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Preparation of freestanding graphene-based laminar membrane for clean-water intake via forward osmosis process†

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Here, a facile strategy is reported to prepare thin freestanding graphene oxide (GO) laminar membranes through thermally-induced chemical cross-linking process. The resultant freestanding cross-linked GO (F-CGO) membranes show controllable inter-laminar distance and hydraulic stability in aqueous media. Semi-permeability tests through a forward osmosis (FO) system show that the water permeation rate of the F-CGO membranes is up to $35.5 \text{ L m}^{-2} \text{ h}^{-1}$ when the draw solution is 2.0 mol L^{-1} NaCl solution. This value is much higher than that of standing cross-linked GO (S-CGO) membranes, which indicates that the internal concentration polarization (ICP) effect has been alleviated by the F-CGO membrane. Furthermore, clean-water was taken in from various wastewaters, but without any detectable residual pollutants by using the F-CGO membrane as a separator. This work demonstrates that the F-CGO membrane is promising for water purification applications through the FO process, especially for portable water purification systems utilized during emergency situations.

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Introduction

Recently, graphene oxide (GO) has emerged as an excellent membrane material, drawing huge attention from separation fields. Nair *et al.* demonstrated that sub-micrometre-thick laminar GO membranes allow for the un-impeded permeation of water, whereas they block everything else in the liquid or vapour form, and even gases including helium.¹ They attribute these anomalous observations to the low-friction monolayer flow of water through the two-dimensional capillaries formed by the non-oxidized regions of the GO sheets. These capillaries block other molecules because they are narrowed or reversibly clogged with water. This work was then immediately expanded to explore the penetration behaviour of ions through GO membranes. Joshi *et al.*² found that laminar GO membranes act as molecular sieves in aqueous solutions, blocking all solutes with a hydrated radii larger than 0.45 nm, while the permeation rates of smaller ions through these membranes are thousands of times faster than those in simple diffusions. For the same reason, the network of

nanocapillaries only accepts species that fit in, and a capillary-like high pressure acts on ions inside the capillaries, accelerating their diffusion behaviour.³ The manifested size-exclusion properties make GO laminar membranes promising materials for ion-selective separation,⁴⁻⁶ organic-water separation,⁷⁻⁹ water desalination¹⁰⁻¹² and purification,¹³⁻¹⁵ and so on. However, owing to the highly hydrophilic and negatively charged oxygen-containing groups of the GO sheets,¹⁶ the inter-layer distance (*d*-spacing) of laminar GO membranes will be enlarged, causing the possible disintegration of the two-dimensional capillaries when they are immersed in aqueous media for long time periods.¹⁷ That is why a pure GO laminar membrane cannot effectively perform its size-exclusion function during membrane separation processes due to its low mechanical stability.^{16,18,19} Therefore, to enhance their stability in aqueous systems, laminar GO membranes need to be modified by processes such as chemical cross-linking,²⁰⁻²² partial reduction^{23,24} and intercalation.²⁵

As an emerging membrane technology, forward osmosis (FO) has attracted much attention because of its wonderful advantages over other traditional membrane separation techniques, including low energy consumption,²⁶ low membrane fouling²⁷⁻²⁹ and low equipment requirement, and so on. Nevertheless, the application of FO technique still suffers from some problems, especially, internal concentration polarization (ICP),²⁶ which stems from the bilayer structure of FO membrane, thin selective layer and porous supporting layer.³⁰ The severe ICP effect during FO process would induce the

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driving force decrease, further reduce the membrane flux. In principle, FO process is based on the natural osmosis phenomenon, in which water is drawn across a semipermeable membrane from a feed solution into a draw solution by an osmotic pressure difference ($\Delta\pi$) between the two solutions, while the solutes are left behind in the feed solution. The FO membrane is just required to stand the osmotic pressure, which is much lower than membranes for reverse osmosis (RO), nanofiltration (NF), or ultrafiltration (UF) process. Therefore, freestanding FO membrane, without the supporting layer, has been proposed to eliminate ICP effect in recent.²³

Taking into consideration the excellent size-exclusion, water permeation, and outstanding mechanical strength properties of laminar GO membranes,^{31,32} they are promising materials for the fabrication of freestanding semipermeable membranes for FO processes. For instance, ultrathin laminar GO membranes (40–200 nm), reduced by hydrogen iodide (HI) vapor, exhibited a high enough tensile strength (170–270 MPa) for FO processes. In addition, the obtained freestanding reduced GO membranes showed a higher salt rejection and water permeation than commercial cellulose triacetate (CTA) FO membranes.²³ Sun *et al.* prepared a freestanding hybrid membrane consisting of anionic GO, cationic Co–Al (or Mg–Al) layered double hydroxide (LDH) that exhibited robust mechanical strength, and a highly selective charge-guided ion transport in concentration gradient-driven systems.²⁵

In this work, we report a facile strategy to prepare thin freestanding graphene-based laminar membrane through thermally-induced chemical cross-linking process, named as freestanding crosslinked GO (F-CGO) membranes. The sensitivity of water to the inter-laminar distance of the F-CGO membranes was evaluated by X-ray diffraction (XRD), and terahertz time-domain spectroscopy (THz-TDS). The salt rejection performance and ICP effects of these F-CGO membranes were investigated in a custom-made FO system, using NaCl solution as the draw solution and ultrapure water as the feed solution. In addition, the F-CGO membrane was used to intake clean-water from various wastewaters that contained heavy metal ions or organic dyes.

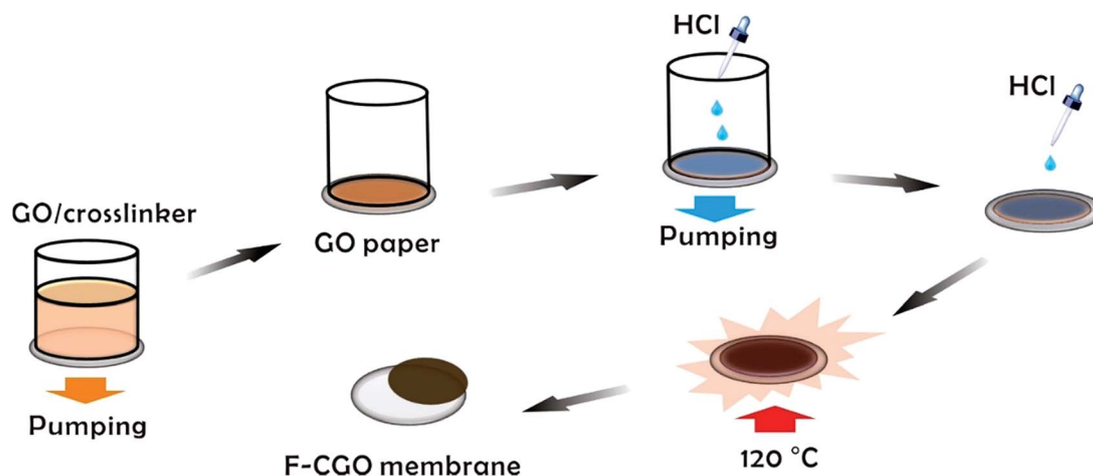
Experimental

Preparation of F-CGO membranes

The monolayer GO sheets were prepared using modified Hummer's method as reported in our previous works.^{33–35} Details are shown in the ESI.† The scheme of the F-CGO membranes preparation was shown as Scheme 1, and the details could be described as follows. 10 mL of GO (0.5 mg mL⁻¹)/PA (propane-dioic acid) mixtures, with different weight ratios of GO/PA, were prepared and mildly shaken for a few minutes. Then, these solutions were poured into the glass cup of a vacuum filter with a CA microfiltration membrane (pore size, 0.22 μm). When a filtration cake was formed on the CA membrane by continuous pumping, several droplets of HCl solution ($\sim 9.6 \text{ mol L}^{-1}$) were distributed on the cake, and then pump-filtrated until dry. After that, the CA membrane coated with GO sheets was taken off from the filter, and infiltrated with another several droplets of the HCl solution. Then, to make sure the cross-linking reaction was carried out, the GO sheets coated CA membrane was put in an oven at 120 °C for 30 min. The cross-linking reaction involves the esterification of the carboxylic acid on the PA with the hydroxyl groups of the GO sheets. After cross-linking, this standing cross-linked GO layer on CA membrane was named S-CGO membrane, while the F-CGO membrane was obtained by mechanically peeling off the cross-linked GO layer from the CA microfiltration membrane. As a contrast, a GO membrane without PA addition was also prepared by the same procedure and labelled as GO-120 membrane. The areal loading of GO sheets was about 0.35 mg cm⁻² in all of here prepared thin freestanding GO-based membranes.

Semi-permeation tests

The semi-permeation tests were carried out on a custom-made FO permeation cell as show in Fig. S1.† The permeation cell consisted of two glass compartments, separated by the F-CGO membrane, with a 19.6 mm² contact area. One compartment was filled with ultrapure water as the feed solution, and the other compartment was filled with an aqueous NaCl solution, at



Scheme 1 Schematic diagram of the F-CGO membranes preparation process.



a pre-set concentration, as the draw solution. The feed solution was circulated between the corresponding compartment of FO permeation cell and an Erlenmeyer flask using a peristaltic pump with a 45 mL min⁻¹ flow rate, and so was the draw solution. The initial volume of the feed solution was equal to that of the draw solution. The Erlenmeyer flask-stored draw solution was weighed on an electronic balance during the permeation tests, and this weigh was used to calculate the water permeation rate of the membrane using the following equation:

$$R_w = \frac{W_{t+1} - W_t}{S_m \Delta t}, \quad (1)$$

where R_w is the water permeation rate; W_t and W_{t+1} are the weight of the draw solution at sampling time points (t) and ($t + 1$), respectively; Δt is the time interval between two sampling points; and S_m is the effective contact area of the membrane between the two compartments of the permeation cell. At the same time, a few millilitres of the feed solution were sampled and determined by Inductively Coupled Plasma Atomic Emission Spectrometry (ICP-AES) to measure the reverse salt rate, which was calculated as follows:

$$R_s = \frac{C_{t+1}V_{t+1} - C_tV_t}{S_m \Delta t}, \quad (2)$$

where R_s is the reverse salt rate; C_t and C_{t+1} are the concentrations of Na⁺ at sampling time points (t) and ($t + 1$), respectively; V_t and V_{t+1} are the volume of the feed solution at sampling time points (t) and ($t + 1$), respectively. To have an indication of the membrane's Na⁺ rejection performance, we calculated the Na⁺ selective permeation coefficient across the membrane based on the water and Na⁺ ion permeation rates, as follows:

$$S = \frac{C_d V_d}{C_f V_f}, \quad (3)$$

where S is the selective permeation coefficient of Na⁺; C_d and V_d are the Na⁺ concentration and volume of draw solution after 4 h permeation, respectively; and C_f and V_f are the Na⁺ concentration and volume of feed solution after 4 h permeation period, respectively. Therefore, the rejection of Na⁺ could be calculated as follows:

$$R_j = \frac{S}{S + 1}, \quad (4)$$

where R_j is the Na⁺ rejection of the F-CGO membrane after 4 h permeation.

Clean-water intake from wastewaters

The setup and procedure of this test are similar to those of the semi-permeation test, except for the feed and draw solutions. For this test, the feed solution was composed of a 10 ppm methylene orange (MO), rhodamine B (RB), or Cu²⁺ aqueous solution; and the draw solution consisted of a 2.0 mol L⁻¹ NaCl solution. The water permeation rate in the different feed solutions was determined by measuring the weight change of the draw solution. The draw solution was monitored by UV-Vis spectroscopy (MO or RB feed solution), or with ICP-AES instruments, to detect the leaking pollutants during the test.

Results and discussion

Preparation of F-CGO membranes

The GO sheets prepared using modified Hummer's method, were monolayer and had a basal size of about 10 μm (Fig. S2†). PA was used as the cross-linker because of its acid-assisted esterification with the hydroxyl groups on the GO sheet.²¹ As the Scheme 1 shows, after HCl solution wetting and a 120 °C heat treatment, the GO-PA paper was easily released from the supporting cellulose acetate (CA) membrane (Fig. 1a). The obtained F-CGO membranes with different cross-linking degrees were prepared by adjusting the PA/GO weight ratio and labelled as F-CGO x membranes, where the x denotes the PA/GO weight ratio in the corresponding GO/PA mixture dispersion used for membrane preparation. The thermogravimetric analysis (TGA) curves of the F-CGO membranes show the weight loss from 130 °C to 210 °C was increased as the PA/GO weight ratio increasing (Fig. S3†), which confirms the cross-linking degree of F-CGO membrane increased, because of the poorer thermal stability of cross-linker than that of GO sheets. We also controlled the thickness of the F-CGO membranes by adjusting the dosage of the GO dispersion. As it is shown in the insert of Fig. 1a, the resultant F-CGO3 membrane exhibits semi-transparency, and the cross-sectional SEM image shows its thickness to be about 1 μm, with a noticeable regular laminar structure (Fig. 1b). Fig. 1c shows that the surface of the F-CGO membrane appears to have a ridge-like morphology, which can be explained by the unevenly stress applied on the GO sheets during their pumping-assistance assembly. This deduction is supported by the reverse surface morphology and optical microscopy images of the F-CGO membrane (Fig. S4†), in which the observed radial trends of the ridges reflect that the strong suction at the micro-pores of the supporting membrane resulted in the uneven assembly of the GO sheet.

The FT-IR spectra of different GO membranes show that the C=O (~1730 cm⁻¹) and C-O (1300–1050 cm⁻¹) absorption bands increased after cross-linking (Fig. 1d). This behaviour is explained by the esterification of the GO hydroxyl groups with the carboxylic acid of the PA.²¹ Investigating the C1s XPS spectra of the F-CGO membrane and GO paper after their drying at room temperature (25 °C) and 120 °C (labelled as GO-25 and GO-120, respectively), we found that the intensity of peak at ~284.6 eV (C-C/C-H) increased obviously, and the peak at ~288.8 eV (C=O) decreased after 120 °C treatment whether PA addition or not (Fig. 2), which demonstrates the GO sheets were partially reduced during the thermal treatment. Comparing with the GO-120 membrane, as shown in the Fig. 2c and d, the intensity of the peak at ~286 eV, assigned as the C-OH signal,³⁶ decreased sharply; while the intensity of the C-O-C=O/C-O-C (~287.2 eV)³⁶ peak increased. These results confirm that chemical cross-linking happened between the PA and GO sheets *via* esterification. Moreover, the higher area ratio of C-O-C=O/C-O-C signal in F-CGO3 membrane (Fig. 2d) indicates the (Fig. 2c) higher cross-linking degree than that of F-CGO1 membrane, which is evidenced by the pyrolysis behaviour of as-prepared F-CGO membranes from different PA/GO



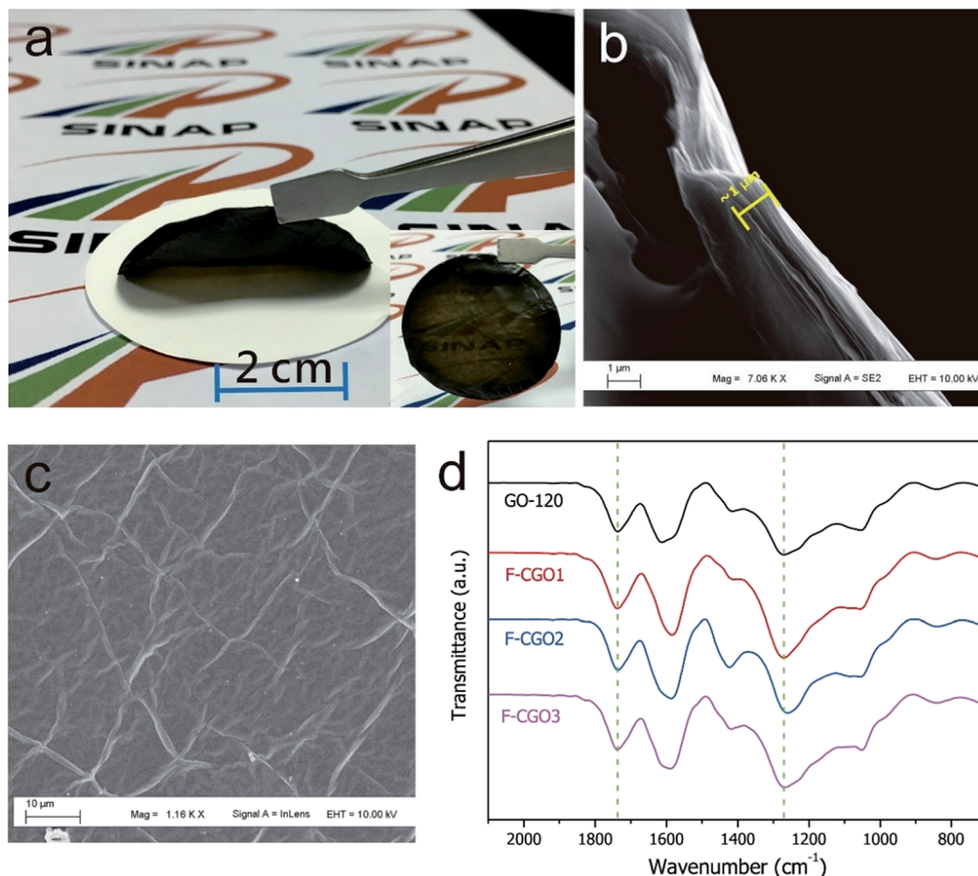


Fig. 1 (a) Digital pictures of the F-CGO3 membrane and its removal from the supporting membrane. (b) Cross-section and (c) surface micromorphology of the F-CGO3 membrane. (d) FT-IR spectra of the GO-120 membrane and different F-CGO membranes, which were prepared using different cross-linker dosages (the numbers denote the weight ratio of PA/GO in the mixture dispersion), which were tested on transmission mode. The scale bar in (b) and (c) was 1 μm and 10 μm, respectively.

dispersions (Fig. S3[†]). The Raman spectra of the GO-120 and F-CGO3 membranes show that the I_D/I_G ratio increased slightly after cross-linking, which is in accordance with that was previously reported, indicating that only small distortions were introduced on the laminar structure after cross-linking (Fig. S5[†]).³⁷ Interestingly, the I_{2D}/I_{S3} ratio was slightly increased from 0.609 of GO-120 to 0.694 of F-CGO3, where the S3 peak is a second-order peak due to the D–G combination, and 2D refers to the graphene 2D peak,³⁸ which indicates that less topological disorder within GO sheets in F-CGO3 membrane.³⁹ In other words, the defect density of GO sheets in the F-CGO3 membranes was further reduced during the thermal treatment at 120 °C with the assistance of cross-linker,³⁸ which was accompanied by a higher intensity of C–C/C–H peak compared to the GO-120 membrane (Fig. 2). Additionally, it would also indicate the cross-linking between adjacent GO sheets enhanced the structure order of GO-based membrane, which gives another reason for the light change of D peaks in Raman spectra after cross-linker introduction.

Water sensitivity and hydraulic stability

In Fig. 3a, the water contact angle of the pristine GO membrane (air-drying at 25 °C, GO-25) was ~40°. This value increased to

>70° after the membrane was oven-dried at 120 °C or cross-linked with PA, which indicates the partial reduction of GO sheets in the membranes after these treatments. This result is in accordance with the Raman spectra of the GO-120 and F-CGO3 membranes. The partial reduction and increased hydrophobicity of the F-CGO3 membrane could enhance its structural stability in aqueous media. The XRD patterns of the dry membranes (Fig. 3b and S6a[†]) show that the (002) diffraction peak increased from 10.2° (GO-25, d -spacing 0.861 nm) to 10.4° (GO-120, d -spacing 0.852 nm) after the heat treatment, indicating the achievement of a more compact laminar structure. It is noted that the diffraction peaks were further increased to 10.5° (d -spacing 0.844 nm), 10.8° (d -spacing 0.814 nm) and 10.9° (d -spacing 0.81 nm), as the PA/GO weight ratio increased. This suggests that the cross-linking between PA and GO does not expand, but narrow the interlayer distance instead. This behaviour could be explained that the small size of cross-linker molecule and low cross-linking degree of PA (as shown by the TGA curves in Fig. S3[†]), are unable to undermine the GO sheets stacking, which happens during the polymer decoration of GO sheets.³⁶ Through comparisons of the XRD patterns from the wet state of the membranes (Fig. 3b and S6b[†]), it was observed that the GO-25 diffraction broadens and its 2θ angle shifts to



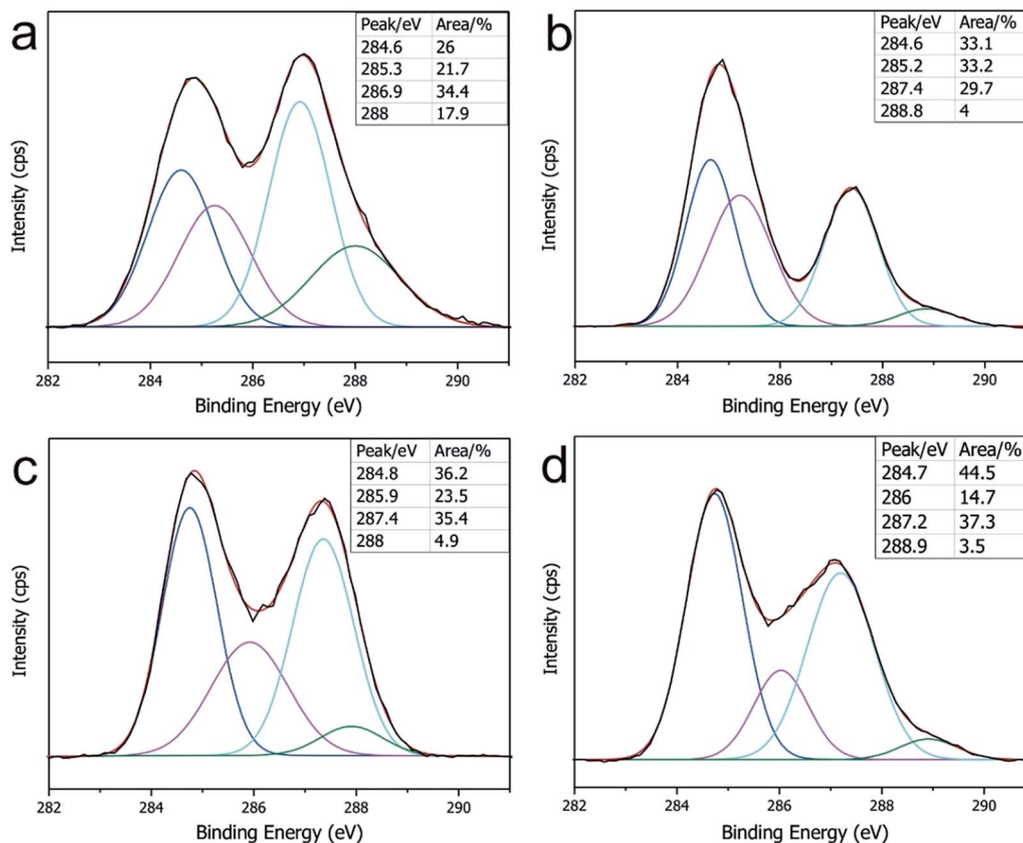


Fig. 2 The C1s XPS spectra of the (a) pristine GO membrane (GO-25), (b) GO-120 membrane, (c) F-CGO1 membrane and (d) F-CGO3 membrane.

6.6° (d -spacing 1.34 nm), while the other membranes still keep sharp peak curves, even though their 2θ angles shift to at least 7.0° (d -spacing 1.25 nm).

The effect of water on the structure of the membranes was monitored by terahertz time-domain spectroscopy (THz-TDS), which is a novel and promising tool to characterize the weak interactions of molecules, such as hydrogen bond and van der Waals force, because of its unique time-resolved feature and high signal to noise (SNR) in THz frequency range (wavelength in 0.03–3 mm).⁴⁰ As a polar liquid, water strongly absorbs THz radiation, which enables the THz-TDS to determine the absorption coefficient and refractive index of aqueous sample in THz range.^{41,42} The THz-TDS spectra of GO-25, GO-120 and F-CGO3 membranes containing different water contents were recorded and shown in Fig. 3c, d and S7a.† Due to the highly sensitive to water, the THz-absorbance of the GO-25, GO-120 and F-CGO3 membranes (Fig. S7b–d†) from 0.5 to 3.5 THz demonstrates a positive relationship with the water contents in the GO-25, GO-120 and F-CGO3 membranes. The THz-TDS spectra show that the transmission intensity of the dry F-CGO3 membrane is slightly lower than that of the dry GO-25 and GO-120 membrane because of the introduction of cross-linker into the F-CGO3 membrane, which would be THz-sensitive and thus bring about the transmittance decreasing of incident THz radiation. This result is very consistent with those

from FT-IR, Raman and TG analyses. A noticeable time delay can be observed in the THz-TDS spectra when the membrane was at different wet states. The time delay was obvious higher for the F-CGO3 membrane than that for the GO-25 membrane (Fig. 3c and d). In principle, the time delay is the function of the refractive index and thickness of sample material, and can be defined mathematically as follows:⁴³

$$\Delta t \approx \frac{(n_2 - n_1)d}{c}, \quad (5)$$

where Δt is the time delay degree; c is the speed of light; d is the thickness of the sample; and n_2 and n_1 are the refractive indexes of the material and background, respectively. Dry air was taken as the background in our experiment.

As eqn (5) shows, the time delay degree is positively related to the refractive index and thickness of the membrane. After being wetted with water, the refractive index and thickness of the GO membranes simultaneously increase. Given the higher water stability of the F-CGO3 membrane, the thickness of the wet F-CGO3 membrane should not be larger than that of the wet GO-25 membrane. Thereby, the refractive index of the wet F-CGO3 membrane should be larger than that of the wet GO-25 membrane. In fact, we calculated the refractive index of GO-25 and F-CGO3 membrane at dry and wet state by the TAS7400 system software and the arithmetic library provided from the THz instruments manufacture (Advantest



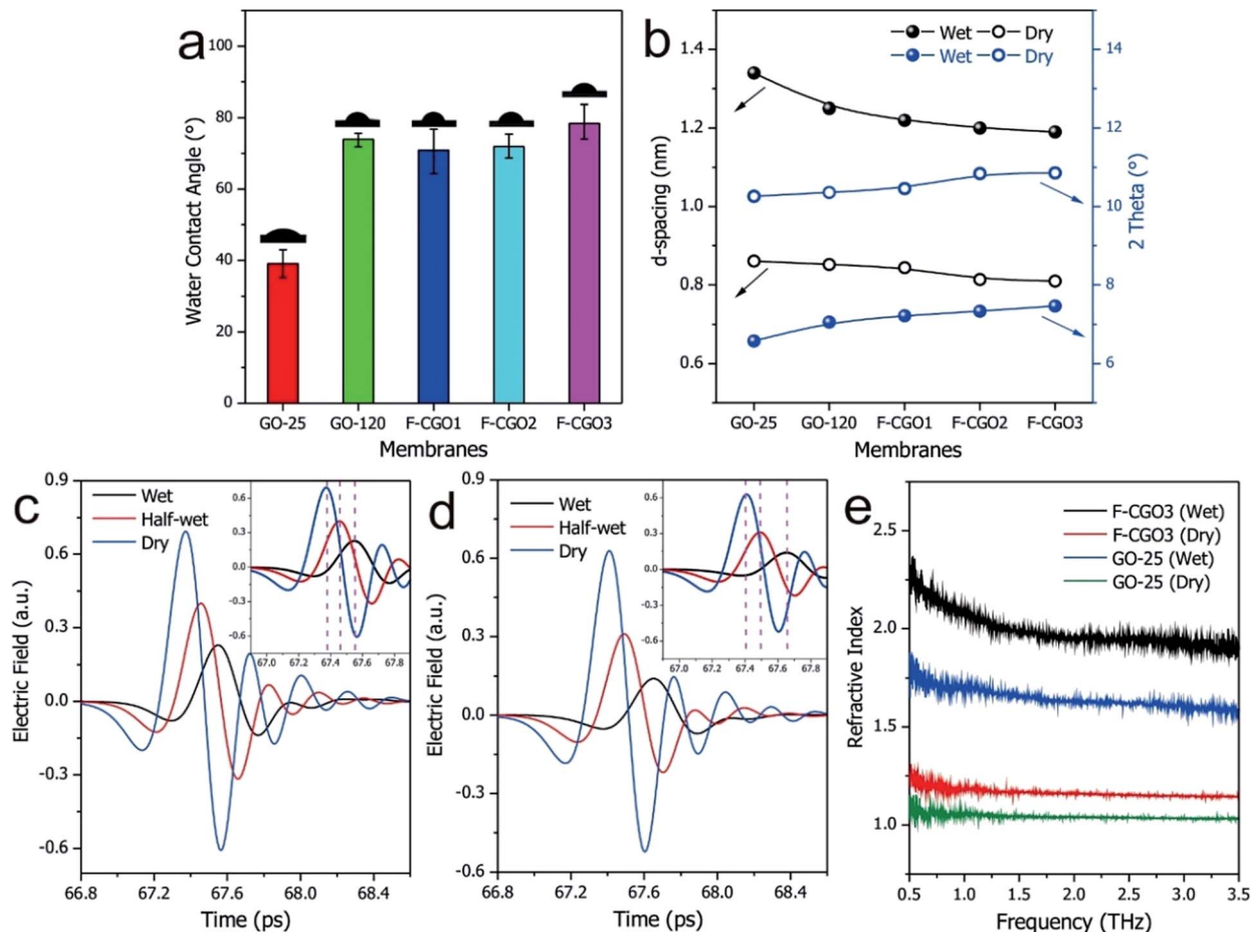


Fig. 3 (a) Water contact angle on different GO membranes surface; (b) comparison of the *d*-spacing of different GO membranes at the dry and wet states (immersed in water for 1 h); and THz-TDS spectra of (c) GO-25 and (d) F-CGO3 membranes at different wet states. (e) The refractive indexes of the GO-25 and F-CGO3 membranes at wet and dry states. *Dry state*: stored in 25 °C desiccator overnight; *wet state*: picked out from ultrapure water after immersing 1 h and blotted with paper; *half-wet state*: leaving in sample chamber of THz spectrometer for 10 min.

Corporation, Japan). As Fig. 3e shows, the refractive indexes of both GO-25 and F-CGO3 membranes are increased obviously after water wetting, but the refractive index of wet F-CGO3 membrane is really larger than that of wet GO-25 membrane. As we know, the refractive index is relevant for the material density.^{44,45} Therefore, our results indicate that the density of wet F-CGO3 membrane is higher than that of the wet GO-25, which is a result of the cross-linking treatment having enhanced the GO sheets stacking, and thus producing a more stable membrane in water.

The obtained F-CGO3 membrane was cut into 0.8×3.0 cm strips, and hung on the bracket of a retort stand. The strips exhibited good strength as they could withstand at least a 20 g weight and 2 g aids (Fig. S8†). To assess the hydraulic stability of the membranes visually, the maximum hydraulic pressure that the membranes could resist in the FO setup was also measured. The F-CGO membranes with 1 μm thickness (attached on a silicone pad with an aperture of 19.6 mm^{-2} , as shown in Fig. S9a†) were able to withstand static hydraulic pressures up to ~ 9 kPa, much higher than that of the GO-120 membrane (Fig. S10b†), which indicates the cross-linking treatment

enhanced the hydraulic stability of GO membrane obviously. It also was shown that the hydraulic stability of F-CGO membranes is proportional to their crosslinking degree, which means that the GO sheets in the membrane were tightly connected together by cross-linker, and formed a consubstantial entity. This result demonstrates the F-CGO membranes would be able to withstand osmotic pressure during FO process.

Semi-permeability of F-CGO membranes

Taking into consideration the high osmotic potential and nontoxicity of NaCl, it is an ideal and promising candidate as a hydration draw reagent for the FO setup. Here, we mainly investigated the Na^+ permeability of F-CGO membranes by using a custom-made osmotic cell, as shown in Fig. S1.† In this set-up, one side of the osmotic cell was ultrapure water (the feed solution), and the other side was the NaCl solution (the draw solution). Fig. 4a shows that when the concentration of the NaCl solution was 0.5 mol L^{-1} , the Na^+ permeation decreased as the cross-linking degree increased, which is in accordance with the *d*-spacing changes observed after cross-linking (Fig. 3b). We



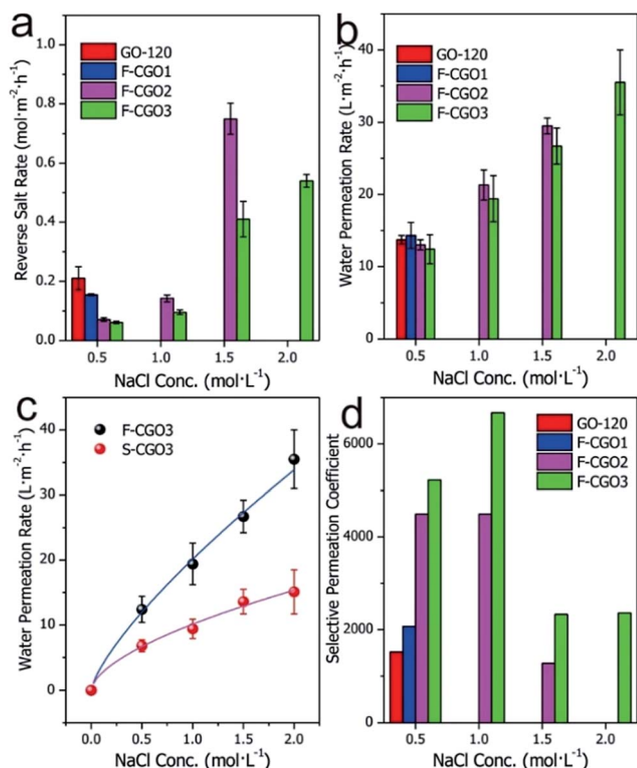


Fig. 4 (a) Reverse salt diffusion and (b) water permeation results of different GO membranes; (c) comparison between the water permeation of F-CGO3 and S-CGO3 membranes; (d) selective Na⁺ permeation coefficient across different GO membranes. The feed solution was ultrapure water, and the draw solution was a 2.0 mol L⁻¹ NaCl solution. The active layer of the S-CGO membrane faced the feed solution. The reverse salt diffusion test lasted 4 h, while the water permeation rate was obtained from a 0.5 h permeation period.

also found that F-CGO membranes with different crosslinking degrees exhibited different swelling durability in high salinity solutions. When a 1.0 mol L⁻¹ NaCl solution was used as the draw solution, the GO-120 and F-CGO1 membranes disintegrated after several minutes of testing. As the NaCl concentration increased to 2.0 mol L⁻¹, the F-CGO2 membrane also experienced disintegration. This behaviour is explained by the high osmotic pressure induced on the GO membranes by the highly concentrated draw solution. These results indicate that the higher crosslinking degree of F-CGO membranes not only hinders the diffusion of Na⁺, but it also provides the membrane with a good structural stability in highly saline water, which is in accordance with the static hydraulic pressure test results (Fig. S9b[†]).

Based on the comparison of Na⁺ permeation through cross-linked GO membranes with and without a supporting CA microfiltration membrane (Fig. S10[†]), it was found that the Na⁺ permeation of the F-CGO3 membrane was higher than that of the S-CGO3 membrane. Given the same structure and thickness of the active layer of the S-CGO3 membrane with that of the F-CGO3 membrane, the effect of the porous supporting membrane is accountable for the differences in the Na⁺ permeation. As previously mentioned, the porous supporting membrane decreases the osmotic gradient decrease because it

concentrates the solutes in the supporting layer (*i.e.* ICP effect). Thus, a decreasing driving force results in a decreasing Na⁺ permeation rate through the crosslinked GO membrane, which indicates that the permeation behaviour of the S-CGO membrane suffers from serious ICP effects, in comparison to the F-CGO membrane, because of its porous supporting layer.

The water permeation of different membranes is presented in Fig. 4b, where it can be observed that the water permeation of F-CGO membranes is inversely proportional to their cross-linking degree, although it does not differ much for the different samples. Even when a highly concentrated draw solution was used, there was only a slight difference between the water permeation values of the F-CGO2 and F-CGO3 membranes, which may be due to differences in the *d*-spacing of GO membranes after they were cross-linked under different PA dosages. This indicates that cross-linking the GO sheets has little impact on the water permeation of the membrane. Fig. 4c shows that the water permeation of the F-CGO3 membrane is higher than that of the S-CGO3 membrane, particularly when using a highly concentrated draw solution. This result confirms that the freestanding membrane (F-CGO) is better for water permeation than the composite membrane (S-CGO) because, even though it has a faster cross-membrane diffusion of the draw reagent (Fig. S10[†]), the ICP effects on the freestanding membrane were efficiently alleviated or eliminated. We also noted that the water permeation rate of the F-CGO3 membrane was ~35 L m⁻² h⁻¹ when using a 2.0 mol L⁻¹ NaCl solution as the draw solution, and ultrapure water as the feed solution. This value is about three times that of commercial cellulose triacetate (CTA) FO membranes produced by HTI Inc. under the same testing conditions.²³

To further reveal the semi-permeability of the F-CGO membrane, we calculated the selective Na⁺ permeation coefficient across the membranes based on the reverse salt rate and water permeation rate. This parameter is defined as the molar ratio of Na⁺ in the draw solution to that in the feed solution after permeation testing; and it indicates the Na⁺ rejection performance of the membranes under different conditions. Fig. 4d shows that when the NaCl concentration of the draw solution is 0.5 mol L⁻¹, the Na⁺ rejection performance increases as the cross-linking degree increases, which is a trend in agreement with the reverse salt diffusion behaviour of different membranes (Fig. 4a). The F-CGO3 membrane showed the highest Na⁺ rejection value of ~6700 (1.0 mol L⁻¹ NaCl solution as the draw solution). Even at the highest saline solution, the Na⁺ selective permeation coefficient was higher than 2000, which means that the F-CGO3 membrane rejected 99.95% of Na⁺ by retaining it on the side of the draw solution during the osmotic-pressure-driven permeation process. This high Na⁺ rejection ability makes the membrane beneficial for applications such as water desalination, and for the removal of toxic ions and organic pollutants from water through forward osmosis processes.

Clean-water intake from wastewaters

Inspired by its good Na⁺ rejection and water permeation, the F-CGO3 membrane was used to intake clean-water from solutions



contaminated with organic dyes, such as methylene orange (MO) and rhodamine B (RB). As Fig. 5a shows, a noticeable liquid level difference appeared in the U-shaped tube between the NaCl solution (2.0 mol L^{-1}) and the wastewaters, which indicates the intake of water by the NaCl solution through the F-CGO3 membrane. The MO and RB pollutants were rejected by the membrane, as evidenced by the colourless NaCl solution after testing. In addition, after the membrane was immersed for 15 days in the high saline water and polluted solutions, the F-CGO3 membranes were able to withstand the high liquid level difference between the draw and feed solutions, showcasing their excellent durability and hydraulic stability. The water intake efficiency (water permeation ratio) of the F-CGO3 membrane was determined using the same setup as the semi-permeability tests shown in Fig. S1,[†] where a 2.0 mol L^{-1} NaCl solution and wastewater were used as the draw and feed solutions, respectively.

Fig. 5b shows that the initial membrane water permeation rates in MO and RB solutions were 12.2 and $19.3 \text{ L m}^{-2} \text{ h}^{-1}$, respectively. These rates decreased as the permeation time was prolonged, and reached stability after a 3 h permeation period.

This is due to the interaction forces and steric effects of the pollutants with the F-CGO3 membrane. As it is known, MO and RB have conjugated aromatic structures (Fig. S11[†]), and are attracted to GO sheets *via* the π - π interaction. Additionally, coulombic interactions may also contribute to the water permeation decline because of the negatively charged GO sheets and positively charged RB. Although RB has a larger conjugated aromatic structure and more favourable coulombic interactions with the GO sheets, its water intake efficiency is higher than that of the MO solution at the same feeding weight concentration. This might be caused by the smaller molecular weight of MO, which results in a higher osmotic potential at the same weight concentration. On the other hand, the smaller molecular size of the MO molecule allows it to easily access the inlets of the inter-laminar membrane channels and block the permeation of water. Furthermore, to evaluate the influence of the supporting layer on the water intake efficiency of the membranes, in Fig. 5b the water permeation rates of the F-CGO3 and S-CGO3 membranes in various wastewaters are compared. The figure shows that when the S-CGO3 membrane was used as the separator, the water permeation rates from feed solutions was

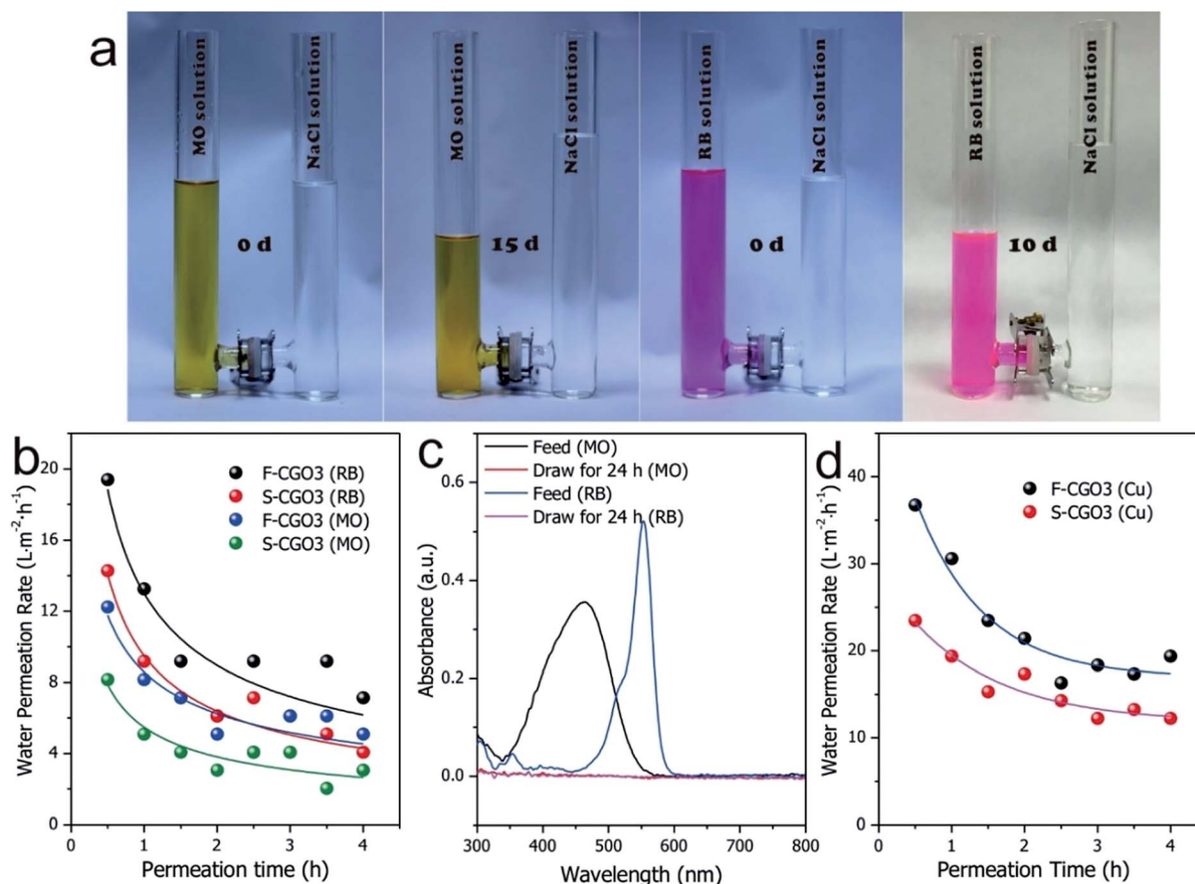


Fig. 5 (a) Digital pictures of the water permeation behaviour from a 10 ppm MO solution (left two) and a 10 ppm RB solution (right two) to a 2 mol L^{-1} NaCl solution through the F-CGO3 membrane, which was fixed in the tunnel of a glass communicating vessel. (b) Water permeation rates of the F-CGO3 and S-CGO3 membranes in the MO or RB solution. (c) UV-Vis spectra of the original MO or RB solutions (diluted 2 times), and the corresponding draw solutions after a 24 h permeation period performed with the F-CGO3 membrane. (d) Water permeation rate of the F-CG and S-CGO3 membranes in the 10 ppm Cu^{2+} solution. The feed solutions were 10 ppm MO, RB or Cu^{2+} aqueous solutions, while the draw solution was a 2 mol L^{-1} NaCl solution.



lower at the beginning and during the stable interval. These results are consistent with those in Fig. 4c, thus indicating the presence of ICP effects on the permeation of water during a concentration gradient driven by a membrane separation process, such as FO. We also detected by UV-Vis spectroscopy that pollutants leaked into the draw solution. The UV-Vis spectra in Fig. 5c indicate that the absorption of MO and RB molecules are both nil, which confirms that the intake from the wastewaters is clean-water.

We also investigated the water intake efficiency of membranes from heavy metal ions solutions using Cu^{2+} as a model (Fig. 5d). From the results, the initial water permeation rate in a Cu^{2+} solution was found to reach $36.7 \text{ L m}^{-2} \text{ h}^{-1}$, and after 3 h, this value stabilized to about $20 \text{ L m}^{-2} \text{ h}^{-1}$. This decrease in the water permeation rate could be owing to the strong coordination interactions between Cu^{2+} and the oxidized region of the GO sheets, which causes the Cu^{2+} to concentrate around the inlets of the inter-laminar channels of the F-CGO3 membrane, and results in a high water permeation resistance across the membrane, similar to what was reported by Shi *et al.*⁴⁶ However, both the initial and stable water permeation rates obtained for the Cu^{2+} solution are larger than those obtained for the MO and RB solutions. This reflects the higher water intake efficiency from inorganic wastewater by the F-CGO membrane. The draw solution after permeation testing was also examined using an ICP-AES instrument. In fact, there was no Cu^{2+} was detected in the draw solution after permeation testing, which indicates that the heavy metal ions, *i.e.* Cu^{2+} , were completely blocked by the F-CGO3 membrane. Therefore, we can conclude that the F-CGO3 membrane can be used to intake clean-water from wastewater by effectively rejecting heavy metal ions and organic pollutants. This performance is useful for the on-site water purification from natural or polluted waters, and therefore the fabricated membrane would be a promising candidate to serve as the functional membrane of some portable water purification devices, such as a military hydration bag for its use during rough missions.

Conclusions

Through the thermally-induced chemical cross-linking process, we successfully prepared a thin freestanding GO-based FO membranes with controllable inter-laminar distance and good hydraulic stability. Comparisons between the permeation performances of the membranes confirmed that the water permeation rate of the freestanding membrane is far higher than that of standing membrane, which suggests that the ICP effect induced by the FO process can be alleviated, and even eliminated, by removing the porous supporting layer. The water intake tests from water contaminated with heavy metal ion and organic dyes demonstrated that the thin freestanding GO-based FO membrane can be used to intake clean-water from wastewater by effectively rejecting heavy metal ions and organic pollutants. These properties make the fabricated membrane a promising material for the assembly of portable water purification systems that can be used by field workers and the military, or to supply drinking water during emergency situations.

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Notes and references

- 1 R. R. Nair, H. A. Wu, P. N. Jayaram, I. V. Grigorieva and A. K. Geim, *Science*, 2012, **335**, 442–444.
- 2 R. K. Joshi, P. Carbone, F. C. Wang, V. G. Kravets, Y. Su, I. V. Grigorieva, H. A. Wu, A. K. Geim and R. R. Nair, *Science*, 2014, **343**, 752–754.
- 3 P. Sun, H. Liu, K. Wang, M. Zhong, D. Wu and H. Zhu, *Chem. Commun.*, 2015, **51**, 3251–3254.
- 4 P. Sun, F. Zheng, M. Zhu, Z. Song, K. Wang, M. Zhong, D. Wu, R. B. Little, Z. Xu and H. Zhu, *ACS Nano*, 2014, **8**, 850–859.
- 5 P. Sun, M. Zhu, K. Wang, M. Zhong, J. Wei, D. Wu, Z. Xu and H. Zhu, *ACS Nano*, 2013, **7**, 428–437.
- 6 J.-Q. Huang, T.-Z. Zhuang, Q. Zhang, H.-J. Peng, C.-M. Chen and F. Wei, *ACS Nano*, 2015, **9**, 3002–3011.
- 7 K. W. Putz, O. C. Compton, M. J. Palmeri, S. T. Nguyen and L. C. Brinson, *Adv. Funct. Mater.*, 2010, **20**, 3322–3329.
- 8 L. Huang, Y. Li, Q. Zhou, W. Yuan and G. Shi, *Adv. Mater.*, 2015, **27**, 3797–3802.
- 9 T. M. Yeh, Z. Wang, D. Mahajan, B. S. Hsiao and B. Chu, *J. Mater. Chem. A*, 2013, **1**, 12998–13003.
- 10 H. M. Hegab and L. Zou, *J. Membr. Sci.*, 2015, **484**, 95–106.
- 11 B. Liang, W. Zhan, G. Qi, S. Lin, Q. Nan, Y. Liu, B. Cao and K. Pan, *J. Mater. Chem. A*, 2015, **3**, 5140–5147.
- 12 P. Lu, S. Liang, T. Zhou, X. Mei, Y. Zhang, C. Zhang, A. Umar and Q. Wang, *RSC Adv.*, 2016, **6**, 56599–56609.
- 13 Y. Han, Z. Xu and C. Gao, *Adv. Funct. Mater.*, 2013, **23**, 3693–3700.
- 14 H. Huang, Z. Song, N. Wei, L. Shi, Y. Mao, Y. Ying, L. Sun, Z. Xu and X. Peng, *Nat. Commun.*, 2013, **4**, 2979.
- 15 J.-G. Gai, X.-L. Gong, W.-W. Wang, X. Zhang and W.-L. Kang, *J. Mater. Chem. A*, 2014, **2**, 4023–4028.
- 16 C.-N. Yeh, K. Raidongia, J. Shao, Q.-H. Yang and J. Huang, *Nat. Chem.*, 2015, **7**, 166–170.
- 17 R. K. Joshi, S. Alwarappan, M. Yoshimura, V. Sahajwalla and Y. Nishina, *Appl. Mater. Today*, 2015, **1**, 1–12.
- 18 L. J. Cote, J. Kim, Z. Zhang, C. Sun and J. Huang, *Soft Matter*, 2010, **6**, 6096–6101.
- 19 S. Sun, C. Wang, M. Chen and M. Li, *Chem. Phys. Lett.*, 2013, **561**, 166–169.
- 20 Z. Jia and W. Shi, *Carbon*, 2016, **101**, 290–295.
- 21 Z. Jia and Y. Wang, *J. Mater. Chem. A*, 2015, **3**, 4405–4412.
- 22 S. Park, K.-S. Lee, G. Bozoklu, W. Cai, S. T. Nguyen and R. S. Ruoff, *ACS Nano*, 2008, **2**, 572–578.
- 23 H. Liu, H. Wang and X. Zhang, *Adv. Mater.*, 2015, **27**, 249–254.
- 24 P. Sun, Q. Chen, X. Li, H. Liu, K. Wang, M. Zhong, J. Wei, D. Wu, R. Ma, T. Sasaki and H. Zhu, *NPG Asia Mater.*, 2015, **7**, e162.



- 25 P. Sun, R. Ma, W. Ma, J. Wu, K. Wang, T. Sasaki and H. Zhu, *NPG Asia Mater.*, 2016, **8**, e259.
- 26 P. Nasr and H. Sewilam, *Clean Technol. Environ.*, 2015, **17**, 2079–2090.
- 27 M. Xie, J. Lee, L. D. Nghiem and M. Elimelech, *J. Membr. Sci.*, 2015, **493**, 748–754.
- 28 S. Zhao, L. Zou, C. Y. Tang and D. Mulcahy, *J. Membr. Sci.*, 2012, **396**, 1–21.
- 29 B. Mi and M. Elimelech, *J. Membr. Sci.*, 2010, **348**, 337–345.
- 30 Y. Huang, H. Jin, H. Li, P. Yu and Y. Luo, *RSC Adv.*, 2015, **5**, 106113–106121.
- 31 M. Daly, C. Cao, H. Sun, Y. Sun, T. Filleter and C. V. Singh, *ACS Nano*, 2016, **10**, 1939–1947.
- 32 Y. Gao, L.-Q. Liu, S.-Z. Zu, K. Peng, D. Zhou, B.-H. Han and Z. Zhang, *ACS Nano*, 2011, **5**, 2134–2141.
- 33 C. Yu, B. Zhang, F. Yan, J. Zhao, J. Li, L. Li and J. Li, *Carbon*, 2016, **105**, 291–296.
- 34 J. Zhao, B. Zhang, C. Yu, Y. Liu, W. Wang and J. Li, *Carbon*, 2016, **109**, 487–494.
- 35 S. Xie, J. Zhao, B. Zhang, Z. Wang, H. Ma, C. Yu, M. Yu, L. Li and J. Li, *ACS Appl. Mater. Interfaces*, 2015, **7**, 17558–17564.
- 36 B. Zhang, Y. Zhang, C. Peng, M. Yu, L. Li, B. Deng, P. Hu, C. Fan, J. Li and Q. Huang, *Nanoscale*, 2012, **4**, 1742–1748.
- 37 G. Jiang, M. Golezdzinowski, F. J. E. Comeau, H. Zarrin, G. Lui, J. Lenos, A. Veileux, G. Liu, J. Zhang, S. Hemmati, J. Qiao and Z. Chen, *Adv. Funct. Mater.*, 2016, **26**, 1729–1736.
- 38 H. Wang, J. T. Robinson, X. Li and H. Dai, *J. Am. Chem. Soc.*, 2009, **131**, 9910–9911.
- 39 B. Zhang, L. Li, Z. Wang, S. Xie, Y. Zhang, Y. Shen, M. Yu, B. Deng, Q. Huang, C. Fan and J. Li, *J. Mater. Chem.*, 2012, **22**, 7775–7781.
- 40 P. Y. Han, M. Tani, M. Usami, S. Kono, R. Kersting and X. C. Zhang, *J. Appl. Phys.*, 2001, **89**, 2357–2359.
- 41 C. Ronne and S. R. Keiding, *J. Mol. Liq.*, 2002, **101**, 199–218.
- 42 J. T. Kindt and C. A. Schmittenmaer, *J. Phys. Chem.*, 1996, **100**, 10373–10379.
- 43 B. Hussain, M. Ahmed, M. Nawaz, M. Saleem, M. Razzaq, M. A. Zia and M. Iqbal, *Appl. Opt.*, 2012, **51**, 5326–5330.
- 44 H. N. Ritland, *J. Am. Ceram. Soc.*, 1955, **38**, 86–88.
- 45 D. Mergel, D. Buschendorf, S. Eggert, R. Grammes and B. Samset, *Thin Solid Films*, 2000, **371**, 218–224.
- 46 J. Liu, G. S. Shi, P. Guo, J. R. Yang and H. P. Fang, *Phys. Rev. Lett.*, 2015, **115**, 164502.

