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Estimation of phase separation temperatures for polyethersulfone/solvent/non-solvent systems in RTIPS and membrane properties

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ABSTRACT:

Phase separation temperature estimations, based on Hansen solubility parameters for poly(ethersulfone) (PES)/solvent/non-solvent systems, were carried out to study the control of phase separation temperature in a reverse thermally induced phase separation (RTIPS) process. Four membrane-forming systems were studied, namely PES/N,N-dimethylacetamide (DMAc)/diethylene glycol (DEG), PES/DMAc/polyethylene glycol 200 (PEG200), PES/DMAc/PEG300 and PES/DMAc/PEG400. The effects of PES molecular weights, PES concentrations, PEG molecular weights and ratios of non-solvent/solvent on phase separation temperature are investigated, and the theoretical Hansen solubility parameter calculation is used to establish a prediction equation for phase separation temperature. A linear relationship between the experimental data and the difference in the

solubility parameters between PES and the mixed solvent was observed. When the membrane-forming temperature was higher than the cloud point, membranes with a bi-continuous structure were acquired and showed a higher pure water permeation flux than that of membranes prepared with the non-solvent induced phase separation (NIPS) process. The pure water permeation flux and the mean pore size of membranes prepared with the RTIPS process decreased in line with an increase of PES molecular weight. When the membrane formation mechanism was the RTIPS process, the mechanical properties were better than those of the corresponding membranes prepared with the NIPS process.

Keywords:

polyethersulfone, membrane, phase separation temperature, reverse thermally induced phase separation

1. Introduction

Polyethersulfone (PES) consisting of rigid phenyl and flexible ether groups, aligned with a conjugated structural unit, chemical resistance, good molecular stability and excellent mechanical properties, and has been widely used for membrane materials [1-3].

Many studies have been successfully prepared PES membranes by employing non-solvent induced phase separation (NIPS) [4-6]. In the NIPS process, the phase separation driving force involves mass transfer between the solvent in the casting solution and the non-solvent in a coagulation bath. As a result of the fast mass transfer, the morphology of PES membranes formed by the NIPS method often shows a dense skin layer and finger-like pores, which usually result in a low permeation flux and poor mechanical properties [7, 8]. Modifications to the NIPS process for better PES membranes have been

investigated extensively [9-13].

Commonly employed approaches include the addition of additives to the casting solutions or coagulations [14, 15], the introduction of an extra process [16, 17] and chemical modification [18, 19]. Nevertheless, these modification methods result in complicated membrane-forming processes, which subsequently lead to an increase in production costs.

As a new membrane formation technology, Castro first introduced the thermally induced phase separation (TIPS) method in 1981 [20]. In the TIPS process, an upper critical solution temperature (UCST) system is used to prepare polymer membranes. The phase separation driving force is heat transfer. Compared to the NIPS method, there are fewer variables to control [21-26], and the prepared membranes have a porous skin layer and bi-continuous pores, which usually results in a high permeation flux and good mechanical properties. Crystalline polymers, such as polypropylene (PP) [28], polyethylene (PE) [29], poly(vinylidene fluroride) (PVDF) [30], poly(ethylene-co-vinyl alcohol) (EVOH) [31], ethylene-acrylic acid copolymer (EAA) [32], polyphenylene (PPS) [33], poly(ether ether ketone) (PEEK) [34], poly(oxymethylene) (POM) [35], poly(ethylene chlorotrifluoroethylene) (PECTFE) [36], poly(4-methyl-1pentene) (PMP) [37] and poly(L-lactic acid) (PLLA) [38], as well as amorphous polymers, such as poly(vinyl butyral) (PVB) [39], poly(methyl methacrylate) (PMMA) [40] and polystyrene (PS) [41], have been prepared successfully by using the TIPS process. Moreover, some researchers have focused on simulating the phase separation dynamics of membrane formation in TIPS [42, 43]. Despite its versatility and success, a lack of suitable diluents is one of the main reasons for the TIPS process being challenging for the large scale industrial production of many polymer membrane materials. Moreover, the high processing temperature also limits the wide use of the TIPS method.

Thus, several new improvements to the TIPS method have been reported [44-48]. Matsuyama et al. [44], for instance, first reported a new technology combining TIPS and NIPS to prepare PMMA microporous membranes. Methanol was used as non-solvent, resulting in the TIPS process and NIPS process occurring simultaneously. As result, a membrane with small pores, which were formed by TIPS near the bottom surface, was prepared. Li et al. [45] achieved the transition of the membrane structure from a typical finger-like morphology to a leaf-like network structure by mixing the solvent N,N-dimethylacetamide (DMAc) and the diluent dibutyl phthalate (DBP), while Cha et al. [46] realised a change in α -type crystallite PVDF to β -type crystallite PVDF by introducing a solvent (DMAc) into the coagulation bath. This method was named "modified TIPS" (m-TIPS). Water flux and the mean pore size of the prepared PVDF membrane increased, but mechanical strength decreased. Complex TIPS (c-TIPS) was proposed by Zhu et al. [47], the highlight of which lies in a complex diluent that is mixture of a water-soluble good solvent and a water-soluble additive. PVDF and polystyrene (PS) membranes with high strength and water flux can be obtained. Low TIPS (L-TIPS) was presented by Lu et al. [48], whereby a self-synthesised additive was utilised and the spinning dope temperature of L-TIPS was lower than that of traditional TIPS. A PVDF membrane with sponge-like pores in the middle was obtained, but a few finger-like pores still co-existed.

Our earlier studies [49-52] have posited a reverse thermally induced phase separation (RTIPS) technology. The employed casting solution features a low critical solution temperature (LCST) and the driving force for phase separation is induced by heat transfer, albeit in an opposite manner of TIPS. The RTIPS technology combines a low membrane-forming temperature with fast heat transfer. PES [49] and polysulfone (PSF) [52] membranes can be successfully attained via RTIPS, and the prepared membranes with porous skin have shown high water flux and good mechanical properties. The key

factor affecting PES membrane formation mechanism is the phase separation temperature of the LCST casting solutions. However, there are no previous studies about the theoretical estimation of the phase separation temperature of PES membrane-forming systems via RTIPS. In this study, therefore, the phase separation temperature of PES/solvent/non-solvent ternary systems with LCST was examined, and the theoretical prediction was achieved through Hansen solubility parameters. DMAc was used as a solvent, and diethylene glycol (DEG) and polyethylene glycol (PEG) with molecular weights of 200, 300 and 400 were used as non-solvents. Moreover, the effects of PES concentration, PEG molecular weight, and membrane-forming temperature on the PES membrane structure and USCK performances are discussed in detail.

2. Experimental

2.1. Materials

Three different polyethersulfones (PES) were used as membrane material in this study. The average molecular weights (stated in units of g/mol) were 51,000 (E6020P, BASF Co. Ltd., German), 62,000 (VERADEL 3000P, Solvay Specialty Polymers USA, LLC) and 77,000 (Jilin Jida High Performance Materials Co. Ltd., China). The polymers are abbreviated herein as PES51000, PES62000 and PES77000, respectively. All three polymers were dried at 60°C before use. N,N-dimethylacetamide (DMAc), diethylene glycol (DEG), polyethylene glycols with molecular weights of 200, 300 and 400, respectively (PEG200, PEG300 and PEG400), and ethanol were purchased from Shanghai Chemical Reagent Co. Ltd. Bovine serum albumin (BSA, average molecular weight of 67,000) was obtained from Shanghai Lianguan Biochemical Engineering Co., Ltd.). All chemicals were used without further purification.

2.2. Preparation of the casting solutions

The compositions of the PES/solvent/non-solvent casting solutions are shown in Table 1 and Table 2. DMAc is a solvent for PES, and DEG and the three types PEG are non-solvents for PES.

Homogeneous casting solutions were obtained by stirring for 48h at 25°C, following which the

solutions were degassed under atmospheric pressure for one day at room temperature.

		Casting so	Mass ratio of		
Casting solutions	Types of PES	DES	DMAc	DEG	DEG/DMAc
		ГĽЭ	DMAC	DEG	(wt. /wt.)
MDEG1-10		10.0	41.5	41.5	1.0:1.0
MDEG1-12		12.0	41.5	41.5	1.0:1.0
MDEG1-14		14.0	41.5	41.5	1.0:1.0
MDEG1-15.5		15.5	41.5	41.5	1.0:1.0
MDEG1-17a	PES51000	17.0	41.5	41.5	1.0:1.0
MDEG1-17b		17.0	42.6	40.4	0.95:1.0
MDEG1-17c		17.0	43.7	39.3	0.9:1.0
MDEG1-17d		17.0	46.1	36.9	0.8:1.0
MDEG1-17e		17.0	47.4	35.6	0.75:1.0
MDEG1-17f		17.0	48.8	34.2	0.7:1.0
MDEG2-17	PES62000	17.0	41.5	41.5	1.0:1.0
MDEG3-17	PES77000	17.0	41.5	41.5	1.0:1.0

Table 1 Compositions of the PES/DMAc/DEG casting solutions.

 Table 2 Compositions of the PES/DMAc/PEG casting solutions.

		Casting s	Casting solution compositions (wt.%)			
Casting solutions	Types of PES	PES	DMAc	PEG	PEG/DMAc (wt. /wt.)	
MPEG200-1		17.0	34.6	48.4	1.4:1	
MPEG200-2		17.0	38.5	82.6	1.6:1	
MPEG200-3	DES51000	17.0	27.7	55.3	2.0:1	
MPEG200-4	PES51000	17.0	25.1	57.9	2.3:1	
MPEG200-5	.0	17.0	20.75	62.25	3.0:1	
MPEG200-6		17.0	15.1	67.9	4.5:1	
MPEG300-1		17.0	20.75	62.25	3.0:1	
MPEG300-2		17.0	16.6	66.4	4.0:1	
MPEG300-3	DES51000	17.0	13.8	69.2	5.0:1	
MPEG300-4	FES51000	17.0	11.9	71.1	6.0:1	
MPEG300-5		17.0	10.4	72.6	7.0:1	
MPEG300-6		17.0	9.2	73.8	8.0:1	
MPEG400-1		17.0	25.1	57.9	2.3:1	
MPEG400-2	DES51000	17.0	20.75	62.25	3.0:1	
MPEG400-3	ress1000	17.0	15.1	67.9	4.5:1	
MPEG400-4		17.0	10.0	73.0	7.3:1	

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MPEG400-5		17.0	9.2	73.8	8.0:1

2.3. Characterisation of the casting solutions

2.3.1 Cloud point determination

The cloud point (T_c) , i.e. the lower critical solution temperature (LCST), is referred to as the "phase separation temperature" of the casting solutions. Transparent homogeneous casting solution was poured into a test tube and heated slowly in a water or oil bath, as shown in Fig. 1. As the temperature increased, phase separation occurred at a certain temperature. The onset of turbidity was considered as an indication of phase separation.



Fig. 1. Schematics of the test of cloud point

2.3.2 Viscosity

The viscosities of the casting solutions in various compositions were measured with a DV-IIPRO Digital Viscometer (Brookfield, USA) at 25°C. The recorded data are the curve of viscosity to the shear rate. The reported data were the average value.

2.3.3 Light transmittance measurement

To investigate what is the major driving force during membrane-forming process, light transmittance experiments were carried out. The method was reported by Li et al [9]. The light

transmittance indicates the phase separation rate of the casting solution, and the intensity of the light transmittance through the membrane was recorded as a function of immersion time.

2.4. Hansen solubility parameters

The Hansen solubility parameter (δ_t , total cohesion solubility parameter) [53] is composed of three parameters (δ_d , δ_p and δ_h) to account for nonpolar interactions (_d), dipolar interactions (_p) and hydrogen bonding interactions (_h), respectively. The Hansen solubility parameter is defined by:

$$\delta_t^2 = \delta_d^2 + \delta_p^2 + \delta_h^2 \tag{1}$$

The solubility parameter of mixed solvents is defined as:

$$\delta_{i,s} = \frac{X_1 V_1 \delta_{i,1} + X_2 V_2 \delta_{i,2}}{X_1 V_1 + X_2 V_2} \tag{2}$$

where X is the molar fraction, V is the molar volume, $\delta_{i,1}$ (i=d, p, h) is the δ_t of the solvent and $\delta_{i,2}$ (i=d, p, h) is the δ_t of the non-solvent.

The difference in the solubility parameters between the mixed solvent and PES can then be described by [49]:

$$\Delta \delta_{s-p} = \left[\left(\delta_{d,s} - \delta_{d,p} \right)^2 + \left(\delta_{p,s} - \delta_{p,p} \right)^2 + \left(\delta_{h,s} - \delta_{h,p} \right)^2 \right]^{1/2}$$
(3)

where $\delta_{i,s}$ (i=d, p, h) are the Hansen solubility parameters of the mixed solvent and $\delta_{i,p}$ (i=d, p, h) are the Hansen solubility parameters of PES. The values of the PES, solvent and non-slovent are listed in Table 3.

Components	$\delta_{\rm d} \\ ({\rm MPa}^{1/2})$	$\frac{\delta_p}{(MPa^{1/2})}$	δ_{h} (MPa ^{1/2})	$\frac{\delta_t}{(MPa^{1/2})}$	$\Delta \delta_{s-p}$ (MPa ^{1/2})
PES	17.60	10.40	7.80	21.90	-
DMAc	16.80	11.50	10.20	22.70	2.76
DEG	16.60	12.00	20.70	29.12	13.04
PEG200*	17.97	6.06	17.30	25.67	10.45
PEG300*	17.91	4.73	15.09	23.89	9.24
PEG400*	17.88	4.00	13.85	22.96	8.81

Table 3 Hansen solubility parameters for PES, solvent and non-solvent.

*Solubility parameters of PEG200, PEG300 and PEG400 were calculated by the group contribution method [54].

2.5. Preparation of the PES membranes

Flat sheet membranes were prepared by employing the RTIPS method. During the membrane formation process, deionised water was used as a coagulation bath, the temperature of which is shown in Table 4, and adjusted to the cloud point. The casting solution was cast onto a clean glass plate, using a casting knife with a gap of 300 μ m, and then immersed into a pure water bath quickly, in order to form a membrane. The residual mixed solvent in the prepared membranes was extracted with ethanol for 2 days. The ethanol was renewed every 12 h. These wet membranes were then dried for 2 days at room temperature, to produce dry membranes for testing.

Mambrona number	Turnes of DES	Water bath temperature
Memorane number	Types of PES	(°C)
MDEG1-10-20	PES51000	20
MDEG1-10-50	PES51000	50
MDEG1-12-20	PES51000	20
MDEG1-12-50	PES51000	50
MDEG1-14-50	PES51000	50
MDEG1-15.5-50	PES51000	50
MDEG1-17a-50	PES51000	50
MDEG2-17-50	PES62000	50
MDEG3-17-50	PES77000	50
MPEG200-4-20	PES51000	20
MPEG200-4-80	PES51000	80
MPEG300-4-50	PES51000	50
MPEG300-4-80	PES51000	80

Table 4 Temperature of the membrane formation.

2.6. Characterisation of the PES membranes

2.6.1. Morphology

The morphologies of the membranes were investigated by scanning electron microscopy (SEM; S-3400 II, Hitachi High-Technologies, Japan). The cross-sections of the prepared membranes were fractured in liquid nitrogen and sputtered with gold in a vacuum.

2.6.2. Permeation performance

Pure water and BSA aqueous solution (300 mg·L⁻¹) were used as the feed solutions. The testing system was a dead-end stirred cell, which has been reported by Li et al. [9]. All tests were carried out at room temperature with a feed pressure of 0.1 MPa. Before the pure water permeation flux (J_w) and the rejection rate (R) of BSA aqueous solution were measured, the modules were pre-pressured at 0.1MPa with pure water for 0.5 h. The BSA contents of the feed and the permeate solutions were

determined by an UV spectrophotometer (UV-300, Shimadzu, Japan), and then the J_w and R for each membrane sample were determined three times and averaged. The J_w and the R are defined as follows [49]:

$$J_{w} = \frac{V}{A \times T} \tag{4}$$

$$R = (1 - \frac{C_P}{C_F}) \times 100\% \tag{5}$$

where J_w is the pure water permeation flux $(L \cdot m^{-2} \cdot h^{-1})$, V is the permeate pure water volume (L), A is the effective area of the membrane (11.34 cm²), T is the permeation time (h). R is the rejection rate of BSA (%) and C_P and C_F are the BSA contents of the feed and the permeate solution (wt. %), respectively.

2.6.3. Porosity and pore size

Membrane porosity ε (%) is the ratio of the total volume of pores in the membrane to the bulk volume of the membrane. In this paper, it was determined by the dry-wet weight method [55]. The porosity was calculated as:

$$\varepsilon = \frac{(m_w - m_D)/\rho_{ethanol}}{(m_w - m_D)/\rho_{ethanol} + m_D/\rho_p}$$
(6)

where m_W is the weight of the wet membrane, m_D is the weight of the dry membrane, ρ_p is the density of PES (1.370 g·cm⁻³), and $\rho_{ethanol}$ is the density of ethanol (0.790 g·cm⁻³).

Mean pore size (r_m) was calculated via the filtration velocity method. As described by the Guerout-Elford-Ferry equation [56], r_m can be expressed by the following equation:

$$r_m = \sqrt{\frac{(2.9 - 1.75\varepsilon) \times 8\eta hQ}{\varepsilon \times A \times \Delta P}}$$
(7)

where ε is membrane porosity, η is the viscosity of water (8.9×10⁻⁴ Pa·s), Q is pure water flux (mL·s⁻¹), h is membrane thickness (mm), A is the effective area of the membrane (11.34 cm²) and ΔP is the feed

pressure (0.1 MPa).

2.6.4. Mechanical properties

The tensile strength, Young's modulus and elongation at break of the prepared membranes were measured by a material tensile testing machine (QJ-210A, Shanghai Qingji Instrumentation Science and Technology Co. Ltd., China). The tensile loading speed was 50 mm·min⁻¹, and the gauge length was 50 mm. Each membrane sample was measured at least three times and then averaged.

3. Results and discussion

3.1. Phase diagram of PES/DMAc/DEG system

The effects of PES molecular weight on phase separation temperature and viscosity are shown in Table 5. The cloud point showed similar values when PES with significant molecular weights was used. Phase separation of the casting solutions with LCST is mainly induced by decreasing hydrogen-bonding interactions between PES and mixed solvent with an increase in the temperature of the casting solution. The difference in PES molecular weight is not attributed to the hydrogen-bonding interactions. Furthermore, the experimental results confirm that phase separation temperature has little to do with PES molecular weight. In addition, the viscosities of the three casting solutions were shifted to higher viscosities, as the PES molecular weight increased as expected.

The phase diagram and the viscosity for PES/DMAc/DEG systems with different PES contents are shown in Fig. 2, in which it is evident that the cloud point decreased as PES concentration increased. The result can be explained by the intermolecular interaction forces between PES and mixed solvent (DMAc/DEG). In spite of the incompatibility between PES and non-solvent DEG, PES can dissolve in the mixed solvent (DMAc/DEG) and maintain stability at room temperature, which is due to the good compatibility between PES and solvent DMAc. The casting solution is LCST system, meaning

that hydrogen-bonding interactions change poorly between DMAc and DEG, in line with increasing the temperature of the casting solution. Correspondingly, the apparent concentration of DEG (non-solvent), which does not bond to solvent (DMAc), increases until it is higher than a critical concentration at a certain temperature. At this point, the poor interactions between PES and DEG dominate the casting solution, while DEG separates out as a non-solvent rich phase and PES precipitates from the casting solution.

This indicates that it is easier to induce phase separation at lower temperature, when PES concentration increases. The viscosities are illustrated in Fig. 2, indicating that the viscosity shifts higher as PES concentration increases.

Casting solutions	Types of PES	$T_{c}(^{\circ}C)$	Viscosity (mPa·s)
MDEG1-17a	PES51000	27	1535
MDEG2-17	PES62000	26	1689
MDEG3-17	PES77000	26	3361

Table 5 Cloud points and viscosities of the PES/DMAc/DEG casting solutions.



Fig. 2. Phase diagram and viscosity of the PES/DMAc/DEG system. Types of PES: PES51000,

DEG/DMAc (wt./wt.): 1:1.

3.2. Estimation of the cloud point for the PES/DMAc/DEG system

The cloud points of different PES/DMAc/DEG systems were estimated. From equations (1), (2) and (3), the difference in solubility parameters ($\Delta \delta_{s-p}$) between the mixed solvent and PES was calculated and is listed in Table 6. It is evident that the δ_{H} increases when increasing the mass ratio of DEG/DMAc, whereas δ_{D} and δ_{P} change very little, thereby indicating that the interactions between PES and mixed solvent are influenced mainly by a hydrogen-bonding interaction. Fig. 3 shows the experimental cloud point temperatures for the casting solutions of PES in DEG/DMAc, and it is notable that the cloud points decrease in line with the increase in $\Delta \delta_{s-p}$. In addition, there is a good linear relationship (r²=0.9980) between the experimental cloud points and the $\Delta \delta_{s-p}$, and the linear regression of the data gives the relation:

$$T_c = 89.5 - 9.1 \Delta \delta_{s-n}$$

where T_c is the cloud point.

From Fig. 3, the experimental results show fair agreement with the linear regression line, thereby suggesting that the mass ratio of DEG/DMAc can be conveniently adjusted to control the phase separation temperature. Based on the experiments and estimations, it is evident that the mass ratio of DEG/DMAc can be applied as a key processing parameter for controlling the thermodynamic properties of casting solutions and provides the necessary theoretical basis for PES membrane preparation via the RITPS method.

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(8)

PES and mixed solvent	DEG:DMAc (wt.:wt.)	$\delta_D\!/MPa^{1/2}$	$\delta_P/MPa^{1/2}$	$\delta_{\text{H}}/\text{MPa}^{1/2}$	$\delta_t\!/MPa^{1/2}$	$\Delta \delta_{s\text{-}p}/\text{MPa}^{1/2}$
PES	-	17.60	10.40	7.80	21.90	-
DEG/DMAc	0.7:1	16.75	11.62	12.82	24.08	5.23
DEG/DMAc	0.75:1	16.74	11.64	13.14	24.26	5.55
DEG/DMAc	0.8:1	16.74	11.65	13.44	24.42	5.84
DEG/DMAc	0.9:1	16.73	11.68	13.89	24.68	6.28
DEG/DMAc	0.95:1	16.72	11.69	14.22	24.87	6.61
DEG/DMAc	1:1	16.72	11.70	14.48	25.03	6.86

Table 6 Solubility parameters for PES and mixed solvent.



Fig. 3. Relationship between $\Delta \delta_{s-p}$ and cloud points for the PES/DMAc/DEG system. The

concentration of PES is 17wt.%.

3.3 Cloud points of PES/DMAc/PEG systems

The effects of PEG molecular weight and mass ratio of non-solvent/solvent (PEG/DMAc) on cloud point are shown in Fig. 4. Firstly, for PES/DMAc/PEG200, PES/DMAc/PEG300 and PES/DMAc/PEG400 systems, the cloud points showed similar trend of decreased cloud point with an increase of mass ratio of PEG/DMAc. The reason for this is the same as that of the PES/DMAc/DEG

systems. Due to good compatibility between PES and DMAc, PES can dissolve in the mixed solvent (DMAc/PEG) and keep stability at room temperature in spite of the incompatibility between PES and PEG (non-solvent). When the amount of PEG increases, the interactions between PES and the mixed solvent become weaker, and consequently the casting solution is easier to induce phase separation within at lower temperature. This indicates that the cloud point can be controlled by the mass ratio of non-solvent/solvent.

Secondly, it is obvious that the cloud point increases with increasing the PEG molecular weight. From the data in Table 3, it can be seen that the $\Delta \delta_{s-p}$ between PES and PEG decreases when the PEG molecular weight increases. This means that the intermolecular interaction forces between PES and mixed solvent become stronger as PEG molecular weight increases. Thus, phase separation of the casting solution will occur at higher temperature.



Fig. 4. Cloud points of different PES/DMAc/PEG systems.

3.4. Estimation of the cloud point for the PES/DMAc/PEG system

The cloud points of PES/DMAc/PES200, PES/DMAc/PEG300 and PES/DMAc/PEG400 systems were estimated, respectively. $\Delta \delta_{s-p}$ between the mixed solvent (PEG200/DMAc, PEG300/DMAc and

PEG400/DAMc) and PES was calculated and is listed in Table 7. From the data in Table 7, it can be seen that δ_H increases and δ_P decreases when increasing the mass ratio of PEG200/DMAc, whereas δ_D changes only a little. The PEG/DMAc/PEG300 and PES/DMAc/PEG400 systems have the same pattern as the PES/DMAc/PEG200 systems. These observations indicate that the interactions between PES and mixed solvent are controlled by dipolar and hydrogen interactions.

Fig. 5 shows the experimental cloud points for the PES/DMAc/PES200, PES/DMAc/PEG300 and PES/DMAc/PEG400 casting solutions, and it is evident that the cloud points decrease when increasing $\Delta \delta_{s-p}$. There is a good linear relationship between the experimental cloud point and $\Delta \delta_{s-p}$, and the linear regression equations of the data are listed in Table 8. These linear relationships indicate that the mass ratios of PEG200/DMAc, PEG300/DMAc and PEG400/DMAc in a simple way can be used to control the cloud point. It is evident that the mass ratio of non-solvent/solvent (PEG/DMAc) is a key processing parameter for controlling the thermodynamics of the casting solutions and is applied as a theoretical estimation of phase separation temperature for the PES membrane preparation via RITPS method.

Furthermore, it can be seen that the linear regression line shifts to higher cloud points as the PEG molecular weight increases from 200 to 300. When the PEG molecular weight further increases from 300 to 400, the shift of the linear regression line is not obvious. The difference is, however, attributed to the difference of $\Delta \delta_{s-p}$ between PES and PEG200, PEG300 and PEG400. From the data in Table 3, it can be seen that δ_H and δ_P decrease when increasing the PEG molecular weight, whereas δ_D changes only a little. This observation indicates that the molecular interactions between PES and PEG are controlled by dipolar and hydrogen interactions. As shown in Table 7, the difference of $\Delta \delta_{s-p}$ is 1.21 when the PEG molecular weight increases from 200 to 300, and the difference of $\Delta \delta_{s-p}$ is 0.43 when

the PEG molecular weight increases from 300 to 400. This means that the effects of PEG molecular weight on intermolecular interaction forces between PES and mixed solvent become weaker as PEG molecular weight increases. Thus, the shift of the linear regression line is an interplay between several factors.

PES and mixed	PEG200:DMAc	$\delta_{\rm D}/{\rm MPa}^{1/2}$	$\delta_{\rm P}/{\rm MPa}^{1/2}$	$\delta_u/MPa^{1/2}$	$\delta/MPa^{1/2}$	$\Delta \delta_{\rm m} / MPa^{1/2}$
solvent	(wt.:wt.)	0 <u>D</u> / 1011 a	0p/1 111 d	0 _H /1011 u	o _t / Will a	Δ0 _{S-p} / 1 v Π α
PEG200/DMAc	1.4:1	17.27	9.29	13.08	23.58	5.41
PEG200/DMAc	1.6:1	17.36	8.90	13.60	23.78	5.99
PEG200/DMAc	2:1	17.49	8.29	14.39	24.12	6.92
PEG200/DMAc	2.3:1	17.57	7.91	14.89	24.35	7.51
PEG200/DMAc	3:1	17.70	7.34	15.63	24.72	8.41
PEG200/DMAc	4.5:1	17.83	6.71	16.46	25.18	9.42
PEG300/DMAc	2:1	17.34	8.18	12.60	22.94	5.29
PEG300/DMAc	3:1	17.55	6.87	13.55	23.21	6.75
PEG300/DMAc	4:1	17.67	6.12	14.08	23.41	7.60
PEG300/DMAc	5:1	17.74	5.69	14.39	23.55	8.10
PEG300/DMAc	6:1	17.78	5.43	14.58	23.63	8.41
PEG300/DMAc	7:1	17.81	5.26	14.71	23.69	8.61
PEG300/DMAc	8:1	17.83	5.14	14.79	23.73	8.75
PEG400/DMAc	2.3:1	17.33	7.80	12.00	22.48	4.95
PEG400/DMAc	3:1	17.47	6.86	12.46	22.53	5.85
PEG400/DMAc	4.5:1	17.65	5.59	13.08	22.67	7.14
PEG400/DMAc	7.3:1	17.78	4.71	13.51	22.82	8.06
PEG400/DMAc	8:1	17.79	4.60	13.56	22.84	8.18

Table 7 Solubility parameters of PES and mixed solvent.



Fig. 5. Relationship between $\Delta \delta_{s-p}$ and cloud points for the PES/DMAc/PEG system. The

concentration of PES is 17wt.%

System	Linear regression equation	r^2
PES/DMAc/PEG200	T _c =202.6-17.8Δδ _{s-p}	0.9981
PES/DMAc/PEG300	T _c =208.5-16.6Δδ _{s-p}	0.9924
PES/DMAc/PEG400	$T_c = 175.3 - 11.5 \Delta \delta_{s-p}$	0.9984

Table 8 Linear regression equations of different PES/DMAc/PEG systems.

In addition, the viscosities of different PES/DMAc/PEG systems are shown in Fig.6, showing that the viscosity increases as PEG molecular weight and $\Delta \delta_{s-p}$ (mass ratio of non-solvent/solvent) increase. Combining with Fig. 4 and Fig. 6, it can be seen that if the molecular weight of PEG continues to increase, the cloud point and viscosity of the casting solution will increase and the membrane formation process is difficult to operate.



Fig. 6. Viscosities of different PES/DMAc/PEG systems.

In general, based on the above discussion, it is evident that by changing the mass ratio of a non-solvent/solvent or PEG molecular weight, the phase separation temperature can be estimated, which in turn can provide theoretical guidance for PES membrane preparation via the RITPS method. The method allows for a quick estimation of the cloud point.

3.5. Membrane morphology

The SEM micrographs of PES flat sheet membranes, prepared by the NIPS and RTIPS processes, are shown in Fig. 7. The cloud point of the casting solution MDEG1-12 is 31.7°C. When the water bath temperature was 20°C, which was lower than the cloud point, the membrane formation process was dominated by the NIPS process. The membrane MDEG1-12-20, with finger-like pores, was formed in the cross-section, and instantaneous demixing was induced by the inflow of pure water, as shown in Fig.8. When the water bath temperature was 50°C, which is higher than the cloud point, the membrane formation process was dominated by the RTIPS mechanism. Membrane MDEG1-12-50 with a bi-continuous structure was prepared, representing the typical morphology of membranes with

high flux, as shown in Fig.8. Since the membrane formation process is induced by the formation of a translucent membrane via mass traansfer or heater transfer, the velocity of light transmittance changes is directly related to the phase separation rate. To understand the difference between the heat transfer rate and mass transfer rate, light transmittance experiments were conducted. As shown in Fig. 9, the light transmittance of casting solution MDEG1-12 at 20°C changed more rapidly than that of the casting solution MDEG1-15.5 at 20°C. This is due to the water bath temperature being 20°C, which was lower than the cloud point, and the casting solution MDEG1-12 possesses the low viscosity (as shown in Fig. 2), which means fast mass transfer. Thus the NIPS process is the dominating membrane formation process and finger-like pore was formed. Meanwhile, when the water bath tempeature was 50°C, which is higher than the cloud point, the slope of the initial part of the light transmittance curve for the casting solution MDEG1-12 and MDEG1-15.5 increase rapidly and there is no slope difference between the two casting solutions. If the NIPS is the dominating process, the phase separation rate of MDEG1-15.5 should be lower than that of MDEG1-12. The observation indicates that the phase rate of heat transfer is independent of the viscosity. This comfirmed that the heat tranfer rate is much faster than the mass transfer rate and RTIPS is the dominating process. Thus, heat transfer is the major driving force of membrane formation when the temperature is higher than the cloud point, which results in the formation of a bi-continuous structure.



Fig. 7. SEM micrographs of PES flat sheet membrane cross-sections.



Fig. 8 Schematic diagram of membrane formation for NIPS and RTIPS method.



Fig. 9. Light transmittance curves of casting solutions.

SEM images of PES membranes prepared with different PES contents via the RTIPS process are shown in Fig. 10. As shown in Fig. 2, the cloud points of the casting solutions with different PES contents are between 27°C and 33°C. When the membrane formation temperature is 50°C, which is higher than the cloud points, the membrane formation process is controlled by the RTIPS mechanism, because of the main driving force of phase separation being heat transfer instead of mass transfer. All

the membranes are presented as a bi-continuous structure, as shown in Fig. 10, which confirms the membrane-forming mechanism is RTIPS process [48,51].



Fig. 10 SEM micrographs of PES flat sheet membrane cross-sections prepared via the RTIPS process.

(L) Cross-section and (R) enlarge cross-section.

The SEM micrographs of PES membranes prepared with different PEG molecular weights via the NIPS and RTIPS processes are shown in Fig. 11. As shown in Fig. 4, the cloud point of the casting solution MPEG200-4 is 67.4°C. When the water bath temperature was 20°C, which was lower than the cloud point, NIPS was the dominating process and the membrane MPEG200-4-20 with finger-like pores was formed in the cross-section. When the water bath temperature was 80°C, which was higher than the cloud point, RTIPS was the dominated process and the membrane MPEG200-4-80 with bi-continuous structure was presented. The cloud point of the casting solution MPEG300-4 is 67.0°C. Similarly, when the membrane formation temperature is 50°C, which is lower than the cloud point, the membrane formation temperature is 80°C, which is higher than the cloud point, the membrane formation temperature is 80°C, which is higher than the cloud point, the membrane formation temperature is 80°C, which is higher than the cloud point, the membrane formation temperature is 80°C, which is higher than the cloud point, the membrane formation temperature is 80°C, which is higher than the cloud point, the membrane formation temperature is 80°C, which is higher than the cloud point, the membrane formation temperature is 80°C, which is higher than the cloud point, the membrane MPEG300-4-80 with a bi-continuous structure was formed.



Fig. 11. SEM micrographs of PES flat sheet membrane cross-sections prepared with different PEG

molecular weights.

In summary, based on the above SEM results, it is evident that a change in the membrane formation

mechanism and the control of membrane structure have been relised by changing only the membrane-forming temperature.

3.6. Permeation properties, porosity and pore size

The determined permeation parameters, the porosity and mean pore size of PES membranes prepared with different PES concentration, water bath temperature and mixed solvent, are shown in Fig. 12 and Fig. 13. Firstly, for MDEG1-12-50, MPEG200-4-80 and MPEG300-4-80, the membrane formation mechanism is the RTIPS process, and J_w is higher than for the corresponding membranes (MDEG1-12-20, MPEG200-4-20 and MPEG300-4-50) prepared by the NIPS process (see Fig. 12a and Fig. 12d). As shown in Fig. 13(a), the MDEG1-12-50 membrane has a smaller r_m than that of the MDEG1-12-20 membrane, so the rejection rate of the membrane MDEG1-12-50 is a higher value (see Fig. 12a). From Fig. 13(d), it is notable that the MPEG200-4-80 and MPEG300-4-80 membranes, which are prepared by the RTIPS process, have a larger r_m and rejection rate (R) than the corresponding membranes (MPEG200-4-20 and MPEG300-4-50) prepared with the NIPS process. Based on these experiments, it can be seen clearly that PES membranes prepared by the RTIPS method can be applied as a new tool for preparing membranes with high flux and a high rejection rate, thereby reflecting the advantages of RTIPS method.

However, for the MDEG1-10-50 membrane, J_w is 126 L·m⁻²·h⁻¹, which is less than the value of 299 L·m⁻²·h⁻¹ of the MDEG1-10-20 membrane, as shown in Fig. 12(a). According to the experimental results, it is evident that the membrane MDEG1-10-50 (prepared by the RTIPS process) has lower pure water flux than that of MDEG1-10-20 membrane prepared by employing the NIPS process. The result is inconsistent with the advantages of the RTIPS method for preparing high flux membranes. According to the membrane-forming mechanism of RTIPS, when the membrane-forming temperature

 $(50^{\circ}C)$ is higher than the cloud point $(33^{\circ}C)$ of the casting solution, the membrane formation mechanism should be dominated by the RTIPS process. Membrane MDEG1-10-50 with a bi-continuous structure was prepared indeed (Fig. 10). In reality, the PES concentration of the casting solution MDEG1-10 is low (10 wt.%,), the tensile strength of the membrane MDEG1-10-20 is poor (0.4 MPa, as shown in Fig. 13). The finger-like pore membrane prepared at low PES concentration is more vulnerable to damage under the test pressure. Therefore, the membrane MDEG1-10-20 showed higher apparent J_w and r_m.

Secondly, the J_w of the membranes prepared by the RTIPS process reaches peak value for the MDEG1-14-50 membrane following an increase in PES concentration (see Fig. 12b), because the viscosity of the MDEG1-14 casting solution is great enough for phase separation via RTIPS and lower than the viscosities of the casting solutions MDEG1-15.5 and MDEG1-17. The phase separation rate between PES-rich phase and PES-poor phase is influenced by the viscosity of the casting solution and reaches equilibrium at 14 wt.%. Thus, the MDEG1-14-50membrane has good pore connectivity, which is embodied in a maximum r_m value (see Fig. 13d). In addition, the porosity of the membranes prepared by the RTIPS process decreases in line with an increase in PES concentration, as shown in Fig. 13(b), due to an increase in solid content.

Thirdly, the J_w and r_m of membranes prepared via the RTIPS process in Fig. 12 and Fig. 13 present a downtrend in line with an increase in PES molecular weights in the following descending order: MDEG1-17a-50, MDEG2-17-50 and MDEG3-17-50. The reason for this can be attributed mainly to viscosity differences in the casting solutions. Higher viscosity means a lower rate of phase separation. Accordingly, r_m is small. From the results in Fig. 13(c) it can be seen clearly that the porosity (ϵ) remains almost constant.



Fig. 12. Pure water permeation flux and rejection rate of the prepared PES membranes.



Fig. 13. Porosity and mean pore size of the prepared PES membranes.

3.7. Mechanical properties

The mechanical properties of PES membranes prepared with different PES concentration, water bath temperatures and mixed solvent are presented in Fig. 14. For MDEG1-10-20, MDEG1-12-20, MPEG200-4-20 and MPEG300-4-50, the membrane formation mechanism is the NIPS process, in which the tensile strength, elongation at break and the Young's modulus are lower than for the corresponding membranes (MDEG1-10-50, MDEG1-12-50, MPEG200-4-80 and MPEG300-4-80), which are prepared with the RTIPS process (see Fig. 14a and Fig. 14d). As shown in Fig. 7, Fig. 10 and Fig. 11, the membrane morphology changes from a finger-like pore structure to a bi-continuous structure when the water bath temperature is higher than the cloud point, and these results in an improvement to the mechanical properties of the membrane.

As Fig. 14(b) shows, the tensile strength and the elongation at break of the membranes prepared with the RTIPS process increase in line with PES content in the following ascending order: MDEG1-10-50, MDEG1-12-50 and MDEG1-14-50. The membrane structure becomes dense following an increase in PES content, which results in an increase in tensile strength and elongation at break. The tensile strength and the elongation at break reached their peak values at 17 wt.% PES content. Furthermore, Fig. 14(c) shows that the tensile strength and the Young's modulus of the membranes MDEG1-17a-50, MDEG2-17-50 and MDEG3-17-50 increase in line with an increase in PES molecular weight. This indicates that the strength and the stiffness of the membranes were enhanced with the increase in PES molecular weight, which is common behavior for polymers.

The membrane MDEG1-17a-50, which was prepared from a mixed solvent of DEG/DMAc, showed higher tensile strength and elongation at break than the membranes MPEG200-4-80 and MPEG300-4-80, which were prepared from PEG200/DMAc or PEG300/DMAc mixed solvent (seen

Fig. 14d). This result may be due to the molecular weights of PEG200 and PEG300, which are much greater than DEG. A small amount of PEG200 or PEG300 embedded in the membrane may result in a decrease in tensile strength and elongation at break due to plasticisation effects.



Fig. 14. Mechanical properties of PES membranes.

4. Conclusions

An estimation of phase separation temperature, based on the Hansen solubility parameter, was performed, and then PES flat membranes were prepared via an RTIPS process. As for casting solutions, DMAc was used as the solvent, while DEG, PEG200, PEG300 and PEG400 were used as the non-solvent. The PES/solvent/non-solvent casting solutions were LCST systems, and the cloud point was shown to be independent of PES molecular weight. With the increase in PES content, the cloud point of the prepared casting solutions decreased and the viscosities increased. When the PEG molecular weight increases, the cloud point and the viscosities of the casting solution increased. The calculated Hansen solubility parameters show that there is a good linear relationship between the experimental cloud points and the $\Delta \delta_{s-p}$. The composition and membrane-forming temperature of the casting solution are the key casting parameters for PES membrane preparation via the RITPS process, and they can be adjusted according to the requirements.

On the one hand, when the membrane-forming temperature was lower than the cloud point, the membrane formation mechanism was the NIPS process. A finger-like pore structure was present in the membrane. On the other hand, when the membrane-forming temperature was higher than the cloud point, the membrane formation mechanism was mainly dominated by the RTIPS process, and a bi-continuous structure was found in the membranes. Those membranes prepared by employing the RTIPS process exhibited higher pure water permeation flux than that of the membranes prepared with the NIPS process, except for the MDEG1-10-50membrane. The porosity of the membranes prepared by the RTIPS method decreased with an increase in PES concentration, while pure water permeation flux and the mean pore size of membranes prepared by the RTIPS process decreased in line with an increase in PES method are reflected by the relatively

high flux and high rejection rate compared to the NIPS method.

With the increase in PES molecular weight and content, membrane strength improved. Membranes prepared by the DEG/DMAc mixed solvent, showed higher tensile strength and elongation at break than for membranes produced by a PEG200/DMAc or PEG300/DMAc mixed solvent. When the membrane formation mechanism was the RTIPS process, tensile strength, the elongation at break and the Young's modulus were higher than for the corresponding membranes prepared by the NIPS process. It is concluded that the advantage of the RTIPS process is the easy preparation of membranes with symmetrical bi-continuous morphology resulting in good mechanical properties.

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Highlights

- > A group contribution estimate of phase separation temperature for RTIPS was proposed.
- > Phase separation temperatures were linearly dependent on Hansen solubility parameters.
- > PES membranes with bi-continuous structure were acquired from RTIPS process.

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> PES membranes prepared by RTIPS showed good permeation and mechanical properties.