Study of Polysulfone and Polyacrylic Acid (PSF/PAA) Membranes Morphology by Kinetic Method and Scanning Electronic Microscopy

Chamekh Mbareck1* and Quang Trong Nguyen2
1Université des Sciences, de Technologie et de Médecine; Faculté des Sciences et Techniques, B.P. 5026, Nouakchott, Mauritanie
2P.B.S. UMR 6270 CNRS - Université de Rouen, 76821 Mont-Saint-Aignan, France

Abstract

This work focuses on the study of the morphology forming of Polysulfone (PSI) and polyacrylic acid (PAA) membranes which are prepared by mixing both of the polymers in DMF and the obtained blend is precipitated in water (non-solvent). The precipitation kinetic and the effects of polysulfone concentration, drying-time in free-air and proportions of both polymers are investigated.

Based on the SEM technique, the kinetic and viscosity measurements, and the visual observations, this study brings to the fore the different steps of the morphology forming of PSF/PAA membranes: formation of finger-like structures, sponge-like-structures, inner pores, superficial pores and the craters. Appearance of these structures is governed by the exchange process, between casting polymer-solution and the precipitation-bath (coagulation), which is controlled by operating conditions. This work constitutes a great tool in understanding the mechanisms of the morphology forming of PSF/PAA membranes and to improve their performances: preparation of tailored membrane.

Keywords: Polysulfone; Poly (acrylic acid); Membrane morphology; SEM; Blends; Macrorvoids

Introduction

Polysulfone is a hydrophobic polymer with excellent chemical and mechanical properties. However, the hydrophobic behaviour prevents its use in several areas such as in water treatment. In order to overcome this drawback, some researchers have proposed the incorporation of hydrophilic groups, to increase the polymer hydrophilicity which improves the membrane water- permeability and reduces membrane-fouling [1,2].

In literature, the synthesis of carboxylated polysulfone membranes was realized through different ways. Membranes were prepared by (i) substitution with functionalized groups, (ii) by polymer graft modification or (iii) by surface modification of prefabricated membranes [2-9].

In a previous work we proposed the preparation of carboxylated polysulfone membranes by mixing PSf and PAA in dimethyl formamide (DMF) solvent and, the obtained blend, was precipitated in water-bath (non-solvent) [10]. Incorporation of carboxylic groups in PSf matrix imparted the obtained membrane high ionic-exchange and complexation capacities. It also increased the membrane hydrophilicity which enhanced the resistance to organic fouling and made membrane suitable for many applications. As a result, it was found that, the membrane properties depended on: (i) uniform distribution of PAA chains in the network formed by the PSf matrix, (ii) rate of immobilization of PAA, and (iii) morphology of the membrane. The PSf/PAA membrane has shown good performances for the ultrafiltration of different heavy metal solutions and dye molecules [10,11]. Here, our interest will be focused on the PSF/PAA membrane morphology in order to learn much about the mechanisms of the morphology forming and to know how to improve the membrane performances in view to prepare tailored membranes.

SEM technique, visual observation, precipitation speed, viscosity measurement are used to characterize the morphology of the PSF/PAA membranes.

Experiment

Polymer properties

Polysulfone (PSF) and Poly (acrylic acid) (PAA) were supplied by Aldrich and dimethyl formamide (DMF) by Prolabo. All polymers and chemical products were used as supplied, without any further purification. Molecular weights of PAA and PSF were 450 000 and 26 000 g/mol, respectively.

Membrane preparation

Polysulfone (PSF) and Poly (acrylic acid) (PAA) were dissolved, separately, in dimethyl formamide (DMF) in a glass reactor equipped with a mechanical stirrer and thermostated at 90°C for over 3 hours. The PSF concentration was 17 (or 19) wt. % and that of PAA was 5 wt. %. Afterwards, both solutions were mixed together in known proportions, stirred for 30 min and de-bubbled. Such de-bubbled casting-solution was casted on a glass plate with a lab-made Gardner knife, dried in free-air for a known time (generally 20 seconds) and finally immersed in a coagulation-bath containing a sufficient volume of MilliQ water (18.2 MΩcm) at 18°C. Membranes were thoroughly washed with water, and stored in a dilute sodium azide solution till their use.

Kinetic measurement

Five polymer-solutions with the same mass (2 g) and different composition (PSF/PAA 100/0, 96/04, 92/08, 89/11 and 83/17) were poured on a glass plate to get exactly a same surface area. Afterwards,
polymer casting-solution is immersed in equal volume of water coagulation-bath (2 liters).

Kinetic measurements were used to determine the time and the speed of PSf/PAA precipitation. The time of precipitation corresponds to the period separating the moment of immersion of casting-solution in the water-bath and the moment at which the formed membrane peels spontaneously off from the glass plate. The speed of precipitation was calculated according to the following formula: 

$$S_p = \frac{m}{t_p}$$

Where, $S_p$ is the precipitation speed of the polymer solution (in gr/sec); $m$ is the mass of casting-solution (in gram); $t_p$ is the time of precipitation (in second).

**Viscosity measurement**

Rheological experiments were carried out using a controlled shear stress (advanced rheometer: AR 2000 from TA instruments). A double gap concentric cylinder device was used. Measurements were done at $25 \pm 0.1^\circ C$ with the shear rates ranging from 0.1 to 660 s$^{-1}$ and shear stress from 0.01 to 622 Pa.

**Scanning Electronic Microscopy**

Membrane surfaces and sections were analyzed by a Zeiss EVO 40 EP microscope with the Secondary Electron Backscattered Detector (CE-BSD). Samples were frozen in liquid nitrogen at -190°C, fractured and vacuum coated with a thin gold film. The SEM observations were carried out at various magnifications while the electron beam energy was fixed at 10 keV.

**Results and Discussion**

**Kinetic measurement**

Kinetic measurements aim at characterizing the behaviour of PSf/PAA membranes during precipitation step. Therefore, precipitations time and speed were determined. It is interesting to note that precipitation time may be also regarded as time of total exchange of solvent and non-solvent between casting-solution and precipitation-bath.

As shown in Table 1, the high precipitation speed of PSf solution-film, compared with those of PSf/PAA solution-films, is the consequence of larger difference in the solubility parameter values of PSf (12.9 cal$^{1/2}$/cm$^{3/2}$) and water (23.4 cal$^{1/2}$/cm$^{3/2}$) [12]. The addition of PAA (14.6 cal$^{1/2}$/cm$^{3/2}$) on the PSf solution - to form a blend - decreases the difference in solubility between polymer-solution and water; and increases the precipitation time of PSf/PAA blends as shown in Figure 1 [13].

It’s well known that PAA is a hydrophilic polymer which can develop strong interactions with water. Then, the increase of PAA content in the casting-solution reduces chemical driving forces of exchange with non-solvent and such a thing delays the precipitation of the casting-solution [14,15].

As a result, the precipitation of PSf/PAA blends depends on the rate of the solvent expulsion from casting-solutions. This expulsion is governed by the difference in the solubility parameter or the affinity (interactions) between polymer and coagulation-bath.

**Effect of the PSf concentration in the casting-solution**

Figure 2a and b presents the SEM micrographs of PSf membranes obtained from the casting-solution containing, respectively, 17% and 19 wt. % of PSf in DMF. These micrographs show that with the increase in the PSf concentration, the membrane morphology changes from finger-like structures to sponge-like structures. Some researches assign the appearance of finger-like structures to instantaneous precipitation of PSf-film [15-17]. Here, suppression of macrovoids, within membrane with higher percent (19 wt % PSf), is mainly due to the increase of the viscosity of casting-solution as both of the casting-solutions have practically the same precipitation time [15,18].

At higher polymer content, growth of the nucleus of poor-polymer phase is disrupted by the high viscosity of the liquid medium; the vitrification of the rich-polymer matrix surrounding the nucleus would occur more easily. However, the macrovoid suppression is accompanied by a change in the porosity, pore size and pore distribution in the membrane [19-21].

**Effect of drying time in free air (pre-coagulation) ($t_d$)**

Three PSf/PAA 96/04 membranes are used to study the effect of drying time on the membrane morphology. Two samples are,
In fact, it is difficult to give a single value to the pore size as the majority is composed of different porous structures as shown in Figure 4. The SEM image of membrane surface shows that each superficial pore contains smaller inner-pores at different levels. For instance, the average size of the superficial pore in the PSf/PAA 92/08 membrane was ca. 680nm, while that of the first inner-pores level was 130nm and the second inner-pores level was 25 nm. It appears that the pore sizes in the membrane active-layer decrease progressively from the upper surface towards the inner section. Progressive decrease of these pore sizes and good performances of these membranes for ultrafiltration of heavy metal ions, published previously [11], suggest the high membrane tortuosity from one face to another.

The exposure of PSf/PAA/DMF film in free-air provokes an exchange between the solvent in casting-film (DMF, b.p. 152.8°C) and the atmospheric water vapour (at 40% relative humidity) [22]. Due to the low solvent volatility and low water content in air, the exchange rate of solvent and nonsolvent (water) is lower in free-air than in water-bath. A short drying time (a rapid quenching) of the casting liquid-film in the coagulation bath induces a rapid film vitrification (phase separation) due to further solvent outflow. This phenomenon provokes high growth of rich-polymer phase in the sub-layer and growth of poor-polymer phase in the bottom layer. These circumstances favor the formation of finger-like structure forms. For long drying time, the exchange of solvent and non-solvent is slow, as well as, the solution-film vitrification. This behavior leads to a more homogeneous structure (more sponge-like structure).

The precipitation of casting-solution in free-air is very slow in comparison with the precipitation in water-bath which is fast. In fact, the high boiling-point of solvent and low inflow rate of water-vapour respectively, dried during 10 and 300 seconds before immersing in water-bath whereas the third is completely dried in free air (water vapour 40%).

This experiment shows that the size of superficial pore increases with the increase in drying time (Figure 3), and macrovoids in the section are completely suppressed for t> higher than 300 seconds (Figure 3C). The size of superficial pores increases from 100 nm at a short time to 10 µm at a long drying time.

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The precipitation of casting-solution in free-air is very slow in comparison with the precipitation in water-bath which is fast. In fact, the high boiling-point of solvent and low inflow rate of water-vapour
in casting-solution reduced sensibly the exchange driving forces between casting-solution and water. Thus, the vitrification of solution-film progresses slowly leading to homogeneous matrix with inner and superficial pores.

These results show again the role of the driving forces arising from solvent and non-solvent exchange: rapid exchange of solvent and non-solvent favors the formation of finger-like structures, whereas the slow exchange enhances the formation of sponge-like structures.

Effect of the blend composition

Figure 5 shows the cross-section and surface morphologies of PSf/PAA 100%, 96/04, 92/08 and 89/11, respectively. The SEM micrographs show that the morphology of PSf 100% is mainly composed of finger-like structures which progressively disappear in favor to sponge-like structures with the increase in PAA content.

The progressive suppression of finger-like structure is in agreement with the previous results (section 3.1). The addition of PAA increases the intermolecular interactions between casting-solution and the non-solvent (water). Such a thing delays the precipitation of casting-solution and enhances the homogeneous structures forming. These results are in agreement with those of Wu et al. [23] who suggest that weak interactions between polyvinylidene fluoride and poly (ethersulfone) chains tend to create a large distance between the polymers chains which induces the formation of macroporous structures.

Visual observation of casting-solution precipitation

Visual observation of the precipitation of PSf/PAA/DMF casting-solution shows a strong film-contraction and thin turbulence-liquid streams which escape rapidly from the top-face of the nascent-membrane. These liquid-streams appear at numerous superficial points. Afterwards, the film peels off from the glass plate pushed by a large amount of solvent which emerges from the membrane bottom-face. This phenomenon strength decreases with the increase of PAA percentage.

The simple analysis of this phenomenon leads to admit that, the immersion of PSf/PDF casting-solution in non-solvent-bath creates strong repulsive forces between PSf chains (hydrophobic) and water molecules (hydrophilic). These forces provoke the rapid retraction of polymers’ chains and the formation of rich-polymer domains and rich-liquid domains. The progressive retraction of polymers’ chains pushes the rich-liquid domains to escape through the polymer matrix inducing the formation of inner and superficial pores, on the one hand, and the formation finger-like structures and sponge-like structures, on the other hand. The increase of PAA percentage reduces the repulsive forces and encourages the homogeneity of membrane morphology.

Interpretation of PSf/PAA morphology Mechanism

Under the driving forces of exchange between casting-solution and non-solvent (water), large fluctuations in concentration appear at the beginning of the phase separation, leading to large stresses between the poor-polymer and the rich-polymer phases, as illustrated in Figure 6a. Transient gel-clusters would be formed due to the dynamic asymmetry in stresses between the system components. Afterwards, the clusters aggregate into large gel domains [24]. Chemical potential changes induce an osmotic pressure which causes contraction of the elastic gel domains and liquid expulsion as illustrated in Figure 6b.
Figure 6: The mechanism of the formation of macrovoids structure and surface craters. Evolution in time of the structure of the solution of polymer film cast on a glass plate:

a) in the first step of polymer gelling into a surface gel skin, just after immersion in the water coagulation bath;

b) seconds later, in the next step of gel domain formation in the sub-gel skin layer;

c) in the step of formation of a surface crater by solvent ejection under gel collapse pressure through weak points in the gel skin, followed by a self-sealing of gel skin at the break point;

d) in the step of formation of a macrovoid by water-rich solvent flow towards the bottom face under gel collapse pressure, and subsequent preferential phase separation and gel domain formation on the flow path.
superficial pore size of PSf/PAA membranes as a function of PAA percentage.

Table 2: Zero-shear viscosity values of polymer casting-dopes and Average superficial pore size of PSf/PAA membranes as a function of PAA percentage.

<table>
<thead>
<tr>
<th>PAA%</th>
<th>0</th>
<th>0.04</th>
<th>0.08</th>
<th>0.11</th>
<th>0.15</th>
</tr>
</thead>
<tbody>
<tr>
<td>η₀ (Pa S)</td>
<td>0.6</td>
<td>1.3</td>
<td>3.6</td>
<td>5.5</td>
<td>14.6</td>
</tr>
<tr>
<td>Average superficial pore size (µm)</td>
<td>0</td>
<td>2.36</td>
<td>0.49</td>
<td>0.42</td>
<td>--</td>
</tr>
</tbody>
</table>

Gel-collapse is physically described by Tanaka et al. [24] who propose that the water penetration induces stresses (osmotic stresses) throughout the different parts of gel structures and the liquid squeezing out of gel. As a result, rich polymer-domains and rich liquid-domains are formed across the gel. If the gel domain is closer to the surface, it will be easier for the squeezed liquid to escape through the weak spots on the nascent gel layer [25]. The ejected DMF-water mixture causes turbulence streams in the water-bath that we observed in the regions surrounding the weak points on the surface, as illustrated in Figure 6c. As early as, the stress is released and the solvent escaped at the weak points on the surface of the nascent gel layer, small pores would be formed at low layer viscosity and craters at high layer viscosity. This behavior is confirmed by the decrease of average pore size and the increase of zero-shear viscosity, with the increase of PAA fraction, as shown in Table 2. The round crater observed on the upper layer is due to a local collapse of the elastic skin after the escape of the liquid from the casting-film.

When there are no breakable weak points in the gel skin, the rich-water liquid would be pushed downwards to the bottom face by the collapsing gel domains. As the water transport, from the liquid-sublayer towards the bottom face, is much faster than that of molecular diffusion, the separation of the polymer-poor phase occurs faster on the path of the squeezed water-rich liquid leading to finger-like structures (macropores) in the membrane section. Bottom parts of these structures are generally larger than the top parts. Such morphology can also be explained on the basis of viscoelastic properties: larger the distance from the skin layer, lower the medium viscosity. Thus, the poor-polymer phase tends to expand more in the lower part of the nascent macropores, as illustrated in Figure 6d.

The addition of PAA increases the viscosity of polymer-solutions (as shown by Table 2) and decreases the intensity of turbulence streams observed above the nascent membrane surface. It also induces the decrease of precipitation time. All these results show that with the reduction of solvent expulsion forces or the delay of precipitation, the morphology tends to be more homogeneous and the sponge-like structures dominate as shown by Figure 5.

Higher affinity of the casting-solution to non-solvent due to the presence of hydrophilic PAA makes the polymer-gel domains more swollen and weaker. As a result, the breaking of weak spots in the gelled skin is still possible, and the deformation of the softer-gel domains is larger at higher PAA contents, leading to larger craters on the surface.

Exploring the morphology of the membrane prepared from casting solutions of polysulfone (PSf), polyvinylpyrrolidone (PVP) and N-methylpyrrolidone (NMP), Han and Nam [26] report that a membrane with a low PVP content (5%) shows larger macro pores than that with a high PVP content (20%). They suggest that the introduction of a low PVP content enhanced sufficiently the demixing ability of the casting solution to provoke a rapid collapse of polymer chains and to induce the formation of macropores. On the contrary, at high PVP content, the increase in the viscosity of the polymer solution due to the enhanced entanglement of polymer chains hinders the solvent-non-solvent exchange, delays the demixing of casting-solution and induces a suppression of macropores. These results may be interpreted by the mechanisms that we propose to explain the morphology of PSf/PAA membrane.

**Conclusion**

This work has elucidated many points of the morphology mechanisms forming of PSf/PAA membrane morphology. Generally, these membranes present two morphologies: sponge-like structure and finger-like structure. The appearance of these morphologies is governed by the driving forces of solvent and non-solvent exchange. These forces provoke the retraction of polymers' chains which push the solvent to escape through the porous matrix inducing the formation of different morphologies: inner and superficial pores, finger-like structures and sponge-like structures. The decrease of these driving forces favors the appearance of sponge-like structure whereas their increase favors the appearance of finger-like structure.

**References**