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Development of an Asymmetric Ultrafiltration Membrane from Naturally Occurring Kaolin Clays: Application for the Cuttlefish Effluents Treatment

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Abstract

This work concerns to the development and characterization of support and ultrafiltration membranes from naturally occurring- kaolin clays as principal components. The preparation and characterization of porous tubular supports, using kaolin powder with corn starch as poreforming agent, were reported. It has been found that the average pore size was about 1 μ m while the pore volume was 44% for supports sintered at 1150°C with a flexural strength of about 15 MPa. The deposition of the active layer was performed by slip casting method. The rheological study of various coatings with different concentration of kaolin powder, polyvinyl alcohol (PVA) and water under different conditions regarding temperature and stirring time was done. After drying at room temperature for 24 h, the membrane was sintered at 650°C. The average pore diameter of the active layer was 11 nm and the thickness was around 9 μ m. The determination of the water permeability shows a value of 78 $l/h.m^2.bar$. This membrane can be used for crossflow ultrafiltration. The application of the cuttlefish effluent treatment shows an important decrease of turbidity, inferior to 1.5 NTU and chemical organic demand (COD), retention rate of about 87%. So, it seems that this membrane is suitable to use for wastewater treatment.

Keywords: Kaolin clays; Sintering; Ceramic supports; Ultrafiltration membrane; Cuttlefish effluent

Introduction

Membrane separation processes extend more and more every day in industrial uses, with new requirements concerning the materials and preparation procedures. Due to their potential application in a wide range of industrial processes such as water and effluent treatments [1-5], drink clarification [6,7], milk pasteurization [8-11], biochemical processing [12], inorganic membrane technology grows in importance.

The use of ceramic membranes has many advantages such as high thermal and chemical stability, pressure resistance, long lifetime, good resistance to fouling, and ease of cleaning [13-17]. The main process to prepare ceramic membranes includes first the obtaining of a good dispersion of small particles and then, the deposition of such dispersion on a support by a slip casting method.

Unfortunately, ceramic membrane fabrication, even though commercially available, still remains highly expensive from a technical and economic point of view due to the use of expensive powders such as alumina [18-22], zirconia, titania and silica [23,24]. To reduce the cost of ceramic membrane fabrication, recent research works are focused on the use of cheaper raw materials such as apatite powder [25], natural raw clay [26-29], graphite [30], phosphates [31,32], dolomite, kaolin [33-37] and waste materials such as fly ash [38-42].

To increase the reliability of the tenacity of these membranes having an asymmetric structure composed of at least two layers which are the support and the active layer, there is still a need for improvement in rheological properties of dip solution required for the preparation of active layer. The control of the rheological parameters of the dip solution is of significant importance for optimizing the final composition of the suspension. Indeed, the rheological properties depend on the physico-chemical characteristics of the raw material used and on the conditions under which the dip solution was prepared. Besides, the active layer thickness and microstructure of the membrane are mainly controlled by the viscosity of the dip solution which depends

mainly on the particle size, the nature of the raw material, the addition of polymer and on temperature [43,44].

The properties of the kaolin suspensions in water are largely modified by the presence of polymers. Many studies reported in the literature, treated as subject the rheological behavior of systems like "water- apatite- additive" and "water-clay-additive" where the additives were often a polymer. Most of these studies were devoted to the colloidal and/or rheological properties of these suspensions and on the effect of polymeric additives in order to obtain the optimal composition of the suspension [45-47].

In order to decrease the membrane cost, the present work describes the preparation of both porous support and active layer based on naturally occurring-kaolin clays (chemical structure: Al₂O₃.2SiO₂.2H₂O [48,49]), to elaborate a new asymmetric ceramic ultrafiltration membrane. This membrane is fabricated in different steps.

The first step is the preparation of porous tubular ceramic support using kaolin clays. Corn starch powder was added as pore-forming agent to produce sufficient porosity with acceptable mechanical property. The properties of porous support were discussed as a function of sintering temperature in order to optimize the preparation conditions. Their structural and functional properties are determined by different techniques. The most important parameters used in the characterization

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of these substrates are: surface and internal morphology, mean pore size, pore size distribution, porosity, mechanical and chemical stability.

The second step is the characterization of the rheological properties of a dip solution which will be used to elaborate an ultrafiltration membrane by deposition using slip casting process of the active layer supported by the kaolin support. The adhesion of the layer to the macroporous ceramic support is achieved by capillary suction. The thickness of the layer depends on the physico-chemical properties of medium, as well as the viscosity of the slip. The study reveals also that the thickness of the active membrane layer can be controlled by the percentage of the kaolin powder added to the suspension and the deposition time.

The last step of this work concerns the application of the ultrafiltration elaborated membrane to the treatment of cuttlefish conditioning wastewater. Figure 1 shows a schematic representation of the sequential steps of the support, active layer, and the procedure of asymmetric ceramic membrane preparation.

Materials and Methods

Materials

In this study, both the supports and membranes were prepared from clay. The clay used in the present study is a kaolin Codex (notes as K), it was recommended by the L.P.M Cerina (Laboratoire des Plantes Medicinales, Tunisia). Corn starch powder was used as a pore former.

Characterization of the kaolin powder

The chemical composition of the kaolin powder was determined by spectroscopic techniques, as Xray fluorescence for metals and by atomic absorption for alkaline earth metals.

Phase identification was performed by XRD analysis (Philips X'Pert X-ray) diffractometer) with Cu K α radiation (λ =1.5406 A°), and the crystalline phases were identified by reference to the International Center for Diffraction Data cards.

Fourier transforms infrared (FTIR) spectra by the KBr method using an IR spectrometer [Perkin-Elmer spectrum BX]. The spectra were collected for each measurement over the spectral range 600-4000 cm $^{-1}$ with a resolution of 4 cm $^{-1}$.

Thermogravimetric analysis (TG) and differential thermal analysis (DTA) were carried out from ambient temperature to 1300°C at a rate of 10°C min⁻¹ under air, using a setaram SETSYS Evolution 1750.

The particle size distributions of kaolin (K) were determined by the Dynamic Laser Scattering (DLS) technique using water as dispersing medium (Mastersizer 2000, Malvern Instruments).

Supports elaboration

The different steps for preparing ceramic materials is the choice of the nature and particle size of the ceramic powder, the choice of organics additions, the optimal volume of water, the aging time paste, shaping by extrusion, and the time of drying and sintering. The preparation of the porous membranes involved the formulation of a plastic and homogeneous paste that contained the clay ceramic powder mixed with organic additive and water.

The optimal formulation of the ceramic precursor paste was carried out in an empirical way. The selected composition of powders used for the plastic paste preparation was 90 wt.% Kaolin (K) and 10 wt.% Starch as a plasticizer. To dos this, a quantity of 540 g of kaolin and 60

g of starch powders were uniformly dry mixed using a rotary mixer at a speed of 65 rpm for 10 min. The powder mixture was aged with a progressive addition of water (300 ml). Water content was adapted according to the type of clay material used in the formulation. As a result, more homogeneity in the final substrate structure was acquired. Before the extrusion phase, an aging stage of the aqueous suspension is necessary to obtain a good homogeneity and to favor the formation of porosities. This step is required to prepare a paste with rheological properties allowing the shaping by extrusion. To this end, the excess of liquid was eliminated and the obtained paste was kept in a closed plastic bag for 24 h under high humidity environment to avoid premature drying and ensure a homogenous distribution of water and organic additives. After aging, the paste was extruded into the tubular specimens trough an extruder and then the wet pieces are set on stems at room temperature during 24 h to ensure a homogenous drying and to avoid twisting and blending.

Finally, a thermal treatment was carried out in a programmable furnace at different final temperatures. The Type 30400 Automatic and Programmable furnaces are general laboratory and heat treating furnaces. Two steps have been determined. Subsequently, the membrane was heated up to 400°C for 2 h at a heating rate of 2°C/min in order to eliminate the organic additive used as pore-forming agent. In a second step, the support was sintered at a temperature

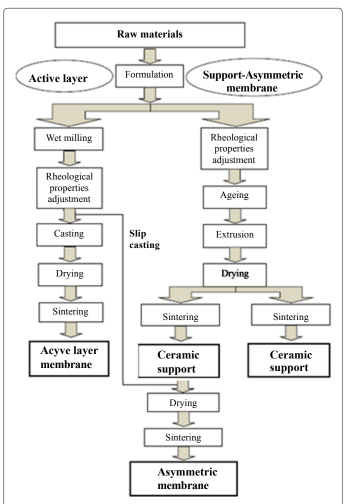


Figure 1: Schematic representation of support, active layer and asymmetric ceramic membrane preparation.

ranging from 1100 to 1250°C with a ramping rate of 5°C min⁻¹ in order to avoid the formation of cracks during sintering of the samples. The final temperature was kept constant for 1 h. After that, the consolidated support was allowed to cool until the environmental temperature was reached.

The temperature-time schedule not only affects the pore diameters and porous volume of the final product but also allows obtaining the final morphology and mechanical strength. Tubular supports were elaborated with external/internal diameter of $16/11~\mathrm{mm}$ and the length of $150~\mathrm{mm}$.

Supports characterization

The evolution of densification and surface quality of the supports sintered at different temperatures were determined by scanning electron microscopy.

Porosity and pore size distribution were measured by mercury porosimetry. This technique relies on the penetration of mercury into a membrane's pores under pressure [34-36,50,51]. The intrusion volume is recorded as a function of the applied pressure and then the pore size was determined.

The mechanical resistance tests were performed using the three points bending method (LLOYD Instrument) to control the resistance of the membranes fired at different temperatures.

The corrosion tests were carried out using aqueous solutions of nitric acid (pH=2.5) and sodium hydroxide (pH=12.5) at 45 and 80°C, respectively. All the samples were ultrasonically rinsed in distilled water, dried at 110°C and stored in a dryer. The degree of corrosion was characterized by the percentage of the weight loss.

Ultrafiltration membrane elaboration

To decrease the probability of cracking or fissures during the sintering process of the membrane, in the preparation of the active layer the same raw materials of the support were used.

The active ultrafiltration layer from kaolin was prepared by a slip casting process on kaolin support from a mixture of the inorganic powder and a 12 wt.% aqueous solution of polyvinylique alcohol (PVA) (Rhodoviol 25/140 (Prolabo)), which is used as binder and plasticizer. The slip casting process [42,52] was applied to coat the support tube to elaborate ultrafiltration membrane as described in Figure 1.

In the first step, a stable suspension or slip has to be obtained by putting in suspension the mineral powder in water. Then, 12 wt.% aqueous solution of PVA was mixed to the preceding suspension. Desagglomeration of the mineral powder and homogenization of the coating formulation was ensured by mechanical stirring using an ordinary magnetic stirrer at its maximum speed. The coating formulation was poured inside the support for a few minutes at room temperature while the tube was closed at one end. A layer is then formed on the inner side of the porous tube due to capillary suction. Afterwards, the excess was drained out. Drying was realized at room temperature for 24 hours. The sintering temperature was fixed at 650°C for 3 hours. A heating temperature at 250°C for 2 hours is necessary in order to eliminate completely the PVA, present in great quantity in the slip [3]. Relatively slow temperature rate (2°C/min) was needed in order to avoid the formation of cracks on the layer.

Composition and characterization of the slip

Different coating formulations B.n differing by PVA concentration

(1<n<3) and kaolin powder (4<n<8) were used to determine the optimal composition of the final layer.

At the first time, the kaolin powder quantity was kept at 4% when the PVA aqueous solution percentage was varied and then, at the second time, the PVA aqueous solution percentage was fixed to 40% when the kaolin powder percentage was varied (Tables 1 and 2).

The viscosity is an important parameter to determine the slip optimal composition. It was measured just before slip casting by using an Antoon Paar rheometer model MCR301.

Filtration pilot

The laboratory pilot used for the filtration experiments was equipped with a cross-flow filtration system implementing tubular ceramic membrane of 15 cm length. The tubular membrane takes place in a stainless steel carter. The transmembrane pressure (TMP) can reach 8 bars wich corresponds to the ultrafiltration range. It was controlled by an adjustable valve on the retentate side. Temperature was kept at 25°C by a thermal exchange system. The membrane was conditioned by immersion in pure deionized water for a minimum of 24 h before filtration tests. The determination of the water membrane permeability was performed with distilled water.

Effluent characterization

Wastewater samples were taken from the effluents produced by a sea-product freezing factory located in Sfax, Tunisia. The dark colour in this effluent was due to the presence of sepia ink (containing melanin) as suspension particles which have a size range between 56 and 161 nm [31].

A large number of analyses were conducted on each sample and the following parameters were measured: turbidity, Chemical Oxygen Demand (COD), temperature, pH and conductivity. The COD values of raw effluents from the production process ranged between 2000 and 3000 mgL $^{\rm -1}$ with an average concentration of 2615 mgL $^{\rm -1}$. The turbidity measured for the raw effluent presents a very high value which is in the range of 335 NTU (Table 3).

The techniques used to analyze collected samples of feed, retentate and permeate are reported below:

- Turbidity: Using a HACH «2100 N Turbidimeter» turbidimeter.

Coating formulation	% of PVA ^a	% of water
B.1	20	76
B.2	30	66
B.3	40	56

Table 1: Composition of coating formulation prepared with different percentage of PVA.

solution added to the powder-water dispertion.

Coating formulation	% of kaolin	% of water/PVA aqueous solutions ^b
B.4	2	94
B.5	4	92
B.6	6	90
B.7	8	88
B.8	10	86
b: mixture of 12 wt % PVA	aqueous solution and wate	r

Table 2: Composition of coating formulation prepared with varying percentage of kaolin.

Turbidity (NTU)	COD (mg.L ⁻¹)	Conductivity (mS.cm ⁻¹)	рН
335	2615	204	7

Table 3: Turbidity, COD, conductivity and pH of raw effluents.

- Dissolved organic carbon: Using a «REHROTEST TRS 200 NF T 90-101» COD Analyser.
 - Conductivity: Using a «Consort K 911» conductimeter.

Results and Discussion

Powder characterization

The chemical composition of kaolin is given in Table 4, where the main impurities are CaO, K2O, TiO_2 and Fe_2O_3 . It reveals that the major components were silica (SiO₂: 47.85%) and aluminium oxide (Al₂O₃: 37.60%).

Phase identification is of great importance before any membrane manufacturing. For example, Figure 2 presents the XRD patterns of the raw kaolin. It can be seen that kaolinite (K) was the major mineral component with a small amount of quartz (Q) and illite (I) impurities. No other components were observed, because the impurities are in so tiny quantity (Table 4) and most of them are probably incorporated into the crystal structure of kaolinite.

The results of FTIR spectroscopy of the starting clay (Figure 3) show the kaolin characteristic bands: Si-O (at around 993, 1024, and 1112 cm⁻¹), Si-O-Al (at around 525, 750, and 795 cm⁻¹), Al-OH (at around 910 cm⁻¹) and OH (at around 3684, 3668, 3651, and 3618 cm⁻¹) [48,49,53].

The particle size distribution of kaolin was determined by the Dynamic Laser Scattering (DLS) technique, and the results shown in Figure 4. This method gave an average particle size in the order of 4 μm .

A total weight loss is observed by TGA to be about 15% of kaolin (Figure 5). In fact, the weight loss consists of two distinct stages: The first one is considered as a slight weight loss between room temperature and 150°C, because of the dehydration of the clay. The second mass loss detected between about 400 and 700°C is mainly due to the phenomenon of dehydroxylation of kaolinite confirmed by DTA which shows an endothermic peak at 560°C leading to the transformation of kaolinite to metakaolinite. A third stage, wich is characterized by an exothermic reaction appeared at about 975°C without any weight loss. The exothermic peak corresponds to the metakaolin-mullite transformation [54].

Supports characterization

For the development of high-quality supports, the following properties are of major importance: pore size distribution, porosity, surface texture, mechanical properties and chemical stability.

Supports morphology: Figure 6 illustrates SEM pictures for the supports sintered at the four different temperatures considered in this work. The optimal sintering temperature was determined by comparing the texture of the different obtained samples.

The ceramic substrates sintered at lower temperature (1100°C) show highly porous structure. Below 1100°C, we detect the presence of intergranular contacts which are large enough to ensure the ceramics cohesion (beginning of sintering). We observe that the formation of grain boundaries is achieved within a narrow temperature range of 1150°C-1200°C. The aspect of the surface is homogeneous and does

not present any macro defect (cracks, etc.). Beyond 1250°C, the densification process (shrinkage) dominates and a relatively open structure is still observed.

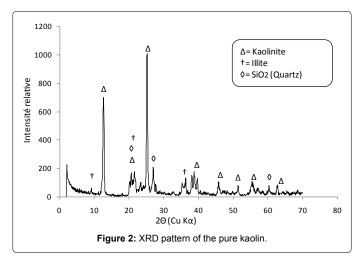
The morphology as well as the microstructure of optimal support fired at 1150°C is shown in Figure 6 for the cross section and the surface views. A smooth inner surface is observed which will allow the deposit of a homogeneous membrane layer.

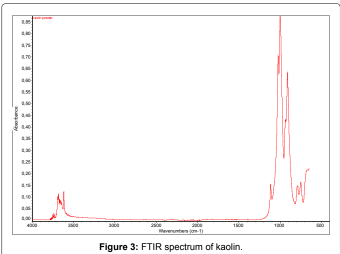
Open porosity and pore size distribution for supports: The evolution of the average pore diameter and the porosity with the temperature of sintering reveals that the porosity decreases from 49 to 35% between 1100°C and 1250°C, while the pore diameter increased from 0.81 to 1.4 μm (Figure 7). The beginning of the material densification occurred when the temperature increases [25,33,36,37,41,55].

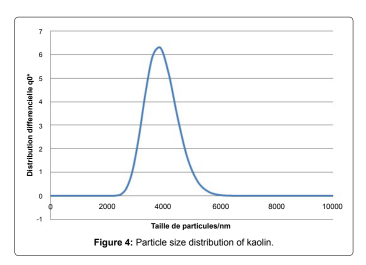
At 1150° C, the characteristics of the support are an average pore diameter of 1 μm and a porosity of 44%. Consider Figure 8 which presents a Single (mono) Modal of pore size distribution or

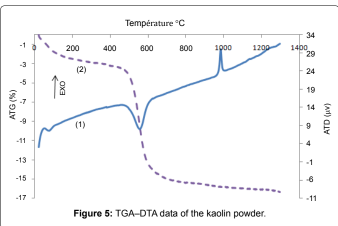
	SiO ₂	Al ₂ O ₃	Fe ₂ O ₃	MgO	K ₂ O	CaO	TiO ₂	LOI*
Kaolin (%)	47.85	37.60	0.83	0.17	0.97	0.57	0.74	11.27
* LOI: Loss on Ignition at 1000°								

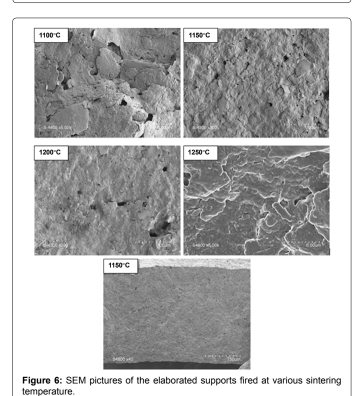
Table 4: Chemical composition of the used kaolin (wt %).











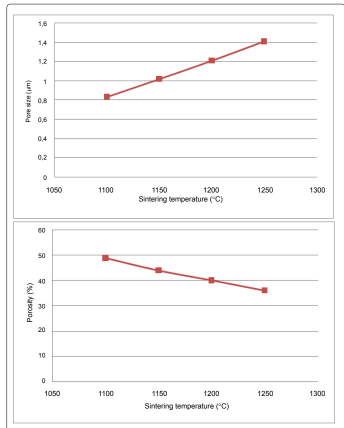
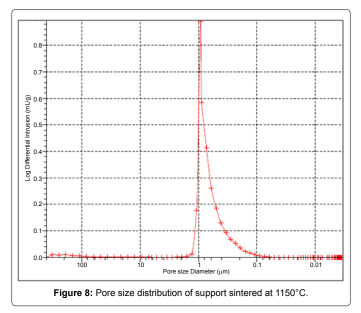
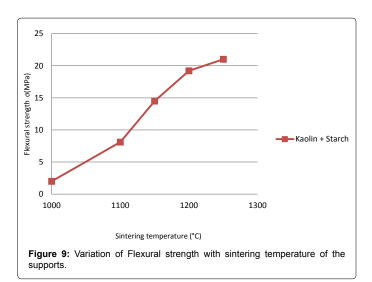


Figure 7: Effect of sintering temperature on average pore size and porosity of supports.



homogenous pore distribution. This is necessary for a good integrity of the membrane.

Mechanical resistance: Figure 9 shows the variation of flexural strength with sintering. In accordance with the SEM pictures and the porosity values, the increase of the sintering temperature is accompanied with a densification phenomenon and consequently an



increase in the tensile strength from 3 MPa at 1000°C to 21 MPa at 1250°C.

Chemical corrosion: The weight loss due to the corrosion of acids and alkali is shown in Figure 10. It can be seen that the membrane shows a better acid corrosion resistance, since its mass loss is much lower than those of supports after alkali corrosion. Therefore, the observed results in weight loss during corrosion tests suggest that the prepared support possesses a good chemical corrosion resistance and it is suitable for applications involving acidic and basic media.

Choice of the support: Finally the best condition to prepare the support are established for a firing temperature of 1150°C, at this condition the average pore diameter is 1 μm and the porosity is 44%. Moreover this support presents good mechanical resistance having a tensile strength of 15 MPa, and also good chemical properties towards the acid and basic solutions. Thus, membrane support sintered at 1150°C for 1 hour has been selected as substrates for filtration tests.

Rheological characterization of the suspension

For a good adhesion on the macroporous support, viscosity must be sufficient to facilitate the coating and to prevent that the support does not absorb solvent too quickly. Many parameters were studied in order to optimise the slip composition (percentage mass in PVA, percentage mass in kaolin, temperature, etc.).

Influence of temperature: The influence of the temperature on the suspension viscosity was studied using different concentrations of PVA (20%, 30% and 40%) and kaolin (2%-10%). The mineral study used for coating was tested at varying temperatures ranging from 25 to 80°C. As expected, a rise of temperature leads to a decrease of the suspension viscosity (Figures 11 and 12) according to an exponential relationship whose expression is given by Dufauda [56]:

$$\eta = A.e^{Ea/Rt}$$

Where A is the pre-exponential factor, Ea is the activation energy for viscous flow, R is the gas constant (8.314 J mol^{-1} K⁻¹), and T is the absolute temperature. The logarithm of the viscosity plotted against the inverse of temperature bits well aspect Arrhenius relation.

Figures 11a and 12a show that the viscosity increases considerably with increasing the concentration of kaolin powder and PVA but

decreases with the increase of the temperature in the range 25-80°C. The influence of temperature on the composition and viscosity can be described using Arrhenius Law. The determination of the activation energy Ea shows a pronounced sensitivity of the viscosity to temperature changes [57,58].

Figures 11b and 12b show an evolution accorded to Arrhenius law for the different prepared suspensions with an increase of the logarithmic viscosity with the percentage of loading of PVA and kaolin in the temperature range between 20°C and 70°C. A straight line was obtained using Arrhenius model which fitted reasonably well with the experimental data and gave high R2 equal to 0.99 with 99% confidence level.

Table 5 lists the corresponding flow activation energy ΔEa and the

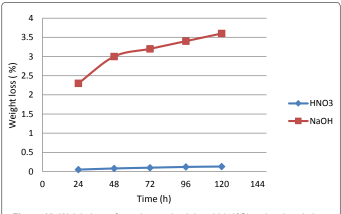


Figure 10: Weight loss of membranes in nitric acid (45°C) and soda solutions (80°C) as a function of time.

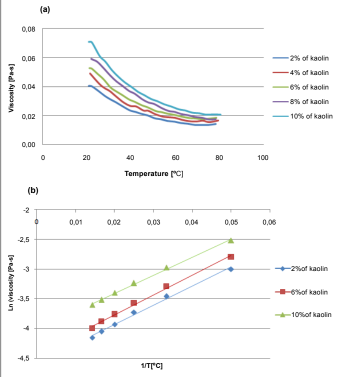


Figure 11: Evolution of viscosity with the temperature for different kaolin concentrations.

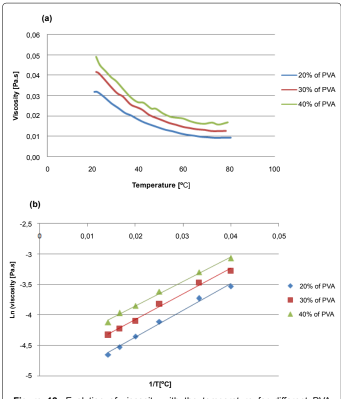


Figure 12: Evolution of viscosity with the temperature for different PVA concentrations.

related fit stability index R2 for different suspensions. With increasing concentration of kaolin powder and PVA, a pronounced increase of the flow activation energy can be observed. Such results are in agreement with the values given in the literature [57,58] and are due to the increasing interaction of the particles with the polymer.

In slip casting process, capillary suction is the driving force that causes the layer deposition on the inner side of the support. Then, to obtain a good and homogenous layer, the suspension does not have to be very viscous nor fluid. The effect of PVA and mineral powder percentages on the viscosity of the suspension is due to the fact that increasing concentration would have a direct effect on the fluid internal shear stress while the temperature effect is obviously due to a weakening temperature of inter-particle and inter-molecular adhesion forces [59].

The PVA and the mineral powder percentage, have a direct influence on the suspension viscosity. This influence is more important for low temperatures. A temperature of 25°C to 35°C will be retained since the layer deposition process is technically possible only at low temperatures. The PVA percentage has the stronger effect on the viscosity, so, it is of interest to study its effect on the rheological behavior of the suspension.

Influence of PVA percentage: Figure 13 shows that the viscosity of the suspension increases slightly depending on the stirring period. However, the increase of PVA percentage leads to a considerable increase of the viscosity. Such rheological behavior is explained by the different types of interactions susceptible to be established between PVA used as Binder and kaolin particles which depend on the concentration of both of them [57]. In conclusion, a stirring time of 90 minutes is a medium duration that can be used when preparing the suspension to be used.

From Figure 14, it can be noticed that dynamic viscosity decreases slightly according to the speed of shearing, but this variation remains insufficient for the determination of the nature of the behavior of such suspension. For identifying the rheological behavior, the variation of the shear stress (τ) according to the speed of shearing (D) was determined

Figure 15 shows that the behavior of dip solution (B.1, B.2 and B.3) is of Newtonian type. The shear rate increases according to the speed of shearing D. These samples show a Newtonian behavior generated by an increase in inter-particle interactions due to increasing the concentration of PVA and formation of complex flocculated structures. Suspension with a PVA percentage comprised between 20 and 40 % can be apparently used to deposit the layer.

Finally, to obtain a suspension with a viscosity of 0.042 Pa.s and having a Newtonian behavior at room temperature of 25 to 30°C, the suspension should have a composition of 4% of kaolin and 40% of PVA and stirred for 90 minutes.

	PVA ^b
17.50	18.64
0.992	0.995
	0.992

Table 5: Estimation of the flow activation energy Ea for kaolin-PVA.

b: Kaolin powder % was fixed at 4%

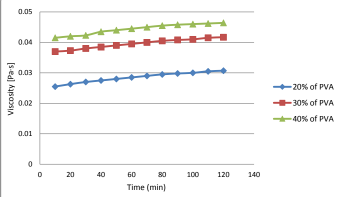


Figure 13: Evolution of the viscosity with the time of stirring for different PVA concentrations.

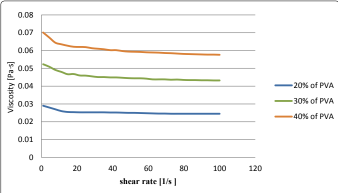
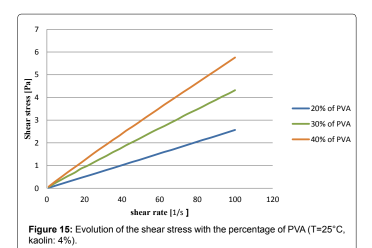


Figure 14: Evolution of the viscosity with the percentage of PVA (T=25°C and 4% of kaolin).



Membrane characterization

Scanning electron microscopy (SEM): Influence of PVA percentage: Three layers were cast on clay tubular supports using three different suspension compositions (B.1, B.2 and B.3). The same casting time was used during the slip casting operation (6 min). The sintering conditions, previously mentioned, were respected.

Figure 16, which shows SEM pictures for surface and cross-section of the obtained layers, give information about the thickness and texture. According to the SEM analysis, it can be noticed at first, that there is a good adhesion between the layer and the clay support. No detachment is observed on the different cross section photographs.

The optimal composition for coating suspension is obtained by using 40% PVA (B.3). This composition enables us to achieve a thickness of 8 μm for ultrafiltration and homogeneous surface free from defects (such as cracks, etc.). On the other hand, the composition corresponding to 20% of PVA (B.1) enables us to obtain a thickness of 3 μm . So, B.3 composition was retained for the preparation of an ultrafiltration membrane.

Influence of deposition time: The deposition of the slip B.3 on the clay support was performed by slip casting using a deposition time between 2 and 20 min. Different ultrafiltration membranes with different layers thickness (between 3 and 30 $\mu m)$ were prepared. SEM (scanning electron microscopy) images of the resulting membranes are shown in Figure 16.

B.3 composition was retained for the preparation of an ultrafiltration membrane. The casting time was fixed to 6 min.

Mercury porosimetry: The average pore size of the optimized membrane was determined by mercury porosimetry. The pore diameter measured was centered near 11 nm (Figure 17), which confirms that an ultrafiltration layer was achieved.

Determination of membrane permeability: The membrane was initially characterized by the determination of water permeability. It can be seen that the pure water flux increases linearly with increasing the applied pressure (Figure 18). The membrane permeability was found to be equal to 78 l/h.m².bar.

Application to the treatment of the cuttlefish effluents

The elaborated ultrafiltration membranes have been applied to the cuttlefish effluents treatment. Figure 19 gives the variation of permeate

flux with transmembrane pressure. Permeate flux increased linearly with transmembrane pressure until 6 bar and then became pressure independent. This behavior can be explained by the formation of a concentrated polarization layer. Beyond 6 bar, the flux value is about 64 l/h.m².

The variation of the permeate flux with time at different transmembrane pressures (TMP) from 2 to 6 bar and a temperature of 25°C, as shown in Figure 20. Permeate flux decreases with the time and stabilizes after the first 20 min whatever the pressure value. This behavior can be attributed to the fouling due to the interaction between membrane material and the waste water solution. It is important to notice that fouling is not very important since the flux decline did not exceed 15% for the different pressures used. This is a specific behavior of

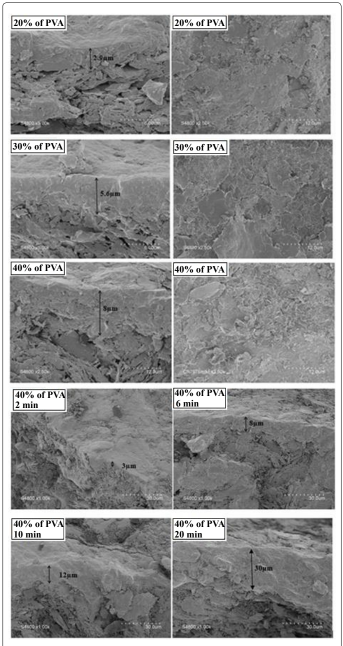


Figure 16: SEM micrographs (cross section and surface) of active layer obtained with different slip compositions and deposition time.

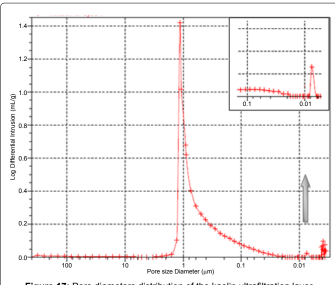
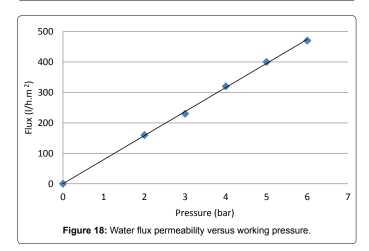
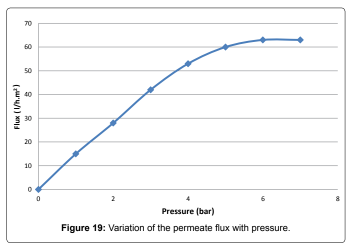


Figure 17: Pore diameters distribution of the kaolin ultrafiltration layer.



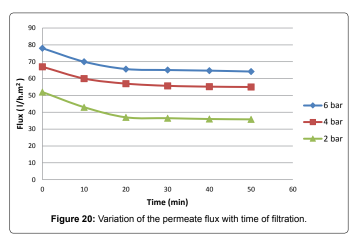


the UF membrane compared to MF membrane that showed generally an important decrease of flux due to pores blocking by retained particles [3,42].

Ultrafiltration performances: Table 6 gives the main physicochemical parameters analyzed for permeate obtained by UF kaolin membrane. These analyses show variability in the turbidity and COD values. The turbidity of the ultrafiltrated effluent was 0.62, 0.95 and 1.36 NTU respectively for 2, 4 and 6 bar. The values of COD (from 350 to 530 mg L⁻¹ with TMP from 2 to 6 bar). The conductivity values were usually in the range of 140-148 ms.cm⁻¹.

Percentage reduction of turbidity and COD as a function of TMP has been shown in Figure 21. Both the turbidity and COD reduction have been found to increase with decrease in TMP, which could be attributed due to the higher rejection at lower TMP. As pressure increases, more melanin permeates through the membrane leaving most of solutes to through the pores of the membrane by increasing transmembrane pressure and subsequently decreased rejection.

Turbidity of permeate was found to get reduced by 99.8% when UF was carried out at a TMP of 2bar, whereas, COD was reduced by only 87%. However, the conductivity in the permeate decreases slightly and is pressure independent. In fact, at all the TMP level, turbidity



	Pressure (bar)	Turbidity (NTU)	COD (mg.L ⁻¹)	Conductivity (mS.cm ⁻¹)
Raw effluents		335	2615	204
	2	0,62	350	140
Filtrate	4	0,95	460	146
	6	1,36	530	148

Table 6: Characteristics of the effluent before and after filtration on kaolin membranes

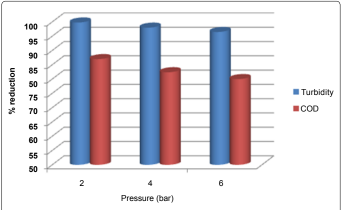


Figure 21: Effect of transmembrane pressures in ultrafiltration on turbidity and COD reduction.

reductions were found to be more than the corresponding COD reduction on percentage basis. As the COD is caused by the presence of low molecular inorganic chemicals also, which might pass through the membrane, may give less (%COD).

Evaluations of resistances: The volume flux in a pressure driven membrane process depends on the hydraulic resistance of the used membrane and the pressure drop over the membrane. This is generally expressed by the following formula:

$$J_{\omega} = \Delta P / \eta_{\omega} R_{\omega} J_{\omega} \tag{1}$$

where, $R_{\rm m}$ is the intrinsic hydraulic resistance of the membrane and $\eta_{\rm w}$ is the viscosity of water.

The permeate flux (J_s) after filtration of the solution can be expressed by the resistance-in-series model.

$$J_{s} = \Delta P / \eta_{s} R_{t} J_{s} \tag{2}$$

in which η_s is the permeate viscosity and \boldsymbol{R}_t is the total resistance that can be defined as:

$$R_t = R_m + R_c \tag{3}$$

where, R_c is the total resistance fouling.

Experimentally, the intrinsic membrane resistance R_m was calculated by measuring the pure water flux J_w and viscosity ηw .

Total resistance R_t was estimated from the solution flow rates under operating conditions using Equation (2):

$$R_{s} = \Delta P / \eta_{s} J_{s} \tag{4}$$

The intrinsic membrane resistance and the total fouling resistance deduced from these Equations for kaolin membrane are presented in Table 7. The major part of the total resistance was due to fouling $((R_p + R_p)/R_p)$ for different working pressure. Intrinsic membrane resistance (R_m/R_p) represented a small portion of the total resistance.

Membrane regeneration

After each experiment, the membrane must be regenerated. The efficiency of the using protocol is verified by the measurement of water flux. The regeneration of the membrane was carried out by firstly, the membrane was exposed to an acid-basic washing sequence (Table 8), and secondly, by leaving the membrane in distilled water. The used protocol appears sufficient because we obtained the value of the initial permeability of the membranes (Figure 22).

Conclusion

In this work, a comprehensive study on the fabrication and characterization of an asymmetric ultrafiltration membrane from naturally occurring- kaolin clays were performed. Ceramic supports have been obtained by extrusion using kaolin powder with corn starch as poreforming agent. It has been concluded that the supports sintered at 1150°C is the optimum supports allowed for depositing the membrane layers. The mechanical and structural properties of the supports are satisfying in terms of porosity and pore diameter.

Pressure (bar)	R _t (10 ¹²) (m ⁻¹)	R _m (10 ¹²) (m ⁻¹)	R _c (10 ¹²) (m ⁻¹)	R _m /R _t (%)	R _c /Rt (%)
2	20.11	4.61	15.5	22.92	77.08
4	26.18	4.61	21.57	17.6	82.4
6	33.75	4.61	29.14	13.66	86.34

 Table 7: Resistance values for kaolin membrane.

Sequence	Agent	Concentration	T (°C)	Pressure (Bar)	Time (min)
Rinsing then evacuation	water		RT	2	
Basic washing	NaOH	5-10 g L ⁻¹	60-65	2	30
Rinsing until neutrality	water		RT	2	
Acidic washing	HNO₃	5 ml L ⁻¹	40-50	2	30
Rinsing until neutrality	water		RT	2	
*RT: Room Temperature					

Table 8: Acido-basic washing sequence.

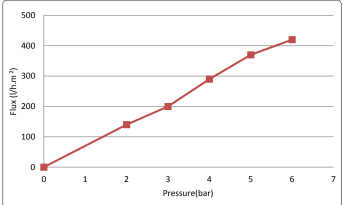


Figure 22: Variation of the permeate flux with pressure for Kaolin membrane after regeneration.

The UF layer, deposited on the supports, was obtained by the slip-casting technique. The composition of the slip was optimized using a rheological study. An excellent link between the support and the ultrafiltration layer was obtained. The thickness of the ultrafiltration layer is about 9 μm ; it can be controlled by the percentage of the PVA powder added in the slip suspension and the duration of the deposition time. The obtained membrane has an average pore size diameter of about 11 nm and a water permeability of 78 l/h.m².bar. This result may allow the novel membrane to be used in water treatment in the ultrafiltration range.

The application of this membrane to the washing cuttlefish effluent treatment exhibited an acceptable stabilized permeate flux of almost 64 l/h.m² at 2 bar and 25°C with a very important decrease of different pollutants in terms of turbidity (99%) and COD (87%). These membranes can also be used as a support for nanofiltration layer.

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