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Eprints ID: 15580

**To link to this article**: DOI:10.1016/j.memsci.2013.12.049


**To cite this version**: Emin, Clélia and Remigy, Jean-Christophe and Lahitte, Jean-Francois *Influence of UV grafting conditions and gel formation on the loading and stabilization of palladium nanoparticles in photografted polyethersulfone membrane for catalytic reactions.* (2014) Journal of Membrane Science, vol. 455. pp. 55-63. ISSN 0376-7388

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Influence of UV grafting conditions and gel formation on the loading and stabilization of palladium nanoparticles in photografted polyethersulfone membrane for catalytic reactions

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Abstract

This paper reports innovative coupling for the development of catalytic membrane reactors. Photografting for the modification of membrane surfaces has been extensively studied and we used the results to optimize metal nanoparticle loading. Polyethersulfone hollow fibers were modified by poly(acrylic acid) grafting and the impact of a cross-linked network and a polymer gel grafted layer on metal loading was examined. Transmission electron microscopy images showed well dispersed metal nanoparticles close to the surface. The catalytic activity of the different hollow fibers was assessed by the reduction of p-nitrophenol into p-aminophenol and a conversion yield of close to 100% was found. We demonstrated the possibility of obtaining various reactors by varying the residence time through alteration of the grafted layer.

Keywords:
Hollow fiber membrane
UV photografting
Functionalization
Palladium nanoparticles
Catalytic membrane reactor

1. Introduction

Polymers have been widely studied in order to modify their properties. Various techniques such as blending, coating or chemical modifications have been successfully developed during the last decade [1–3]. UV irradiation graft polymerization is one of these techniques [4,5], which presents many advantages including low energy costs, simplicity and cleanliness of the method and long-term chemical stability [6]. In membrane science, this technique has mainly been reported for the reduction of fouling on various polymer membranes [7,8]. Polyethersulfone (PES) membranes are widely used in water treatment and are well known for their mechanical strength, thermal and chemical stability. Moreover, the advantage of poly(arylsulfone) membranes is their intrinsic photosensitivity to UV irradiation provided by the phenoxypyphenyl sulfone chromophore in the backbone of the polymer chain leading to the production of free radicals [9,10]. In the presence of vinyl monomers, e.g. acrylic acid, grafting occurs with covalent bonds being formed between the membrane support and graft chains [11]. The grafted chains remain at the surface without affecting the bulk properties. Thus, with this technique, it is possible to obtain various specific surface properties depending on the nature of the monomers.

Currently, a great interest is focused on nano-scale materials due to their unique properties (e.g. high specific surface areas, special optical properties) that differ from those of the bulk material. More specifically, metal nanoparticles (MNP) such as those of platinum group metals are considered as an interesting choice for catalytic reactions [12]. In the nanoparticle state, palladium is well known to have the ability to adsorb hydrogen into its lattice and dissociate it. The unique affinity of this metal towards hydrogen makes it a good catalyst for various hydrogenation reactions [13]. However, without stabilizing agents such as polymers or surfactants [14], nanoparticles tend to agglomerate in solution forming aggregates and thus reducing their high efficiency. The functionalization of membrane surfaces with charged groups can stabilize nanoparticles in the polymeric matrix [15]. These functional groups are able to retain metal ion nanoparticle precursors. Once the precursor is immobilized at the membrane surface, MNP can grow by chemical reduction of metal ions. MNP thus become stabilized inside the grafted layer.

Modified membranes can then be used for an innovative application in terms of water depollution. Currently, water treatment is ineffective for many polluting molecules like pesticides or endocrine disruptors. These compounds are a major concern due their high toxicity in groundwater and drinking water and a combination of traditional membrane processes and catalysis could be an opportunity to deal with them. Catalytic membrane
reactors (CMR) containing MNP can perform the reaction and separation steps simultaneously in the same unit. However, the very low concentrations of the target molecules and the toxicity of the by-products can affect the viability of such processes. In the chemical industry, membrane processes involving catalysts can be an alternative to the synthesis of chemical compounds. For example, Astruc [16] mentions numerous reactions catalyzed by transition metal nanoparticles. However, the use of nanoparticles in reactors such as fluidized beds involving cohesion forces could lead to aggregation of the nanoparticles [17] due to interaction forces such as London–van der Waals or electrostatic interactions. Additionally, recovery of the nanoparticles in a continuous process could pose further problems. Catalytic membrane reactors appear to offer good prospects to overcome these limitations, especially with MNP.

Research with various membrane modifications for the stabilization of MNP in the polymer matrix have already shown good results. Ouyang et al. [18] reported the catalytic reduction of 4-nitrophenol with NaBH₄ using catalytic hollow fiber membranes prepared using layer-by-layer polyelectrolyte deposition and gold nanoparticles. Smuleac et al. [19] effectively modified poly(vinylidene fluoride) (PVDF) by in situ polymerization of acrylic acid for dechlorination applications. This functionalized membrane contained Fe or core-shell Fe/Pd nanoparticles formed by reduction with NaBH₄. Doménech et al. [20] used polymeric membranes made with sulfonated PES (PES-Cardo) and nanoparticles were prepared using an intermatrix synthesis methodology. The catalytic activity of the reduction of p-nitrophenol has also been studied with Pt nanoparticles in a thin film of poly (2-vinylpyridine) [21].

In a previous paper [22], we already demonstrated the feasibility of stabilizing MNP in a grafted polymer. In the present work, we studied the impact on stabilized noble nanoparticles of modifying hollow fiber by photografting. The catalytic activity of the different membranes obtained was assessed. The influence of the nature and the thickness of graft chains was studied on palladium loading and on the reactivity of the nanoparticles. The catalytic performance of the membranes was evaluated by the reduction of p-nitrophenol to p-aminophenol, p-Nitrophenol, which is produced during the synthesis of paracetamol, is frequently used as a model pollutant. Reactions were performed in two different configurations [23]: flow through membrane contactor and in a batch reactor for comparison.

2. Materials and methods

2.1. Materials

Microfiltration PES hollow fibers MicroPES® were purchased from Membrana (Wuppertal, Germany). This membrane has a nominal pore size of 0.2 μm, an inner diameter of 300 μm and a wall thickness of 100 μm. Sodium p-styrene sulfonate, acrylic acid, NN-methylenebisacrylamide, 4-hydroxybenzophenone, sodium chloride, sodium borohydride, tetraamminepalladium (II) chloride monohydrate, toluidine blue O and p-nitrophenol were purchased from Sigma Aldrich (France). Sodium hydroxide, hydrochloric acid and acetic acid were purchased from Carlo Erba (France) and nitric acid from VWR (France). All compounds were used without further purification and solutions were prepared with deionized water.

2.2. Methods

2.2.1. Functionalization of hollow fiber surface by photografting polymerization

Hollow fibers were functionalized using a continuous UV photografting polymerization setup already described in a previous paper [24] with slight modification. Before the polymerization process, membranes were water-soaked to prevent overheating of the fiber. The membranes were then dipped into a monomer solution containing acrylic acid (AA) as monomer, 4-hydroxybenzophenone as photoinitiator and NN-methylenebisacrylamide as cross-linker. The photoinitiator ensured the creation of free radicals and the cross-linker ensured formation of a polymer network. The concentration of AA was 15 wt% and 25 wt% in water. Concentrations of cross-linker and photoinitiator are expressed in mol% of monomer. The membrane was passed through two industrial UV polychromatic lamps (UVA-PRINT LE, doped halogen lamps, lᵣ = 5520–10,080 mW cm⁻², Hoene UV France, Lyons, France). The operating rate was adjusted to between 8 and 20 m min⁻¹, corresponding to an irradiation time (IRT) of between 3.75 and 1.50 s. The filter was made of quartz. In our conditions, the thickness of the liquid film coating was determined by the Landau and Levich equation [25] and the mode of entrainment characterized by a visco-inertial mode as demonstrated by Goma-Bilongo et al. [26] in a numerical simulation of the photografting setup. For this series of experiments, the hollow fiber was irradiated in the first lamp in order to create radicals at the surface. The experiment was performed under nitrogen atmosphere as oxygen is an inhibitor for the polymerization [27,28]. Then, the fiber was passed through the monomer solution and polymerization took place under the second lamp. After grafting, all membranes functionalized with carboxylic groups (–COO–H⁺) were washed with, and subsequently stored in, deionized water to remove excess monomer and any homopolymer not covalently bound to the membrane. The water flux was measured on pristine and grafted membranes in dead-end filtration mode using laboratory-made modules. The structure and the thickness of the graft polymer of membranes washed then dried in an oven for several days were determined under a scanning electron microscope (SEM, TM-1000 Tabletop Microscope, Hitachi). Surface analysis of the membranes was performed by attenuated total reflection Fourier transform infrared spectroscopy (ATR-FTIR, Thermo-Nicolet Nexus 670). The nominal incident angle was 45°. The thickness of the membranes was determined by X-ray photoelectron spectroscopy (XPS). The concentration of palladium in the palladium complex [Pd(NH₃)₄Cl₂]⁻ according to Eq. (1). Palladium nanoparticles were then formed on reduction of the Pd²⁺ following immersion of the membrane in a 0.1 M aqueous solution of NaBH₄. Formation of the nanoparticles shifts the ion exchange equilibrium in Eq. (2) to the right since Pd²⁺ cations are consumed to form Pd⁰ [31]. Consequently, intermatrix synthesis enables the ionogenic functions, leading to another cycle of metal exchange and reduction which further increases metal loading.

\[ \text{COO}^{-} + \text{H}^{+} + [\text{Pd(NH}_3)_4\text{Cl}_2]^{2-} \rightarrow (\text{COO}^{-})_2[\text{Pd(NH}_3)_4\text{Cl}_2^{2+} + 2\text{H}^{+}] \] (1)

\[ (\text{COO}^{-})_2[\text{Pd(NH}_3)_4\text{Cl}_2^{2+} + 2\text{Na}^{+} \rightarrow 2\text{R}^{-} \text{COO}^{-} \text{Na}^{+} + [\text{Pd(NH}_3)_4\text{Cl}_2^{2+}] \] (2)

\[ [\text{Pd(NH}_3)_4\text{Cl}_2^{2+} + 2\text{H}_2\text{O} \rightarrow \text{Pd}^{0} + 7\text{H}_2 + 2\text{B(OH)}_3 + 4\text{NH}_3 \] (3)

In order to evaluate the effects of pH on ion exchange equilibrium, a set of poly(acrylic acid) (PAA) grafted membranes were immersed in a solution of sodium chloride (5.2 M) at pH 10 for at least 6 h prior to ion exchange with the palladium salt to convert –COO⁻H⁺ to the –COO⁻Na⁺ form as described by Smuleac et al. [19].
2.2.3. Quantification of palladium

Palladium loading of the membrane was determined by inductively coupled plasma optical emission spectrometry (ICP-OES, Ultima 2, Horiba Jobin Yvon). 1 cm² of membrane sample was dissolved in aqua regia and diluted in water. This diluted solution was analyzed using ICP. Measurements were repeated three times using different membrane samples on different positions along the hollow fibers. The palladium concentration in permeates was also determined in order to quantify leaching during filtration.

2.2.4. Characterization of palladium nanoparticles

The presence of Pd was confirmed by energy-dispersive X-ray spectroscopy (EDX) analysis. Images of nanoparticles were obtained by transmission electron microscopy (TEM, JEOL Jem 1400). Samples of membranes were first embedded in resin and then thin slides were cut with a microtome. The analyses provided the size and the distribution of the MNP.

2.2.5. Catalytic conversion of p-nitrophenol

The catalytic reduction of p-nitrophenol to p-aminophenol was performed in the presence of sodium borohydride solution (NaBH₄) in convection mode and batch reactor. In the batch reactor system, 8.6 cm² of membrane were immersed in 40 ml of a mixture of p-nitrophenol (0.72 mM) and NaBH₄ (14.38 mM). We assumed that the carefully mixed solution was homogeneous and only small samples were taken in order to keep the reaction volume almost constant. In flow through membrane configuration, a solution containing p-nitrophenol and NaBH₄, at the same concentration, was filtered across the membrane at room temperature using hollow fiber modules containing two fibers corresponding to a total surface area of 8.6 cm². The reducing compound was in excess (20:1), so the kinetics of the reduction could be assimilated to a pseudo first order reaction. The concentration of p-nitrophenol was monitored by UV-visible spectroscopy (λ = 400 nm, Uvilit, XTD5) versus time in order to follow the conversion.

3. Results and discussion

3.1. UV-induced membrane modification

As demonstrated in previous articles [2,4], UV photografting polymerization is an efficient method to modify a polymeric membrane surface. Fig. 1 clearly shows that altering the radiation energy received by the membrane surface (determined by the irradiation time and the power supply of the UV lamp) modifies the weight of polymer grafted onto the membrane surface. The greater the energy received by the membrane the greater the weight of polymer grafted. Three zones, with different slopes on the plot can be identified in Fig. 1. In the first zone from 0 to 12.3 J cm⁻², the energy is mostly consumed by the inhibitor/oxygen present in the monomer solution (used without purification). After this induction time, all the inhibitor is destroyed and the dissolved oxygen consumed thus the weight polymer grafted rises almost linearly with the received energy from 12.3 to 21.4 J cm⁻². Then, for energies higher than 21.4 J cm⁻², the evolution tends to become asymptotic. This result is in agreement with the observation of Goma-Bilongo et al. [26] who determined, by modeling that a total consumption of the monomer occurred for high energies received by the membrane, maximizing the amount of polymer grafted.

The pure water flux of the grafted membranes decreases with the weight of grafted polymer to reach almost zero for 0.55 g m⁻² grafting. Grafting reduces pore size and thus decreases permeability. No permeability decrease was observed with a pristine membrane irradiated in the same conditions but without reactant, indicating that, in our conditions, UV radiation did not degrade the membrane surface significantly and mass transfer properties were maintained. Permeability is plotted versus energy received by the membrane for different concentrations of 15 wt% and 25 wt% AA (Fig. 2). For both AA concentrations, a similar trend was found but with 15 wt% the permeability reached zero for a higher energy (28.5 J cm⁻²).

ATR-FTIR analysis was used to examine the change in the chemical composition of hollow fiber surface and to confirm the presence of grafted polymer. Fig. 3 shows spectra for pristine PES hollow fiber and different weights of PAA grafted onto membranes. The two bands at 1578 and 1486 cm⁻¹ are characteristic of the aromatic rings of PES. In the region below 3000 cm⁻¹, the two bands at 3060 and 3049 cm⁻¹ are attributed to aromatic C–H vibration of the membrane support. The analysis performed on grafted PAA hollow fibers showed a new absorption band at 1702 cm⁻¹ corresponding to the stretching of the carboxylic group (C=O) of PAA. The intensity of this band increased with the weight of grafted polymer, as seen from the spectra of modified membranes (b)–(d) indicating an increase of functionalized groups (Table 1). Below 2700 cm⁻¹ a broad peak corresponds to the asymmetric vibration of O–H stretching for the grafted membranes (not visible on pristine PES). Although the membranes were dried in
The variation was not linear and after 0.53 g AA m\(^{-2}\) the layer increased with the weight of grafted polymer. However, as shown in Fig. 5, the thickness of the grafted layer can also be clearly seen by TEM images. A higher porosity, as observed.

Membranes were studied by SEM to visualize surface modifications and to estimate the thickness of the grafted layer. Thicknesses of the grafted polymer layer are reported in Table 1.

After a certain degree of grafting corresponding to a grafted weight of 0.53 g AA m\(^{-2}\), the pores disappeared, and the surface becomes smoother leading to a dramatic change in the appearance of the surface with an increase of the degree of grafting. The cross section allows the grafted polymer layer (indicated by the arrow on the picture) to be clearly distinguished and its thickness measured. It appears uniform and dense (see Table 1).

The thickness of the grafted layer was plotted versus the weight of grafted polymer (Fig. 5). As expected, the thickness of the layer increased with the weight of grafted polymer. However, the variation was not linear and after 0.53 g AA m\(^{-2}\) the slope became steeper. The theoretical thickness of the corresponding dense polymer layer can be calculated from the weight of grafted polymer assuming the density of the grafted PAA to be equal to that of pure dense PAA. This calculated thickness was significantly lower than the measured one indicating that the grafted layer is porous. The calculated porosity (defined here as the ratio thickness of polymer layer/thickness of equivalent dense layer) increased with the degree of grafting and two porosities were observed: for a degree of grafting below 0.53 g AA m\(^{-2}\), the calculated porosity was around 0.7 (as seen on the calculated thickness for a porosity of 0.7) whereas above 0.65 g AA m\(^{-2}\) the calculated porosity was around 0.8.

The use of a cross-linker during the polymerization leads to formation of a polymer network. The hydrated 3D polymer network formed can then collapse during the drying step before microscopy. This phenomenon will depend on the degree of cross-linking. In the experimental conditions used, the degree of cross-linking increased with the UV energy received by the membrane. At low UV energy, polymerization was incomplete and the polymer layer only partially cross-linked. For a higher UV energy, the degree of polymerization increased and the gel point was crossed. A chemical gel was obtained. This leads to two different kinds of layer, a partially cross-linked polymer and a chemical polymer gel. After drying, both layers become porous, but the less cross-linked one will collapse more than the gel leading to a lower porosity, as observed.

The formation of a grafted layer at the membrane surface leads to a decrease of membrane permeability. This decrease is due to greater resistance to water transfer and to smaller pore diameters, the pores being progressively blocked by the grafted layer of hydrated cross-linked polymer (Fig. 6).

### 3.2. Palladium nanoparticle synthesis in the functionalized PES membranes

As shown in Fig. 7, palladium loading increased with the amount of grafted polymer. Measured values were higher than those theoretically expected for stoichiometric ion exchange between Pd\(^{2+}\) and polyelectrolyte counter ions. We observed a large increase in palladium loading after 7 meq m\(^{-2}\) of grafted PAA due to the transition of the polymer layer structure from partially cross-linked to a chemical polymer gel. As seen in Fig. 5, the chemical polymer gel is more porous than the partially cross-linked layer, allowing easier access of the Pd\(^{2+}\) to polymeric charge. This porous volume can also sterically entrap Pd\(^{2+}\) solution without need of a specific charge effect.

Pre-treating grafted membranes with a concentrated solution of sodium chloride facilitated the ion exchange capacity which slightly increased the Pd loaded into the membrane (cross sign). The results show that Pd nanoparticles can also be synthesized in a pristine membrane without any grafting at the surface, certainly due to limited adsorption of Pd salt at the membrane surface. However, the amount of Pd included in this condition is significantly lower (1.7 meq Pd m\(^{-2}\)).

TEM analysis confirmed the nanoparticle state of the Pd. Fig. 8 shows a cross section of hollow fibers analyzed by TEM. The first image corresponds to the pristine PES membrane and the other is an example of PAA grafted membrane (PES-PAA5). In the figures, arrows show the external surface of the membrane. It appears that the grafted layer can also be clearly seen by TEM images. A higher magnification of the square zone is shown in Fig. 9. The palladium detected in the pristine membrane was found to mainly occur as nanoparticles but aggregates are also clearly visible. In grafted membranes, the size of the nanoparticles was found to be smaller than 5 nm. The nanoparticles were clearly located in the grafted polymer layer, at the surface of the membrane, homogeneously and without noticeable aggregation.

In all experiments, a second loading increased the concentration of metal (Table 2). The TEM analyses, reported in Fig. 10, demonstrate the effective incorporation of an additional quantity of palladium after this second cycle. TEM observation indicated that the average diameter of nanoparticles increased from 4.7 to 15 nm, and without noticeable aggregation.

<table>
<thead>
<tr>
<th>Membrane</th>
<th>[\text{E} \text{ (J cm}^{-2})]</th>
<th>Weight of grafted polymer ((g \text{ AA m}^{-2}))</th>
<th>Thickness of the grafted polymer ((\mu m))</th>
</tr>
</thead>
<tbody>
<tr>
<td>PES</td>
<td>0</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>PES-PAA1</td>
<td>12.3</td>
<td>0.19 ± 0.01</td>
<td>0.7 ± 0.1</td>
</tr>
<tr>
<td>PES-PAA2</td>
<td>15.3</td>
<td>0.27 ± 0.02</td>
<td>0.8 ± 0.1</td>
</tr>
<tr>
<td>PES-PAA3</td>
<td>17.1</td>
<td>0.42 ± 0.05</td>
<td>1.2 ± 0.1</td>
</tr>
<tr>
<td>PES-PAA4</td>
<td>20.4</td>
<td>0.53 ± 0.08</td>
<td>1.7 ± 0.1</td>
</tr>
<tr>
<td>PES-PAA5</td>
<td>21.3</td>
<td>0.55 ± 0.05</td>
<td>2.2 ± 0.1</td>
</tr>
<tr>
<td>PES-PAA6</td>
<td>28.5</td>
<td>0.63 ± 0.01</td>
<td>3.2 ± 0.1</td>
</tr>
<tr>
<td>PES-PAA7</td>
<td>42.7</td>
<td>0.72 ± 0.07</td>
<td>3.6 ± 0.1</td>
</tr>
<tr>
<td>PES-PAA8*</td>
<td>20.4</td>
<td>0.27 ± 0.12</td>
<td>0.8 ± 0.1</td>
</tr>
</tbody>
</table>

\* \([\text{AA}] = -15 \text{ wt}%; [\text{PI}] = 0.03 \text{ mol}%; [\text{cross-linker}] = 2.7 \text{ mol}%.\)
8.4 nm corresponding to a drop in specific surface area from 106 to 59 m^2 g^{-1}. For single loading, the weighted mean diameter calculated as a function of the percentage frequency with 1 nm intervals tends to a Gaussian distribution (white diamonds). The average diameter (for a second loading) was higher than the values obtained for the heterogeneous primary nucleation. The distribution shifted to the right due to favorable secondary nucleation kinetics. Thus, the first nanoparticles incorporated in the membrane act as precursors for the second load, as described for “core–shell” nanoparticle synthesis. No or few new particles were created. The average diameter calculated from the number of particles generated during a primary nucleation and the actual Pd loading for a second cycle was around 6.8 nm, lower than the measured one. However, this value is within the margin of error.

3.3. Leaching of MNP

One of the many advantages of catalytic membranes is that as the catalyst is supported by the membrane, it does not have to be removed from the solution at the end of the process. The leaching of MNP incorporated inside the membrane was determined in order to ascertain if MNP were well trapped in the grafted layer. This conditions the viability of the process. Deionized water was filtered through the membrane, at constant pressure of 0.1 bar and the permeate was analyzed by ICP. We found a concentration in the permeate below the limit of quantification (LOQ), 15 ppb. Thus, the values shall only be expressed qualitatively. The concentration detected was slightly above the limit of detection (LOD), 3 ppb, indicating leaching close to zero. These low values for pristine and grafted membrane indicate that no leaching occurred in our membranes i.e. the MNP present inside the membrane were stabilized and well secured.
3.4. Catalytic reduction of p-nitrophenol

The reduction of p-nitrophenol into p-aminophenol is described by a Langmuir–Hinshelwood mechanism [32]. After a certain reaction time, the solution changes color, turning from yellow to colorless for total conversion, indicating the complete reduction of the nitrophenol. Moreover, for all essays, the absorbance of p-aminophenol was followed at 297 nm and a decrease of the p-nitrophenol peak was found to occur concomitantly with an increase of the p-aminophenol peak, indicating the reduction of the substrate. Thus, conversion was expressed as absorbance of p-nitrophenol.

In the experiment, sodium borohydride was used in excess, thus the apparent kinetic constant \( k_{\text{app}} \) is defined by

\[
\frac{dC_{\text{RNO}_2}}{dt} = -k_{\text{app}}C_{\text{RNO}_2}
\]

where \( k_{\text{app}} \) is the apparent kinetic constant and \( C_{\text{RNO}_2} \) is the concentration of p-nitrophenol.

3.4.1. Batch reactor

In batch reactor conditions for a reaction with first order kinetics, plotting \( C_{\text{RNO}_2} \) versus time gives access to the kinetic constant \( k_{\text{app}} \).

\[
C_{\text{RNO}_2} = C_{\text{RNO}_2}^{0}\exp\left(-k_{\text{app}}t\right)
\]

where \( C_{\text{RNO}_2}^{0} \) is the initial concentration of p-nitrophenol and \( t \) is the time.

To take into account the palladium loading of the membranes, the apparent kinetic constant \( k_{\text{app}} \) was defined as a new constant \( k \) corresponding to the ratio \( k_{\text{app}}/\text{mass of Pd} \). Experiments were performed in batch reactors for a single and double loads of Pd (Table 3).

The constant \( k \) obtained for pristine catalytic membrane was higher than that obtained for grafted membranes. This can be explained by a difference of catalyst accessibility for the reactant, nanoparticles simply and directly adsorbed at the membrane surface would be more accessible than nanoparticles entrapped inside a polymer layer. Comparing the effect of the number of cycles, results indicated better activity with single loading. This could be explained by the difference of reactivity of the nanoparticles. The MNP presented a specific surface area of 106 m\(^2\) g\(^{-1}\) for a single load leading to greater catalytic activity g\(^{-1}\) MNP than with a second loading where the specific surface area was 59 m\(^2\) g\(^{-1}\).

The strategy to obtain the best catalytic efficiency should therefore be to load a large number of small diameter nanoparticles.

3.4.2. Flow-through membrane contactor

Reactions were performed in a flow-through membrane contactor. The solution was filtered through the membrane in a dead-end filtration configuration and the permeate was analyzed after a single pass. The catalytic membrane was assimilated to a plug flow tubular reactor. Thus, assuming that the flow does not differ at the entry and at the end of the reactor, the kinetics for a first order reaction can be described as follows:

\[
C_{\text{RNO}_2} = C_{\text{RNO}_2}^{0}\exp\left(-k\tau\right)
\]

Table 2

<table>
<thead>
<tr>
<th>Membrane</th>
<th>Weight of grafted polymer (g m(^{-2}))</th>
<th>“Single loading”</th>
<th>“Second loading”</th>
</tr>
</thead>
<tbody>
<tr>
<td>PES</td>
<td>–</td>
<td>0.009 ± 0.001</td>
<td>0.030 ± 0.003</td>
</tr>
<tr>
<td>PES-PAA1</td>
<td>0.19 ± 0.01</td>
<td>0.032 ± 0.002</td>
<td>0.065 ± 0.003</td>
</tr>
<tr>
<td>PES-PAA3</td>
<td>0.42 ± 0.05</td>
<td>0.062 ± 0.009</td>
<td>0.183 ± 0.011</td>
</tr>
</tbody>
</table>

3.5. TEM images of (a) aggregate in pristine polyethersulfone and (b) external cross section of poly(acrylic acid) grafted membrane (PES-PAA5).
where $\tau$ is the residence time. In order to take into account the catalyst load, an apparent rate coefficient, $k_{app}$ was also defined as $k=k_{app}/\text{mass of Pd}$.

For each experiment, the transmembrane pressure was adjusted in order to change the permeate flux and thus the residence time. For a given pressure (i.e. given residence time), the concentration was followed as a function of the time. No decrease of conversion was observed indicating a steady state but also no loss of reactivity (which can occur in the case of catalyst poisoning). Conversion of p-nitrophenol in the permeate is plotted versus permeate flux (Fig. 11). By decreasing the flow, the p-nitrophenol concentration decreased in the permeate until quasi-total conversion. The corresponding residence time was between 0.02 and 0.09 s in the grafted reactor layer. The curve calculated from Eq. (6), was fitted to the experimental points and an apparent rate coefficient $k$ of 84 s$^{-1}$ g$^{-1}$ was found. Compared to the value obtained on the same membrane in batch conditions ($k_{\text{batch}}=3.6$ s$^{-1}$ g$^{-1}$), this constant is five orders of magnitude higher. In batch conditions nanoparticles can be considered to be located in the total volume of treated solution whereas in flow through filtration, the catalyst is concentrated in the grafted polymeric layer. This high concentration enhances the kinetics and explains such a high value of $k$. It is thus possible to obtain a good conversion yield by varying the process parameters. However, an increase of the residence time involves a reduction of

<table>
<thead>
<tr>
<th>Membrane</th>
<th>$k$ ($s^{-1} \cdot g^{-1}$)</th>
<th>Single load</th>
<th>Second load</th>
</tr>
</thead>
<tbody>
<tr>
<td>PES</td>
<td>3.9</td>
<td>-</td>
<td>-</td>
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<tr>
<td>PES-PAA1</td>
<td>2.7</td>
<td>1.8</td>
<td>-</td>
</tr>
<tr>
<td>PES-PAA3</td>
<td>1.8</td>
<td>0.7</td>
<td>-</td>
</tr>
<tr>
<td>PES-PAA8</td>
<td>3.6</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 3: Kinetic constants $k$ obtained in batch reactor for pristine and grafted membranes with palladium nanoparticles.

![Fig. 10. TEM images of the cross section of poly(acrylic acid) grafted membrane (PES-PAA3) for a single (a) and a second loading (b) and images at higher magnification for a single (c) and a second loading (d).](image)

![Fig. 11. Conversion of p-nitrophenol in the permeate versus the flux obtained after a single pass on PSS-PAA8 (palladium loading: 0.33 mg).](image)

![Fig. 12. Conversion of p-nitrophenol into p-aminophenol as a function of palladium loading for a pristine and for differently grafted membranes (PES, PES-PAA1, PES-PAA3, PES-PAA8).](image)
treated solution. Thus, for a constant flux we performed the reaction using different membranes with various thicknesses of grafted polymer.

Fig. 12 shows the conversion for a pristine membrane and 3 different grafted membranes with different thicknesses and thus different catalyst loads, at identical flux kept to 28 L h\(^{-1}\) m\(^{-2}\). We assume that the Pd load is proportional to the reactor volume due to the different thicknesses of the grafted polymer layer. Consequently, the kinetics was determined as

\[
R_m = -\frac{dQ_{\text{NO}_2}}{dt}
\]

Consequently, the kinetics was determined as

\[
C_{\text{NO}_2} = C_{\text{NO}_2}^0 \exp \left(-k_{\text{app}} \frac{k_1}{Q_v} t\right)
\]

where \(k_1\) is the proportionality coefficient between the volume of the reactor and the Pd loading determined as being equal to 110–6 L mg\(^{-1}\), \(m\) is the mass of catalyst in the membrane (mg) and \(Q_v\) is the flux expressed in L s\(^{-1}\).

The conversion rate was thus seen to increase with the mass of catalyst and follows the model for grafted membranes (black squares), expressed by the calculated conversion determined by Eq. (8). The different catalytic membranes behave as a series of plug-flow tubular reactors or similar to a plug flow tubular reactors with the dimension of all grafted membrane reactors. The rate coefficient \(k_{\text{app}}\) was estimated to be 36 s\(^{-1}\). As demonstrated before, an increase of the residence time, between 0.05 and 0.12 s\(^{-1}\) for this set of experiments, led to better conversion yields.

In terms of reaction rate, the second system (with variation of the reactor volume due to the different thicknesses of the grafted polymer) compared to the first system (with the variation of the flux) seems to be slightly better. In fact, we obtained, from theoretical correlation, constants equal to 36 s\(^{-1}\) and 28 s\(^{-1}\) respectively for both systems. Increasing the thickness of the grafted polymer layer could therefore be a good route for increasing the efficiency of catalytic membrane reactors.

4. Conclusions

Catalytically efficient hollow fibers were prepared by photo-grafting polymerization and intermatrix synthesis. The weight of grafted polymer has a clear impact on the final property of the membrane in terms of permeability, nanoparticle loading, catalytic efficiency, and so on. Photografting was found to be a suitable tool to functionalize the membrane surface in order to incorporate MNP acting as catalyst. The greater the functionalization, the higher the MNP incorporation. However, we highlighted the presence of two different types of grafted polymer network: a partially cross-linked polymer and a chemical polymer gel and we correlated the impact of the structure of the polymer network to palladium loading. The nanoparticles created in the grafted layer are in the nanometer size range and are dispersed throughout with no aggregation. They are well trapped in the grafted layer avoiding losses due to the operating flux. The different catalytic tests show good catalytic activity for the reduction of nitrophenol into p-aminophenol for very short residence times (lower than 0.15 s). The residence time in the membrane reactor, directly connected to the conversion, can be controlled by the permeate flow but also by the thickness of the grafted layer. The hollow fiber membrane reactor can be considered as a series of plug flow tubular reactors or as a plug flow tubular reactor with the dimension of all grafted membrane reactors summed. Catalytic membrane synthesis using this versatile technique ensures the availability of various tunable materials for numerous applications, from flow-through catalytic reactors to gas/liquid contacts.

Acknowledgments

We would like to gratefully acknowledge ‘Le Ministère Français de l’Enseignement Supérieur et de la Recherche’ for providing a scholarship.

References


