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Fabrication of Porous Matrix Membrane (PMM) Using Metal-Organic Framework as Green Template for Water Treatment

Jian-Yuan Lee, Chuyang Y. Tang, & Fengwei Huo

Pressure-driven membranes with high porosity can potentially be fabricated by removing template, such as low water stability metal-organic frameworks (MOFs) or other nanoparticles, in polymeric matrix. We report on the use of benign MOFs as green template to enhance porosity and interconnectivity of the water treatment membranes. Significantly enhanced separation performance was observed which might be attributed to the mass transfer coefficient of the substrate layer increased in ultrafiltration (UF) application.

With the increasing world population, the demand for clean drinking water also increases exponentially. One of the most challenging global problems affecting many countries around the world is clean water shortage and thus growing concerns over the access to safe drinking water. To meet this challenge, membrane-based water treatment processes have become one of the most energy efficient and most reliable technologies for supplying water. Compared to conventional thermal distillation processes, membrane-based water treatment processes have relatively lower energy consumption, improved cost effectiveness, and ensuring a more sustainable production of drinking water in the future. Hence, membrane-based water treatment processes have been employed in a wide range of applications such as oil/water separation, wastewater treatment, water purification, seawater desalination, food processing, and valuable product recovery.

In pressure-driven membrane processes, such as ultrafiltration (UF), nanofiltration (NF), and reverse osmosis (RO), an external pressure is applied on the solution at one side of the semi-permeable membrane, which serves as a driving force to separate water effectively from the pollutant in the feed solution. The water permeability coefficient is mainly determined by the membrane structure, such as porosity, pore diameter, and tortuosity. In theory, an ideal membrane is expected to have highly porous but very thin and low tortuosity structure. Therefore, higher porosity of membrane will result in higher water permeability coefficient and therefore the higher flux of the membrane processes. Previous studies showed that a variety of additives such as LiCl, polyvinylpyrrolidone (PVP), polyethylene glycol (PEG) were added into the polymer dope solution in order to promote the pore formation. However, all these additives can easily dissolve in the polar organic solvent and was removed during the non-solvent induced phase inversion (NIPS), which resulted in lower porosity. Alternatively, mixed matrix membrane (MMM) is an emerging strategy to enhance several intrinsic membrane properties of pressure-driven membranes. For example, water permeability, anti-fouling property, thermal and mechanical stability of pressure-driven membranes can be potentially enhanced by incorporating nanomaterials such as zeolite, silica nanoparticle, titanium oxide nanoparticle, and silver nanoparticle. However, poor dispersion of the filler in the polymer matrix and incompatibility issue between the polymer and filler might result in poor selectivity of the MMMs. Herein, we proposed the use of benign metal-organic frameworks (MOFs) as green template for the formation of macropores to enhance porosity and interconnectivity of the water treatment membranes.

Our basic idea is to construct a porous matrix membrane (PMM) using MOF particles as template, which initially cannot dissolve in the polar organic solvent but can be easily washed away by water. We believe that our strategy might potentially further improve the porosity of the membranes after removal of these particles. In addition, compared to conventional etching methods, such as the removal of silica nanoparticle by hydrofluoric acid or the removal of gold nanoparticle by potassium iodide, our strategy uses water as the...
etching solution, which is low cost, environmental-friendly and easily available. Herein, we report on the use of MOFs as green template to enhance porosity and interconnectivity of the water treatment membranes. MOFs were chosen as the templates because some of MOFs have relatively low water stability compared to zeolite and silica gel but relatively high stability in organic solvents. Firstly, we studied the dispersion of MOF as filler by embedding less water soluble MOF (F300, iron benzene-1,3,5-tricarboxylate) into polyacrylonitrile (PAN) as a negative control. Secondly, we chose two types of hydrophilic MOF (A100, aluminium terephthalate and C300, copper benzene-1,3,5-tricarboxylate), which is relatively lower water stability than the F300 as the green template. Lastly, the morphology and structure of MOF-based MMMs and PMMs were systematically characterized by field emission scanning electron microscopy (FESEM) and energy-dispersive X-ray spectroscopy (EDX). We speculated that the performance enhancement of pressure driven membrane process is due to the mass transfer coefficient of the PMM increased in ultrafiltration (UF) application.

Results
Firstly, the morphology information and water stability of MOF (A100, C300, and F300) in pure water were shown in Figure 2. A100 is a grey powder while F300 is an orange powder as observed by naked eyes. Both A100 and F300 showed as irregular shaped fine particle under the observation of FESEM. On the other hand, C300 is a blue powder and stone-like particle under the observation of FESEM. Both A100 and C300 showed relatively low water stability and can be completely dissolved in pure water compared to F300, which showed relatively high water stability and can only partially dissolved in aqueous solution.

The preparation of PMMs is outlined in Figure 3. MOFs were firstly dispersed in the DMF by ultrasonication of the solution for 1 hour. Polymer and LiCl were then added into the DMF solution followed by stirring for at least 24 h at 60°C until a homogeneous solution was obtained. A casting knife was used to cast the polymer solution onto a clean glass plate. The polymer film was then immersed into the tap water bath at room temperature followed by soaking in continuous flow water bath to completely remove the MOF as well as the excess solvent inside the membrane. The advantages of our strategy including: 1) MOFs were used as green template, which can be easily removed by water without using toxic chemical, strong acid or base; 2) the fabrication process is under mild condition: neutral pH, at room temperature and under atmospheric pressure; 3) the processes is environmental-friendly, which
minimizes the use of chemicals and reduces the impact on the environment; 4) our strategy is compatible with the current membrane manufacturing processes without extra steps or modification; 5) MOFs is commercial viable, low cost and could be easily scale-up.

Figure 4 showed the FESEM and EDX images of cross sections of PMMs and MMMs. The separation performance and the porosity of the PMMs compared with the control and MMM were shown in Figure 5. The cross-section images of a control, which represent the commercial UF membrane and a negative control (F300), which represent MOF-based MMM were shown in Figure 4 (a) and (b). The presence of MOF (F300) can still be observed using FESEM and further confirmed by EDX analysis (Figure 4 (c) and (d)). These results proved that the MOF were successfully incorporated into the polymer matrix and good dispersion was achieved after sonication. Correspondingly, the marginal increased in water permeability and porosity were observed in Figure 5 (a) and (b). This might be explained by the partial dissolution of F300 under neutral pH condition.

**Discussion**

More importantly, the successful removal of MOF particle was confirmed by FESEM as shown in Figure 4 (e) and (f). Significant amount of macropores were created in the polymer matrix. As a result, a highly porous PMM with porosity of 83±2% based on gravimetric measurement was obtained (Figure 5 (b)). Generally, a membrane substrate with high porosity is generally more favored due to increase the mass transfer of water in pressure-driven membrane processes. Indeed, the highly porous matrix ultrafiltration membranes (PMM and PMC) generally showed higher water permeability without compromising the rejection of dextran. The highest water permeability of 260.5 L/m² h bar was achieved for PMM, which probably due to the combination of enhanced porosity and interconnectivity of the ultrafiltration membranes. It is important to note that the dextran rejection did not significantly change between control and highly porous PMM after removal of MOF in the substrate layer, which means the membrane selectivity remain unchanged. Our overall results clearly show that the removal of MOFs in the polymer matrix substantially enhanced the porosity and is a promising strategy for better separation performance of pressure-driven membranes.

In order to determine the residual amount of MOFs inside the PMM membrane, the PMM samples were digested with a mixture of concentrated nitric acid and concentrated hydrochloric acid in the ratio of 3:1 for 6 hours followed by topping up the concentrated acid solution to 10 mL with water. Precipitates were observed at the bottom of the bottles prior to inductively coupled plasma (ICP) analysis, which can determine metal ions in aqueous solutions down to part per millions (ppm) or even part per billions (ppb) level. Indeed, the PMM has the highest residual percentage of iron (24.0±6.9 wt%). In comparison, PMMA has significantly lower residual percentage of aluminium (8.4±0.6 wt%) and PMMC has the lowest residual percentage of copper (3.8±0.4 wt%) among the three PMM samples. It is noteworthy that the order of membrane water permeability (F300, A100, C300) coincides with the (reverse) order of water stability and residual percentage of MOF (F300>A100>C300). Thus, one reasonable explanation for the highest water permeability of PMC is the higher water solubility of C300. The enhanced pore connectivity in addition to improved porosity is likely responsible to the marked increased in membrane:

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**Figure 4 |** FESEM micrographs of cross-section of (a) control, (b) PMM, (c) PMM, inset: enlarged image of F300, which cannot be completely remove under neutral condition, (d) EDX image of the enlarged image, (e) PMMA, (f) PMMC.

**Figure 5 |** Effect of different types of MOF on the performance of pressure-driven ultrafiltration membranes (a) the pure water permeability and dextran rejection of the ultrafiltration membranes; (Testing conditions: DI water or dextran solution as the feed solution; applied pressure: 5 bars) and (b) membrane porosity.
water permeability. However, the authors admitted that other important parameters, such as the loading, shape, and size of MOF particles, water stability of MOFs, and other types of water soluble MOFs or other kinds of nanomaterials should be optimized to further enhance the performance of PMM membranes.

In summary, MOF-based MMMs and PPM were systematically synthesized and characterized in this study. PPMs can significantly enhance water permeability in pressure driven membrane processes probably due to the removal of MOF particles in the polymer matrix, which enhanced membrane porosity and interconnectivity of the membrane. More importantly, the current study demonstrates which enhanced membrane porosity and interconnectivity of the membrane. More importantly, the current study demonstrates that the MOF particles, water stability of MOFs, and other types of water soluble MOFs can improve the performance of membrane. The enhanced membrane porosity and interconnectivity of the membrane. More importantly, the current study demonstrates that the MOF particles, water stability of MOFs, and other types of water soluble MOFs can improve the performance of membrane.

Typical procedure for fabrication of porous matrix membrane (PMM). To achieve good dispersion of the particles, 1.0 wt.% of MOF particles were added into DMF followed by ultrasonicating the solution for 1 h. PAN (18 wt.%) and LiCl (2 wt.%) were then added into the DMF solution followed by stirring for at least 24 h at 60 °C until a homogeneous solution was obtained. The polymer solution was then cooled down to room temperature overnight without stirring to completely remove any gas bubbles. A casting knife (Elcometer Pte. Ltd., Asia) was used to spread the polymer solution onto a clean glass plate at a gauge height of 150 μm. The plate was then immersed into tape water at room temperature for 10 min till the PAN substrate was separated from the glass plate. The resulting membrane substrate was soaked in tap water bath followed by degoned (DI) water rinsing and the resulting membrane was designated as (Control, PMMDM, PMMDM, and PMMDM).

Membrane characterization. The surface and cross section morphological structures of the prepared PAN membranes were characterized by field emission scanning electron microscopy (FESEM, JSM-7600F, USA). A freeze-dryer was used to dry all the membranes samples at room temperature for at least 12 h followed by coated with a uniform platinum layer before observation. Element mapping was detected with the FESEM microscope equipped with energy-dispersive X-ray spectroscopy (EDX, Oxford Instrument, UK). The methods for determining intrinsic separation properties such as pure water permeability and dextran rejection can be found elsewhere. 

Gravimetric measurement. The membrane porosity, \( \varepsilon \), was defined as the volume of the pores divided by the total volume of the membrane. It can be determined by gravimetric method, measuring the weight of water contained in the membrane pores. By measuring the dry mass (\( m_{dry} \)) and wet mass (\( m_{wet} \)) of membrane samples, the membrane porosity (\( \varepsilon \)) can be calculated according to following equation (1):

\[
\varepsilon = \left( \frac{m_{dry} - m_{wet}}{m_{dry}} \right) \times 100\%
\]

where \( m_{dry} \) and \( m_{wet} \) are the density of water (1.0 g/cm³) and the density of PAN (1.18 g/cm³). It was assumed that all the pores in the membrane and silica gel were completely filled with water.


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Author contributions
J.-Y. L. conceived the idea, designed and carried out the experiments, analyzed the data and co-drafted the manuscript. F.H. and C.Y.T. supervised the project, helped design the experiments and co-drafted the manuscript. All the authors contributed to the interpretation of the results and writing the manuscript.

Additional information
Competing financial interests: The authors declare no competing financial interests.


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