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Breaking through permeability-selectivity trade-off of thin-film composite membranes assisted with crown ethers

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Abstract

In this study, we deployed a modified interfacial polymerization process to incorporate multifunctional crown ethers (CEs) into thin-film composite (TFC) polyamide membranes. These CE additives acted as both the phase-transfer catalyst and co-solvent to facilitate the diffusion of amine monomers into the organic phase and also enhanced the free volume content of the selective layer, facilitating water transport and inhibiting the diffusion of draw solutes. Various characterization techniques were employed to elucidate the modification mechanism as a function of CE chemical and physical properties on the microstructure of resultant TFC membranes and consequently separation performances. Compared to TFC membranes produced from traditional interfacial polymerization method, CE-modified membranes exhibited a 146% water flux enhancement and 59% lower reverse salt fluxes with a suitable draw solution. CE-modified membranes also exhibited improved antifouling performance with a lower flux drop (34% decline) and a higher flux recovery ratio (38% improvement).

Keywords: Thin-Film Composite Membrane; Polyamide; Crown Ether; Interfacial Polymerization; Permeability-Selectivity Trade-off
1. INTRODUCTION

By 2050, the global population will increase by 2 billion, reaching 9.7 billion people.¹ Key to sustaining this burgeoning growth is overcoming water scarcity for anthropogenic activities.² Various water treatment technologies have been exploited in the past few decades to produce clean water.³⁻⁶ For example, membrane-based separation techniques such as ultrafiltration,⁷,⁸ nanofiltration,⁹⁻¹¹ reverse osmosis¹²,¹³ and forward osmosis (FO)¹⁴⁻¹⁶ have been implemented at various levels for desalination, wastewater and water treatment. The heart of the successful implementation of these technologies is a high-performance membrane with superior separation efficiency and excellent fouling resistance. Since the commercialization of thin-film composite (TFC) membranes in the 1970s,¹⁷ such membranes are preferred in the industry for their high fluxes, reasonable selectivity and ease of fabrication. TFC membranes are generally produced by interfacial polymerization (IP) of aromatic amine monomers dissolved in an aqueous solution and an organic solution containing acyl chloride monomers to form a dense aromatic polyamide (PA) layer on a porous support layer.¹⁸⁻²¹ Both the reaction rate and degree of the polymerization are determined by the solubility of the reactive monomer in the opposite phases i.e. the diffusion rate of amine monomers from the aqueous phase into the organic phase.²² Hence the diffusion-limited growth of PA chains coupled with the extremely fast polymerization kinetics between both highly reactive monomers typically result in poor control over bulk properties and microstructure of the formed PA layer, consequently impacting on the separation performances of resulting TFC membranes.

The IP process can be tailored via various approaches to enhance separation properties from an optimized PA microstructure. For example, the addition of catalytic
additive in the monomer solution or a co-solvent system altered diffusion rates of amine monomers into the organic phase and IP kinetics that consequently optimized bulk properties and enhanced separation performances. Alternatively, effervescence i.e. formation of nanosized gas bubbles in the amine solution via ultrasonication or addition of NaHCO₃ to the amine solution also impacted on the morphology and separation performances of TFC membranes. Similarly, we recently showed that ultrasonication during IP was effective for facilitating efficient mixing of two monomer phases that optimized free volume size and content in the PA layer. Additionally, the thickness, roughness and chemical composition of the PA layer could be controlled by molecular layer-by-layer deposition method reported recently, to yield maximum separation performances, where molecular transport was not impeded by kinetic and mass transfer limitations associated with the traditional IP process. It also can be achieved by employing an intermediate layer of nanostrands that regulated the release of amine monomers at the water-hexane interface, providing precise control over the PA morphology and thickness. Besides, 3D printed PA membranes developed by electrospraying of monomers on to a substrate also provided precise control over film thickness and roughness. Recently, Wang et. al. reported a nanostructure-mediated IP process to prepare rough PA layers with extensively crumpled nanoscale structures. The combination of these unique nanoscale membrane topologies led to unprecedented permeances, which were realized by preloading sacrificial templating nanoparticles on a substrate.

Different from these studies, here we incorporated crown ethers (CE) in the monomer solution to tailor the formation of aggregate and network pores within PA selective layers of TFC membranes that were crucial for addressing the tradeoff relationship between permeability and selectivity (Figure 1). These membranes were
applied in FO for the first time. CE, a cyclic polyether, is miscible with both water and organic solvents\textsuperscript{31} and is an excellent chelating agent with intrinsic cavities that are selective towards metal ions (especially for alkali metal ions).\textsuperscript{32} Therefore, it is widely used as the phase-transfer catalyst in organic reactions to transfer inorganic reaction reagents into the organic phase, accelerating chemical reactions.\textsuperscript{32} Moreover, the inner and outer of the CE cavity are hydrophilic and lipophilic, respectively.

Ascribing to these unique characteristics of CE, we hypothesized that CE-assisted IP could significantly impact on the microstructure, morphology as well as FO performances of resulting TFC membranes. This hypothesis was underpinned by two main attributes. First, the miscibility of CE with organic solvents such as hexane and the ability to form hydrogen bonds with amine monomers enable CE to act as a co-solvent and phase-transfer catalyst to enhance amine diffusion into the organic phase, contributing to a more complete IP reaction. Second, the hydrophilicity of CEs could be harnessed to improve the water permeance of resultant PA films during FO whilst chelating metal ions of draw solutes (NaCl and KCl used in this study), contributing to less draw solute leakage. We systemically verified these hypotheses using a series of complementary characterization techniques to elucidate the effects of various modification routes (adding in the aqueous or organic phase), CE type and concentration, as well as the draw solute type are investigated systematically in this study.
FIGURE 1 Schematic illustration of PA networks with CE15 and CE18 incorporated

2. EXPERIMENTAL

2.1 Materials

M-phenylenediamine (MPD, 99.5%) and 1, 3, 5-benzenetricarbonyl trichloride (TMC, 98%) were bought from Aladdin and kept in a refrigerator before use. Polysulfone (PSf) (Mw = 800,000 Da) was obtained from Beijing HWRK Chem co. Ltd. (China). PSf was dried overnight in a vacuum oven at 80 °C prior use. Polyethylene glycol 400 (PEG 400, CP), N-methyl pyrrolidone (NMP, anhydrous, ≥99.5%), n-hexane (≥97%, anhydrous), 18-crown-6 (CE18, 99%, powder), 15-crown-5 (CE15, 99%, liquid), sodium alginate (SA, Mw: 98.11), potassium dihydrogen phosphate (KH$_2$PO$_4$, 99.5%), magnesium sulfate (MgSO$_4$, 99%), sodium sulfate (Na$_2$SO$_4$, 99%), sodium bicarbonate (NaHCO$_3$, 99.5%), ammonium chloride (NH$_4$Cl, 99.5%), calcium chloride (CaCl$_2$, 96%) and sodium chloride (NaCl, ≥99.5%) were purchased from China National Medicine Corporation.
2.2 Preparation of TFC membranes

The PSf substrate membrane was fabricated using the non-solvent induced phase separation method.\textsuperscript{33,34} Briefly, a degassed dope solution (PSf/PEG-400/NMP: 18/16/66 wt.\%) was poured onto a clean glass plate and cast with a casting knife (3520-8, Elcometer, UK) to obtain at thickness of 100 μm. The PSf substrate was immediately immersed into a coagulation bath at room temperature to start the phase inversion process. To remove trapped solvent molecules in the PSf substrate, the water was changed every 12 hours for 2 days. These substrates were stored in deionized (DI) water that was changed every 12 h to remove residual NMP.

CE-modified TFC membranes were fabricated by adding CE into the aqueous (1 – 9 wt.\% of CE15 or 3 wt.\% of CE18) or organic phase (3 wt.\% of CE15 or CE18) prior IP. Detailed compositions of the two monomer solutions studied here are listed in Table S1. Depending on the type of CE used, 18-crown-6 or 15-crown-5, CE-modified membranes were denoted as CE(A/O)-18/15-\(X\) where “A” and “O” represents the deployment of CE in the aqueous or organic phase, respectively, and “\(X\)” stands for the CE content. The miscibility of CE15 and solubility of CE18 in water and hexane (Figure S1 and Table S2) were exploited here to underpin in-situ modification of PA selective layers of as-fabricated TFC membranes. The intrinsic cavities of CE18 molecules ranged from 2.6 – 3.3 Å in diameter (Figure 2),\textsuperscript{32,35-37} which are ideal for permeating water molecules (dynamic diameter of 2.6 Å).\textsuperscript{22} Meanwhile, the cavity diameters of CE15 molecules are significantly smaller (1.7-2.2 Å) and are impermeable for water molecules.\textsuperscript{32,35-37} These differences in cavity diameters also favor stable chelation of various cations where CE15 and CE18 molecules preferentially chelate Na\(^+\) (1.90 Å) and K\(^+\) (2.66 Å) ions, respectively.
Pristine PSf membranes were immersed in an aqueous MPD/crown ether (3.4/0 – 9 wt.%) mixture for 2 min firstly. Excess MPD solution was removed carefully from the substrate surface using a rubber roller. Next, the TMC/crown ether/hexane solution (0.15/0 – 3/100 w/w/v) was poured on to the membrane top surface and left for 1 min before draining off. The newly formed TFC membranes were washed with DI water and stored in DI water before use.

![Figure 2](image.png)

**FIGURE 2** Molecular structures and intrinsic cavity diameter of (a) CE15 and (b) CE18

### 2.3 Characterizations of TFC membranes

Changes in the chemical structures of TFC membranes were characterized using Fourier Transform Infrared spectroscopy in the Attenuated Total Reflectance mode (ATR-FTIR, Brucker, VERTEX-70) and X-ray Photoelectron Spectroscopy (XPS, VG Multilab 2000, Thermo VG Scientific, UK) where a monochromatic Al Ka X-ray source was employed. The free volume content of the PA layer micro-structure was characterized using Positron Annihilation Spectroscopy (PAS, National University of Singapore) via a variable mono-energy slow positron beam. The inter-chain distance ($d$-spacing) of the PA layer was examined using X-ray diffractometer (XRD,
SmartLab-SE, Rigaku, Japan) with $2\theta$ ranging from 10° to 60°. Surface hydrophilicity of TFC membranes was evaluated using the water contact angle (WCA) measured by a Contact Angle Goniometer (DSA 25, KRÜSS, Germany). Zeta potentials of TFC membrane were measured at 25 °C by a Zeta Potential Analyzer (SurPASS™ 3, Anton Paar, Austria) using 0.001M KCl aqueous solution with pH of 2–10. The surface morphology and topology of TFC membranes were observed by a Scanning Electron Microscope (SEM, VEGA3, TESCAN, Czech) and Atomic Force Microscope (AFM, SPM9700, Shimadzu, Japan) respectively.

### 2.4 Evaluation of separation performance

A lab-scale FO apparatus was employed to perform FO tests around 22±0.5 °C, using DI water and 2.0 M NaCl aqueous solution as the feed and draw solutions, respectively. All membrane samples were tested for at least three times under both FO mode (active layer facing feed solution, AL-FS orientation) and PRO mode (active layer facing draw solution, AL-DS orientation). Each test was stabilized for 30 minutes before data collection. Weight changes in the draw solution side were detected by a digital weight balance (FX3000-GD, AND, Japan). Additionally, concentration changes in the feed solution were monitored using a conductivity meter (FE30, Mettler Toledo, Switzerland). Water flux ($J_v$) and reverse salt flux ($J_s$) were determined to evaluate FO performance of TFC membranes, as defined by Eqs. (1) and (2).

$$J_v = \frac{\Delta V}{A_m \Delta t} \quad (1)$$

$$J_s = \frac{\Delta (C V_f)}{A_m \Delta t} \quad (2)$$

where $\Delta V$ is the volume change in the draw solution side over a predetermined time.
\[ (\Delta t), A_m \text{ is the effective membrane area in FO test (3.87 cm}^2\text{), } C_i \text{ and } V_i \text{ are the salt concentration and volume of the feed solution, respectively.} \]

### 2.5 Dynamic fouling tests

Following well-established protocols described in our previous works,\textsuperscript{36, 37} we performed dynamic fouling tests in the FO mode at 22±0.5 °C using synthetic wastewater and 2.0 M NaCl aqueous solution as the feed and draw solutions, respectively. Fresh membrane samples were stabilized by using DI water as both feed and draw solutions for 30 min. The initial FO flux was measured after reaching steady state by replacing DI water with 2 M NaCl solution as the draw solution. Next, the fouling process was conducted by replacing DI water in the feed solution with synthetic wastewater for 18 h (The fouling process time is 18 h using synthetic wastewater as the feed solution). The flow rate in above stages were maintained at 0.3 L/min. Fouled membranes were flushed with DI water as both feed and draw solutions at a flow rate of 0.6 L/min for 30 min. Finally, the FO fluxes of cleaned membrane were measured again.

### 2.6 Evaluation of CE stability in modified PA layers

CE stability in the modified PA layer was evaluated by a long-term FO test and vigorous agitation. For the long-term (72 h) FO test, a large amount of draw solution (4 L 2 M NaCl or KCl solution) and feed solution (4 L DI water) were employed to mitigate dilution effects of draw solutions and the concentration effects of feed solutions. The vigorous agitation treatment was conducted by immersing fresh
membrane samples (25 cm²) into a centrifuge tube with 50 mL ultrapure water, which
was fixed in an oscillator for 7 days, with daily water changes. The amount of CE
released in the ultrapure water after various treatment durations were characterized
using a Total Organic Carbon analyzer (TOC, Vario, Elementar, Germany).

3. RESULTS AND DISCUSSION

3.1 Modification mechanism

The impact of CEs on the physicochemical structure of PA was characterized by FTIR
analyses and validated by XPS (Figure 3). From the FTIR spectra of membranes
studied here (Figure 3a), we observed peaks centered at 1624 and 1548 cm⁻¹ that
were characteristic to CO-NH groups i.e. amide bonds.⁴⁰ The presence of CE in
modified PA selective layers was validated by additional peaks centered at 1085 and
986 cm⁻¹ that are correlated to the ether functional group⁴¹,⁴² that are unique to CE
across all membranes studied in this work. Interestingly, the intensity of the peak
centered at 3460 cm⁻¹ (stretching vibration of -OH) was reduced in CE-modified
membranes. We hypothesize that the more complete IP reactions were facilitated by
CE-assisted amine diffusion/dissolution in the organic phase that resulted in efficient
mixing of both reactive monomers. This contradicts the principles of IP where amine
molecules are typically insoluble in organic phases.¹⁸,²² This hypothesis was validated
from FTIR analyses where hydrogen bonding between amine molecules and hexane
in the presence of CE were observed (Figure S2).

Quantitative evidence of CE presence in PA selective layers was observed from
XPS analyses (Figure 3b). OII (*O-C/H-*O-C/C-*O-C, BE=532.2 eV)⁴³ and CII
(*C-OR, BE=285.4 eV\textsuperscript{44}) peaks that corresponded to ether functional groups were only present in CE-modified PA membranes where both the OII\% and CII\% (peak area ratio) of the CE15-modified membrane were larger than those modified by CE18 molecules (Figure S3). Based on our hypothesis, as more CE molecules are present, the IP reaction would be more complete. This was also observed from the larger increase in O/N ratio (Table S3 and Figure S4) where N is unique to PA and the increase in O content was primarily due to CE and higher intensity ratio of I\textsubscript{OIII}/OI (OIII corresponded to O atoms in carboxylic acid groups hydrolyzed from residual TMC acyl chloride groups and OI was correlated to unreacted TMC carbonyl). Both O/N and I\textsubscript{OIII}/OI ratios of CE15-modified membranes were higher than those of CE18-modified membranes. Attributing to a smaller molecular weight that facilitated better diffusion of CE15 into hexane, the O/N ratio in PA membranes modified by CE15 molecules was larger than that of membranes modified by CE18. TGA results also revealed that weight losses of CE-modified PA powders in the second stage (200-800 °C, caused by the thermochemical decomposition of PA chains) were lower than those of pristine PA powders, possibly due to the intermolecular hydrogen bond interaction between PA chains and embedded CEs, indicating the stable incorporation of CEs in the PA network (Figure S5).

The deployment of CE as co-solvent or phase-transfer catalysts during IP also led to the formation of rougher PA layers as confirmed by SEM and AFM images in Figures 3c-d. Figure 3c also revealed that pristine PA membranes were thinner with more nodular-like surfaces, while thicker CE-modified membranes comprised obvious leaf-like and flake-like structures. Accordingly, as shown in Figure 3d, the surfaces of CE-modified membranes were rougher than pristine membranes, and the average roughness (Ra) increased with higher CE concentration or smaller CE
molecular weight (Figure 3e). However, the WCAs of CE-modified membranes were lower than those of pristine membranes, contradicting the $Ra$ trends (Figure 3e). This was ascribed to the intrinsic hydrophilicity of CEs and rougher surface of CE-modified PA layers. Additionally, the lower carboxyl content in CE-modified PA i.e. higher IP conversion rates also attributed to higher zeta potential values at the same pH condition and the corresponding higher isoelectric points in CE-modified membranes (Figure 3f).
FIGURE 3 (a) FTIR spectra, (b) OII and CII peak area ratios, (c) SEM, (d) AFM, (e) water contact angles (WCA) and average roughness (Ra), (f) zeta potentials of the pristine and CE-modified membranes.

Structural changes induced by CE in the resulting PA layers were characterized by PAS and XRD (Figure 4). PAS results based on Doppler Broadening Energy Spectrum providing qualitative information about free volume, can be reflected by $S$ parameter, which is defined as the ratio of total annihilation counts at central region (511 keV). Generally, $S$ measures free volumes between 0.1 and 1 nm based on $2\gamma$
annihilation. Higher $S$ values typically indicate the more and/or larger free volume cavities in the polymer matrix, resulting in a looser structure.\textsuperscript{45} In this study, $S$ value at around 2 keV indicating the top PA layer was applied to estimate the free volume. $S$ values in CE-modified membranes were larger than those of the pristine membranes (Figures 4a and S6), indicating the formation of looser PA layers with larger free volume content. The $S$ value also increased as a function of CE15 content, CE molecular weight and intrinsic cavity size. Clearly, the effects of 9 wt.% of CE15 on microstructure and porosity could be achieved with 3 wt.% of CE18. The increases in free volume content and size were due to the enlargement of inter-PA chain distances caused by the addition of CE, where additives typically affect chain packing density.\textsuperscript{22,47} The interchain distance increased from 4.834 Å to 5.014 Å as CE15 concentration increased from 1 to 9 wt.% (Figures 4b and S7). Meanwhile, the incorporation of 3 wt.% of CE18 could achieve the same effect as 9 wt.% of CE15, propping PA chains apart by 5.001 Å.

![Graph](image_url)
323 (b)

FIGURE 4 (a) S and (b) d-spacing values of the pristine and CE-modified membranes

3.2 Separation performance, antifouling property, and long-term stability

Compared to pristine PA membranes, the larger free volume, higher surface hydrophilicity and rougher surface (larger transport area for water molecules) of the PA layers containing 1 – 7 wt.% CE15 enhanced water flux by 164% during FO operation were (Figure 5). However, as CE15 content increased to 9 wt.%, the thickness of the selective layer increased to 389.47±28.69 nm that consequently reduced the water flux by 14% and 17% in FO and PRO modes respectively. Different from typical membrane modifications that increase reverse salt fluxes, the addition of CE15 reduced reverse salt fluxes as a function of higher CE content during modification. This was due to the combination of thicker PA layers, the chelation of CE15 with sodium ions and the repulsion effect of trapped NaCl molecules.

Alternatively, we added both CEs into the organic phase during IP. This approach was impractical as unstable membranes that discolored after IP were formed (Figure S8). Membrane discoloration observed here could be due to the dehydration of
amine-saturated substrates caused by CE attraction. Additionally, PA layers modified with CE in the organic phase became yellow and shrank after freeze drying, and delaminated from the discolored PSf substrate (Figure S8). PA delamination resulted from the inability to achieve polymer intrusion into the pores during IP, leading to less anchoring sites. The combination of delamination and poor chemical resistance in PA membranes modified with organic phase CE led to significantly higher water fluxes when compared to those of pristine membranes (Figure 5b). However, the reverse salt fluxes of these CE(O) membranes were also 11-15 times higher than that of pristine membranes.

(a) (b)
with different CE15 loadings added in the aqueous phase; (b) with different CEs added in the organic phase; with different CEs added in the aqueous phase using (c) NaCl or (d) KCl as the draw solution.

To investigate the chelation ability of the CE15 and CE18 with Na\(^+\) and K\(^+\) ions, respectively, we employed draw solutions that contained 2M NaCl or KCl draw solution during FO separation (Figures 5c-d). The water fluxes of CE-modified TFC membranes in both FO and PRO modes increased as a function of CE cavity size i.e. water fluxes of CE-18 modified membranes were higher than that of CE-15 modified membranes. The osmotic pressure of these two draw solutions were nearly equal,\(^{49}\) hence resulting in 4-7% change in water flux, regardless membrane modification. However, the reverse salt fluxes of these TFC membranes with the two different draw solutions were vastly different. With NaCl draw solutions, the reverse salt flux of pristine PA membranes was 23% and 21% lower than that with KCl draw solution under FO and PRO mode. This was due to the smaller hydration radius of KCl. The reverse salt flux of PA membranes modified with CE15 was reduced by 57%, from 23.29 gMH to 10.03 gMH when NaCl draw solutions were employed. The reduction in reverse salt flux was only 24% CE15-modified PA membranes when KCl draw solutions were employed. The lower reverse NaCl flux of CE15-modified membranes was attributed to the stronger chelation ability between CE15 and Na\(^+\) ions, and smaller hydration radius of KCl. However, this was not the case for CE18-modified PA membranes where the reverse salt flux was lower when KCl draw solutions were deployed during FO and PRO. This was in spite of the smaller hydration radius of
KCl when compared to NaCl molecules. Clearly, the chelation ability CE18 with K\(^+\) ions played a more dominant role here. Compare to CE15-modified membranes, the reverse salt flux of CE18 membranes towards KCl draw solution was 31% lower, even when the cavity size of CE18 molecules were larger. EDX characterization were performed to quantify metal ion content in the membranes after FO tests (Table 1 and Figure S9). After FO test with NaCl draw solution, the Na content trapped in CE15-modified membranes was 2.27%, 6 times higher than that in CE18-modified PA membranes. Meanwhile, after FO test with KCl draw solution, K content in CE18-modified membranes were 139% higher than that in CE15-modified membranes, confirming the stronger chelation ability of CE18 with K\(^+\) ions.

**TABLE 1** Elemental composition of the pristine and CE-modified membranes after FO tests using different draw solutes

<table>
<thead>
<tr>
<th>Membrane code</th>
<th>C</th>
<th>N</th>
<th>O</th>
<th>Cl</th>
<th>Na</th>
<th>K</th>
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<td>Pristine-NaCl</td>
<td>58.34</td>
<td>11.86</td>
<td>28.38</td>
<td>0.87</td>
<td>0.55</td>
<td>0.00</td>
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<tr>
<td>Pristine-KCl</td>
<td>58.61</td>
<td>10.18</td>
<td>30.16</td>
<td>0.63</td>
<td>0.00</td>
<td>0.42</td>
</tr>
<tr>
<td>CE(A)-15-3-NaCl</td>
<td>58.73</td>
<td>9.27</td>
<td>28.14</td>
<td>1.58</td>
<td>2.27</td>
<td>0.00</td>
</tr>
<tr>
<td>CE(A)-15-3-KCl</td>
<td>59.22</td>
<td>8.95</td>
<td>31.28</td>
<td>0.05</td>
<td>0.00</td>
<td>0.51</td>
</tr>
<tr>
<td>CE(A)-18-3-NaCl</td>
<td>58.75</td>
<td>9.50</td>
<td>31.36</td>
<td>0.09</td>
<td>0.30</td>
<td>0.00</td>
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<tr>
<td>CE(A)-18-3-KCl</td>
<td>58.35</td>
<td>6.53</td>
<td>32.74</td>
<td>1.17</td>
<td>0.00</td>
<td>1.22</td>
</tr>
</tbody>
</table>

The fouling resistance of polymer membranes is crucial for determining the quality of processed water and membrane lifespan.\textsuperscript{50} Anti-fouling properties of
membranes studied here were investigated using synthetic wastewater containing SA and calcium ions in the feed solution. Calcium ions acted as the bridge to crosslink SA molecules, forming a SA gel layer on membrane surfaces. The formed SA gel layer could enhance both transport resistance that reduces water permeation, and osmotic pressure to significantly reduce the effective transmembrane osmotic pressure difference.

Upon fouling, the water flux of pristine PA membranes was reduced by 60%, reaching 6.5 LMH as shown in Figures 6a and S10. There as a 40% loss in water flux even after physical cleaning as the water recovery ratio only reached 60%. Meanwhile, the impact of fouling on the water flux of CE-modified membranes was less prominent where water fluxes were reduced by 24% and 30% for CE15- and CE18-modified membranes respectively. However, we were able to recover 95% of the water flux after physical washes. The improved anti-fouling properties of CE-modified membranes were ascribed to surface hydrophilicity and less reactive carboxylic acid groups. The predominant factor was surface hydrophilicity where the oxygen atoms in CE behaved as hydrogen bond acceptors, attracting water molecules onto the membrane surface to form a hydration layer that prevented the adsorption of SA molecules. Additionally, the more complete IP reaction in CE-modified membranes contained less carboxylic acid groups on the PA layer that typically function as complexation sites to chelate calcium ions.
FIGURE 6 (a) Dynamic fouling test results of the pristine and CE-modified membranes; (b) long-term FO test results of CE-modified membranes; (c) FO performance comparison with recently reported TFC membranes with various modifications (DI water and 2 M NaCl used as the feed and draw solutions, PRO mode).

We also investigated the stability of embedded CE within the modified PA layers via long-term FO tests and vigorous shaking treatments. CE-modified membranes exhibited stable water fluxes over continuous 72 h FO tests (Figures 6b and S11). Additionally, the reverse salt fluxes of CE-modified membranes increased by only...
9-12%, which was possibly due to the concentration effect of the feed solution. The above results demonstrated that the FO performance of CE-modified TFC membranes could be maintained in long term operation, inferring the stability of CE modifiers in modified PA layers during FO operation. PAS and XRD characterization of membranes after long term FO operation revealed that the $S$ parameter value decreased by 0.10-0.11% and the $d$-space decreased by 0.25-0.28% Å (Table 2 and Figure S12). These slightly smaller values suggested that the decrease in the free volume content of modified membranes was mainly underpinned by a reduction in intrinsic cavity free volume of CEs. It also could indirectly reflect that the increment in the free volume of CE-modified PA layer was mainly contributed by the larger aggregate and network pores (i.e. the enlarged spatial distance)\textsuperscript{22}. We also deployed a TOC analyzer to characterize the amount of CE released from the membranes after vigorous shaking (Figure S13). The average CE release rates of CE15-membranes and CE18-membranes during the 7-day shaking treatment are 0.064±0.02 and 0.055±0.01 ppm/cm$^2$·day, respectively, which further confirmed the stable CE-driven modification of the PA selective layer. The stability of CE modifications, creation of additional porosity whilst facilitating complete IP reactions underpinned the superior separation performances of PA membranes containing 7 wt.% of CE15 molecules. The FO performance of this membrane was more superior than those of various state-of-the-art TFC-PA membranes as displayed in Figure 6c,\textsuperscript{23,33,53-60} highlighting the feasibility of our approach.
TABLE 2 θ and 2 theta values of CE-modified membranes before and after long term FO tests

<table>
<thead>
<tr>
<th>Membrane code</th>
<th>S</th>
<th>d-spacing (Å)</th>
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<tbody>
<tr>
<td>CE(A)-15-3</td>
<td>0.4784</td>
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<td>CE(A)-15-3-NaCl</td>
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<td>4.856</td>
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<td>CE(A)-18-3</td>
<td>0.4867</td>
<td>5.001</td>
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<tr>
<td>CE(A)-18-3-KCl</td>
<td>0.4864</td>
<td>4.987</td>
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</table>

4. CONCLUSIONS

In the present study, we investigated the use of two identical CEs, CE15 and CE18, as additives in the aqueous monomer phase to modify the PA selective layer of resultant TFC membranes. Miscibility with hexane and the ability to hydrogen bond with MPD molecules render CE as an efficient phase-transfer catalyst and co-solvent that facilitated the diffusion of MPD molecules into the organic phase during IP. Consequently, PA layers with higher crosslinking degrees and rougher surfaces were achieved. Additionally, the incorporation of CE into the PA layer imbued higher membrane surface hydrophilicity and higher free volume content. This was ascribed to the polar ether groups and intrinsic cavities in CE as well as the enlarged spatial distance of PA chains. Moreover, in comparison with pristine PA membranes, the water fluxes of CE-modified membranes were enhanced by 164% and increased with higher CE concentration or larger CE variants. Crucially, the use of CE during IP lowered reverse salt fluxes of CE-modified membranes. We also demonstrated that chelation abilities with Na⁺ or K⁺ ions could be harnessed to tailor higher solute rejections of CE15- and CE18-modified membranes with suitable draw solutions. CE-modified membranes were less prone to fouling because of their higher surface
hydrophilicity and lower carboxylic content in the PA layer. The stable embedment of CE into the modified PA layer was also testified by long-term FO test. More importantly, our approaches of CE modification and CE-assisted IP produced FO membranes with state-of-the-art separation performances in a facile manner. This modification approaches could be extended to produce polymer membranes for nanofiltration and gas separations, potentially impacting other important separation applications.

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