



Cite this: *RSC Adv.*, 2017, 7, 53907

A novel gas conductor–gas barrier (GC–GB) blending membrane with adjustable gas separation capacity

Qi He,^a Xiaoyue Xiao,^b Wenxia Wang,^a Yanbo Huang,^a Jiapeng He^a and Kaijun Xiao^{*a}

A promising avenue to preserve harvested fruits and vegetables can be achieved by designing a “breathable” membrane named as equilibrium modified atmosphere packaging (EMAP). In this work, such “breathable” packaging was prepared using a gas conductor –gas barrier (GC–GB) blending membrane (BM). The miscibility between the GC and GB in the BM was displayed using structural scanning and thermal analysis, and the physicochemical properties of the BM were characterized by a series of assays. By changing the SPEEK content in the BM, CO₂ permeabilities of the BM in pure gas can be adjusted from 105 to 1494 kPa, while the O₂ permeabilities can be adjusted from 19 to 297 kPa. Thus, the CO₂/N₂ and O₂/N₂ ideal selectivity of the BMs were 20.48–33.95 and 5.38–6.75, respectively. On this basis, the membrane was used to preserve oranges and cauliflower. The equilibrium internal atmospheres included 4.6–7.9% of CO₂ + 7.8–4.4% of O₂ (p/p) and 3.6–8.7% of CO₂ + 9.2–4.2% of O₂ (p/p), which were respectively recommended to preserve the stored products. The results indicated such adjustable membranes will have promising applications to preserve different kinds of fresh products.

Received 6th September 2017
Accepted 31st October 2017

DOI: 10.1039/c7ra09953c

rsc.li/rsc-advances

1. Introduction

Recently, increasing attention has been paid to developing an effective approach to preserve harvested vegetables or fruits.¹ Since preserved produce is alive, it still needs to consume nutrients and exchange gases through respiration behavior after it is harvested.² Thus, more CO₂ and less O₂ in the storage atmosphere are usually advantageous because these conditions can significantly reduce respiratory intensity and nutrient consumption of the products. However, minimal O₂ should be kept in case the living products “suffocate” to death.²

A promising strategy can be achieved by designing a “breathable” membrane named as equilibrium modified atmosphere packaging (EMAP).³ Its internal atmosphere can be kept in a relatively stable state by balancing the gradient-induced gas transport through the membrane (O₂ entering the packaging and CO₂ leaving) against the gas exchange caused by respiration of the products (O₂ uptake and CO₂ production).⁴ On this basis, the major goal to design an EMAP is to achieve ideal selectivity and permeability. It can be achieved by selecting suitable membrane materials according to the chemical structure, crystallinity and morphology of materials.⁵

Extensive research has focused on tailoring polymeric membranes to achieve superior separation properties, especially some approaches to improve polymers that can hardly fulfill the required properties. For example, Tin *et al.*⁶ performed a room-temperature chemical cross-linking modification on Matrimid as gas separate membrane. Results showed that the permeability of CO₂ through the membrane in mixed gas (40% CO₂ in CH₄, 35 °C and 1.01 × 10⁶ Pa) reduced from 6.72 bar to 3.26 bar after 7 days of treatment. Ammar *et al.*⁷ loaded 0–3 wt% of C30B into polysulfone/polyimide mixed film. The O₂ permeation of the prepared films at 1 atm and 23 °C roughly reduced by half. Khan *et al.*⁸ aimed to gas separation using polysulfone acrylate–zeolite mixed matrix membranes. The mixed gas selectivity for H₂/CO₂ separation increased from 1.53 for pure polysulfone to 3.57 at 40% zeolite loading.

Poly vinylidene fluoride (PVDF) is generally considered as a promising gas barrier (GB) polymer because of its remarkable economic feasibility, chemical compatibility, thermal stability, mechanical strength but low gases permeability.^{9,10} However, its performance is often limited by its hydrophobic nature that associates with pervasive fouling.¹¹ To counter these challenges, this study used sulfonated poly ether ether ketone (SPEEK) as a charged gas conductor (GC) additive into PVDF. A series of assays revealed that the introduction of SPEEK as additive agent in PVDF had at least three advantages compared to other competitors. The first one is miscibility, represented by that SPEEK had free compatible with SPEEK at free percentage. Secondly, the gas separation performance of the composite

^aSchool of Food Science and Technology, South China University of Technology, Guangzhou, China. E-mail: fekjxiao@scut.edu.cn; Tel: +86 20 87113843

^bDepartment of Water Resource and Environmental Engineering, Tamkang University, Taipei, China



membranes had widely adjustable gas selectivity, so that they can be applied as EMAP for different preserved products.^{12,13} Last but not least, traditional PVDF based materials as hydrophobic polymers are prone to be fouled, while the introduction of $-SO_3$ groups in SPEEK can significantly enhance its hydrophilicity.¹⁴ As the results, the composite membranes will be a promising candidate as EMAP to preserve different harvest products.

2. Experimental

2.1. Materials

PEEK (Victrek450G) was obtained from Victrek, UK. PVDF (FR904) was obtained from New Materials Co. Ltd., China. H_2SO_4 solution (98 wt%) and *N,N*-dimethylformamide (DMF) were obtained from Donghong Chemical Plant, China. Microbiology agars were obtained from Huankai Microbe Company, China. *Escherichia coli*, *Pseudomonas aeruginosa* and *Staphylococcus aureus* strains were obtained from Guangdong Province Bacteria Collection Center, China. Chemicals in this study were used without further purification.

2.2. PEEK sulfonation

PEEK was sulfonated by thoroughly dissolving into 98 wt% H_2SO_4 (10 g/200 mL).¹² The mixture was heated to 60 °C and kept under mechanical stirring at nitrogen atmosphere for 16 h. Then, the resulting solution was slowly injected into a glass tank with ice water. The raw SPEEK material was obtained by the precipitate. It was adequately washed and dried under vacuum at 60 °C for 48 h.

Functional groups in the SPEEK were investigated using a Fourier transform infra-red spectrum (FTIR) analyzer (Vector 33, Bruker Company, Germany).¹⁵ Ion exchange capacity (IEC) and the degree of sulfonation (DS) of the prepared SPEEK were determined by H^+ release amount. For this, minced SPEEK (~1 g) was suspended in 2 M NaCl solution (100 mL) for 24 h. The resulting solution was titrated using 0.1 M NaOH solution with phenolphthalein as indicator. The results were calculated by eqn (1) and (2):¹⁶

$$IEC \text{ (mequiv. g}^{-1}\text{)} = \frac{V_{NaOH} - V_{NaCl}}{1000W_{dry}} \quad (1)$$

$$DS = \frac{M_0 IEC}{1000 - M_{SO_3Na} IEC} \quad (2)$$

where V_{NaOH} and V_{NaCl} were the volume of used NaOH and NaCl, respectively; W_{dry} is the weight of dry PEEK; M_0 (288 g mol⁻¹) and M_{SO_3Na} (103 g mol⁻¹) were the molar mass of the initial polymer and the $-SO_3Na$ group, respectively.

2.3. Preparation and characterization of the membranes

Pure PVDF and SPEEK solutions were respectively prepared by dissolving ~15 wt% of corresponding polymer into DMF solvent. On this basis, the casting solutions of blending membrane (BM) were prepared by mixing the pure solutions in different ratio (for the samples of BM25, BM50, and BM75,

25 v/v%, 50 v/v% and 75 v/v% of SPEEK were blended into PVDF, respectively). Each solution was centrifuged at 4000 rpm for 3 min to eliminate foam. Then they were casted on a clear glass plate (360 mm × 300 mm) at ~20 °C and ~75% relative humidity. The casted glass plates were dried at 60 °C for 24 hours to evaporate solvent and form membranes.¹²

Each prepared membrane was characterized by a series of assays. Micro-structure was scanned using a scanning electron microscope (SEM, Nova Nano 450, US) operated at 20 kV and 80 mA.¹⁷ Thermal behavior was reflected using a differential scanning calorimeter (DSC, TGA/DSC-1/1100, Mettler-Toledo Co. Ltd., Switzerland).¹⁸ Hydrophilicity was determined using a contact angle (CA) goniometer (OCA15, Data-Physics Company, Germany).¹⁹ Anti-biofouling capability was assessed through the inhibiting ring on inoculated agars.¹⁸ Porosity was measured by a Brunauer–Emmett–Teller (BET) nitrogen adsorber (JW-BK222, JWGB, China).²⁰ Internal void space was examined *via* water-absorption porosities values (ϵ).²¹

2.4. Gas permeation measurements

An air-tight system (as Fig. 1a) was established for gas analysis.²² The gas selectivities and permeabilities of the membrane were determined by comparing the gas composition at both sides of the membranes using a compact gas chromatograph (CGC, Interscience, Fran). Using pure gases, gas permeability (P) and ideal selectivity (S) were determined from the rate of pressure increase (dP/dt) obtained when permeation reached steady state, as the following equations:²²

$$P_1 = \frac{273.15 \times 10^6 y_1 V L}{760 \times \frac{76}{14.7} A T x_1 P_f} \left(\frac{dP}{dt} \right) \quad (3)$$

$$S_{12} = \frac{P_1}{P_2} \quad (4)$$

where P_1 and P_2 are the gas permeability of component 1 and 2, respectively, y_1 and x_1 are the mole fraction of component 1 at the feed and permeated sides respectively, V is the downstream volume (cm³), L is the membrane thickness (μ m), A is the membrane area (cm²), T is the operating temperature (K), P_f is the pressure of the feed gas (Pa).

When it came to mixed-gas, the selectivity (α) was calculated by the ratio of mole fraction of the gases, as eqn (5):²²

$$\alpha_{12} = \frac{y_1/y_2}{x_1/x_2} \quad (5)$$

where y_1 and y_2 are the mole fractions of components 1 and 2, respectively at the permeated side, x_1 and x_2 are the mole fractions of components 1 and 2, respectively at the upstream.

2.5. Actual applications

The evaluation of the applicability and feasibility of gas separation membranes requires the performance testing under actual conditions. In this study, navel orange (*Citrus sinensis*)



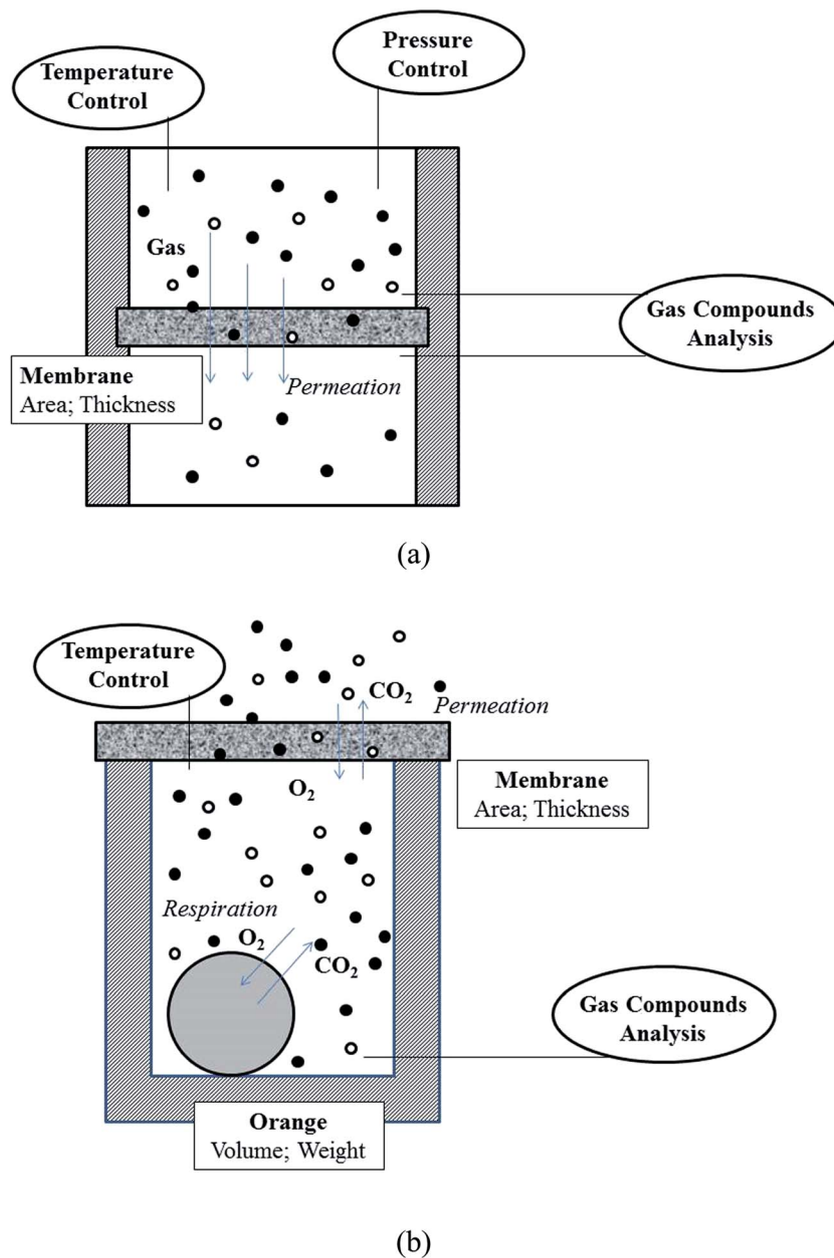


Fig. 1 Schematic for the experimental setup in this study.

and cauliflower (*Brassica oleracea* L. var. *botrytis*) obtained from local farmers were used as experimental samples. Their average individual weight was ~ 300 g. Each sample was assayed within three hours after harvested.

A cylindrical measure chamber ($\varnothing 15$ cm \times 12 cm) was set up to preserve sample in the assay (as Fig. 1b). Its wall and bottom were made by glass that can thoroughly isolate air, while its opened bottom tightly attached a prepared membrane (pure PVDF, BM25, BM50, BM75, or pure SPEEK) for analysis. Additionally, two control groups were set up for comparison. C1 used a glass cover on the chamber bottom that can thoroughly isolate air. C2 let the bottom of the chamber expose to the air without a shield.

3. Results and discussion

3.1. Sulfonated behaviors of PEEK

PEEK sulfonation can be considered as a process in which $-\text{SO}_3\text{H}$ groups are grafted into available $-\text{H}$ sites.²³ During this reaction, DS of SPEEK showed increasing trends, but such growth would slow down gradually and tended to equilibrium with increasing time.

Fig. 2 illustrated FTIR results. The peaks presented at ~ 1600 cm^{-1} can be attributed to vibration of the aromatic ring skeleton, whereas that around ~ 1200 cm^{-1} was primarily due to the presence of aryl oxide.²⁴ Meanwhile, the sulfonation behavior can be represented by the characteristic peaks of



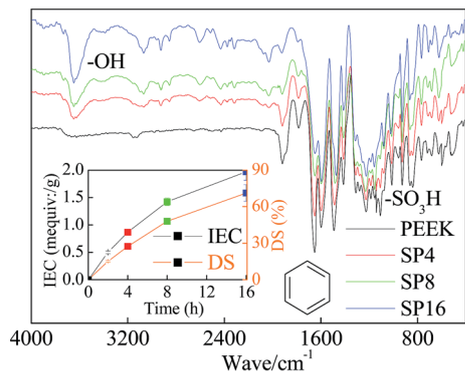


Fig. 2 FTIR spectra of the SPEEK with different DS.

$-\text{SO}_3\text{H}$ around $\sim 1100\text{ cm}^{-1}$. Another noteworthy peak was around $\sim 3200\text{ cm}^{-1}$ caused by $-\text{OH}$.²⁴ It is a typical sign caused by hydration, suggesting the hydrophilicity of materials was significantly improved during the sulfonation process.²⁵

3.2. Blends miscibility

PVDF has a neatly arranged main chain alternating $-\text{CH}_2-$ and $-\text{CF}_2-$. In this chain, H ($1s^1$) atoms are prone to lose its only electron, and F ($1s^2 2s^2 2p^5$) atoms tend to absorb one electron to fill their empty $2p$ -orbital. Meanwhile, SPEEK has an aromatic backbone contains a lot of oxidative phenyl O ($1s^2 2s^2 2p^5$) and reductive phenyl H ($1s^1$). The graft of $-\text{SO}_3\text{H}$ groups increased the number of activated O atoms further.¹² When these two components were mixed, possible interactions between two components are shown in Fig. 3a.^{23,26} As it shown, H atoms from the aromatic ring of SPEEK can lose its only electron to occupy the empty $2p$ -orbital of F from PVDF. O atoms from SPEEK can also form 2 lone-pairs coordination bond with H atoms from PVDF.²⁷ On this basis, the blending materials can maintain a stable state.

To verify the miscibility between PVDF and SPEEK, an initial indication was that all casting solutions and prepared polymers were transparent and homogeneous.²² More obvious details were illustrated by SEM analysis (as Fig. 3b). No individual matrix or mixed interface can be found in the surface and cross-section structure of the BMs. Additionally, further criterion was given by DSC results that obtained the existence of single glass transition temperature (T_g), by using the Fox equation, given as:²²

$$T_g = \frac{1}{\frac{W_1}{T_{g1}} + \frac{W_2}{T_{g2}}} = \frac{T_{g1}T_{g2}}{W_1T_{g2} + W_2T_{g1}} \quad (6)$$

where W_1 and W_2 are the mass fractions of the components 1 and 2 in the blend, and T_{g1} and T_{g2} are the glass transition temperatures of the components 1 and 2 in the blend. Fig. 3c shows the theoretical and experimental comparison of the blend systems with respect to their composition. The comparison showed an excellent agreement between the experimental and theoretical results.

3.3. Antifouling and porosities

Surface hydrophilicity can enhance membranes' fouling resistance, because it can improve membranes' wettability to

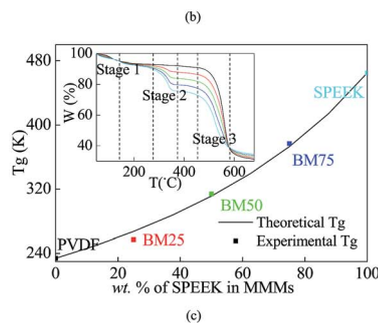
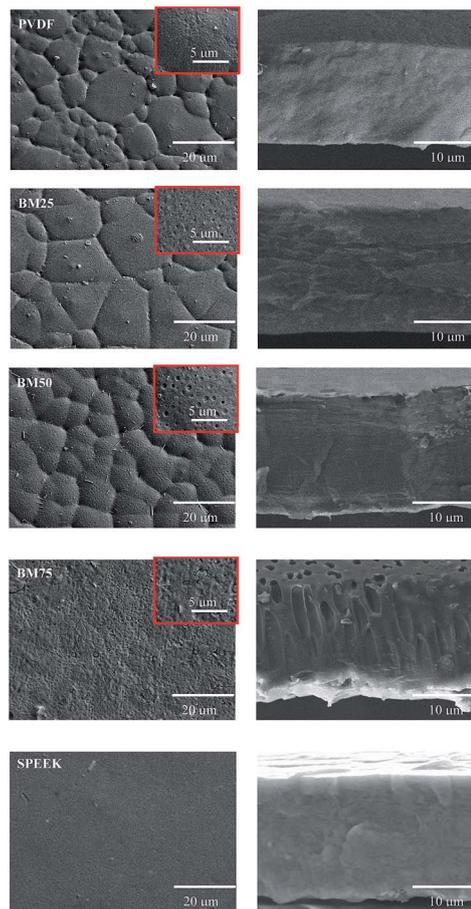
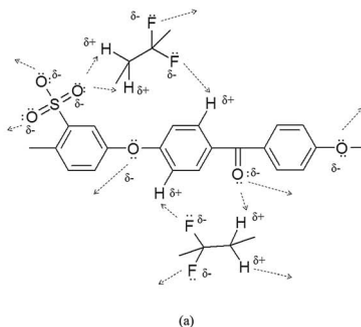


Fig. 3 Miscibility of the BMs: (a) possible interactions; (b) SEM morphology; (c) T_g and TGA.

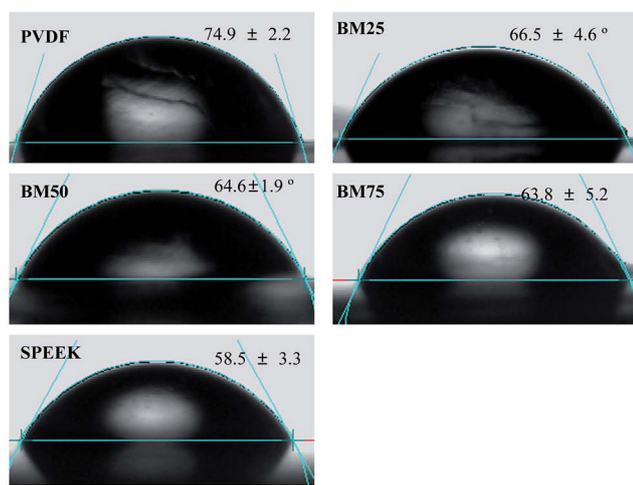
suppress the effects of concentration polarization.⁵ PVDF, as a polymer formed by nonpolar monomers, is a relatively hydrophobic material. Therefore, as shown in Fig. 4a, the CA of



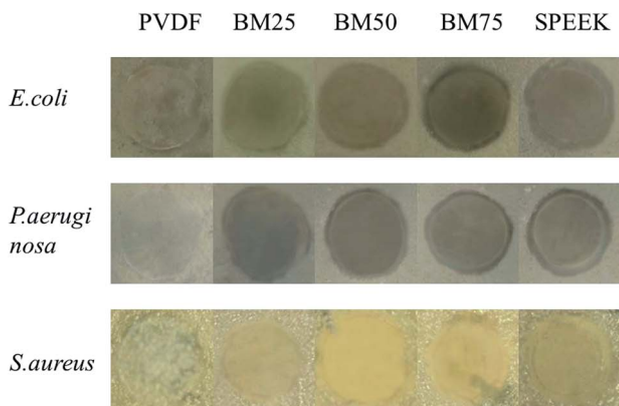
pure PVDF was measured as $\sim 75^\circ$. By comparison, the CA of pure SPEEK was $\sim 60^\circ$, with a lot of hydrophilic $-\text{SO}_3\text{H}$ groups on its main chain.²⁸ On this basis, the CA of the BMs can be theoretically set up to a range from 60° to 75° .

Additionally, an important effect that was effective in the BM's anti-biofouling capacity was the antimicrobial capacity of SPEEK, attributed to the $-\text{SO}_3\text{H}$ groups.²⁹ As shown in Fig. 4b, the pure PVDF cannot make any effects toward the growth of each microbial species, but the BMs with introduction of SPEEK showed inhibiting rings around the membrane disks. Obviously, the SPEEK concentration in the BMs dictated the width of the rings.

When it came to the porosities, Fig. 5a exhibited the results of BET analysis. As it shown, each membrane showed similar N_2 adsorption-desorption isotherms with a convex upward that matches the type I isotherm.³⁰ Compared to SPEEK, PVDF has much sharper N_2 curve with relatively smaller average pore diameter. Thus, more SPEEK compounds in the BMs can slightly improve the average pore size (as Fig. 5a), as well as total pore volume and water-absorption porosities values ϵ (as Fig. 5b).

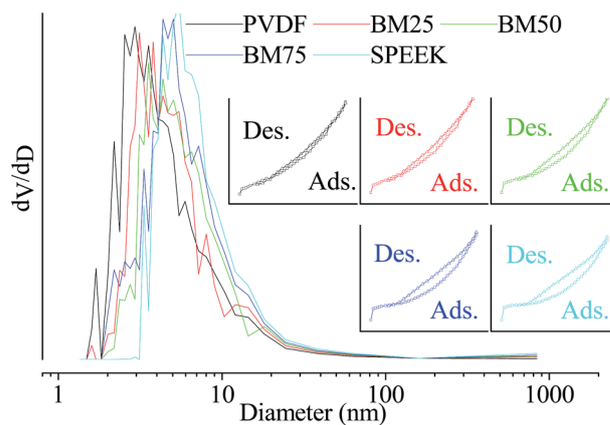


(a)

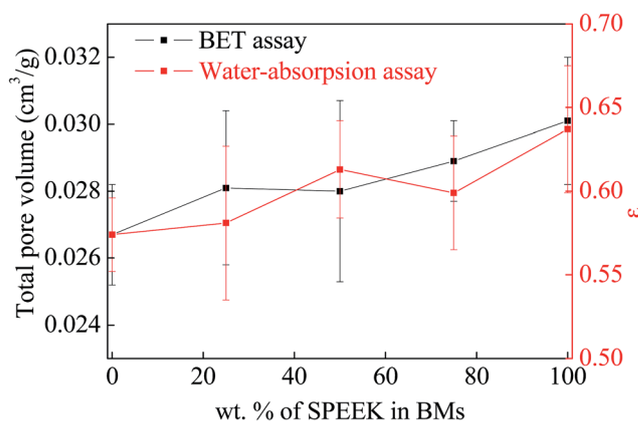


(b)

Fig. 4 Antifouling capacities of the BMs: (a) hydrophilicity and (b) antibiofouling.



(a)



(b)

Fig. 5 Porosities of the BMs: (a) pore diameter distribution and (b) total pore volume.

3.4. Pure gas transport

Generally, major component in internal atmosphere of a packaging is N_2 . It is often considered as a chemically inert gas. The followed gases are O_2 and CO_2 , both of which determine the respiration rate of the stored products. Table 1a summarizes permeabilities and ideal selectivities of pure SPEEK membrane with different DS for pure gas separation. It displayed that the increase of DS can improve the permeabilities of all gases. The reason can be attributed to the introduction of the polar $-\text{SO}_3$ groups in the polymeric chains can improve the solubility coefficients of the gas particles.¹² Compared to N_2 or O_2 , CO_2 had a larger quadrupole moment so that it showed more obvious increase in solubility coefficient, resulting in that the ideal selectivity increased more significantly.¹²

Additionally, Table 1b revealed the gas separation properties of the BMs. Unlike SPEEK, pure PVDF has an apolar main chain that can hardly produce interaction with gas particles. Thus, PVDF can hardly permeate almost all gases so



Table 1 Pure gas permeabilities and ideal selectivities of (a) SPEEK membranes with different DS; and (b) the prepared BMs^a

(a)						
Membr.	DS (%)	Permeability (kPa)			Ideal selectivity	
		CO ₂	O ₂	N ₂	CO ₂ /N ₂	O ₂ /N ₂
PEEK	0	430 ± 63	113 ± 14	21 ± 5	20.48	5.38
SP4	27.4	6650 ± 65	147 ± 12	27 ± 2	24.63	5.44
SP8	48.0	1018 ± 107	212 ± 47	35 ± 6	29.09	6.06
SP16	71.2	1494 ± 207	297 ± 54	44 ± 9	33.95	6.75

(b)					
Membr.	Permeability (kPa)			Ideal selectivity	
	CO ₂	O ₂	N ₂	CO ₂ /N ₂	O ₂ /N ₂
PVDF	105 ± 39	19 ± 5	5 ± 1	21.00	3.80
BM25	602 ± 68	121 ± 20	23 ± 6	26.17	5.26
BM50	942 ± 106	185 ± 24	33 ± 7	28.55	5.61
BM75	1300 ± 72	249 ± 33	41 ± 4	31.71	6.07
SPEEK	1494 ± 207	297 ± 54	44 ± 9	33.95	6.75

^a The study was operated at 293 K and 1000 kPa of feed pressure.

that it is often considered as a gas barrier. In this study, the incorporation of SPEEK in the BMs improved their polar, in addition to the dissolution towards the penetrant molecules. As a result, both permeabilities and ideal selectivities increased with more SPEEK concentration (DS = 71.2%) in the BMs.

3.5. CO₂ plasticization

As a highly condensable gas, CO₂ can plasticize the polymer matrix and lead to a drastic turning point in permeability at a certain partial pressure named as plasticization pressure.³¹ During the gas particles go through a polymeric membrane, CO₂ plasticization will make significant influences to evaluate a membrane's performance.²² This effect can be suppressed by chemical cross-linking,⁶ thermal treatment³² or blending with less plasticizable polymers.³³

In this study, the change in CO₂ permeation with respect to CO₂ partial pressure was analyzed according to the study of Khan *et al.*²² As shown in Fig. 6a, the initial CO₂ permeability decreased with increasing feed pressure, which was in agreement with the permeation of gases through glassy polymeric membranes. As the pressure exceeded plasticization pressure, the gas permeability started to increase with pressure exhibiting CO₂ plasticization. On this basis, SP4 membrane with the lowest DS (27.4%) showed a slight decrease in the CO₂ permeability with increase in pressure until ~600 kPa. SP8 with higher DS (48.0%) improved the plasticization performance by shifting the plasticization pressure to ~1200 kPa. The SP16 (DS = 71.2%) had the highest plasticization pressure of ~3500 kPa. The increase of plasticization pressure can be possibly attributed to increase in chain stiffness induced by the higher content of sulfonic groups.¹²

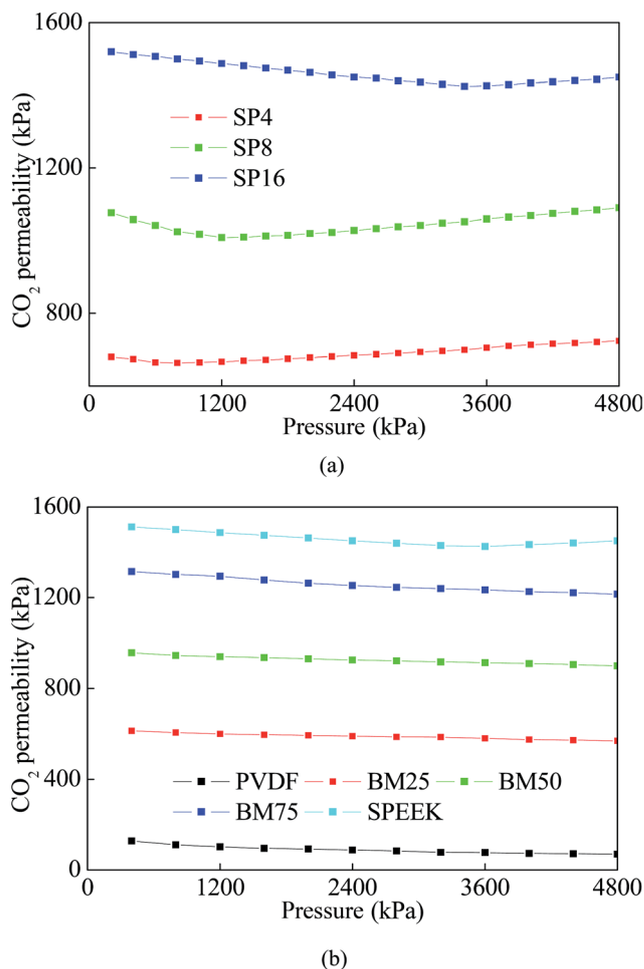


Fig. 6 CO₂ plasticization of (a) the SPEEK and (b) the BMs.

On this basis, plasticization effects in the prepared BMs were demonstrated in Fig. 6b. As it shown, no obvious sign of plasticization can be found in the each membrane until 50 bar. Further investigation at higher pressures was not possible due to limitations of the applied gas permeation system. These results indicated that blending with PVDF could significantly help in suppressing the CO₂ plasticization by increasing the plasticization pressure.

3.6. Binary mixed gas permeations

When using a membrane for mixed gas, an important effect that should be addressed is the competitive sorption of the penetrant gas molecules. It dictates the difference between the ideal and mixed gas selectivities, especially when the gas has a high sorption in the polymer.³⁴ Table 2 displayed the permeabilities of the BMs for different kinds of binary mixed gases (1 : 1). As the results, permeability of each gas in the mixed gas assays was lower than that in the pure gas measurements. Relatively, the decrease in CO₂ permeability was more significant in comparison to that of N₂, resulting in decreasing CO₂/N₂ selectivity. This decrease can be ascribed to competitive sorption of the slower permeating N₂.



Table 2 CO₂/N₂ and O₂/N₂ mixed gas (1 : 1) permeabilities and selectivities of the prepared BMs^a

Membr.	Permeability (kPa)			Permeability (kPa)		
	CO ₂	N ₂	α	O ₂	N ₂	α
PVDF	99 ± 16	5 ± 0	19.80	14 ± 3	4 ± 1	3.50
BM25	532 ± 53	23 ± 6	23.13	106 ± 18	22 ± 3	4.82
BM50	817 ± 76	32 ± 3	25.53	142 ± 14	28 ± 3	5.07
BM75	1048 ± 120	39 ± 2	26.87	196 ± 30	37 ± 6	5.30
SPEEK	1243 ± 87	43 ± 4	28.91	230 ± 31	42 ± 5	5.48

^a The study was operated at 293 K and 1000 kPa of feed pressure.

3.7. Ternary simulated gas permeations

Respiration effect of stored products will consume O₂ and release CO₂ in the packaging, while a simultaneous effect was gas exchange through the membrane caused by difference in gas concentration. Generally, the internal atmosphere of an ideal EMAP will show a trend that reducing O₂ composition is replaced by N₂ and CO₂ until the internal atmosphere get into equilibrium. As a result, O₂ concentration in the atmosphere would decrease from ~21% to ~5% (p/p). N₂ would increase from ~78% to ~90% (p/p). CO₂ would increase from ~0.4% to ~6% and then decrease to ~5% (p/p).¹

In this study, 4 ternary mixed gases (MG1, MG2, MG3 and MG4) were prepared to simulate the atmosphere in different stage of products storage to analyze CO₂ and O₂ selectivities. As shown in Fig. 7, because difference in the compounds of simulated gases was slight, the difference in different gases is not very obvious. However, such difference reasonably plays an important role for equilibrium atmosphere in the packaging during the products storage.

3.8. Modified atmosphere assays

As shown in Fig. 8a, using the prepared BMs to preserve orange, CO₂ concentration inside the packaging got into an equilibrium state after 11.1–23.8 h. The final concentration reached and

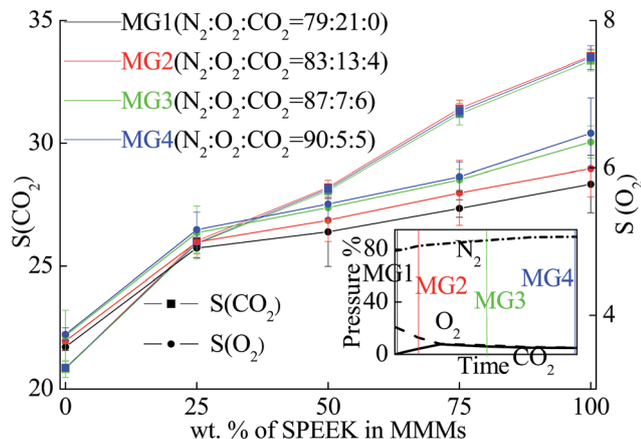
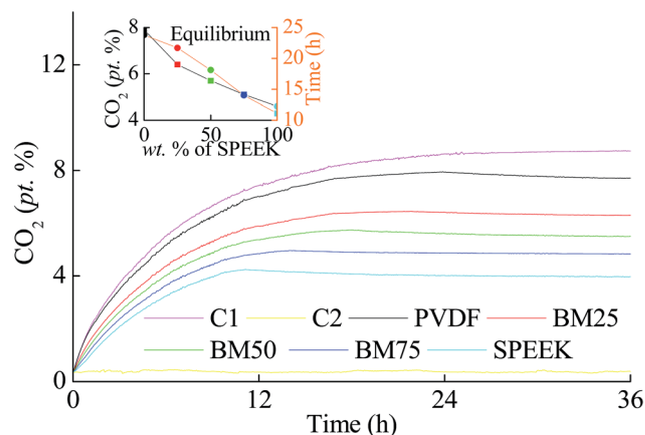


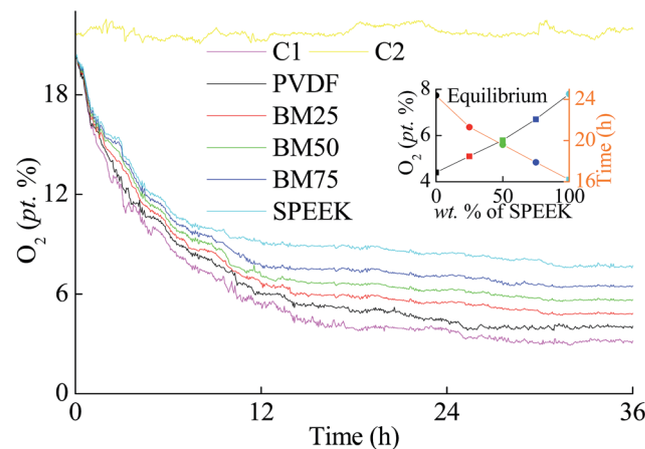
Fig. 7 CO₂ and O₂ selectivities of the BMs in simulated ternary mixed gas at 293 K and 1000 kPa of feed pressure.

kept at 4.6–7.9% (p/p). Meanwhile, O₂ concentration got into the equilibrium concentration of 4.4–7.8% (p/p) after 16.2–24.4 h (as Fig. 8b). Similarly, when the BMs were used to preserve cauliflower, the CO₂ concentration arrived at 3.6–8.7% (p/p) after 3.7–9.1 h of storage (as Fig. 9a), while the O₂ concentration for cauliflower arrived in 4.2–9.2% (p/p) after 5.5–9.6 h (as Fig. 9b). The results indicated a phenomenon that more SPEEK compound in BMs led to shorter equilibrium time, with an equilibrium atmosphere included lower CO₂ concentration and higher O₂ concentration.

Stored vegetables or fruits need to respire as long as there are gases available. Suitable stored atmosphere can significantly prolong the shelf life of stored products by reducing their loss in stored energy reserves through lowering the respiration rate. Because of the biological difference, the optimum O₂ and CO₂ concentrations of different product species vary a lot. As reported, Table 3 shows the recommended concentration of CO₂



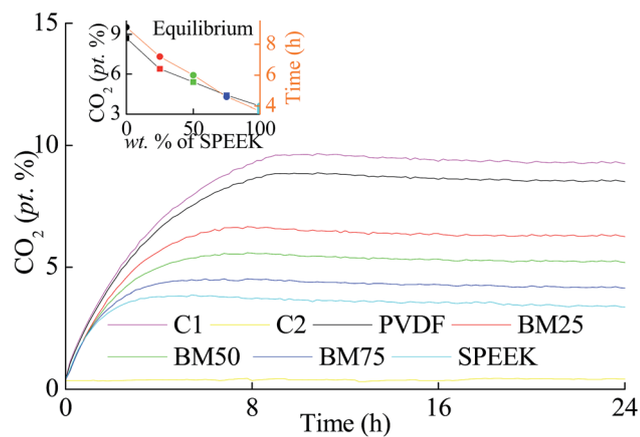
(a)



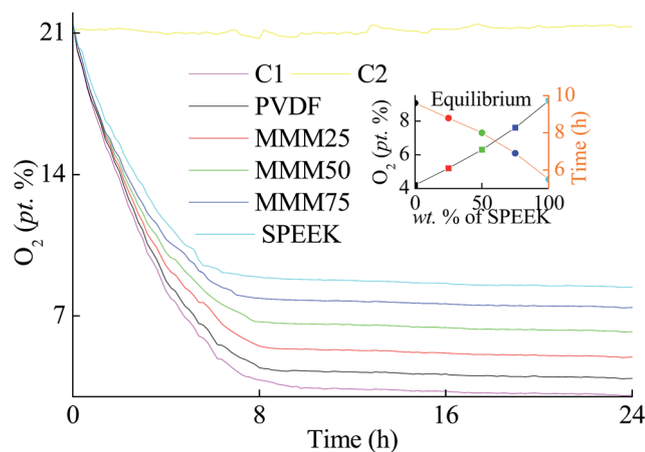
(b)

Fig. 8 The variations in (a) CO₂ concentration and (b) O₂ concentration during the storage of orange using the BMs.





(a)



(b)

Fig. 9 The variations in (a) CO_2 concentration and (b) O_2 concentration during the storage of cauliflower using the BMs.

and O_2 in storage atmosphere that is suitable for many common fruits and vegetables.¹ Accordingly, the BMs packaging aren't only suitable for both orange and cauliflower, it can be also offers a suitable atmosphere to preserve many other fruits and vegetable species.

Last but not least, another notable effect for harvested fruits and vegetables is the transpiration effect, by which they will be detached from the source of water. Actually, a loss in weight of only 5% will cause many vegetables to appear wilted or shriveled. Without the correct packaging, this can happen in a few hours.² Luckily, both PEEK and PVDF have relatively low water vapour transmission rate,^{13,35,36} so that both of them can effectively stop the fluids loss from products' transpiration. Reasonably, the prepared BMs, which are mixed those two fluids barrier, will be effective in the fresh-keeping of the stored products by keeping their moisture. This effect will be studied in further studies.

Table 3 Recommended CO_2 and O_2 concentration for fruits and vegetable's storage²

Fruits	O_2 (%)	CO_2 (%)	Vegetables	O_2 (%)	CO_2 (%)
Apple	1–2	1–3	Artichoke	2–3	2–3
Apricot	2–3	2–3	Snap beans	2–3	5–10
Avocado ^a	2–5	3–10	Broccoli	1–2	5–10
Banana ^a	2–5	2–5	Sprouts	1–2	5–7
Grape	2–5	1–3	Cabbage ^a	2–3	3–6
Grapefruit ^a	3–10	5–10	Carrot ^a	5	3–4
Kiwifruit	1–2	3–5	Cauliflower ^a	2–5	2–5
Lemon ^a	5–10	0–10	Chili pepper ^a	3	5
Mango ^a	3–7	5–8	Sweet corn ^a	2–4	10–20
Orange ^a	5–10	0–5	Cucumber	3–5	0
Papaya ^a	2–5	5–8	Lettuce	1–3	0
Peach	1–2	3–5	Mushroom ^a	3–21	5–15
Pear	2–3	0–1	Spinach	Air	10–20
Pineapple ^a	2–5	5–10	Tomato	3–5	0
Strawberry	5–10	15–20	Onion	1–2	0

^a These vegetable and fruit species may be suitable for storage using the BMs.

4. Conclusions

In this study, a novel GC (SPEEK)–GB (PVDF) blending membrane with adjustable gas separation performance was prepared. The miscibility between SPEEK and PVDF was evidenced by structural scan and thermal analysis. SPEEK can significantly enhance antifouling and optimize the porosities of the BM, while PVDF can effectively suppress the CO_2 plasticization. The blending polymer was used for gas permeations. Its CO_2 and O_2 permeabilities in pure gas can be adjusted at the ranges of 105–1494 kPa and 19–297 kPa, respectively. Thus the ideal selectivity of CO_2/N_2 and O_2/N_2 were 20.48–33.95 and 5.38–6.75, respectively. Similarly, in binary mixed gas with 50% of N_2 , the CO_2 and O_2 permeabilities can be adjusted at the ranges of 99–1243 kPa and 14–230 kPa, respectively. Thus the CO_2/N_2 and O_2/N_2 selectivity were 19.80–28.91 and 3.50–5.48. When the blending polymer was used for orange storage, the atmosphere inside the packaging finally got into an equilibrium state that included 4.6–7.9% of CO_2 and 7.8–4.4% of O_2 (p/p). Meanwhile, the equilibrium state for cauliflower storage included 3.6–8.7% of CO_2 and 9.2–4.2% of O_2 (p/p). The results indicated this packaging can offer recommended atmosphere for both orange and cauliflower. The atmosphere was also suitable for many other fruits and vegetable species. Consequently, the prepared blending polymer will have a promising application as MA packaging to preserve fresh products.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors are grateful for financially supported by the financial support by Guangzhou Science and Technology Program Key Projects (201508020086), Science and Technology



Equipment Mobilization Project of Guangdong Province (x2skB2160440), Applied Science and Technology Research and Development Special Project of Guangdong Province (2015B020230001), the Project of Technology Research Center for Lingnan Characteristic Fruits & Vegetables Processing and Application Engineering of Guangdong Province (No. [2015] 1487) and the Project of Food Science Innovation Team of Guangdong Higher Education Institutes (2016KCXTD020).

References

- 1 A. Aspelund and K. Jordal, *Int. J. Greenhouse Gas Control*, 2007, **1**, 343–354.
- 2 Sandhya, *LWT-Food Sci. Technol.*, 2010, **43**, 381–392.
- 3 T. Dong, S. X. Song, M. Liang, Y. Wang, X. J. Qi, Y. Q. Zhang, X. Y. Yun and Y. Jin, *J. Food Sci.*, 2017, **82**, 97–107.
- 4 D. Rico, A. B. Martin-Diana, J. M. Barat and C. Barry-Ryan, *Trends Food Sci. Technol.*, 2007, **18**, 373–386.
- 5 P. Zhang, J. W. Qian, Y. Yang, Y. X. Bai, Q. F. An and W. D. Yan, *J. Membr. Sci.*, 2007, **288**, 280–289.
- 6 P. S. Tin, T. S. Chung, Y. Liu, R. Wang, S. L. Liu and K. P. Pramoda, *J. Membr. Sci.*, 2003, **225**, 77–90.
- 7 A. Ammar, A. Elzatahry, M. Al-Maadeed, A. M. Alenizi, A. F. Huq and A. Karim, *Appl. Clay Sci.*, 2017, **137**, 123–134.
- 8 A. L. Khan, A. Cano-Odena, B. Gutierrez, C. Minguillon and I. F. J. Vankelecom, *J. Membr. Sci.*, 2010, **350**, 340–346.
- 9 A. Bagheri, M. Javanbakht, H. Beydaghi, P. Salarizadeh, A. Shabanikiac and H. S. Amolid, *RSC Adv.*, 2016, **6**, 39500–39510.
- 10 Y. Q. Zhang and X. Lv, *J. Membr. Sci.*, 2016, **520**, 54–65.
- 11 E. A. Feijani, A. Tavasoli and H. Mahdavi, *Ind. Eng. Chem. Res.*, 2015, **54**, 12124–12134.
- 12 A. L. Khan, X. F. Li and I. F. J. Vankelecom, *J. Membr. Sci.*, 2011, **372**, 87–96.
- 13 L. Jia, X. Xu, H. Zhang and J. Xu, *J. Appl. Polym. Sci.*, 1996, **60**, 1231–1237.
- 14 W. R. Bowen, T. A. Doneva and H. B. Yin, *Desalination*, 2002, **145**, 39–45.
- 15 I. Sawada, R. Fachrul, T. Ito, Y. Ohmukai, T. Maruyama and H. Matsuyama, *J. Membr. Sci.*, 2012, **387–388**, 1–6.
- 16 A. Rahimpour, S. S. Madaeni, S. Ghorbani, A. Shockravi and Y. Mansourpanah, *Appl. Surf. Sci.*, 2010, **256**, 1825–1831.
- 17 Q. He, J. L. Dai, L. Zhu, S. F. Li, K. J. Xiao and Y. R. Yin, *Sci. Adv. Mater.*, 2016, **8**, 1878–1886.
- 18 H. Basri, A. F. Ismail, M. Aziz, K. Nagai, T. Matsuura, M. S. Abdullah and B. C. Ng, *Desalination*, 2010, **261**, 264–271.
- 19 C. Fang, Y. Jing and Z. X. Lin, *Int. J. Adhes. Adhes.*, 2017, **73**, 1–7.
- 20 Q. He, Y. Shen, K. J. Xiao, J. Y. Xi and X. P. Qiu, *Int. J. Hydrogen Energy*, 2016, **41**, 20709–20719.
- 21 H. Basri, A. F. Ismail and M. Aziz, *Desalination*, 2011, **273**, 72–80.
- 22 A. L. Khan, X. F. Li and I. F. J. Vankelecom, *J. Membr. Sci.*, 2011, **380**, 55–62.
- 23 Q. He, Z. P. Zhu, H. Dong and K. J. Xiao, *RSC Adv.*, 2017, **7**, 37000–37006.
- 24 A. Muthumeenal, S. S. Pethaiah and A. Nagendran, *Renewable Energy*, 2016, **91**, 75–82.
- 25 H. L. Wen, C. S. Song, Y. F. Tong, L. Chen and X. L. Liu, *J. Appl. Polym. Sci.*, 2005, **96**, 489–493.
- 26 H. Yong, H. H. C. Park, Y. S. Kang, J. Won and W. N. Kim, *J. Membr. Sci.*, 2001, **188**, 151–163.
- 27 L. Han, R. Wang, D. Yuan, B. Wu, B. Lou and M. Hong, *J. Mol. Struct.*, 2005, **737**, 55–59.
- 28 N. Tan, G. Xiao and D. Yan, *Chem. Mater.*, 2010, **22**, 1022–1031.
- 29 S. Sami and E. Rahimpour, *Orient. J. Chem.*, 2015, **31**, 2195–2200.
- 30 Q. He, J. L. Dai, L. Zhu, K. J. Xiao and Y. R. Yin, *J. Alloys Compd.*, 2016, **687**, 326–333.
- 31 Y. Liu, R. Wang and T. S. Chung, *J. Membr. Sci.*, 2001, **184**, 231–239.
- 32 H. Kawakami, M. Mikawa and S. Nagaoka, *J. Membr. Sci.*, 1996, **118**, 223–230.
- 33 A. Car, C. Stropnik, W. Yave and K. Peinemann, *Sep. Purif. Technol.*, 2008, **62**, 110–117.
- 34 R. Wang, S. L. Liu, T. T. Lin and T. S. Chung, *Chem. Eng. Sci.*, 2002, **57**, 967–976.
- 35 T. Ahmadzada, D. R. McKenzie, N. L. James, Y. B. Yin and Q. Li, *Thin Solid Films*, 2015, **591**, 131–136.
- 36 F. F. Yang, Y. Li, X. Yu, G. G. Wu, Y. Xia, J. Y. Yu and B. Ding, *RSC Adv.*, 2016, **6**, 87820–87827.

