Soft Matter

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Cite this: *Soft Matter*, 2024, 20, 6834

Received 16th April 2024, Accepted 15th July 2024

DOI: 10.1039/d4sm00449c

rsc.li/soft-matter-journal

Ionic nanoporous membranes from selfassembled liquid crystalline brush-like imidazolium triblock copolymers†

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There is a need to generate mechanically and thermally robust ionic nanoporous membranes for separation and fuel cell applications. Herein, we report a general approach to the preparation of ionic nanoporous membranes through custom synthesis, self-assembly, and subsequent chemical manipulations of ionic brush block copolymers. We synthesized polynorbornene-based triblock copolymers containing imidazolium cations balanced by counter anions in the central block, side-chain liquid crystalline units, and sidechain polylactide end blocks. This unique platform comprises: (1) imidazolium/bis(trifluoromethanesulfonyl)imide (TFSI) as the middle block, which has an excellent ionexchange ability, (2) cyanobiphenyl liquid crystalline end block, a sterically hindered hydrophobic segment, which is chemically stable and immune to hydroxide attack, (3) polylactide brush-like units on the other end block that is easily etched under mild alkaline conditions and (4) a polynorbornene backbone, a lightly crosslinked system that offers mechanical robustness. These membranes retain their morphology before and after backbone crosslinking as well as etching of polylactide sidechains. The ion exchange performance and dimensional stability of these membranes were investigated by water uptake capability and swelling ratio. Moreover, the length of the carbon spacer in the imidazolium/TFSI central block moiety endowed the membrane with improved ionic conductivity. The ionic nanoporous materials are unusual due to their singular thermal, mechanical, alkaline stability and ion transport properties. Applications of these materials include electrochemical actuators, solid-state ionic nanochannel biosensors, and ion-conducting membranes. **PAPER**
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1. Introduction

Well-designed ion-conducting polymer membranes move charge via the transport of ions rather than electrons and function as $(poly)$ electrolytes,¹ electrochemical actuators,² and ion exchange membranes for applications in demineralization of salt solutions, 3 desalination, 3 industrial waste treatment, 4 fuel cells,⁵ selective separation of different anions by electrodialysis,⁶ and high pollution environments.⁷ An essential strategy to obtain high ion conductivity and excellent performance includes having densely functionalized ionconducting groups on a hydrophobic polymer matrix creating ion channels and dimensional stability.^{8,9} Tethering the ion

exchange on regularly spaced flexible side-chains improves ion conductivity rather than attaching the ion-exchanging head group directly to the polymer backbone. $8,10-12$ In principle, lengthening the spacer group should improve the ionic conductivity by supporting the formation of phase-separated morphology.^{13,14} This distinct phase separation occurs due to the enthalpy related to the demixing of the hydrophobic polymer backbone with the hydrophilic ionic side chains, typically observed in block copolymer architectures.

Block copolymers containing ion-conducting groups in the main chain and side chains enable the creation of ion conducting membranes or ionic nanochannels (10–100 nm) through control of their microstructures due to clustering of ionic groups and microphase segregation of incompatible domains.15–17 Furthermore, the composition of the block copolymers, the type and concentration of ionic groups, the presence of spacers, and the morphology and alignment of the ion-conducting domains will impact the thermal properties, mechanical properties, ionic conductivity and ion exchange capacity of these ion-containing block copolymers.¹⁸⁻²⁰ Also,

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[†] Electronic supplementary information (ESI) available. See DOI: [https://doi.org/](https://doi.org/10.1039/d4sm00449c) [10.1039/d4sm00449c](https://doi.org/10.1039/d4sm00449c)

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in this architecture, the polymer backbone being protected by the hydrophobic polymer matrix is stable against alkaline $media.^{21,22}$

Another promising approach towards the design of nanochannels with high ion conductivity is to improve ionic mobility. The self-assembly of liquid crystalline polymers is a versatile approach that improves conductivity by creating well-defined and ordered ionic channels.²³⁻²⁶ Although the incorporation of liquid crystals (LC) to obtain an ordered alignment of ionic clusters is somewhat limited, initial success on designing ion transporting self-assembled soft materials forming 1D, 2D, and 3D ionic pathways has been recently reported.^{24,27–29} Thus, the design and structure–property investigation of ion-containing liquid crystalline block copolymers, currently unknown, will be a unique platform for the preparation of ion-conducting materials due to (i) formation of ionic domains from ion-containing groups (imidazolium and pyrimidinium groups), (ii) liquid crystalline ordering, anisotropic ordering and responsive properties and (iii) microphase segregation of block copolymer architecture.

In our previous studies, we used sequential ring opening metathesis polymerization (ROMP) of norbornene monomers to synthesize liquid crystalline brush-like block copolymers (LCBBCs) and explored their phase behavior.³⁰⁻³² The advantage of these LCBBCs over conventional liquid crystalline block copolymers is that the introduction of the brush-like side results in tailored meso and micro-structured morphology of these polymers, reduced entanglement and viscosity as well as enhancement of alignment kinetics with or without directed field assemblies. The majority block is composed of polynorbornene bearing side-chain liquid crystalline cyanobiphenyl units, which facilitate the formation of a smectic mesophase. $33-35$ The minority block is composed of polynorbornene bearing poly(D,L-lactide) (PLA) brush-like units, which can be selectively etched by hydrolysis under gentle alkaline

conditions. The selective removal of PLA blocks results in a functionalized nanoporous surface. 32 The use of the polynorbornene backbone allows the system to be crosslinked and aligned with field-directed assembly, including shear and magnetic alignment, and by using this method, mechanically robust nanoporous membranes could be obtained. Based on this design principle, we recently reported the synthesis and properties of nanoporous membranes from norbornenebackbone-based triblock copolymers containing cyanobiphenyl and polylactide side chain units as end blocks and protected amine or thiol containing monomers in the central block. 32 These amine and thiol groups can ligate to heavy metal ions. The resulting nanoporous membranes are used for heavy metal ion removal applications.^{32,36,37}

In principle, the design strategy developed for liquid crystalline nanoporous membranes for metal ion removal can be applied to create a new brush-like liquid crystalline imidazolium functionalized triblock copolymer platform with stable ionic channels and high ion exchange capacity (Fig. 1). Selfassembly within these compression molded films, followed by thiolene crosslinking chemistry of polynorbornene olefinic backbones and mild alkaline selective etching of PLA, presents nanoporous membranes (illustrated in Fig. 1) containing ionconducting groups with high ion exchange capacity (IEC) and excellent ionic conductivity. The modular platform discussed in this paper is one of its kind: (1) imidazolium/TFSI as the middle block is alkaline stable, has high IEC and ionic conductivy, $38(2)$ inclusion of the majority cyanobiphenyl block not only offers natural alignment capabilities but also creates a steric hindrance effect, hence chemically stable and OH-immune conductive nanochannels are produced and (3) the lightly crosslinked hydrophobic polynorbornene system from which PLA side chains can be easily etched under mild alkaline conditions and create alkaline stable ion-conductive nanochannels for potential use in fuel cells and water electrolysis.37,39–42 Paper

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Fig. 1 Schematic illustration of a new nanoporous membrane with high IEC. The design strategy for this system involves (a) sequential block polymerization of three different norbornene monomers (NBCB12 (liquid crystalline unit), NBIm_mX (imidazolium unit) and NBPLA (brush-like moiety)), (b) resulting self-assembled triblock copolymer system comprising NBCB12 and NBPLA end blocks and NBIm_mX middle block with a crosslinked backbone and (c) etching out PLA to create nanochannels through which ion exchange is realized. The nanoporous membrane produced lacks longrange order and hexagonal arrangements of the pores.

2. Results and discussion

2.1. Synthesis of monomers

The monomers $NBCB12^{23}$ and $NBPLA2K^{32}$ are synthesized according to previously reported procedures (Fig. S1 and S2, ESI†). New imidazole-functionalized norbornene monomers are synthesized using modified literature procedures and details are provided in Scheme 1.⁴³

Briefly, in a dried round-bottomed flask equipped with a magnetic stir bar, 5-norbornene-2- carboxylic acid (3.6 g, 0.026 mol) and 6-bromo-1-hexanol (4 g, 0.022 mol) are dissolved with the help of 30 mL of CH_2Cl_2 . Then EDC (6.35 g, 0.033 mol) and a catalytic amount of DMAP and pyridine mixture are added to the reaction mixture stirring at room temperature for 12 h, then the reaction is quenched by adding water (30 mL). The crude product is extracted three times with $CH₂Cl₂$, and the organic layer is dried with anhydrous $MgSO₄$. Then the organic layer is filtered and concentrated by a rotary evaporator to afford an oily product. The crude product is further purified by column chromatography using hexanes/EtOAc (20 : 1) as the mobile phase and silica gel as the stationary phase to yield a yellow viscous oil, NB ${\rm (CH_2)_6Br}$ (5.04 g, 76%). ¹H NMR (500 MHz, CDCl₃): δ 6.17-5.85 (m, 2H), 4.01 (t, 2H), 3.36 (t, 2H), 3.17 (s, 1H), 2.99–2.85 (m, 2H), 2.13–2.22 (m, 3H), 1.92–1.79 (m, 2H), 1.68–1.21 (m, 7H) (Fig. S3, ESI†). The same steps were followed in the synthesis of $NB(CH_2)_{12}Br$, the only slight change was the use of 12-bromo-1-hexanol in place of 6-bromo-1-hexanol. Soft Matter

2. Results and discussion

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2.1.1. Synthesis of 5-norbornene-2-carboxylate-1-hexyl-3-methyl-imidazolium. $NB(CH_2)_6Br$ (5.0 g, 0.016 mol) and

1-methylimidazole (1.75 mL, 0.024 mol) are refluxed in acetonitrile for 18 h. After cooling to room temperature, the reaction mixture is concentrated using a rotary evaporator to afford yellow viscous oil and then washed with diethyl ether $(3 \times 100 \text{ mL})$ to obtain (NBI₆mBr) (4.46 g, 80% yield) (Fig. S3, ESI[†]). The resulting oil is dissolved in $H₂O$ (100 mL), LiTFSI (4.88 g, 0.017 mol) is added, and the mixture is stirred at room temperature for 12 h.

The yellow oil is extracted with dichloromethane and the organic layer is washed with water and dried over anhydrous $MgSO₄$ to obtain yellow oil, NBIm₆Fs (4.63 g, 75%) (Fig. S4, ESI†). ¹H NMR (500 MHz, CDCl₃): δ = 8.69 (s, 1H), 7.30 (d, 2H), 6.13–5.86 (m, 2H), 4.11 (t, 2H), 3.96 (t, 2H), 3.71 (s, 3H), 3.15 (s, 1H), 1.84 (m, 2H), 1.82 (m, 3H), 1.58–1.55 (m, 2H), 1.35– 1.24 (m, 7H). HRMS: calculated ${M⁺$ 303.43, observed 303.20 (Fig. S6, ESI†). The same procedure was followed in the synthesis of the NBI m_{12} Fs monomer, and it was characterized with ¹H NMR (Fig. S5, ESI†) and HRMS: calculated ${M_1^+}$ 387.59, observed 387.30 (Fig. S7, ESI†).

2.2. Synthesis of triblock copolymers and preparation of crosslinked ionic membranes

Ring-opening metathesis polymerization (ROMP) of norbornene-functionalized monomers is carried out as shown in Scheme 2. In a typical example, the feed ratio of NBCB12 : NBIm_{(6 or 12})X (X = Br or Fs): NBPLA2K is 75:5:20 wt%, which is expected to result in a triblock copolymer with cylinders decorated with imidazolium ions within the polymer

Scheme 1 Synthesis of intermediate compounds and monomers, (NB(CH₂₎₆Br), NB(CH₂₎₁₂Br, 5-norbornene-2-carboxylate-1-hexyl-3-methylimidazolium bromide (NBIm₆Br), 5-norbornene-2-carboxylate-1-dodecyl-3-methyl-imidazolium bromide (NBIm₁₂Br), 5-norbornene-2-carboxylate-1-hexyl-3-methyl-imidazolium bis((trifluoromethyl)sulfonyl)amide (NBIm6Fs), and 5-norbornene-2-carboxylate-1-dodecyl-3-methyl-imidazolium bis((trifluoromethyl)sulfonyl)amide (NBIm₁₂Fs).

matrix.³² NBCB12 (300 mg, 0.6 mmol) dissolved in 10 mL of $CH₂Cl₂$ is added to a round-bottomed flask equipped with a magnetic stir bar and a rubber septum and purged with nitrogen for 10 min. In a scintillation vial, modified Grubbs 2nd generation (12.5 mg, 0.018 mmol) is dissolved in CH_2Cl_2 (5 mL) and purged with nitrogen for 10 min. The solution is then added to the round bottom flask containing $NBCB₁₂$ and the reaction is allowed to proceed for 30 min at room temperature. After the completion of first polymerization, the pre-purged CH_2Cl_2 solution containing NBIm_(6 or 12)X (X = Fs or Br, 24 mg, 0.05 mmol) is added and stirred for a further 2 hours. Once the completion of second polymerization, the pre-purged CH_2Cl_2 solution containing NBPLA2K (106 mg, 0.05 mmol) is added and stirred for a further 30 minutes. Then, the reaction is terminated by adding excess ethyl vinyl ether (EVE). The resulting polymer is precipitated into a methanol–isopropanol mixture, filtered, and dried overnight in a vacuum oven at room temperature for later use (Table 1).

The successful synthesis of brush-like triblock copolymers is a result of the salient features of ROMP such as its living polymerization, precise molecular weight control, low molecular weight dispersity, and tolerance to an adverse range of functional groups, for example in this case imidazolium groups. The molar composition of the block copolymers is confirmed by ¹H NMR spectroscopy. The characteristic proton

Table 1 Sample description (E = etched, $XL = crosslinked$, $XLE =$ crosslinked-etched, subscript indicates the length of the carbon spacer)

absorption peaks for the imidazolium ring appear at 10.2 ppm in NBIm₆Br and 8.9 ppm in NBIm₆Fs (Fig. S8–S10, ESI†). When the norbornene monomer is polymerized, the peak broadens but remains at the same position (5.7–6.3 ppm). The successful synthesis of triblock copolymers is confirmed by ${}^{1}H$ NMR

 a The % composition calculated from ¹HNMR for TBIm_(6 or 12)Br shows a discrepancy from the theoretical value. This could be as a result of the peak used for the case of imidazole at 10.2 ppm for TBIm($_{6 \text{ or } 12}$ Br and 8.9 ppm for TBIm($_{6 \text{ or } 12}$)Fs, respectively. The peak is broad due to the surrounding environment composed of Br– or TFSI–. All other peaks overlap with those of NBCB12 and PLA. ^b Theoretical molecular weight of the synthesized triblock copolymers, TBI($_{6 \text{ or } 12}$)Br and TBI($_{6 \text{ or } 12}$)Fs. ϵ Mn determined by GPC relative to polystyrene standards using the ELSD detector with THF as an eluent. $\frac{d}{dx}$ The molecular weight distribution or the polydispersity index (*Ð*).

spectroscopy. Furthermore, the molar compositions of these triblock copolymers are determined, and the related images and results are included in Table 2 and Fig. S8, S9, ESI.† The target theoretical molecular weight of all the triblock copolymers is set at 25 kDa. The GPC results indicate that the terpolymers possess a high degree of polymerization and number average molecular weights of up to \sim 23 kg mol⁻¹ (Table 3) and a narrow dispersity (D) of 1.1-1.3.

2.2.1. Crosslinking representative protocol. 200 mg of the triblock copolymer (TBImBr), 5 mg of 2-hydroxy-4'-(2hydroxyethoxy)-2-methylpropiophenone, and 12 mg of 1,10 decanedithiol are weighed in separate vials covered with aluminum foil. 1 mL of THF is added to the bottles to dissolve the contents and they are mixed thoroughly. The solvent is then evaporated by purging with nitrogen for a few minutes and vacuum dried overnight. A thin film of the sample is produced by compression molding at 82 \degree C and quenched to room temperature, followed by UV crosslinking for 3 hours resulting in a crosslinked film. The gel fraction of a crosslinked sample in THF is determined to be \sim 42%. We followed our previous reported methods to prepare nanoporous membranes decorated with imidazolium ions whose brief details are described in the supporting information (ref. 32).

2.3. Thermal properties

Thermal properties of both triblock copolymers before and after crosslinking and etching are analyzed by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC), respectively. Both libraries of terpolymers exhibit excellent thermal stability with 5% weight loss from TBIm₆Fs at \sim 297 °C, TBIm₆Br at \sim 260 °C, TBIm₁₂ Br at \sim 262 °C, and TBIm₁₂Fs at \sim 265 °C (Fig. 2 and Fig. S12, ESI†). Interestingly, the crosslinked and etched terpolymers, $TBIm₆Br-XLE$ $(\sim 324 \text{ } ^\circ\text{C}), \quad \text{TBIm}_6\text{Fs-XLE} \quad (\sim 343 \text{ } ^\circ\text{C}), \quad \text{TBIm}_{12}\text{Br-XLE}$ (\sim 338 °C), and TBIm₁₂Fs-XLE (\sim 349 °C) exhibit much higher thermal stability compared to non-crosslinked $(TBIm₆Br,$ TBIm₆Fs, TBIm₁₂Br, and TBIm₁₂Fs), and crosslinked terpolymers, TBIm₆Br-XL (\sim 310 °C), TBIm₆Fs-XL (\sim 316 °C), TBI m_{12} Br-XL (\sim 302 °C), and TBI m_{12} Fs-XL (\sim 311 °C), respectively (Fig. 2 and Fig. S12, ESI†). The thermal stability of the crosslinked and etched terpolymer polymers is promising for membrane applications. Soft Matter **Fraction Content Content Content Content Common Content Content Content Content Common Content Common C**

Thermal mesophase transitions of triblock copolymers before and after crosslinking, and PLA etching are determined using DSC. All the samples are heated up to $150\degree C$ to remove thermal history and the first cooling cycle is used to interpret the data. These copolymers exhibit glass transition

N.B.: all measurements are measured in triplicates and the average values are recorded. E – etched, XL – crosslinked, XLE – crosslinked and etched/ PLA removed.

Fig. 2 Thermogravimetric analysis of triblock copolymers before (TBIm₁₂Br and TBIm₁₂Fs) and after crosslinking (TBIm₁₂Br-XL and TBI m_{12} Fs-XL), and after PLA etching (TBI m_{12} Br-XLE and TBI m_{12} Fs-XLE), respectively.

 $(T_{\rm g})$ temperature around \sim 17–22 °C for the non-crosslinked triblocks (TBIm₆Br, TBIm₆Fs, TBIm₁₂Br and TBIm₁₂Fs), and \sim 25–28 °C for the crosslinked (TBIm₆Br-XL, TBIm₆Fs-XL, $TBIm_{12}Br-XL$ and $TBIm_{12}Fs-XL$), and etched (TBIm₆Br-XLE, $THIm_6Fs-XLE$, $THIm_{12}Br-XLE$ and $THIm_{12}Fs-XLE$) copolymers, respectively (Fig. 3a and b). The liquid crystalline transition temperature, which is also the liquid crystalline clearing temperature (T_{LC}) for both non-crosslinked and crosslinked

triblock copolymers was around \sim 73–77 °C (Fig. 3a and b), and T_{LC} does not change much after crosslinking and etching (Fig. 3a and b), indicating that post polymerization modifications of crosslinking and etching PLA do not significantly alter the microstructures of these systems.

2.4. Mechanical properties

Dynamic mechanical analysis (DMA) was applied to elucidate the effect of crosslinking on the mechanical performances of the ionic nanoporous membranes. Fig. 3c and d illustrate the tan δ (loss factor/tangent) for the non-crosslinked (TBIm₆Fs and $TBIm_{12}Fs$) copolymers and crosslinked copolymers (TBI m_6Fs - XL and $TBIm₁₂Fs-XL$, respectively, as a function of temperature. The tan δ peak maxima values correspond to the thermal properties observed from DSC, the values indicate comparable glass transition temperatures that are consistent with the DSC results. For example, tan δ maxima occurred at \sim 23 °C for TBIm₆Fs, \sim 31 °C for TBIm₆Fs-XL (Fig. 3c), \sim 22 °C for TBIm₁₂Fs, and \sim 30 °C for TBIm₁₂Fs-XL, respectively. The second tan δ peaks observed at \sim 60–70 °C are associated with liquid crystalline clearing temperature (T_{LC}) of the copolymers and the results are in good agreement with the DSC results (Fig. 3a–d). The crosslinked samples exhibit slightly higher tan δ values, an indication of successful crosslinking, superior energy dissipation from the tougher networks and denser chain Paper West Common Source 2022. The the effective companies are considered under the considered under the common state of the common state are common state and children are common state and children are common state and co

Fig. 3 (a) First cooling cycle DSC thermograms for triblock copolymers before (TBIm₆Br and TBIm₆Fs), after crosslinking (TBIm₆Br-XL and TBIm₆Fs-X), and PLA etching (TBIm₆Br-XLE and TBIm₆Fs-XLE membranes). (b) First cooling cycle DSC thermograms for triblock copolymers before (TBIm₁₂Br and TBIm₁₂Fs), after crosslinking (TBIm₁₂Br-XL and TBIm₁₂Fs-XL), and PLA etching (TBIm₁₂Br-XLE and TBIm₁₂Fs-XLE membranes) showing glass transition (T_g) and LC transition temperatures (T_{LC}). (c) and (d) Variation of loss factor, tan δ with temperature for non-crosslinked (TBIm_(6 or 12)Fs) and crosslinked $(TBIm_{(6 or 12)}Fs-XL).$

Fig. 4 (a) Variation of storage modulus and loss modulus with temperature for TBIm₆Fs-XL and TBIm₆Fs as a control sample. (b) DMA curves for TBIm₁₂Fs-XL and TBIm₁₂Fs as control copolymers. (c) Stress–strain curves for TBIm₆Br, TBIm₁₂Br, TBIm₆Br-XL, and TBIm₁₂Br-XL. (d) Stress–strain curves for TBIm₆Fs, TBIm₁₂Fs, TBIm₆Fs-XL, and TBIm₁₂Fs-XL. The stress–strain behaviors of neat TBIm₆Br, TBIm₁₂Br, TBIm₁₆Fs, and TBIm₁₂Fs copolymers were compared as control samples.

entanglements in the systems. The DMA storage and loss moduli curves for TBIm₆Fs, TBIm₆Fs-XL, TBIm₁₂Fs, and $TBIm₁₂Fs-XL$ are shown in Fig. 4a and b, respectively. Both the storage and loss moduli curve trends depict the thermal transitions (T_g and T_{LC}) for the copolymers. All copolymers exhibited a glass transition region and above it there are rubbery plateaus with crosslinked films having a higher rubbery modulus than non-crosslinked, an indication of successful formation of tough chain networks in the systems. In general, there is a slight decrease in the storage modulus in the $T_{\rm g}$ region, a sharp decline at > 50 °C (T_{LC} region) and sample yielding > 75 °C (isotropic region) (Fig. 4a and b). The storage modulus values at low temperatures for $TBIm₆Fs-XL$ samples were slightly higher as compared to those of $TBIm_{12}Fs-XL$. The 12-carbon spacer in $TBIm_{12}Fs-XL$ could have increased the plasticity in the system that led to flexible chains and elastomeric nature in the systems (Fig. 4c and d). This is demonstrated in the tensile tests (stress–strain curves), and the crosslinked samples had higher toughness and stiffer behaviors as compared to non-crosslinked samples (Fig. 4c and d). Both TBIm₆Br-XL and TBIm₆Fs-XL were stiffer as compared to $TBIm_{12}Br-XL$ and $TBIm_{12}FS-XL$ samples (Fig. 4c and d).

As shown in Fig. 4c and d, the non-crosslinked TBIm_{6}Br , and TBIm12Br had a low yield strength of 1.4 MPa and 0.3 MPa, respectively, as compared to crosslinked samples $\text{TBIm}_{6}\text{Br-XL}$ and TBI m_{12} Br-XL which had a yield strength of 3.6 MPa and

3.3 MPa (\sim 157% and 1000% increase in yield strength), respectively. After the introduction of TFSI counterions in both non-crosslinked (TBIm₆Fs and TBIm₁₂Fs) and crosslinked samples (TBIm₆Fs-XL and TBIm₁₂Fs-XL), there was a significant enhancement of the yield strength (Fig. 3c and d). This enhancement can be attributed to additional physical crosslinking from TFSI ion–ion interactions. This in turn reinforces the chemical crosslinking/mechanical properties of the membranes as it can be witnessed in the high yield strength for both TBI m_6 Fs-XL (13.8 MPa) and TBI m_{12} Fs-XL (11.5 MPa) (Fig. 4c and d). Fig. S16 and S17 (ESI†) show the tensile and viscoelastic properties for the etched membranes, $TBIm_{12}Br-XLE$ and TBI m_{12} Fs-XLE, respectively. As indicated in Fig. S16 (ESI†), the tensile strength for the etched membranes was in the range of 0.25–0.4 MPa, and these values are less than those for the unetched crosslinked membranes. We speculate that the introduction of pores in the nanoporous membranes might have influenced the films' uniformity/thickness, thus affecting their mechanical properties. Tensile properties depend entirely on the uniformity of the films and hence, the stretching capacity in the etched membranes was likely reduced as a result of the size and distribution of the pores in the films leading to low yield strength. However, Fig. S17 (ESI†) clearly shows that even after etching, the targeted nanoporous membranes retain their mechanical integrity at temperatures below and above their glass transition temperatures. The Tan delta curves maxima for the etched membranes, $TBIm_{12}Fs-XLE$ and $TBIm_{12}Br-XLE$ occurred at \sim 25 and \sim 29 °C and were comparable to those of unetched crosslinked membranes and those observed from DSC.

In general, the crosslinked copolymers with 12 spacer imidazole monomers and TFSI⁻ ions exhibited a slight decrease in tensile strength, and a slight increase in elongation (large strain break) as compared to both non-crosslinked and crosslinked copolymers with 6 spacer imidazole monomers and Br ions, indicating tougher, robust and more flexible nanoporous membranes. Hence, crosslinking leads to a dense, tough, and continuous network which in turn promotes stress transfer and significant mechanical improvement in the systems. $44,45$

The triblock copolymer composition and architecture are designed in such a way that cylindrical morphology in the targeted membranes is obtained and retained. The use of the norbornene backbone is to allow easy thiolene crosslinking chemistries to provide the required rigidity to support the nanopores in these systems. The PLA block is introduced as a sacrificial block by easily etching it using mild basic conditions leaving behind unaltered morphology in the final membrane. Without crosslinking, the stability of the nanopores produced in these systems would be quite limited because of the Laplace pressure. It is therefore necessary to make the system mechanically robust enough to prevent the pore collapse before the nanopores are generated. When crosslinking LC based systems, one should always aim to obtain a mechanically robust system while ensuring that the system retains their morphologies and liquid crystalline properties.

2.5. Microstructural analysis

The crosslinking chemistry and surface chemistry of robust, crosslinked nanoporous membranes (after PLA removal) are confirmed by SEM imaging (Fig. 5). The porous nature of the

Fig. 5 Comparison of SEM images of gold sputter-coated triblock copolymers (a) before crosslinking (TBIm $_6$ Br) and (b) after crosslinking and PLA etching (TBIm₆Br-XLE), (c) before crosslinking (TBIm₆Fs) and (d) after crosslinking and PLA etching (TBIm₆Fs-XLE).

Fig. 6 FTIR spectrum of the TBIm $_6$ Fs triblock copolymer before crosslinking, after crosslinking, and etching of the crosslinked nanoporous samples.

membrane is revealed by SEM where Fig. 5a and Fig. S13a (ESI†) show unetched samples and Fig. 5b and Fig. S13b (ESI†) show crosslinked etched nanoporous membranes. The pore size for the nanoporous membranes $(TBIm₆Fs-XLE$ and TBIm₁₂Fs-XLE) from SEM was \sim 30 \pm 3 nm and is in good agreement with the SAXS results and our previous studies (Fig. 7).^{32,46} A slight difference exists between the pore size and domain spacing. A similar observation has been noted by Seungwan Cho et al. This may be attributed to the enhanced chain mobility in the elevated temperature during etching, leading to a partial collapse of some LC domains during hydrolysis.⁴⁶ During crosslinking, the vibrational and stretching bands from the PLA moiety are restricted, thus the peak at 1752 cm⁻¹ disappears and the intensity of the peak at 1173 cm^{-1} is reduced relative to other peaks (Fig. 6). The presence of a broad amine peak is indicated by N–H stretching at 3294.3 and 3373.5 cm^{-1} , and N–H bending at 1602 cm^{-1} .³² The accessibility of functional groups on the surface of nanoporous membranes is revealed by the intensity of these peaks (Fig. S14 and S15, ESI†). The persistence of the C–O stretching ester peak at 1247 cm^{-1} indicates the chemical stability of the triblock terpolymer. The CN stretching vibration observed at 2225 cm^{-1} is attributed to the cyano group from the liquid crystalline moiety. Paper Weight exched on 29 and 29 Yuly 2024. Because with a spectral on 2022. The method of the Creative Common and the Creative Common and the Creative Commons are the common and the common and the spectral in gradient co

> Thus, the ionic nanoporous membrane (triblock copolymer) has several unique features: (1) the imidazolium/TFSI as the middle block is alkaline stable, (2) the cyanobiphenyl block is sterically hindered and hydrophobic and resistant to attack from hydroxide anions that are also used to etch PLA and (3) the lightly crosslinked system is easily etched under alkaline conditions, which only removes PLA but leaves all other chemical constituents intact and creates mechanically stable ion-conductive nanochannels.

2.6. Influence of the membrane microstructure on ion exchange capacity

We previously reported that the norbornene-based liquid crystalline diblock brush copolymer, LC73-PLA27, with composition

Fig. 7 SAXS profile for the triblock copolymers showing the retention of morphology before and after crosslinking/etching.

comprising NBCB12 of 73 wt% and NBPLA2K of 27 wt% selfassembles into PLA cylinders within majority LC phase from which cylindrical pore nanoporous membranes are derived upon

Furthermore, the porous structure of the membrane may not be continuous through the thickness of the compressionmolded triblock copolymer before etching and after etching and crosslinking, respectively. The lack of long-range order of the domain and discontinuous nature of the porous structure while the membrane is mechanically robust and elastomeric are hypothesized to be advantageous when ion exchange studies are carried out.

2.7. Water uptake and swelling ratio

Water uptake (WU) and swelling ratio (SR) are significant parameters in evaluating anion exchange membrane performance. The membrane films are initially soaked in deionized water for 24 h and weighed to obtain the hydrated membrane mass. The films are subsequently dried under a vacuum for 24 h at 80 \degree C and weighed to obtain their dry weight. Similarly, the swelling ratios are obtained by measuring the length of the films after soaking in deionized water and after vacuum drying. Water uptake is calculated in percent using the following equation:

Water uptake =
$$
\frac{\text{mass of hydrated membrane} - \text{mass of dry membrane}}{\text{mass of dry membrane}} \times 100\%
$$
 (1)

Swelling ratio is calculated as follows:

Water uptake =
$$
\frac{\text{length of hydrated membrane} - \text{length of dry membrane}}{\text{length of dry membrane}} \times 100\%
$$
 (2)

removal of PLA.36 We also reported the synthesis of triblock copolymers bearing amine and thiol ligands in the central block, which are compression-molded, crosslinked and etched to remove PLA. The nanoporous membrane retains its morphology, an indication that crosslinking does not modify the alignment of the pores. Small angle X-ray scattering analysis of compression molded films of triblock copolymers before (TBIm $_6$ Fs and TBIm_{12} Fs) and after crosslinking/etching (TBIm₆Fs-XLE and TBI m_{12} Fs-XLE, respectively) is shown in Fig. 7.

The triblock copolymers before and after crosslinking and etching show a negligible shift in scattering reflections (q-range positions), which indicates that the microstructure of the liquid crystalline mesophase and the overall morphology of the triblock copolymer are largely unaltered. The SAXS profile of these samples suggests a cylindrical packing of PLA side chains within the smectic layered LC matrix. This is in agreement with the extensive studies that we have previously conducted on these systems and have shown that the full scattering peaks can be observed if the system is aligned under a magnetic field. Using the q^* indicated, the PLA domain decorated on the edges with imidazolium is \sim 29 nm, and the q_{LC1} indicates that the system is smectic A (SmA) consisting of interdigitated bilayers with a *d*-spacing of 5.51 nm where q_{LC2} is a higher reflection of q_{LC1} . Additionally, it can be deduced from the scattering pattern observed that there is no long-range hexagonal ordering of the PLA cylinders in the liquid crystalline domains.

The water uptake and swelling ratio of the membranes etched in 0.5 M NaOH are observed to increase with an increase in the hydrocarbon chain for both (TBIm_(6 or 12)Br and TBIm_(6 or 12)Fs) polymers (Table 3). This could be attributed to the architectural domains which could possibly allow water to percolate through the polymer matrix. Interestingly, TBI m_{12} Fs films exhibited the highest water uptake and swelling ratios, which are attributed to the network of well-connected bis(trifluoromethanesulfonyl)imide (TFSI) hydrophilic domains improving the films interaction within an aqueous medium. The TBI m_6 Br films with hydrophobic polymer domains showed comparatively less water uptake and swelling ratio which are mainly due to lower IEC. Moreover, the water uptake and swelling ratio of the crosslinked/etched films are lower than the etched films but higher than the non-etched films, an indication that crosslinking and etching improved the dimensional stability and the performance of the membranes, respectively. The performance and dimensional stability of the membranes are evaluated based on their water uptake and swelling ratio, thus higher water absorption and less swelling are effective ways to improve conductivity without losing mechanical stability.⁴⁷ Notably, the percent swelling ratio values are less compared to those reported in the literature, an indication that these membranes have remarkable dimensional stability endowed by the presence of cyanobiphenyl liquid crystalline end block, a sterically hindered hydrophobic segment, which is chemically stable and immune to hydroxide attack. The length of carbon spacers was observed to improve the water uptake and swelling ratio.

2.8. Ion exchange capacity (IEC) studies

Ion exchange capacity is a measure of the total ion exchanging sites or functional groups per gram of dry membrane. In liquid crystalline ionic materials, the morphology, domain size (5–10 nm), the counterion on the ion exchange membrane and orientation have a huge impact on ion exchange capacity and/ or ionic conductivity.³⁶ Similarly, block copolymer architecture has been shown to exhibit better ion exchange capacity compared to homopolymers and random copolymers because of the various morphologies that can be achieved through block copolymers which enhances ion transport.^{20,48,49} The incorporation of liquid crystalline units within brush-like block copolymers in this study creates defined ionic channels within a liquid crystalline matrix, which ultimately improves ion exchange capacity.^{50–52} Furthermore, optimal crosslinking chemistry of the polynorbornene backbone in this study allows for the design of ion exchange membranes with sufficient mechanical integrity as illustrated by the mechanical properties discussed in the previous sections while retaining high ion exchange capacity. This has been a technical challenge in conventional ion exchange membranes that this work addresses.10,53 The ion exchange capacity (IEC) of the thin films is determined by calibrated potentiometric measurements (Fig. S11, ESI†) and (Table S1, ESI†) and the IEC value is calculated as follows:

$$
IEC = \frac{\text{no. of moles of ions exchanged (mmol)}}{\text{dry weight of the membrane (g)}} \tag{3}
$$

The highest IEC is observed in membranes that are both etched and crosslinked where both membranes $\text{TBIm}_{12}\text{Br-XLE}$ and TBIm₁₂Fs-XLE are made up of \sim 5 wt% of the ion conducting block by composition. It should also be noted that by having flexible methylene spacers between the side chain moieties, the backbone microphase separation is enhanced and this cushions the membrane against mechanical collapse, especially in alkaline environments.⁵⁴⁻⁵⁷ Li and coworkers have reported an IEC value of 1.85 \times 10⁻³ mol g⁻¹ for quaternary ammonium groups grafted onto the norbornene backbone.⁵⁸ The observed IEC for this triblock copolymer is comparable to the other reported systems taking into account that the IEC values depend on the content of the ionic ion groups incorporated into the copolymer. Thus, in our present brush-like triblock which contains a small percentage of the ionic block (\sim 5 wt%) in the copolymer composition, the IEC observed, in this case, is significant and indicative of the number of ionic exchange sites available, highlighting the potential for use as ion exchange membranes or ion-conducting membranes.⁵⁹⁻⁶¹

The alkaline stability of the nanoporous ionic membrane is promoted by the presence of π -electrons from imidazolium and cyanobiphenyl groups. $62,63$ The chemical stability is further enhanced by the norbornene backbone, the alkyl spacer between the norbornene backbone and imidazolium group.^{64,65}

Our system is particularly intriguing due to the well-defined polymer morphology supporting both ion conductivity and membrane stability. Additionally, the presence of discontinuous pores facilitates the performance of the membrane via the formation of ion channels and increasing surface area for ion exchange. The IEC and ionic conductivity performance of the thin films immersed in an alkaline solution (0.5 M NaOH) for etching compared to the unetched films indicates that these films exhibit alkaline stability behavior.⁶⁶ Mckeown, Guiver and Xu et al. reported that microporous copolymer ionomer based AEM shows high ion conductivity suggesting that the microporosity of the charged copolymer matrix facilitates the ion transport.⁶⁷ Zhang et al. reported that semi-flexible poly(aryl ether sulfone) containing flexible aliphatic side-chains with imidazolium cations at the end shows high hydroxide conductivity (93 mS cm⁻¹ at 60 °C), highlighting the fact that the introduction of flexible side chains increases the mobility leading to better hydrophilic/hydrophobic microphase separation.⁶⁸ In summary, while several strategies have been developed over the years to achieve high conductivity and acceptable alkaline stability, our work presents a novel approach to the molecular engineering of mechanically, thermally and alkaline stable, nanoporous membranes with unusually high IEC despite comprising 5 wt% of the ion exchange moiety in the polymer architecture. Paper
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2.9. Ionic conductivity of thin films

The ionic conductivity of the thin films comprising only 5 wt% of the ion exchange moiety in the polymer architecture is evaluated using the Osilla Four Point Probe. The conductivity of the terpolymer with Br as the counterion increases from 1.41×10^{-5} to 2.32×10^{-5} . On the other hand, the conductivity of the terpolymer, $TBIm₆Fs$ with TFSI as the counterion increases from 2.34 \times 10⁻⁵ to 3.72 \times 10⁻⁵ (Table 3). Etching thin films helps create anchoring points for ions in the polymer matrix,⁶⁹ while crosslinking helps to introduce branched structures that provide a certain free volume to help build ion channels with lower transmission resistance.^{70,71} When compared to the terpolymers with Br as counterions, the terpolymers with TFSI as counterions showed higher ionic conductivity (σ) (Table 3). This could be attributed to the excellent ionexchange ability of the imidazolium/TFSI central block moiety and its hydrophilicity, which improves the membrane interaction with water. Notably, ionic conductivity also increases with the increase in carbon spacer groups due to improved hydroxyl interactions with the polymers' functional groups due to bond flexibility, improving the local mobility of cationic groups.⁷²

3. Conclusions

We describe the synthesis, properties, and high ion exchange capacity of a novel brush like liquid crystalline imidazolium functionalized triblock copolymer platform (as high IEC capacity nanoporous membranes) by ring-opening metathesis polymerization for potential use as an anion exchange membrane

for fuel cells and water electrolysis applications. The majority block contains a side-chain LC block that provides some degree of order that may be further enhanced due to their ability to align under the applied external field, and the PLA brush-like block allows the formation of nanopores decorated with ioncontaining imidazolium cations counterbalanced by anions. The triblock copolymers and subsequent crosslinked ionconducting nanoporous membranes are thermally stable with some nanoporous membranes such as $TBIm_{12}Fs-XLE$ having degradation temperatures at 5% weight loss, T_d , as high as 349 \degree C, well above the 150 \degree C, which is the typical operating temperature for fuel cells and water electrolysis applications. These crosslinked membranes are mechanically robust as has been illustrated by the detailed analysis of mechanical properties. The morphology of the nanoporous membranes was established by SEM and SAXS, and the size of the pores is determined to be \sim 32 nm, with the domain spacing ranging from \sim 28 to 30 nm. The morphological studies indicate that the nanoporous membrane may not be continuous throughout the thickness of the pores and lack long-range order. However, for optimal IEC and ionic conductivity, these morphological characteristics are highly desirable for these applications. IEC is determined using potentiometry, and the highest IEC is observed for films that are both crosslinked and etched, which suggests that the formation of nanopores facilitates ion conductivity and is not impeded by lack of long-range order. Ongoing studies are focused on methods to optimize longrange order using aligning capabilities of the cyanobiphenyl units as well as the construction of continuous nanopores and their impact on IEC and ionic conductivity of these new materials for AEM applications. Soft Matter

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Data availability

All of the data reported in this manuscript are available in tables and figures both in the main manuscript as well as the ESI.†

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

This work was supported by the NSF under grants DMR-1507045 and CMMI-1246804. The central instrumentation facilities at the Institute of Materials Science and Chemistry Department at UConn are acknowledged.

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