Superhydrophobic Hexamethylene Diisocyanate Modified Hydrolyzed Polymers of Intrinsic Microporosity Electrospun Ultrafine Fibrous Membrane for the Adsorption of Organic Compounds and Oil/Water Separation

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ABSTRACT: Polymers of intrinsic microporosity (PIMs) have gained significant research interest because of their successful applications in adsorption and separation. PIM-1 is the first and most studied member of this class because it shows specific interactions with some certain organic species. Chemical modification of PIM-1, which can be achieved by simply hydrolyzing the nitrile groups in the backbone, provides an advantage of tailoring its adsorption and separation performances. In this study, electrospinning of ultrafine fibers from hydrolyzed polymer of intrinsic microporosity (HPIM) and blends of hexamethylene diisocyanate (HMDI)/HPIM was achieved in several different ratios of HMDI/HPIM ranging from 1:9 to 1:1 (w/w). Bead-free and uniform fibers were obtained in the form of self-standing ultrafine fibrous membranes, which were then thermally treated at 150 °C to introduce chemical cross-linking between HMDI units and carbonyl groups of HPIM, resulting in HMDI-modified HPIM fibrous membranes (HMDI/HPIM-FMs). The solubility behavior has been altered by an introduced modification that makes membranes insoluble in all common organic solvents. Chemical cross-linking has been confirmed by using a Fourier transform infrared technique showing urethane linkage between HMDI and HPIM, and it was further supported by X-ray photoelectron microscopy and elemental analysis techniques that show a significant increase in the relative ratio of nitrogen in HMDI/HPIM-FMs compared to HPIM-FM. The average fiber diameters of fibrous membranes were found between 1.38 ± 0.29 and 0.96 ± 0.22 μm depending on the blend compositions and applied electrospinning parameters. Moreover, the water contact-angle value for HPIM-FM increased with the introduced HMDI modification from 140 ± 4° to 159 ± 7°, changing the nature of the membrane from hydrophobic to superhydrophobic. Consequently, HMDI/HPIM-FMs were successfully employed in oil/water separation due to the superhydrophobicity. In addition, the adsorption properties of HPIM-FM and HMDI/HPIM-FMs were explored for common organic solvents. While both HPIM-FM and HMDI/HPIM-FMs show promising results, the structural stability of HMDI/HPIM-FMs in liquids was found to be more stable and reusable with respect to HPIM-FM. Hence, HMDI/HPIM-FMs are more favorable for organic adsorption and separation purposes from an aqueous system.

KEYWORDS: polymers of intrinsic microporosity (PIM-1), electrospinning, superhydrophobic, organic removal, oil/water separation

INTRODUCTION

The issue of water pollution has gained tremendous attention in recent years because it has become a worldwide challenge. Water pollution is mainly caused by industrial actions and oil spill accidents that lead to serious environmental damage. Therefore, oil spill cleanup is an urgent task, and the need for producing cost-effective and environmentally friendly methods and materials is an ongoing demand. Several methods have been introduced for oil spill cleanup including in situ burning of oil on water, chemical degradation, mechanical extraction, and centrifugation. Each has its own limitations and most have low separation efficiency and high operation cost and require complex equipment. Among these cleanup strategies, the most promising technique is considered to be the use of sorbent materials that can selectively separate oil/water mixtures. One of the most required properties for ideal sorbents is maintaining special wettable surfaces, such as superhydrophobic/superhydrophilic surfaces. Superhydrophobic surfaces display water contact angles of greater than 150° and can be prepared by controlling the chemical composition and topological...
structures. Until now, several methods have been employed to obtain superhydrophobic surfaces such as colloidal assembly, layer-by-layer deposition, chemical deposition, template-based techniques, and electrospinning. Electrospinning is one of the most peaked techniques for the fabrication of superhydrophobic polymeric or composite nanofibers. Thus, a number of electrospun materials have been reported for oil/water separation.

In the past few years, great emphasis has been put on the research of the synthesis and applications of microporous materials that were obtained from purely organic components because of their outstanding performances in adsorption and separation. Polymers of intrinsic microporosity (PIMs) have been developed based on this approach. PIMs possess rigid and contorted structures that limit the packing efficiency. Therefore, they have high specific surface area and high chemical and thermal stabilities, which make them suitable for several applications.

PIM-1 is the first and most studied member of the PIM family because it can be synthesized by using commercially available 5,5′,6,6′-tetrahydroxy-3,3,3′,3′-tetrakis(1,1′-spirobisindane and tetrafluoroterephthalonitrile monomers. PIM-1 is soluble in common organic solvents such as chloroform, tetrahydrofuran, and dichloromethane, and thus it can be prepared in the form of powder, membrane, and fiber, showing excellent performance in adsorption and separation applications. Recently, electrospun PIM-1 was introduced, and it has been utilized successfully for adsorption and separation purposes. PIM-1 possesses nitrile units in the backbone, which enables possible post-modification reactions to alter the properties of PIM-1 for the desired applications. Therefore, some efforts have been devoted to the modification of PIM-1. Hydrolysis is the most studied modification among all of them because of the simple reaction conditions. From this standpoint, Zhang et al. studied the post-treatment of an electrospun PIM-1 fiber with a base to obtain a hydrolyzed PIM-1 (HPIM) fiber. The study revealed that the dye removal ability of electrospun PIM-1 toward cationic dyes in aqueous media has been improved by hydrolysis. However, the hydrolysis procedure has caused severe damage to the mechanical and structural integrity of fibers. Recently, we introduced a new strategy to fabricate self-standing, electrospun HPIM ultrafine fibrous membranes as a molecular filter to remove methylene blue from an aqueous solution, showing a high removal efficiency.

In this study, we reported a facile approach to the further modification of HPIM with 1,6-hexamethylene disocyanate (HMDI) in order to obtain superhydrophobic ultrafine fibers using an electrospinning technique. Chemically cross-linked, self-standing, HMDI-modified HPIM fibrous membranes (HMDI/HPIM-FMs) showed improved hydrophobicity with increased water contact angle. After modification, HMDI/HPIM-FMs became insoluble, which enhanced the structural stability of membranes against several organic compounds. Thanks to their superhydrophobicity and insolubility, HMDI/HPIM-FMs were successfully employed in oil/water separation as well as organic removal applications.

**Experimental Section**

**Materials.** 5,5′,6,6′-Tetrahydroxy-3,3,3′,3′-tetrakis(1,1′-spirobisindane (98% (Alfa Aesar) and tetrafluoroterephthalonitrile (98%, Aldrich) were recrystallized as reported previously. Anhydrous potassium carbonate (99.0%, Fisher) was dried in an oven at 110 °C overnight. 1,6-Hexamethylene disocyanate (HMDI, >98%, Aldrich), dimethylformamide (DMF), ethanol (EtOH), dimethylacetamide, methanol (MeOH), toluene, sodium hydroxide, tetrahydrofuran (THF), dimethyl sulfoxide (DMSO), dichloromethane (DCM), hexane, chloroform (CHCl₃), sunflower oil (Yudum, obtained from a local shop), pump oil (Edwards 19, ultragrade vacuum oil), and silicone oil were used as received. The synthesis and hydrolysis of PIM-1 was conducted as reported previously.

**Electrospinning.** For electrospinning of HPIM, HPIM powder (0.35 g) was dissolved in DMF (0.5 mL) to obtain 70% (w/v) polymer concentration, and the HPIM solution was stirred at room temperature for 4 h and then degassed for 15 min. After that, the solution was loaded into a 1 mL syringe fitted with a blunt needle (0.5 mm diameter) and placed horizontally on a syringe pump (KDS101, KD Scientific). Microfibers were collected on aluminum foil and then ground by a metal collector. Electrospinning was performed using a commercial spinner (Nanoweb, Mavi Teknik, Mersin/Turkey). The applied parameters were as follows: flow rate, 0.5 mL h⁻¹; voltage, 11 kV; distance, 16 cm. Then the collected fibers were removed from the aluminum surface by tweezers. Eventually, HPIM fiber was dried in an oven at 150 °C for 2 h. HMDI/HPIM blend solutions were prepared in several different ratios ranging from 1:9 (w/w) (H10) to 1:1 (w/w) (H50) while maintaining the same amount of HPIM (0.35 g in 0.5 mL of DMF; 70% (w/v)). Similarly, a HPIM solution was first prepared, and predetermined amounts of HMDIs were added to this and stirred at room temperature for an additional 1 h. The solution was degassed for 15 min prior to the electrospinning process. The electrospinning parameters of blend mixtures were identical with those of HPIM, and cross-linking of the electrospun HMDI/HPIM-FMs was carried out at 150 °C for 2 h in an oven.

Additionally, H50 has been prepared in the dense membrane (film) form using the same HPIM/HMDI compositions in 2 mL of DMF. The solution was placed in a glass Petri dish and heated to 60 °C for 3 h. Then it was dried at 150 °C in an oven for 2 h.

**Methods.** A Bruker Alpha P spectrometer was used to collect attenuated total reflectance infrared (ATR-IR) spectra of samples that were scanned 64 times at a resolution of 4 cm⁻¹. ¹H NMR spectra of the samples were recorded using a Bruker DPX-400 MHz spectrometer. Samples were dissolved in DMSO-d₆. X-ray photoelectron spectroscopy (XPS) spectra were obtained by a flood gun charge neutralizer system equipped with a monochromated Al Kα X-ray source. Elemental analysis was carried out using a Thermo Scientific Flash 2000 series CHNS-O analyzer. 2,5-[Bis(5-tert-butyl-2-benzoxazol-2-yl)]thiophene and vanadium pentoxide were used as a standard and a catalyst, respectively. The morphology of the samples was studied by scanning electron microscopy (SEM; FEI Quanta 200 FESEM). The samples were coated with 5 nm of gold/palladium prior to imaging. The average fiber diameters were determined using ImageJ software. Thermogravimetric analysis (TGA; Q500, TA Instruments) was utilized to explore the thermal resistance of samples that were heated to 600 °C at a heating rate of 20 °C min⁻¹ under N₂. The thermal properties of the samples were also investigated using differential scanning calorimetry (DSC; Q2000, TA Instruments). The samples were heated to 500 °C at 20 °C min⁻¹ under a N₂ atmosphere. For DSC measurements, the samples were equilibrated at 0 °C and then heated to 500 °C at 20 °C min⁻¹ under a N₂ atmosphere. Water contact-angle data of the samples were obtained by a Dataphysics water contact-angle system (OCA) using deionized water (0.4 μL), which was automatically dropped onto the nanofibrous mats, and a Laplace—Young fitting was applied to the measurements. The measurements were repeated 10 times at different positions for an average value. The mechanical properties of the samples were examined using dynamic mechanical analysis (DMA; Q800, TA Instruments). The thicknesses of the samples were measured using a Forma ProMax digital caliper. Rectangular-shaped samples (8 × 6 × 0.2 mm) were employed to obtain stress–strain curves in controlled force mode with a 0.05 N min⁻¹ force ramp rate by three different samples at room temperature. Young’s modulus was obtained from the linear region of the curves. Similarly, the storage modulus was tested for rectangular-shaped samples (15 × 6 × 0.02 mm) in multifrequency mode with an amplitude value of 5 μm. Samples were heated to 200 °C.
Nitrogen adsorption/desorption isotherms of the samples were obtained at $-196^\circ$C using an Autosorb iQ gas sorption analyzer. The samples ($\sim 0.05$ g) were degassed under high vacuum at $120^\circ$C for 16 h before analysis. A multipoint analysis method was applied to calculate the Brunauer–Emmett–Teller (BET) surface areas.

Adsorption of Organic Compounds and Oils. For performance evaluation, fibrous membranes ($1 \times 1$ cm) were immersed in organic liquids for an hour and then taken out for weight measurements. The membranes' weights before and after adsorption were recorded to estimate the adsorption capacity. The weight measurements were performed immediately after the membranes were taken out of the liquid to avoid any evaporation effect. The percent weight gain was calculated based on eq 1

$$\% \text{ weight gain} = \frac{m_1 - m_0}{m_0} \times 100$$

where $m_0$ and $m_1$ are the weights of the fibrous membranes before and after adsorption, respectively. The regeneration and adsorption/desorption performances of HMDI/HPIM-FMs were tested by H50 ($2 \times 1$ cm, $15$ mg) for 10 cycles using pump oil. After the weight measurements, the membranes were washed with hexane and immediately used for the next adsorption. Comparisons were made by film form of the same polymer.

Separation Experiments. For the oil/water separation experiments, 3 mL each of pump oil and water were mixed and the oil was colored with Sudan III. Free-standing fibrous membranes ($\sim 15$ mg) with dimensions of $2 \times 1$ cm were immersed in the mixture to separate the oil from the mixture. The adsorbed oil was transferred to another sample vial. Following that, the membranes were immersed in hexane, and they were gently rinsed with the help of tweezers to remove adsorbed oil from the fibrous membrane surface. An oil/water separation experiment was performed using sample H50 for 5 continuous cycles.

Results and Discussion

Structural Characterization of PIM-1 and HPIM. The synthesis and characterization of PIM-1 and HPIM are now well-known; thus, detailed structural characterization and purity of PIM-1 and HPIM were performed based on previous studies.43 The ATR-IR and $^1$H NMR spectra and TGA curves of PIM-1 and HPIM powders are provided in Figure S1.

Electrospinning HPMs and HMDI/HPIM-FMs. The properties and applications of powder and film form of PIM-1 have been reported in the literature.30,44,45 The electrospinning of fibers from PIM-1 is a fairly new concept compared to those of powder and film form.31,33 Recently, we reported the electrospinning of HPIMs and their optimized parameters to obtain uniform bead-free fibers along with their dye adsorption abilities based on the degree of hydrolysis.43 In the present study, initially we performed the electrospinning of fully hydrolyzed PIM-1 (HPIM) ultrafine fibers in DMF, which is similar to a previous report.43 Subsequently, mixtures of HMDI/HPIM were prepared in DMF in different compositions (Table 1) for the electrospinning of fibrous membranes (HMDI/HPIM-FMs). Bead-free and uniform ultrafine fibers were obtained under the same electrospinning parameters as pristine HPIM fibers. Consequently, HMDI/HPIM-FMs were heated at $150^\circ$C for 2 h to dry the membranes and to catalyze the cross-linking reaction. The optimization process of membrane production is explained in detail in Figures S2–
The electrospinning process, reaction pathway, and estimated structures of HMDI/HPIM-FMs are displayed in Figure 1.

SEM images and average fiber diameter distributions of HPIM-FM and cross-linked HMDI/HPIM-FMs are displayed in Figure 2. All samples show bead-free and relatively uniform fibrous morphology. Furthermore, the average fiber diameters were found to be between 1.38 ± 0.29 and 0.96 ± 0.22 μm (Table 1), depending on the amount of HMDI added to the HPIM solution for electrospinning. The reduction in the average fiber diameter was explained by the miscibility of HMDI and DMF because they can mix easily without precipitating HPIM. HMDI, which is added to the HPIM solution, acts as a solvent along with DMF. Owing to the increased volume of the solvent, the concentration of HPIM decreased, resulting in a smaller fiber diameter. The solubility of HPIM-FM changed after cross-linking with HMDI, and thus the HMDI/HPIM-FMs became insoluble in common organic solvents (THF, CHCl₃, hexane, diethyl ether, DCM, acetonitrile, MeOH, EtOH, 2-propanol, DMSO, DMF, and N-methyl-2-pyrrolidone), which indicates more stable materials against solvents. The samples were soaked in an electrospinning solvent (DMF) overnight to examine the success of the cross-linking reaction between HPIM and HMDI, and SEM images were obtained before and after treatment, as shown in Figure 2. While HPIM fibers were immediately dissolved, samples with less HMDI content (H10 and H20) showed slight swelling behavior in DMF. However, samples with more HMDI content...
(H33 and H50) exhibited no swelling behavior in DMF, indicating that the structure stability of the fiber was improved by increasing the amount of HMDI used in the preparation of HMDI/HPIM-FMs.

**Structural Characterization of HMDI/HPIMs.** Structural characterization of HMDI/HPIM-FMs was conducted using both Fourier transform infrared (FT-IR) and XPS measurements, and the effect of the HMDI content on the chemical structure was studied by altering the amount of HMDI in the fibrous membrane. The FT-IR spectra of HPIM-FM and HMDI/HPIM-FMs are depicted in Figure 3a. It can be seen that HPIM shows several $\sim$NH peaks in the region 3100~3500 cm$^{-1}$, which gradually turns a single $\sim$NH peak because of the formation of a urea structure. The expanded spectra of this region are also shown in Figure 3b. In addition, the carbonyl peaks of amides 1 and 2 (1600 and 1660 cm$^{-1}$) also disappear after chemical cross-linking and turn into a single peak while shifting slightly to 1680 cm$^{-1}$. Moreover, the signals around 2260 cm$^{-1}$ indicate the presence of NCO functionality. The presence of this peak can be explained by the bifunctional structure of HMDI. It can be observed to be due to either the unreacted side of HMDI or a slight excess of HMDI usage.

The XPS spectra of the corresponding samples are also displayed in Figure 4, and a clear intensity increased can be seen by integration of the N/O ratio, which increases from 0.54 to 1.19 by HMDI modification of HPIM. The nitrogen contents of HMDI/HPIM-FMs are increased by chemical cross-linking compared to HPIM-FM. This was also supported by elemental analysis of electrospun HMDI/HPIM-FMs, which is displayed in Table 1. While the carbon content does not change significantly, the nitrogen content increases proportionally from 5.6 to 10.1%, which highlights the addition of HMDI units to the HPIM structure.

The thermal properties of HMDI/HPIM-FMs were investigated using TGA and DSC techniques. Clear differences in the thermal stability were observed, as displayed in Figure 5a. The HPIM fiber shows a small weight loss (~3%) around 150 °C, which may be due to either the small amount of DMF trapped in the fibers because its powder form does not show this weight loss (Figure S1c) or decomposition of the carboxylate salt present in HPIM. Then the polymer starts to degrade from 240 to 540 °C, which is due to amide functional group and polymer backbone degradation, respectively. HMDI/HPIM-FMs show two significant degradation steps based on the amount of the HMDI content. The first degradation occurs between 240 and 330 °C because of decomposition of the urethane groups. The second one happens because of degradation of the HPIM backbone. Similar degradation behaviors are also observed in the DSC measurements, as presented in Figure 5b.

Furthermore, the mechanical behaviors of HMDI/HPIM-FMs were also investigated by the DMA technique. The mechanical strengths of HMDI/HPIM-FMs were found to be higher than that of pristine HPIM-FM. The stress–strain curves of selected samples are displayed in Figure 6a. Young’s moduli of the samples were found to be 5 ± 0.4, 25 ± 2, and 110 ± 23 MPa for HPIM, H20, and H50 respectively. Cross-linking usually increases the storage modulus of the polymer because of a reduction in the chain mobility. The mechanical strength of HPIM improved after cross-linking with HMDI, and storage moduli of the selected samples are presented as a function of the temperature in Figure 6b. An increase in the amount of the HMDI content of the membrane increased the storage moduli of the samples, which were found to be 0.4, 3.2, and 14.3 MPa for HPIM, H20, and H50, respectively, at 200 °C. Visual representations of the flexibilities of HPIM-FM and, unlike its film form, which has a water contact angle of...
about $60^\circ$, the water contact angle of HPIM-FM was found to be $140 \pm 4^\circ$ (Table 1 and Figure 7a). The result, in general, is in accordance with the previous studies. Mostly, the value of the contact angle depends on the surface roughness, and it increases with an improved surface roughness. Therefore, fibrous membranes are more hydrophobic than the films.\textsuperscript{52,53} In addition, the effect of cross-linking HPIM with HMDI on the hydrophobicity was also studied, and it was found that membranes became superhydrophobic depending on the amount of HMDI unit integrated in the polymer. As shown in Figure S5, both HPIM-FM and HMDI/HPIM-FM (H50) were soaked in organic liquids for 1 h. While HPIM is soluble in DMSO and DMF, H50 is insoluble in any common organic solvent that we tested. The weight-based adsorption capacities of HPIM and H50 fibrous membranes are given in Figure 8a. Also, data are presented in Table S1. While both fibrous membranes show high adsorption capacities against organic compounds, HPIM-FM shows a greater adsorption capacity, which is possibly due to the fluffy characteristic of the sample. Liquids can penetrate the HPIM structure more easily than they can H50, that has a more film-like structure, as provided in supporting video S1 (HPIM) and supporting video S2 (H50). Even though the adsorption capacity of H50 is lower than that of HPIM, H50 is able to adsorb DMSO and DMF better than HPIM-FM thanks to its insolubility. The maximum weight gain can reach up to 3400% for HPIM-FM and 2000% for H50 for CHCl$_3$ adsorption. Similarly, silicone oil adsorption can reach over 2000% for both fibrous membranes. Nevertheless, the insolubility in the organic solvents and the structural stability of H50 make it a better candidate for adsorption studies because the reusability is as important as the capacity for an efficient adsorbent. It is possible to reuse H50 by simply using hexane, but HPIM-FM can lose its physical properties in oils or after removal of oil with hexane, as shown in Figure S6. Therefore, the performance of H50 in adsorption was investigated for pump oil adsorption for 10 cycles. After 10 adsorption/desorption cycles, H50 exhibited 89% removal efficiency in comparison to its first usage, as shown in Figure 8b. SEM images of reused H50 are presented in Figure 8c, showing no structural damage on the fiber morphology.

The prepared HMDI/HPIM-FMs have hydrophobic surfaces along with 3D interconnected pore structures. Hence, it was expected to be a good candidate for the adsorption of organic compounds. Several different oils (silicone oil, pump oil, and sunflower oil) and organic solvents (MeOH, CHCl$_3$, DCM, hexane, toluene, DMF, and DMSO) were tested to investigate the adsorption capacities of HPIM-FM and HMDI/HPIM-FMs for organic compounds. As shown in Figure S5, both HPIM-FM and HMDI/HPIM-FM (H50) were soaked in organic liquids for 1 h. While HPIM is soluble in DMSO and DMF, H50 is insoluble in any common organic solvent that we tested. The weight-based adsorption capacities of HPIM and H50 fibrous membranes are given in Figure 8a. Also, data are presented in Table S1. While both fibrous membranes show high adsorption capacities against organic compounds, HPIM-FM shows a greater adsorption capacity, which is possibly due to the fluffy characteristic of the sample. Liquids can penetrate the HPIM structure more easily than they can H50, that has a more film-like structure, as provided in supporting video S1 (HPIM) and supporting video S2 (H50). Even though the adsorption capacity of H50 is lower than that of HPIM, H50 is able to adsorb DMSO and DMF better than HPIM-FM thanks to its insolubility. The maximum weight gain can reach up to 3400% for HPIM-FM and 2000% for H50 for CHCl$_3$ adsorption. Similarly, silicone oil adsorption can reach over 2000% for both fibrous membranes. Nevertheless, the insolubility in the organic solvents and the structural stability of H50 make it a better candidate for adsorption studies because the reusability is as important as the capacity for an efficient adsorbent. It is possible to reuse H50 by simply using hexane, but HPIM-FM can lose its physical properties in oils or after removal of oil with hexane, as shown in Figure S6. Therefore, the performance of H50 in adsorption was investigated for pump oil adsorption for 10 cycles. After 10 adsorption/desorption cycles, H50 exhibited 89% removal efficiency in comparison to its first usage, as shown in Figure 8b. SEM images of reused H50 are presented in Figure 8c, showing no structural damage on the fiber morphology. An investigation of the adsorption ability of HMDI/HPIM-FMs was performed by comparing dense (film) and fibrous membranes of H50. Note that the production of the film form of H50 was significantly more difficult by the solvent evaporation method. We were able to detach the membrane from the glass surface, but maintaining the membrane in a flat shape was not possible. The membrane was prone to roll by

**Figure 6.** Representative (a) stress–strain curves and (b) storage modulus–temperature graphs of HPIM-FM and HMDI/HPIM-FMs.

**Figure 7.** (a) Water contact angle versus % HMDI for HMDI/HPIM-FMs and (b) a digital image of pump oil (red) and water droplets (yellow) on HPIM and H50. The oil was colored by Sudan III dye, and water was colored by orange II dye.

**Figure 8.** (a) Stress–strain curves and (b) storage modulus–temperature graphs of HPIM-FM and HMDI/HPIM-FMs.
itself as the solvent evaporated (Figure S7). It is now well-known that PIM-1 has high BET surface area ($\sim 770$ m$^2$ g$^{-1}$), and it can adsorb organic molecules from its surroundings. Similarly, HPIM also has a relatively high surface area ($\sim 400$ m$^2$ g$^{-1}$) in powder form, but it loses its surface area in the membrane form ($\sim 20$ m$^2$ g$^{-1}$). The addition of HMDI cross-linking further reduces the surface area of the fibrous membrane (Figure S8). Owing to a reduction in the surface area, the adsorption performance of HMDI/HPIM-FMs can only be explained by the 3D structure of the fibrous membranes. Great hydrophobicity, along with a fibrous nature, enables HMDI/HPIM-FMs to have high adsorption capacity. Comparative pump oil adsorption of the dense and fibrous membranes of H50 has been studied to support this idea. While the dense form can only have 60% weight gain, which is due to the accumulation of oils on the film surface, the fibrous membranes have a higher removal rate.

Figure 8. (a) Adsorption capacities of HPIM and H50 toward different organic liquids and oils, (b) % removal of pump oil by H50 in 10 adsorption/desorption cycles, (c) SEM images of H50 after 10 pump oil higher (inset) and lower magnification adsorptions, and (d) SEM image of an oil-adsorbed H50 fibrous membrane.

Figure 9. Digital images showing the adsorption performance of H50 on organic compounds upon removal of (a) pump oil (red) from the water surface and (b) CHCl$_3$ (red) from the bottom of water as a function of time.
membrane can reach up to 1300% weight because of the fibrous nature. This further provides a slightly dried SEM image of H50 after oil adsorption in Figure 8d. As can be seen, oil can penetrate inside the fibrous membrane and remains there. In addition, the change in the dimensions of fibrous membranes is investigated using isopropyl alcohol. The samples with dimensions of 3 × 3 were employed for HPIM and H50. After an effective equilibrium was reached, both samples were swollen and increased in size. However, HPIM was not a handful material, and weight measurement was not successfully conducted, as experienced in the oil/water separation experiment (Figure S9). On the other hand, H50 maintained its membrane structure (Figure S9), and it had a significant weight gain of around 2200% and a ~10% increase in its dimensions, as shown in Figure S10.

The oil adsorption capacity of HMDI/HPIM-FMs (H50) was found to be approximately between 13 and 25 mg g⁻¹, depending on the type of the oil that was used in the experiment (derived from Figure 8a). Some examples of oil adsorbents and their adsorption capacities along with their structural forms are provided in Table S2. The adsorption capacity of H50 is not the highest data in the literature, but it is greater than those of many other oil adsorbents. In addition, the research related to electrospon PIMs is still in its early stages. It is highly possible that the capacity may be easily improved in the future by using different forms of PIM and cross-linker structure. Nevertheless, within the scope of this study, the reusability and durability of the HPIM fibrous membrane were improved by the introduced modification. The feasibility of HMDI/HPIM-FMs for practical applications was further demonstrated by using H50 to separate two organic compounds (pump oil and CHCl₃) with different intensities from an aqueous system. As shown in Figure 9a, pump oil (colored by Sudan III) can be immediately and selectively adsorbed by a piece of H50 mat as a result of the superhydrophobic surface and 3D interconnected fibrous structure. The membrane was still floating on the water surface after adsorption because of the low density, which may help to remove oil easily from water. In another example, a piece of H50 mat was soaked in water to adsorb CHCl₃ (colored by Sudan III), which was at the bottom of water (Figure 9b). The results clearly show that HMDI/HPIM-FMs could immediately separate organic compounds from aqueous systems regardless of their densities. The rapid separation can be seen in supporting video S3 (oil adsorption) and supporting video S4 (CHCl₃ adsorption). The reusability and adsorption/desorption performances of HPIM and H50 were also investigated in effective oil/water separation experiments several times, as presented in supporting video S5 (HPIM) and supporting video S6 (H50). While H50 can be used five times continuously in oil/water separation experiments without sustaining any damage to its structural stability, HPIM loses its membrane form after 2 cycles of separation. Thus, it is not suitable for practical oil/water separation.

CONCLUSION

HMDI/HPIM-FMs were successfully fabricated by an electrospinning technique prior to thermal treatment. In order to evaluate the effect of modification on the properties of electrospun HPIM-FM, the fiber morphology, mechanical strength, adsorption and separation performances, and thermal resistance were investigated. HMDI/HPIM-FMs were found to be insoluble in common organic solvents. In addition to insolubility, the materials are thermally and chemically stable up to 240°C. The mechanical properties of HPIM-FM were also improved by HMDI modification, in which HMDI/HPIM-FMs have shown higher Young’s and storage moduli compared to pristine HPIM-FM. HMDI/HPIM-FMs have demonstrated superhydrophobicity with water contact angles of up to 159 ± 7°. Therefore, they were successfully employed in adsorption and separation applications. Owing to the mechanical strength, insolubility, and superhydrophobicity, HMDI/HPIM-FMs exhibited notable adsorption and separation performances along with high reusability.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsanm.8b00115.

Characterization of PIM-1 and HPIM, optimization of the membrane production, and organic and oil adsorption (PDF)

Supporting video S1 (HPIM flexibility) (AVI)
Supporting video S2 (H50 flexibility) (AVI)
Supporting video S3 (oil adsorption) (AVI)
Supporting video S4 (CHCl₃ adsorption) (AVI)
Supporting video S5 (HPIM oil/water separation) (AVI)
Supporting video S6 (H50 oil/water separation) (AVI)

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The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors thank Dr. Amaresh C. Pradhan for his practical help with XPS analysis.

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