. Each phenomenon was affected by both others: for instance, the crosslinking could have an influence on phase inversion dynamics by reducing the polymer chain mobility; in the same time, the evaporation could affect the crosslinking due to the progressive increase of the polymer concentration, and phase inversion and the solvent evaporation are also interdependent: on the one hand, evaporation during phase inversion leads to a deviation of the composition path in the phase diagram and on the other hand, the temporal variation of the lean composition phase due to phase separation dynamics affects the evaporation dynamics. To start deciphering this complex interplay, two temperature ramps were tested (2 and 120 °C/min), leading to different membrane formation dynamics in terms of solvent evaporation, crosslinking rate and phase separation dynamics.

3.2.2. Crosslinking

• Closed system (without solvent evaporation)

Crosslinking dynamics was first followed in closed system using a rheometer (Physica MCR 301, Anton Paar). In these conditions, water evaporation was not possible and hence the polymer concentration was kept constant during the experiments. The tests were performed under small amplitude oscillatory shear stress (1%) and a frequency of 1 Hz. Surprisingly, the elastic modulus $G'$ did not cross the viscous modulus $G''$, even after 16 hours of experiments despite the presence of the crosslinking agent and the catalyzer in appropriate proportions. This result exhibited that the crosslinking was not efficient enough in such operating conditions, i.e. the number of bonds between polymer chains was too low for inducing a sol-gel transition. Indeed, the creation of bonds between HPC polymer chains was hindered because of HPC chemical structure: due to the random substitution of -OH groups by -OR groups (Fig. 1), at a polymer concentration of 20 wt% and using
GA, the probability to have a favorable conformation for chemical crosslinking was quite low, thus explaining the low crosslinking efficiency in such closed system.

- **Open system (with solvent evaporation)**

During the membrane preparation, the system was open to atmosphere and solvent evaporation occurred simultaneously to phase inversion and crosslinking. In this case, the polymer concentration progressively increased over time until the membrane structure was fixed (after almost complete water evaporation). The crosslinking efficiency was characterized by checking the membrane stability after its immersion in water during several days. Swelling and re-solubilization ratios were determined for the two different temperature ramps: 2 and 120 °C/min. For those two conditions of temperature ramp, we demonstrated that HPC membrane did not dissolve in water after 40 days of immersion (the re-solubilization was null). Such results obtained in water bath clearly demonstrated the efficiency of the chemical crosslinking reaction performed during membrane formation. The same test was then determined in two aprotic solvents (dimethylsulfoxide and tetrahydrofurane), and two protic solvents (methanol and chloroform). In all case too, no re-solubilization was observed, hence demonstrating that the crosslinking was efficient enough to prevent the dissolution in polar aprotic solvents as well.

### 3.2.3. Membrane morphologies

Classically, the membranes formed by UCST-TIPS process exhibit symmetric morphologies since the heat transfer induced by a temperature quench is fast comparing to the mass transfer, which would create anisotropy in the membrane structure; Indeed it has been previously demonstrated that anisotropic structures could be formed if an evaporation stage is done prior to TIPS process [8-11]. For this work, the SEM photographs of the final membranes prepared by LCST-TIPS method were reported in Fig. 4.

| Membrane HPC/GA/PEG 120 °C/min | Membrane HPC/GA/PEG 2 °C/min |
Top surface

Membrane cross-section

Top cross-section

Middle cross-section

Low cross-section

(a) 1.2 µm

(b) 59.5 µm

(c) 30.0 µm

(d) 50.0 µm

(e) 1.2 µm
The membranes were prepared using the procedure described in the material and methods section and the formulation was the same as the one chosen for the crosslinking experiments: HPC 20 wt%, GA 5 wt%, HCl 1 wt%, PEG200 5 wt%. For a temperature ramp of 120 °C/min, Figures 4a and 4f present the SEM of top and bottom surfaces, respectively, and Figures 4b, 4c, 4d and 4e present the SEM of cross-sections at different locations in the membrane thickness. For a temperature ramp of 2 °C/min, Figures 4g and 4l present the SEM of top and bottom surfaces, respectively, and Figures 4h, 4i, 4j and 4k present the SEM of cross-sections. The top and bottom surfaces of the membranes seem to show a very low apparent porosity, whatever the temperature ramp, meaning that low pore sizes can be expected in both operating conditions. These photographs exhibit that the final membrane thickness in dry state after solvent evaporation is close to 50 µm whatever the temperature ramp, i.e. close to 20% of the initial thickness (250 µm). This indicates that the global porosity of the HPC membrane is quite low. Nevertheless, zooms performed near the top surface (Figures 4c and 4i), at the middle (Figures 4d and 4j) and at the bottom (Figures 4e and 4k) of the membrane show that HPC membranes are porous in the whole thickness, for both temperature ramps. SEM observations in Fig. 4 suggest that the phase separation led to the formation and the coexistence of two interconnected phases, a rich phase composed of concentrated HPC and a lean phase mainly composed of water and GA. This polymeric structure was bicontinuous, reminiscent of spinodal decomposition mechanism; this result was expected for two main reasons: (i) as shown by the phase diagram determined in this work and reported in Fig. 2, the spinodal and binodal curves were shown to be very close, and hence the metastable region is very thin for the HPC/water system.
Furthermore, (ii) the target temperature was significantly higher than the spinodal curve at given polymer concentration, and hence whatever the temperature ramp, the system did not stay enough time in the metastable region to induce nucleation and growth mechanism that would have formed cellular structures in the membrane.

Fig. 4 also shows that although the HPC membrane seemed to be isotropic regarding the global cross-section photographs (Fig. 4b and 4h), the zooms at different locations in the membrane thickness exhibit anisotropy between the top and the bottom cross-section. More precisely, the porosity is higher near the upper interface (air/membrane interface) than below in the membrane thickness, and the more dense structure can be observed just at the middle cross-section. This observation was done whatever the temperature ramp, i.e. 2 or 120 °C/min, but the morphologies obtained at these different temperature ramps are different: the membrane porosity is lower when the membranes were prepared at 2 °C/min comparing to those prepared at 120 °C/min. A difference of evaporation rate combined to different cross-linking efficiencies between 2 and 120 °C/min would be probably the reason that explains those differences. Indeed, since the cross-linking was performed by a chemical reaction between glutaraldehyde and HPC polymer chains, its efficiency directly depends on the temperature, and as a consequence also on the temperature ramp because all phenomena involved in the membrane structuration are coupled. Faster the temperature will be reached during membrane formation, earlier the membrane structure will be fixed by chemical cross-linking. At 120 °C/min, the target temperature is reached after only 20 s (from 20 °C to 60 °C) whereas it is reached after 20 min at 2 °C/min. At the lowest temperature ramp, the evaporation is therefore expected to be almost finished when the membrane structure is fixed and most of the water lefted the system. In this case, the polymer chains keep on moving and collapse during water extraction, thus forming the dense-like structure presented in Fig. 4i, 4j, and 4k. On the contrary, at 120 °C/min, the morphology is fixed much earlier by cross-linking and hence the morphology is fixed whereas the water extraction by evaporation is not finished, thus keeping higher porosity in the membrane structure.
Finally, SEM photographs were obtained at different concentrations of PEG200 (from 0 to 5 wt%), which exhibited that the membrane morphology was not affected by the formulation (figure not shown in the paper).

3.2.4. Water sorption

Water sorption experiments were performed on HPC membranes with and without PEG200 in the initial formulation. The membranes were prepared at the two aforementioned temperature ramps: the membranes were stored during 40 days after their preparation in a water bath at ambient temperature (Table 3). In Table 3, the water sorption is expressed in swelling ratio (SR) defined in section 2.2.4.5.

Table 3. Swelling ratio (SR) of HPC membrane after 40 days storage in water bath.

<table>
<thead>
<tr>
<th></th>
<th>120 °C/min</th>
<th>2 °C/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>HPC/GA</td>
<td>45% ± 4%</td>
<td>87% ± 8%</td>
</tr>
<tr>
<td>HPC/GA/PEG</td>
<td>150% ± 13%</td>
<td>210% ± 19%</td>
</tr>
</tbody>
</table>

The values of water sorption not only depend on the formulation, and more precisely the presence of PEG200, but also on the temperature ramp imposed during the membrane preparation by LCST-TIPS method. The table 1 exhibits that the water sorption was three times higher when the PEG200 was added in the initial formulation, going from 45% to 150%, meaning that the porogen enhances hydrophilicity of the polymeric matrix, thus promoting the water intake. The great difference of water sorption observed depending on the formulation demonstrated that a significant proportion of porogen remained in the membrane after washing. Hydrophilic polymers are known to favor the water intake but the water sorption strongly depends on the cross-linking efficiency as well. Table 1 points out that the chemical cross-linking by GA was clearly more efficient at the highest temperature ramp; indeed the water sorption was divided by two without porogen between
2 °C/min and 120 °C/min and by almost 1.5 when the PEG200 was added to the initial formulation, from 210% to 150%.

Fig. 4. exhibited that the membrane morphology was different depending on the temperature ramp: the structure was denser, i.e. the porosity was lower, in the middle of the membrane thickness at the lowest temperature ramp 2 °C/min. In the same time, the water sorption was shown to be much higher at this temperature ramp, indicating that the membrane was almost a hydrogel in such conditions. On the contrary, at 120 °C/min, the porosity is higher in the membrane thickness and the water sorption is much lower, what confirms that the chemical cross-linking was more efficient at 120 °C/min than at 2 °C/min, probably since the chemical reaction is favored at high temperature and even if the target temperature (60 °C) was the same for both conditions, it was reached earlier at 120 °C/min than at 2 °C/min, i.e. 40 s and 1200 s, respectively.

3.2.5. Mechanical properties

First, the influence of the casting speed was investigated on the mechanical properties of HPC membranes. Table 4 reports the tensile strength, the maximum elongation and the Young modulus for two casting speeds, i.e. 3.4 and 15.0 cm/s.

<table>
<thead>
<tr>
<th>Casting speed</th>
<th>Tensile strength (MPa)</th>
<th>Maximum elongation (%)</th>
<th>Young module (Mpa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.4 ± 0.2 cm/s</td>
<td>1.46 ± 0.2</td>
<td>1.00 ± 0.1</td>
<td>1173 ± 100</td>
</tr>
<tr>
<td>15.0 ± 0.5 cm/s</td>
<td>3.34 ± 0.2</td>
<td>2.15 ± 0.1</td>
<td>1011 ± 100</td>
</tr>
</tbody>
</table>

These results clearly exhibit that the mechanical properties of HPC membranes cast with the higher shear rate were improved compared to those of membranes prepared with the lower one. Rheological conditions during casting and spinning have been proved to be an important parameter influencing the final membrane performance in the case of dense and porous active layer, as well
It is assumed that high shear rate promotes the polymer chain orientation resulting in their closer packing in the bulk matrix. This effect can account for our findings regarding the improvement of mechanical properties. Despite the modest range of casting speed rate studied, a significant increase of tensile strength and elongation at break was obtained. It can then be expected that even much better properties should be reached in fabrication of HPC hollow fibers by spinning in which high shear rates are usually applied. For the rest of this paper, only data obtained with membranes prepared using the maximum casting speed attainable with our set-up will be presented.

The influence of the temperature ramp and of the formulation on the mechanical properties was also investigated and the results are reported in Table 5. As mentioned in the section 2.2.4.4, three successful measurements were used to obtain average values for mechanical properties. For each condition, one characteristic graph representing the tensile stress versus the tensile strain was reported in Fig. 5.

### Table 5. Influence of the temperature ramp on the mechanical properties of HPC membranes (HPC 20%, GA 5%, HCl 1%) prepared by LCST-TIPS method at 60 °C

<table>
<thead>
<tr>
<th>Temperature ramp</th>
<th>120 °C/min</th>
<th>2 °C/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>Membranes HPC/GA</td>
<td>Tensile strength (MPa)</td>
<td>3.34 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>Maximum elongation (%)</td>
<td>2.15 ± 0.1</td>
</tr>
<tr>
<td></td>
<td>Young module (MPa)</td>
<td>1011 ± 200</td>
</tr>
<tr>
<td>Membranes HPC/GA/PEG200</td>
<td>Tensile strength (MPa)</td>
<td>1.7 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>Maximum elongation (%)</td>
<td>0.46 ± 0.1</td>
</tr>
<tr>
<td></td>
<td>Young module (MPa)</td>
<td>553 ± 100</td>
</tr>
</tbody>
</table>
Figure 5. Tensile stress versus tensile strain corresponding to the mechanical properties presented in Table 5.

As can be seen, whatever the formulation, the temperature ramp did not show any drastic effect on the mechanical properties. It was concluded that the polymer chain arrangement seems not to be disturbed too much by the crosslinking at surface or at depth. It should be noted that an expected decreasing trend was observed in the latter case. By contrast, the presence of a porogen (PEG200) in the initial formulation strongly affected the whole mechanical properties. The elongation at break markedly decreased indicating a likely anti-plasticizer effect of PEG200. Although the drop of both
the tensile strength and Young modulus is not in agreement with this assumption, our findings consistent with previous report showing that extruded films made of HPC and 1% PEG400 exhibited instability of mechanical properties with a continuous decrease of elongation at break and tensile strength during storage [38].

At this point, the obtained mechanical properties were not high enough to enable experiments under pressure. For this reason, the HPC polymer dope solutions were cast onto a non-woven PET support prior to further filtration. SEM cross-section of supported membranes is presented in Fig. 6, exhibiting a uniform HPC layer that did not clog the non-woven porosity.

![SEM cross-section of supported membrane](image)

**Figure 6.** Cross-section of HPC membrane onto a non-woven PET support

### 3.2.6. Permeability results

The procedure for the filtration tests was detailed in the material and methods section. A membrane conditioning precedes the filtration tests, corresponding to a gradual increase of pressure from 0 to 3.5 bar each 0.2 bar.

The filtration tests were conducted in this work for different kinds of membranes, i.e. different formulations and different preparation procedures: membranes with or without PEG200 were studied; two temperature ramps were tested (2 and 120 °C/min) during the membrane preparation
by LCST-TIPS process in the Linkam hot stage; for the highest temperature ramp, membranes were prepared under 50% and 100% RH. The results of filtration measurements were reported in Table 6.

**Table 6.** Permeability of HPC membranes for two temperature ramps (HPC 20 wt%, GA 5 wt%, HCl 1 wt%), under 50 % and 100 % RH, with or without porogen agent (PEG200).

<table>
<thead>
<tr>
<th>Temperature ramp</th>
<th>Membrane HPC/GA</th>
<th>Membrane HPC/GA/PEG</th>
</tr>
</thead>
<tbody>
<tr>
<td>120 °C/min (RH 50 %)</td>
<td>0 L/h/m²/bar</td>
<td>12.2 L/h/m²/bar</td>
</tr>
<tr>
<td>2 °C/min (RH 50 %)</td>
<td>0 L/h/m²/bar</td>
<td>1 L/h/m²/bar</td>
</tr>
<tr>
<td>120 °C/min (RH 100 %)</td>
<td>0 L/h/m²/bar</td>
<td>1 L/h/m²/bar</td>
</tr>
</tbody>
</table>

The whole membranes prepared without PEG200 were almost not permeable to water, even for the highest pressures applied and whatever the preparation procedure. By contrast, adding the porogen agent in the initial formulation helped opening the membrane porosity, and in this case the filtration performances were significantly affected by the preparation procedure: at 2 °C/min the membrane permeability was very low and the flux was almost null until 3 bars. At 3 bars, a flux around only 3 ± 0.2 L/h/m² was detected. By contrast, the membranes prepared with PEG200 under 50% RH at 120 °C/min, were permeable. The water filtration flux was around 38 ± 5 L/h/m² at 3 bars in these conditions. This result clearly indicates that the membrane preparation procedure had a significant effect on the membrane permeability, suggesting that the membrane morphology could have been also affected by the membrane formation procedure. The Figure 7 represents the flux versus the pressure for the membranes obtained with PEG200 under 50% RH at 120 °C/min.
At 2 °C/min, the previous results of water sorption exhibited that the water intake within the membrane is high compared to 120 °C/min, suggesting that the membrane could be assimilated to a hydrogel. Such a behavior was related to a poor cross-linking at 2 °C/min leading to the formation of a hydrogel, allowing water sorption in large amounts when immersed in water. The SEM photographs pointed out that the membrane structure was denser at 2 °C/min than at 120 °C/min, in agreement with the permeability results, and they also exhibited that the filtration layer was located in the middle of the membrane, especially at 2 °C/min. In this region, the membrane structure is almost dense, explaining the poor filtration performances of the membranes prepared in these conditions.

On the contrary, at 120 °C/min, the chemical cross-linking is more efficient because the final temperature is reached after only 20 s. Thus, the characteristic length between cross-linking bonds is lower than at 2 °C/min and the water sorption is significantly reduced.
4. Conclusion

This paper deals with the development of a novel porous polymeric membrane from biobased polymer (HPC) and without the use of organic solvent. A thermally induced phase separation process was used to induce the phase inversion, but increasing the temperature above the LCST of HPC/water solution. Using optical transmission methods, the phase diagram of the polymeric system was investigated, and the cloud point curve was plotted in the polymer concentration range [1-30 wt%]. The spinodal curve was determined in the same time and it was demonstrated that the metastable region is very thin in the classical concentration range used to make membranes (10-25 wt% of polymer). SEM photographs exhibited that HPC membranes were porous in the whole cross-section, characterized by a quite low porosity, especially when the temperature ramp is low (2 °C/min). At such a low temperature ramp, it was suggested that the chemical crosslinking reaction using GA was delayed thus forming a hydrogel matrix. At higher temperature ramp (120 °C/min), the membrane showed lower swelling and better filtration performances. The prove of concept was therefore demonstrated: a membrane, stable in water, was made from water-soluble biobased polymer, which may reduce the use of non-renewable resource materials (petroleum) and reduce pollution levels, in agreement with the principles of green chemistry.

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6. References


