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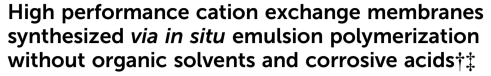


**PAPER** 

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The synthesis of cation exchange membranes (CEMs) usually involves using organic solvents and/or a sulfonation process. In this study, green and scalable synthesis of high performance CEMs is achieved without organic solvents and sulfonation. The synthesis is carried out *via in situ* polymerization of lithium styrene sulfonate in a porous support. Different preparation procedures are developed and optimized. Functional sulfonate groups are successfully loaded onto and into the membrane support, as verified by FTIR. Besides, water plays an important role in membrane synthesis. By reducing the amount of water used, the ratio of functional polymers to the membrane support in the synthesized CEMs is increased. Therefore, the synthesized CEMs show increased ion exchange capacity (IEC). This is significant because it means that high IEC can be achieved without introducing cation exchange resins into the membranes. Finally, the synthesized membranes demonstrate high desalination performance. This new methodology may shed new light on preparing CEMs in an efficient and eco-friendly way.

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# Introduction

Ion exchange membranes (IEMs) have been undergoing prosperous development in recent years. 1-5 A recent statistical study reveals that over 20 000 research papers relating to IEMs have been published since 2001.6 IEMs are widely used in many applications, including but not limited to fuel cells,7-10 electrodialysis, 11-13 and reverse electrodialysis. 14-16 Generally, IEMs are classified into two categories: cation exchange membranes (CEMs) and anion exchange membranes (AEMs). 17-19 In order to meet the growing requirements for IEMs, many synthesis methods have been developed.20-24 Among these methods, solution casting is a very popular one. This method involves casting a polymer solution onto a film, followed by phase inversion to remove the solvent and get solid membranes. 25-28 However, there are several disadvantages of using this method. Firstly, harmful organic solvents such as dimethylformamide (DMF) and dimethylacetamide (DMAc) are often used in this method, and therefore post-disposal of these solvents is inevitable. Secondly, when ion exchange resins are added to the solution, additional procedures such as sonication are usually necessary so as to disperse the resins uniformly and to avoid aggregation of resins. Also, the amount of resins that can be added is usually limited. As a result, the ion exchange capacity (IEC) of the prepared membranes will be affected. Thirdly, the mechanical properties of the membranes are usually not robust and the membranes can be easily broken if bent. Another common method for preparing IEMs with different properties is radiation induced grafting.29-31 However, with this method, high-energy radiation is usually indispensable. In addition, organic solvents are usually required. Furthermore, in some procedures, strong acids are used to introduce functional groups. The sol-gel method is an emerging method to introduce inorganic particles into organic polymers. 32,33 However, it is difficult to control the dispersion of inorganic particles, and therefore the membrane performance will be compromised. This method also inherits the issues and challenges of the solution casting method.

In our previous work, CEMs with high IEC were synthesized using a porous support.<sup>34</sup> Similar to other common polymerization methods for membrane preparation, concentrated sulfuric acid is used to introduce sulfonate groups into the membranes. Since the acid waste is harmful to the environment, sodium hydroxide is needed to neutralize the waste. A similar method which also uses porous supports is pore filling.<sup>35,36</sup> However, with this method, organic solvents and/or sulfonation are usually needed, and therefore post-treatment of solvents and acid wastes is necessary.<sup>37,38</sup> In addition, in some cases, in order to induce crosslinking, additional procedures such as annealing treatment are necessary.<sup>39,40</sup>

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 $<sup>\</sup>dagger$  The raw experimental data and additional images are available from the open repository DOI: 10.5281/zenodo.1622251

 $<sup>\</sup>ddagger$  Electronic supplementary information (ESI) available: Solubility study of LiSS in green solvents, membrane thickness increase in the  $M_{CDX}$  series, theoretical derivations of ion exchange capacity, ED test details, and large-scale production and yield of membranes. See DOI: 10.1039/c9ta06248c

In this work, a new methodology is developed to synthesize high performance CEMs in an effective and environmentfriendly way (Scheme 1). This methodology includes avoiding the use of organic solvents and strong acids, reducing the use of unnecessary chemicals, and realizing rapid synthesis via optimizing preparation procedures. To achieve the above goals simultaneously, in situ polymerization, emulsification, functional monomers, and porous supports are combined together for the first time. To be specific, a monomer containing the functional sulfonate group (e.g., lithium p-styrenesulfonate) is used so sulfonation is not needed. Also, water is used to dissolve the monomer so no organic solvents are needed. The detailed process of choosing water as the solvent is provided in Section 1 of the ESI.‡ In order to homogenize the mixture which mainly contains water, the functional monomer, and the crosslinking agent (e.g., divinylbenzene), a small amount of eco-friendly emulsifier is used. The porous support is used to absorb the liquid mixture and also to ensure the mechanical strength of the membranes. The polymerization reaction takes place within and on the porous support. This effective methodology may open up a new world of possibilities for scalable synthesis of IEMs in an eco-friendly way.

# **Experimental**

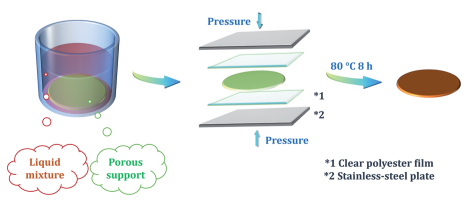
#### Chemicals and materials

Lithium p-styrenesulfonate (LiSS) was provided by Tosoh Organic Chemical. Divinylbenzene (DVB), styrene, benzoyl peroxide (BPO, brand Luperox® A75), sodium chloride, phenolphthalein solution, cation exchange resin (CER, brand Amberlyst® 15 hydrogen form) and emulsifier (brand Kolliphor® EL) were supplied by Sigma Aldrich. Sodium hydroxide and hydrochloric acid were supplied by VWR International. As the porous support, nonwoven fabric Novatexx 2471 was purchased from Freudenberg Filtration Technologies. As provided by the supplier, Novatexx 2471 is a thermal bonding material with a weight of 85 g m<sup>-2</sup>, thickness of 0.18 mm, air permeability of 150 L m<sup>-2</sup> s<sup>-1</sup> at 200 Pa, maximum tensile strength of 270 N/5 cm (machine direction) and 170 N/5 cm (cross direction), and elongation of 25% (machine direction) and 30% (cross direction) at maximum tensile strength. As the spacer for synthesis, clear and flat

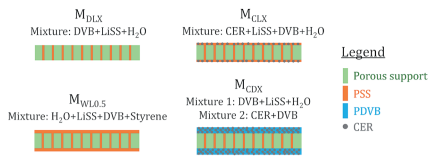
polyester films (model PMX727 with a thickness of 125 microns) were provided by HiFi Industrial Film.

#### Membrane preparation

Membrane preparation procedures are demonstrated in Scheme 1. The general procedures are described as follows. Firstly, LiSS was dissolved in water. Next, a certain amount of DVB or styrene was added into the solution. Then, a small amount of emulsifier was added into the mixture to make the mixture uniform. Then, a certain amount of CER was added into the mixture. Next, a small amount of BPO was added into the mixture. Then, the porous support was immersed into the mixture to adequately absorb the mixture. Then, the support was taken out of the mixture and placed between two clear polyester films, and then the films containing the support were placed between two stainless-steel plates. Finally, the stainlesssteel plates were fastened and placed in an oven to activate the polymerization process. The reaction temperature and time were set to 80 °C and 8 hours, respectively. After polymerization, the membranes were peeled off from the polyester films. As shown in Scheme 2 and Table 1, four series of membranes were prepared. For the  $M_{DLX}$ , the subscript DLX means that the mass ratio of DVB to LiSS is X where X is a number. Different DVB/ LiSS ratios (i.e., X) were studied, including 0.2, 0.4, 0.5, 0.6, 0.8, and 1.0. Meanwhile, the mass ratio of water to LiSS was kept the same, namely 1.2. For the  $M_{CDX}$ , the subscript CDX means that the mass ratio of CER to DVB is X. Different CER/DVB ratios (i.e., X) were studied, including 0.2, 0.4 and 0.6. Meanwhile, the mass ratio of DVB: water: LiSS was kept the same, namely 0.6:1.2:1.0. For the M<sub>CLX</sub>, the subscript CLX means that the mass ratio of CER to LiSS is X. Different CER/LiSS ratios (i.e., X) were studied, including 0.2, 0.4 and 0.6. Meanwhile, the mass ratio of DVB: water: LiSS was kept the same, namely 0.6:1.2:1.0. For the M<sub>WLX</sub>, the subscript WLX means that the mass ratio of water to LiSS is X. Different water/LiSS ratios (i.e., X) were studied, including 0.5 and 1.2. Meanwhile, the mass ratio of DVB and styrene to LiSS to water was kept the same, namely 0.5. It should be noted that MWL1.2 was replaced with M<sub>DL0.5</sub> because they were actually identical. Also, the procedures to prepare the MCDX series were slightly different compared with the other three series. To be specific, for the



Scheme 1 Schematic illustration of membrane preparation procedures.



Scheme 2 Schematic illustration of membrane cross-sections. M is short for the membrane; LiSS is short for lithium styrene sulfonate; DVB refers to divinylbenzene; PSS refers to polystyrene sulfonate; CER refers to cation exchange resin; PDVB refers to polydivinylbenzene.

Table 1 Summary of the acronyms used to describe different membrane series<sup>a</sup>

M series	X meaning	X values	Other parameters
$\mathrm{M}_{\mathrm{DL}X}$	DVB/LiSS ratio	0.2, 0.4, 0.5, 0.6, 0.8, 1.0	Water/LiSS ratio was 1.2
$M_{\mathrm{CD}X}$	CER/DVB ratio	0.2, 0.4, 0.6	DVB: water: LiSS was 0.6:1.2:1.0
$M_{CLX}$	CER/LiSS ratio	0.2, 0.4, 0.6	DVB: water: LiSS was 0.6:1.2:1.0
$M_{WLX}$	Water/LiSS ratio	0.5, 1.2	(DVB + styrene)/LiSS ratio was 0.5

<sup>&</sup>lt;sup>a</sup> M is short for the membrane; DVB is short for divinylbenzene; LiSS is short for lithium p-styrenesulfonate; CER is short for cation exchange resin.

M<sub>CDX</sub> series, two different mixtures were used together for synthesis. The first mixture was mainly composed of DVB, water and LiSS. The second mixture was mainly composed of CER and DVB. The main polymerization reactions occurred during membrane synthesis are shown in Scheme 3.

#### Membrane characterization

Thicknesses of the synthesized membranes were measured using a micrometer. Chemical structures of the synthesized membranes and corresponding materials were investigated via a PerkinElmer Spectrum-100 Fourier-transform infrared spectroscopy (FTIR) spectrometer. IEC was determined using the titration method.20 Briefly, membrane samples were firstly immersed in 1 M HCl solution to convert the Li<sup>+</sup> form into the H<sup>+</sup> form. Next, the samples were washed thoroughly with water and then immersed in 2 M NaCl solution to convert the H<sup>+</sup> form into the Na<sup>+</sup> form. Next, the membrane samples were taken out from the solution and washed with water, and the water was then added into the solution. Next, the solution was titrated using 0.01 M NaOH solution. IEC was calculated using the following equation: IEC =  $C \times V/W$  where C is the concentration of NaOH solution, V is the volume of NaOH solution consumed during titration, and W refers to the dry weight of samples. The loading ratio (LR) was one of the most important parameters in this study. As its name implies, the LR was used to evaluate the amount of functional materials loaded onto and into the porous support after polymerization. To calculate the LR, the weight of the synthesized membrane samples (denoted as  $W_1$ ) and the weight of the corresponding porous support (denoted as  $W_2$ ) were recorded. Then, the following equation could be used to calculate the LR: LR =  $(W_1 - W_2)/W_2$ . As the porous support contained no functional groups (i.e., sulfonate groups), the loaded functional

materials were the only source of sulfonate groups. Obviously, the LR had a direct effect on IEC. To measure the water uptake (or salt solution uptake) of the synthesized membranes, the samples were firstly immersed in 2 M NaCl solution for 24 h, and then excess water on the sample surface was removed, and then the wet weight of the samples was measured using a balance. After that, the samples were dried in an oven at 60 °C. Water uptake was equal to the difference of wet weight and dry weight, divided by the dry weight of the sample. Water contact angles of the synthesized membranes were measured using a standard ramehart goniometer (model 250-U1). Membrane morphology and sulfur distribution were studied using a Jeol JSM-6400 scanning electron microscope (SEM) in combination with an energydispersive X-ray spectrometer (EDS or EDX). Membrane thermal properties were evaluated via thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC). The membrane samples were tested using a Netzsch STA instrument (model 449 F5 Jupiter®). Under a nitrogen flow of 20 cm<sup>3</sup> min<sup>-1</sup>, the samples were firstly heated to 170 °C followed by an isothermal process for 10 min, and then cooled down to 30 °C followed by another isothermal process for 10 min, and then heated to 650 °C. Both the heating rate and cooling rate were kept at 10 °C min<sup>-1</sup>. The second DSC heating curve was used.

#### Membrane desalination performance

Desalination performance measures the efficiency membranes in removing the ions from the salt solution. Desalination performance can be quantified using the following equation:

$$DP = \frac{\left(D_{C_0} - D_{C_t}\right) \times V_D}{t} \tag{1}$$

Scheme 3 Polymerization reactions during membrane synthesis. Synthesis of M<sub>DLX</sub> and M<sub>CLX</sub> mainly involved reaction (a); synthesis of M<sub>WL0.5</sub> mainly involved reaction (c); synthesis of M<sub>CDX</sub> mainly involved reactions (a) and (b). The above reaction equations are for demonstration purposes only, as the ratios of different reactants are not considered in these equations.

where DP is short for desalination performance;  $D_{C_0}$  is the initial salt concentration of the solution in the dilute compartment;  $D_{C_r}$  is the salt concentration of the solution in the dilute compartment after the desalination test with a duration of t; and  $V_D$  is the volume of salt solution in the dilute compartment, which remained unchanged during the desalination test.

Desalination performance was tested using a PCCell ED 64004 electrodialysis cell. Commercial membranes were purchased from PCA GmbH. Five pairs of membranes, including six CEMs and five AEMs, were used each time. In order to compare the performance of synthesized CEMs in this study to that of commercial CEMs, the same AEMs from PCA GmbH were used for all the tests. A BioLogic VSP potentiostat was used to supply power to the cell. Current density was constant during the desalination test. Sodium chloride solution was used in the concentrate compartment and dilute compartment for desalination demonstration. The volume of the concentrate compartment, dilute compartment and electrode compartment was 2.5 L, 2.5 L, and 4.0 L, respectively, and the flow rates were around 0.6 L min<sup>-1</sup>, 0.6 L min<sup>-1</sup> and 2.4 L min<sup>-1</sup>, respectively. The desalination test lasted for sixty minutes and the samples were collected every ten minutes. The conductivity of the samples was measured using a Cole-Parmer TDS meter (model C100). The concentration was calculated based on the relationship between conductivity and concentration (see Section 4 in the ESI‡ for more details).

# Results and discussion

## Membrane synthesis and thickness

The effect of membrane thickness on membrane properties is complicated. An increase in membrane thickness indicates that more functional materials are loaded into and onto the membrane support. When a membrane contains more functional materials (e.g., sulfonate groups for CEM), the desalination efficiency could be increased because more salt ions can pass through the membrane (e.g., sodium ions for CEM) per unit time. Nevertheless, when the membrane thickness is

increased, the membrane resistance will also increase. As a result, the desalination efficiency will be reduced. This contradictory result reveals that membrane thickness has a trade-off relationship with membrane properties. As shown in Fig. 1, the synthesized membranes exhibited different membrane thicknesses under different preparation conditions. When the other conditions were kept the same, DVB/LiSS ratios (i.e., the  $M_{DLX}$  series) did not have an obvious effect on membrane thicknesses (Fig. 1a). This was due to the low loading ratios of functional materials in the membranes. When the porous support was immersed in the liquid mixtures, water competed with functional monomers because of the limited space in the support. As the total liquid uptake capacity of the support did not change much, the loading of functional monomers was affected. During polymerization, water was evaporated gradually, which made more space available. At the same time, the monomers underwent polymerization. However, due to the pressure applied onto the support, most of the liquid mixture on the support was removed before the reaction. As a result, very little monomer was available to fill in the space left by water. Therefore, the loading ratio of functional materials was reduced. Furthermore, the pressure applied on the reactor also showed a negative effect on membrane thickness. Although the loaded functional materials had a positive effect on membrane thickness, the negative effect of the pressure was greater than the positive effect of functional materials when the loading ratio was low. As a result, some synthesized membranes showed decreased thicknesses compared to the thickness of the pure membrane support (Fig. 1).

When cation exchange resin (CER) was introduced, it had a positive effect on membrane thickness. However, its effect was different in different situations. For example, when the ratios of LiSS, water, and DVB were kept the same, CER/LiSS ratios (i.e., the M<sub>CLX</sub> series) had a moderate impact on membrane thicknesses, especially when the CER/LiSS ratio was low (Fig. 1b). Nevertheless, when CER was mixed with DVB, the resulting mixture became very viscous. Moreover, the viscosity increased with increasing the CER/DVB ratio. As a result, the mixture had very poor mobility. When the mixture was applied onto the

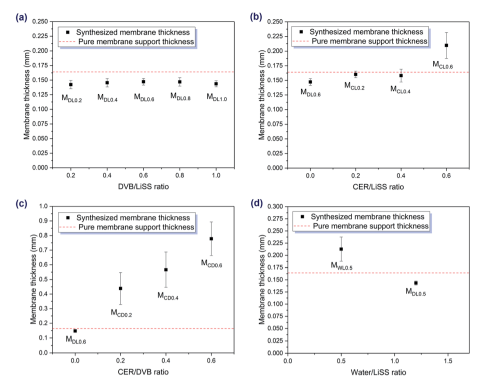


Fig. 1 Thicknesses of different membrane series. (a) Membrane thicknesses under different DVB/LiSS ratios. (b) Membrane thicknesses under different CER/LiSS ratios. (c) Membrane thicknesses under different CER/DVB ratios. M<sub>DL0.6</sub> from (a) is shown in (b) and (c) again for comparison. (d) Membrane thicknesses under different water/LiSS ratios.

support surface, it could not spread out. Even when pressure was applied, the thickness of the mixture did not reduce much. As a result, the synthesized membranes (i.e., the  $M_{CDX}$  series)

showed a dramatic increase in thickness compared to the pure support (Fig. S7‡). Moreover, the higher the CER/DVB ratio, the thicker the membranes (Fig. 1c). Besides, it was interesting to

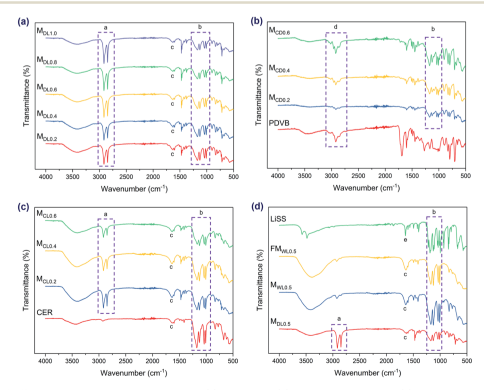


Fig. 2 FTIR spectra of different membrane series and materials. (a)  $M_{DLX}$  series. (b)  $M_{CDX}$  series and PDVB. (c)  $M_{CLX}$  series and CER. (d) LiSS,  $M_{DL0.5}$ ,  $M_{WL0.5}$ , and  $FM_{WL0.5}$ . PDVB refers to polydivinylbenzene; FM refers to the functional material in the membranes.

point out that when CER was introduced into DVB, its color changed to black, so possibly this was not a simple physical mixture. In other words, it was possible that unwanted chemical reactions happened after the two chemicals were mixed together. Therefore, one interesting future research topic is to investigate the interactions between CER and DVB.

The water/LiSS ratio also showed an effect on membrane thickness. As shown in Fig. 1d, reducing the water/LiSS ratio could increase membrane thickness. The reason was that, as water usage was reduced, more space was occupied by functional monomers, and therefore the loading ratio was increased. At the same time, lowering the water ratio also made the mixture more viscous, and thus the pressure applied on the support was less effective in removing the mixture on the surface. As a result, after polymerization, more functional materials were loaded into and onto the support, and therefore the membrane thickness was increased.

#### Membrane chemical structure

Chemical structures of the membranes were investigated by FTIR. The broad concave shape with the shape center at around  $3410 \pm 16 \text{ cm}^{-1}$ , in combination with another peak at  $1632 \pm 3$ cm<sup>-1</sup> (denoted as c in Fig. 2a, c and d), indicated the presence of water in these samples. In contrast, in the spectra of the  $M_{CDX}$ series, these two peaks were not prominent, indicating the low amount of water. This corresponded well to the following water uptake analysis, as the MCDX series exhibited much lower water uptake compared to the other membrane series.

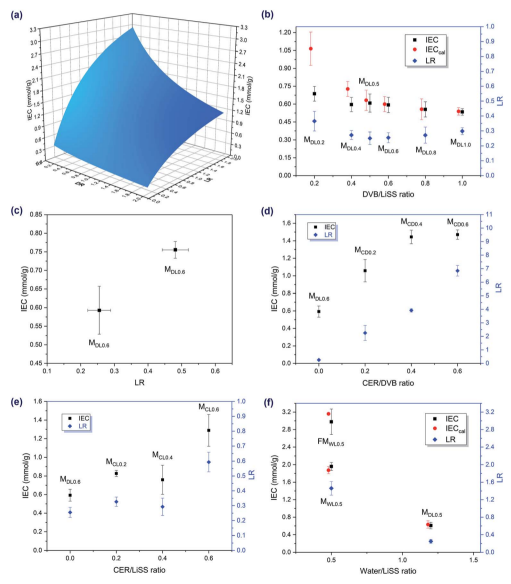


Fig. 3 IEC and LR of different membrane samples and materials. (a) Theoretical effect of DR and LR on IEC. (b) IEC and LR under different DVB/ LiSS ratios. (c) Effect of LR on IEC. (d) IEC and LR under different CER/DVB ratios. (e) IEC and LR under different CER/LiSS ratios. (f) IEC and LR under different water/LiSS ratios. FM refers to functional material in the membranes. The black square points correspond to experimental IEC values and the red circle points correspond to theoretical IEC values.

The peaks at  $1173 \pm 2 \text{ cm}^{-1}$ ,  $1127 \pm 1 \text{ cm}^{-1}$ ,  $1039 \pm 2 \text{ cm}^{-1}$ , and  $1009 \pm 1 \text{ cm}^{-1}$  (denoted as b in Fig. 2) were characteristic peaks of the sulfonate group (salt form). As discussed above, the thicknesses of the M<sub>CDX</sub> series were much higher compared to the thickness of the pure membrane support. As a result, the mass fraction of polydivinylbenzene (PDVB) and the cation exchange resin (CER) in the membranes was much higher than that of the support. That is why these membranes did not show characteristic peaks of the support. Instead, the spectra of these membranes exhibited spectra characteristics of PDVB (denoted as d in Fig. 2b) and CER.

Besides, the two peaks around 2916 cm<sup>-1</sup> and 2849 cm<sup>-1</sup> (denoted as a in Fig. 2a, c and d) were characteristic peaks of the membrane support. Furthermore, M<sub>WL0.5</sub> and FM<sub>WL0.5</sub> exhibited very similar FTIR spectra to each other, indicating the high ratio of functional materials in the membranes. This further revealed the high IEC of the membranes, as discussed below. In addition, these spectra agreed well with those of cross-linked sulfonated polystyrene salts, indicating the successful polymerization of LiSS. Besides, the successful polymerization of LiSS could also be verified via the spectrum of LiSS. To be specific, the sharp peak of the LiSS spectrum at 1647 cm<sup>-1</sup> (denoted as e in Fig. 2d) revealed the presence of the vinyl group. This sharp peak disappeared after the reaction, which indicated that the vinyl group was involved in the reaction.

# Ion exchange capacity and loading ratios

Ion exchange capacity (IEC) is one of the most important properties of IEMs. In this work, preparation conditions demonstrated a great influence on the IEC of the synthesized membranes, mainly by influencing the loading ratio of

functional materials (denoted as LR below) and also the IEC of loaded functional materials. To be more specific, the IEC of the synthesized membranes could be increased by increasing the LR and/or IEC of loaded functional materials. Generally, the LR could be controlled by controlling the immersion time of the porous support in the liquid mixture and/or by controlling the compositions of the liquid mixture. For example, the M<sub>CDX</sub> series demonstrated very high loading ratios compared to the other membrane series. The reason for this was that the mixture of CER and DVB was very viscous. As a result, more CER/DVB mixture was loaded onto the membrane support. Moreover, the IEC of loaded functional materials had a positive relationship with the amount of sulfonate groups in the materials. Therefore, the IEC of loaded functional materials could be increased by increasing the amount of sulfonate groups in it. If assuming all water was removed during membrane preparation and all water-soluble substances were removed during conversion, then the IEC of the synthesized membranes was affected by two parameters, including the LR and mass ratio of DVB and styrene to LiSS in the liquid mixture (denoted as DR below). The following equation was obtained to quantitatively describe the influence of the LR and DR on IEC. Also, Fig. 3a demonstrates this quantitative relationship. Detailed derivation procedures for the following equation are provided in Section 3 of the ESI.‡

$$IEC = \frac{1000 \times LR}{(190.15 \times DR + 206.19) \times (1 + LR)}$$
 (2)

As shown in Fig. 3b, under the same preparation conditions, the LR did not seem to be affected by the DVB/LiSS ratio in the mixture. Furthermore, when the LR was about the same, the

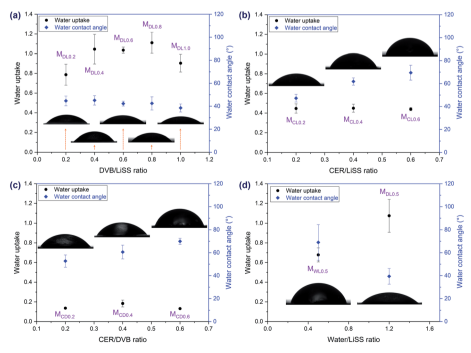


Fig. 4 Water uptake and contact angles of different membrane series. (a) Water uptake and contact angles under different DVB/LiSS ratios. (b) Water uptake and contact angles under different CER/LiSS ratios. (c) Water uptake and contact angles under different CER/DVB ratios. (d) Water uptake and contact angles under different water/LiSS ratios

DVB/LiSS ratio revealed a negative relationship with IEC, though the relationship was not prominent. At high DVB/LiSS ratios, experimental IEC values corresponded well to theoretical IEC values. At low DVB/LiSS ratios, some deviations were observed. This was possibly due to the fact that eqn (2) was obtained based on the assumption that all water was evaporated during the reaction. However, when the DVB/LiSS ratio was low, the amount of water used for synthesis was high, and therefore the above assumption was no longer valid. In other words, the effect of water could not be ignored at low DVB/LiSS ratios. Also, the IEC values of the membrane samples in Fig. 3b are not high, due to the low LRs. One possible way to increase the LR was to increase the amount of liquid mixture in and on the membrane support before the reaction. In addition, as revealed by Fig. 3c,

when the other conditions were the same, the LR had a positive relationship with IEC, which agreed well with theoretical predictions (Fig. 3a).

As revealed by Fig. 3d and e, cation exchange resin (CER) had a positive effect on IEC. Moreover, compared to the other membrane series, increasing the CER/DVB ratio could also dramatically increase the LR (Fig. 3d). As a result, IEC was also increased. In addition, the effect of CER/LiSS ratio on the LR and on the IEC showed a similar trend, due to the positive relationship between the LR and IEC (Fig. 3e). Though CER could increase IEC, it was possible to prepare membranes with comparable or even higher IEC without using CER. One strategy to prepare high IEC membranes without CER was to reduce the amount of water used for dissolving LiSS in the mixtures

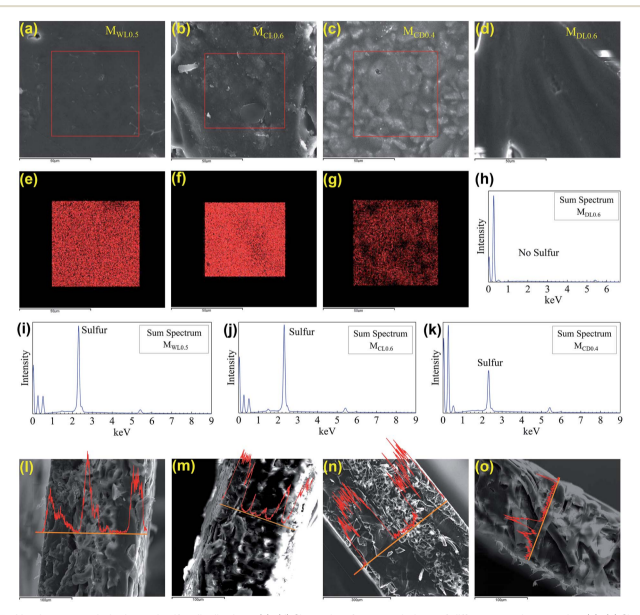


Fig. 5 Membrane morphologies and sulfur distributions. (a)-(d) Showed surface morphology of different membrane series. (e)-(g) Showed surface sulfur distribution (red color) of different membrane series. (h)-(k) Showed the corresponding energy spectrum diagrams of different membrane series. (i)–(o) Showed cross-section morphology of different membrane series. The red curves showed sulfur distribution at a certain position across the membranes (yellow line).

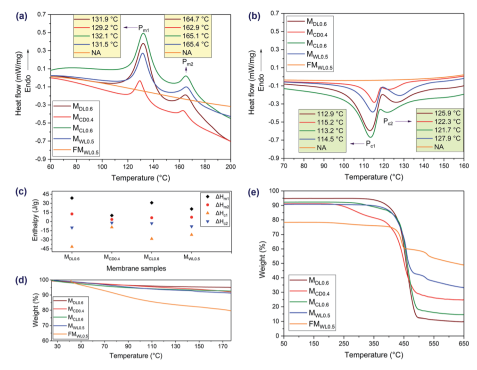


Fig. 6 Thermal properties of different membrane series and materials. (a) DSC heating curves of different membrane series and materials.  $P_{m1}$  and  $P_{m2}$  refers to the first and second melting peak temperature, respectively. (b) DSC cooling curves of different membrane series and materials.  $P_{c1}$  and  $P_{c2}$  refers to the first and second crystallization peak temperature, respectively. (c) Melting and crystallization enthalpies of different membrane series.  $\Delta H_{m}$  and  $\Delta H_{c}$  refers to melting and crystallization enthalpy, respectively. (d) TGA curves of different membrane series and materials (first heating). (e) TGA curves of different membrane series and materials (second heating).

(Fig. 3f). When the water amount was reduced, more functional polystyrene sulfonates could be formed in and on the support, and therefore the LR and IEC also increased. This also agreed well with theoretical calculations.

### Water uptake and contact angles

Water uptake and contact angles were another two parameters revealing electrochemical properties of CEMs. As shown in Fig. 4, all the membrane samples exhibited hydrophilicity but varied in degree. The  $M_{\mathrm{DL}X}$  series demonstrated the highest water uptake and the lowest contact angle compared to the other series. In contrast, due to the thick PDVB layer formed on the support, the  $M_{\mathrm{CD}X}$  series exhibited the lowest water uptake among all the series.

#### Membrane morphology and sulfur distribution

As revealed by Fig. 5a–d, different series of membrane samples exhibited different surface morphologies. As the sulfonate group played an important role in determining IEC and membrane performance, its distribution within the membranes were investigated using the sulfur mapping technique. As shown in Fig. 5e, f, i, and j, both  $M_{WL0.5}$  and  $M_{CL0.6}$  showed rich and uniform sulfur distribution on the membrane surface. As shown in Fig. 5c, the surface of  $M_{CD0.4}$  was rough. To some extent, this uneven surface increased the difficulty to detect sulfur. As a result, the sulfur distribution of  $M_{CD0.4}$  was not as uniform as those of  $M_{WL0.5}$  and  $M_{CL0.6}$ . In contrast, as shown in Fig. 5h,

M<sub>DL0.6</sub> did not show sulfur distribution on the surface due to the low loading ratio of functional materials in the membranes compared to those of the other membrane series. To summarize, the surface sulfur distribution of different membranes shown in Fig. 5 agreed well with the above IEC analysis. In addition, as shown in Fig. 5l-o, sulfur distributions of the membrane cross-sections were not uniform, which was due to the porous structure of the membrane support. For example, the cross-section of M<sub>CD0.4</sub> clearly showed three layers, due to the high loading ratio. The outer two layers did not contain the membrane support while the inner layer contained both membrane support and functional materials. Besides, it seemed that there were some "pores" in the membranes, as shown in Fig. 5l-o. However, it should be clarified that they were not pores. In fact, they were

Table 2 Summary of synthesized membranes and commercial membranes for the desalination test

	Synthesized membrane	Commercial membrane
Supplier	_	PCA GmbH company
ID	$ m M_{WL0.5}$	PC SK
Type	CEM	CEM
Charged group	Sulfonate group	Sulfonate group
Width (cm)	11	11
Length (cm)	11	11
Thickness (mm)	$0.21\pm0.02$	$\textbf{0.18} \pm \textbf{0.02}$

cross-sections of the membrane support, which had a fiber shape. To conclude, the synthesized membranes were semihomogenous, where the fibers contained no sulfonate groups while the functional materials had uniform distribution of sulfonate groups.

#### Membrane thermal properties

As revealed by Fig. 6a, the membranes showed two melting peaks, which were characteristic peaks of the membrane support. Also, all the peaks were there, only shifting slightly. Furthermore, as revealed by the TGA curve (Fig. 6e), during the second heating process, there was no mass loss below 200 °C. These phenomena indicated that no chemical reactions happened to the porous support during membrane preparation. In addition, the membranes also showed two crystallization peaks (Fig. 6b). In contrast, the functional materials did not show any melting or crystallization peaks at relevant temperatures, indicating their amorphous structure.41 Besides, many factors could cause shift of the melting peaks and crystallization peaks, such as the heating rates and cooling rates. 42,43 As revealed by Fig. 6c, M<sub>DL0.6</sub> had the highest melting enthalpies and crystallization enthalpies (absolute value, the same as below) while M<sub>CD0.4</sub> had the lowest melting enthalpies and crystallization enthalpies. The reason lied in that  $M_{\mathrm{DL}0.6}$  had the lowest loading ratio compared to those of the other series, while M<sub>CD0.4</sub> had the highest loading ratio (Fig. 3). In other words, the mass ratio of the porous support was highest in  $M_{\mathrm{DL}0.6}$  and lowest in M<sub>CD0.4</sub>. As a result, M<sub>DL0.6</sub> had the highest enthalpies while  $M_{\mathrm{CD0.4}}$  had the lowest enthalpies. As shown in Fig. 6d, the weight of the samples decreased gradually, which was due to the loss of absorbed water in the membranes. The sulfonate groups in the membranes had a positive role in retaining the water molecules in the membranes, so the release of water molecules was a continuing process.44 The following weight losses in Fig. 6e were due to the decomposition of the sulfonate group and degradation the polymer matrix. In addition, the functional material which had the highest sulfur amount showed the highest residual weight. In contrast, M<sub>DL0.6</sub> which had the lowest sulfur amount showed the lowest residual weight. One possible explanation was that during degradation, a highly thermostable sulfur-bridged polymer was formed.45

# Membrane desalination performance

Membrane desalination performance was demonstrated using M<sub>WL0.5</sub> due to its advantages over the other membrane series. Firstly, M<sub>CDX</sub> had high thicknesses compared to the other membrane series, which indicated that the membrane resistance could be very high. As a result, its desalination efficiency was very low. Secondly, M<sub>DLX</sub> had low IEC compared to the other membrane series, which would also affect its desalination efficiency. Thirdly, although MCLX did not show obvious disadvantages over the other membrane series, CER was used to improve its performance. In addition, it was much easier to get uniform

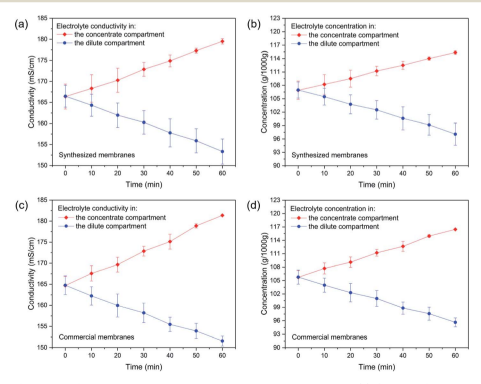


Fig. 7 Desalination performance of synthesized membranes and commercial membranes. (a) Change of electrolyte conductivity in the concentrate and dilute compartment using synthesized membranes M<sub>WL0.5</sub>. (b) Change of electrolyte concentration in the concentrate and dilute compartment using synthesized membranes  $M_{WL0.5}$ . (c) Change of electrolyte conductivity in the concentrate and dilute compartment using commercial membranes. (d) Change of electrolyte concentration in the concentrate and dilute compartment using commercial membranes.

sulfur distribution for  $M_{WL0.5}$  compared to that for  $M_{CLX}$ . Therefore, among all of the membrane series,  $M_{WL0.5}$  stands out.

The basic properties of the synthesized membranes and commercial membranes used for the desalination test are summarized in Table 2. The membranes have the same square shape with an active area of 64 cm<sup>2</sup> (see Section 4 in the ESI‡ for more details). As revealed by Fig. 7, in the desalination process, the conductivity and concentration of electrolyte (i.e., sodium chloride solution) in the concentrate compartment increased while the conductivity and concentration of electrolyte in the dilute compartment decreased. More importantly, the synthesized membranes showed high desalination performance, which was comparable to that of commercial membranes, even given that the synthesized membranes had a higher thickness compared to commercial membranes. Therefore, it is envisioned that when the membrane thickness is reduced while its high IEC is maintained, the desalination performance could be further increased. This is one interesting future research topic. Furthermore, it was possible to realize large-scale production of membranes using this method (see Section 5 in the ESII for more details). Admittedly, as this method is new, there are still many aspects which need further study, such as membrane stability study. Especially, future research should focus on improving the stability of M<sub>WLX</sub> in dilute solutions. Nevertheless, as green synthesis is the future of membrane synthesis, it is envisioned that this environment-friendly methodology could be widely used to synthesize other kinds of ion exchange membranes, and therefore could open up a new era of membrane synthesis.

# Conflicts of interest

There are no conflicts to declare.

# Conclusions

In summary, a new and effective methodology to prepare high performance cation exchange membranes was proposed and studied. Instead of using conventional organic solvents and concentrated sulfuric acid for sulfonation, water was selected as the solvent and also no sulfonation was needed. A series of membranes were synthesized using this methodology and different preparation conditions were investigated in order to optimize the synthesis parameters. Polymerization successfully took place on and within the membranes, and the support did not take part in the reactions. Also, when cation exchange resin was not used, ion exchange capacity (IEC) was determined by two factors, including the mass ratio of divinylbenzene and styrene to lithium p-styrenesulfonate, and the loading ratio of functional materials. In addition, water played an important role in membrane synthesis. IEC could be greatly increased by reducing the amount of water used for synthesis. More importantly, high IEC could be achieved without cation exchange resin, and therefore the synthesis procedure could be more efficient and the synthesized membranes could be more homogeneous. Finally, the synthesized membranes demonstrated high desalination performance. We envision that this

methodology may open up new possibilities for the synthesis of ion exchange membranes in an effective and environmentfriendly way. In order to further improve this methodology, more studies on membrane stability should be carried out in future research.

#### Acronyms

The acronyms below are listed in alphabetical order.

AEMs Anion exchange membranes

BPO Benzoyl peroxide

CEMs Cation exchange membranes
CER Cation exchange resin
DMAc Dimethylacetamide
DMF Dimethylformamide
DP Desalination performance

DR DVB and styrene to LiSS mass ratio
DSC Differential scanning calorimetry

DVB Divinylbenzene

EDS or EDX Energy-dispersive X-ray spectroscopy

FM Functional material

FTIR Fourier-transform infrared spectroscopy

IEC Ion exchange capacity
IEMs Ion exchange membranes
LiSS Lithium *p*-styrenesulfonate

LR Loading ratio
PDVB Polydivinylbenzene
PSS Polystyrene sulfonate

SEM Scanning electron microscope TGA Thermogravimetric analysis

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