Microporous polymers with cascaded cavities for controlled transport of small gas molecules

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In membrane-based separation, molecular size differences relative to membrane pore sizes govern mass flux and separation efficiency. In applications requiring complex molecular differentiation, such as in natural gas processing, cascaded pore size distributions in membranes allow different permeate molecules to be separated without a reduction in throughput. Here, we report the decoration of microporous polymer membrane surfaces with molecular fluorine. Molecular fluorine penetrates through the microporous interface and reacts with rigid polymeric backbones, resulting in membrane micropores with multimodal pore size distributions. The fluorine acts as angstrom-scale apertures that can be controlled for molecular transport. We achieved a highly effective gas separation performance in several industrially relevant hollow-fibrous modular platform with stable responses over 1 year.

INTRODUCTION

The regulation of mass transport at interfaces via molecular engineering of membrane pores is crucial for improving product separation efficiency and productivity (1–3). Recent advances in microporous membrane materials, including polymers (1, 4, 5), graphene (6), zeolites (7), and metal–organic frameworks (8), have been made to meet the growing demands for energy-efficient molecular transport applications. However, the available pore sizes are still discrete and thus limited. As an alternative, the exploitation of hybrid microporous materials using a multistep approach has also been reported (9). In this case, the introduction of microporous materials into ultrathin membranes requires sophisticated techniques for production (10).

We streamlined the refinement and organization of pore size distributions, particularly broad or multimodal distributions typically found at membrane surfaces, via direct fluorination of polymer precursors. The fluorination process has been mainly applied for “nonporous” polymers like poly(phenylene oxide) (11), polysulfones (12, 13), and polyimides (PIs) (14); however, these attempts were unsuccessful to allow the hierarchical refinement of surface pores. There have also been reports on direct fluorination of typical microporous polymers like poly[1-(trimethylsilyl)-1-propyne] (15–17) but never on newer microporous materials developed during the past two decades, including thermally rearranged (TR) polymers (1) and polymers with intrinsic microporosity (PIMs) (18). We specifically targeted several TR polymers and PIMs (hereafter denoted c-TR and c-PIM, respectively; Fig. 1A to C; figs. S1 to S3; and table S1).

One representative nonporous polymer, a PI, was also introduced for comparison. We hypothesized that the surface pore size and its distribution in ready-made microporous membranes can be tailored via substitution of atoms bonded toward the interiors of the micropores, i.e., components of the inner rim of the pore walls. TR and PIM-1 backbones were considered because of their rigid aromatic segments that were directly connected with stiff heterocyclic rings, which hinder intra-chain rotation and thus help permanently maintain the desired microporous structures (4).

RESULTS

During the fluorine exposure, hydrogen atoms with kinetic diameters of 2.89 Å (19) linked to the TR and PIM-1 aromatic polymer backbones were gradually replaced with larger fluorine atoms (3.65 Å) (19). Control over pore sizes and distributions by elemental fluorine resulted in angstrom-scale “apertures” that can be manipulated to regulate mass transport, which is reminiscent of iris diaphragms in the eyes and in microscopes. Note that iris diaphragms in the eyes adjust the amount of light reaching the retina in response to environmental conditions by controlling the diameter of the pupil, from the completely “open” position of the aperture down to the target aperture size. A smaller pupil diameter means that less light passes through the pupils. We demonstrated that, after fluorination, the pore size ($\tau_3$) and pore population ($I_3$) (Fig. 1A and fig. S3) were continuously engineered, particularly at the uppermost TR membrane surface, for example, with radii from 3.36 ± 0.06 Å down to ~2.48 ± 0.07 Å and $I_3$ from 17.22 ± 1.36% to ~7.81 ± 1.39%, respectively, from slow beam positron annihilation lifetime spectroscopy (sb-PALS) (20).  

Pore size and its distributions in polymers are correlated with fractional free volume (FFV), and thus, changes in S parameter [the ratio of the integrated counts between energies of 510.3 and 511.7 keV (S width) to the total counts in Doppler broadening energy spectroscopy (DBES), proportional to FFV] (20) for the fluorinated, microporous, and nonporous membranes are investigated (Fig. 1B and figs. S4 and S5). After exposure, all fluorinated membranes show a decrease in $S$ parameters near surfaces down to ~1 μm, which agrees well with the observed decrease in $\tau_3$ and $I_3$. Apparently, the fluorinated membrane surface domains are composed of two different
sublayers: a deposition layer \( (L_D) \) at membrane top surfaces and a pore-regulated layer \( (L_R) \) at subsurfaces, respectively. Differences in \( S \) parameters in \( L_D \) and \( L_R \) for the fluorinated membranes were ascribed to differences in the probability of fluorine substitution reaction in polymers at the surface of membranes. Molecular fluorine has facile reactions with polymers at the membrane top surfaces, whereas only fluorine that penetrates through the membrane participates in the substitution reaction at the subsurface. This latter process experiences strong mass transfer limitations compared to the external surface reactions, indicating that there is a “cascading” effect on the pore size and distribution from the top surface through the membrane.

Note that pore regulation using molecular fluorine has not yet been realized in the literature due to the poor accessibility of the membrane subsurfaces \( L_R \) to fluorine vapor. The kinetic diameter of molecular fluorine is comparable to or slightly larger than the average nanospatial dimensions in conventional nonporous polymers \((21)\). This fact means that molecular fluorine cannot viably penetrate into nonporous matrices before self-limiting fluorine layers form on the top surface of the membrane \( (L_D) \). For example, nonporous P84 PI membranes had a \( L_R \) thickness \((92 \text{ nm}) \) ~3.8-fold thinner than microporous polymers \((c\text{-TR of } 351 \text{ nm} \text{ and } c\text{-PIM-1 of } 234 \text{ nm}, \text{respectively}; \text{fig. S5})\).

Our pore engineering approach is particularly well suited for microporous TR and PIM-1 polymers due to their peculiar structural features. Specifically, these polymers retain a microporous \( \tau_2 \) larger than molecular fluorine, thereby enabling fluorine penetration into the membrane subsurface where C-H to C-F substitution occurs.
However, the surface-localized molecular fluorine does not appreciably change the physicochemical characteristics of the underlying bulk membrane (e.g., scattering patterns, thermal stability, mechanical toughness, and optical properties; fig. S6). Consequently, c-TR and c-PIM-1 membranes show cascading characteristics in terms of the pore size over a thickness of ~1 μm from the surface into the film (Fig. 1C).

The thickness window of $L_R$ in c-TR gradually changes from 311 nm (30-min exposure) to 874 nm (300-min exposure), while that in c-PIM varies from 209 nm (30 min) to 683 nm (300 min) (fig. S4). In particular, the increment in thickness of the cascaded $L_R$ is more evident with increasing exposure time, which is ascribed to another unique feature of pristine TR and PIM-1 membranes, namely, bottleneck-shaped bimodal pore size distributions consisting of $\tau_3$ and $\tau_4$ ($\tau_3 < \tau_4$) (table S2). During fluorine exposure, $\tau_4$ cavities adjacent to $\tau_3$ “throats” in TR and PIM-1 membranes can facilitate molecular fluorine penetration farther into the matrices.

After subnanopore control by fluorination, the reductions in $\tau$, $I$, and the FFV are amplified in the “throat-like” pores ($\tau_3$) relative to the “cavity-like” pores ($\tau_4$) (Fig. 1D), ultimately reducing the average micropore size on the membrane surface. Computational studies reveal that collisions of TR polymers with molecular fluorine in throat-like pores are more probable than in cavity-like pores (Fig. 1E).

These selectively tuned bottlenecks, which are connected to the intact cavity-like pores in the cascaded microporous membranes, block transport of relatively large molecules and do not affect the mass flux of smaller molecules. This phenomenon leads to a remarkable improvement in molecular differentiation without a loss of mass throughput of the smaller species.

To conceptually elucidate the efficacy of the fluorine-manipulated pore structure in practical molecular separation problems, a molecular dynamics (MD) simulation was conducted on the basis of the pore size information revealed by sb-PALS (Fig. 1F and Supplementary Text). The unprecedented permeation capability of the membrane was first benchmarked by taking helium recovery from natural gas wells as an example. This separation has global importance as helium is a strategic resource (22, 23). Figure 1F presents ~9-fold increases in He/CH$_4$ diffusion selectivities for c-TR membranes, resulting from the substantial decline in diffusion of methane molecules by 95.4% (from $6.01 \times 10^{-7}$ cm$^2$/s to $2.8 \times 10^{-8}$ cm$^2$/s) due to the shrunken throat-like pores.

We further demonstrated that the membranes showed molecular cutoff behaviors by examining single gas permeation of He, H$_2$, CO$_2$, N$_2$, and CH$_4$ (Fig. 2). Upper bounds for perfluoropolymers (24) and nonperfluoropolymers (25), which represent a trade-off relationship between gas permeability and gas selectivity for intrinsic
polymeric membrane materials (26), were also provided to guide the eye. This pore tuning approach evidently enhanced the separation properties of all polymers in this study. However, tailorable separation performances can be achieved specifically in microporous TR and PIM-1 polymers: ~70-fold enhancement in natural gas processing applications (c-PIM-1) and ~38-fold (c-TR) versus ~1.7-fold (P84) but no critical decrease of more than 15% in He permeability for all polymers (Fig. 2B and table S3). Note that the enhanced sieving properties of c-TR and c-PIM-1 membranes for other gas separation applications, including H2/CH4, CO2/CH4, N2/CH4, H2/CO2, H2/N2, or O2/N2, were obtained simply by adjusting the reaction conditions (Fig. 2, C to F, and table S3). On the basis of the series resistance model (27), these results were ascribed to the prevailed enhancement in molecular cutoff behaviors in LR compared to LD (Supplementary Text and table S4). For example, H2/CH4 selectivity values of LD and LR for c-TR-A2-F60 were 102 and 1945, respectively. LR for c-TR-A2-F60, whose H2/CH4 selectivity was 19-fold higher than LD, significantly contributed to the enhancement in H2/CH4 selectivity of c-TR-A2 by 271 from 68. Likewise, H2/CH4 selectivity of LR for the direct fluorinated P84 membranes (973) rendered that of the resulting membrane to be 158 from 94; however, the increment (1.7-fold) was less noticeable than c-TR-A2 (4.0-fold). The difference in the increment was attributed to the shrunken throat-like pores in LR, which can be only observed in the microporous polymer membranes after subnanopore control by fluorination.

In addition, different types of TR membranes derived from diverse precursors [polyamides (28), poly(amide-co-pyrrolone)s (29), PIs (30), and cross-linked PIs (31)] were considered to assess their feasibility in our subnanopore control technique (Fig. 2B). The resulting TR analogs have different pore size distributions depending on the pore size in the precursor membranes (28, 31): cross-linked PIs > PIs > poly(amide-co-pyrrolone)s > polyamides. In particular, TR membranes from polyamides were deliberately investigated because their throat-like pores were even smaller than molecular fluorne. Our concept was successfully realized for all microporous TR membranes after exposure. However, the impact of the pore regulation approach on the tailored transport and separation performance was evidenced for the TR derivatives with larger pore sizes. For example, the highest He/CH4 selectivity of ~790 (3733%) was obtained for TR-cross-linked PI derivatives compared to 21 for the unmodified pristine sample. TR-amide derivatives showed a He/CH4 selectivity of 312 (198%), compared to 70 for the pristine sample, implying the importance of polymer selection. At the same time, an apparent decrease in He permeability for TR-cross-linked PI derivatives (~17%) compared with TR-amide derivatives (~1%) was observed. Likewise, a more modest increase in He/CH4 was observed in PIM-1, in which the throat-like pores are larger than TR polymers (table S2). These results suggest that sophisticated size control over throat-like pores in parent materials can directly regulate molecular differentiation behavior.

Polymer membranes can be fabricated into modules (32, 33), and the pores in TR and PIM-1 hollow fibers are obviously modified for thicknesses of the effective selective layers down to 300 nm (Fig. 3, A and B). Their multimodal pore size distributions can also be observed as those in flat sheet membranes (figs. S7 and S8). Accordingly, c-TR

Fig. 3. Toward practical applications. (A) Schematic of a c-TR hollow fiber membrane module. Each color corresponds to the case of Fig. 1. (B) Electron probe x-ray microanalysis (EPMA) micrographs of cross sections of the c-TR hollow fiber morphology. For c-PIM-1, graded domain would be positioned on shell side. (C to E) Practical feasibility of using pristine (black) and c-TR (red) modules in helium separation applications. Solid ones are aged for 2000 hours. (C) He/CH4 performances of c-TR and c-PIM-1. For PIM-1, black-colored open star indicates the pristine, while blue-colored open diamond is a pore-regulated one. Other hollow-fibrous or asymmetric flat sheet membrane modules made of cellulose acetates (55), PIs (34, 53, 56), perfluoropolymers (35, 53, 57, 58), and carbon molecular sieve materials (34) were also provided for a comparison. Detailed data are given in table S5. (D and E) He recovery and purity dependence in a He/CH4 mixture fed with 1% helium under 20 bar at ambient temperature. Arrows and dotted lines guide the eye to indicate 90% He recovery. Solid lines are from a predictive process model, while solid bullets are from experimental results. The blue pentagons correspond to results from thin-film composite modules made of commercially available perfluoropolymers (fig. S10) (53).
and c-PIM-1 hollow fiber membranes still presented enhancement in separation performances for all of the applications in which we also considered with a flat sheet geometry (Fig. 3C, fig. S9, and table S5). In particular, c-TR hollow fiber modules exhibit orders-of-magnitude higher helium selectivities compared to the same modules made of commercially available PIs (34) and perfluoropolymers (35). We estimated the practical applicability of the c-TR modules, focusing on evaluating the helium purity dependence with recovery in a He/H₂ mixture fed with 1% helium. These experimental results were then compared to the calculated ones from a predictive process model based on a numerical method (Fig. 3, D and E; fig. S10; and table S6), and we successfully achieved reliable and long-term operation until ~9000 hours (or 1 year) without failure (fig. S9C).

We extend this pore-reorganizing concept in other fields including energy-related applications like fuel cells, reverse electrodialyzers, or water splitting where external water management systems are required (36). Focusing on difference in kinetic diameters of water vapor (2.65 Å) versus hydrogen (2.89 Å) or nitrogen (3.64 Å) can be acquired (37). Focusing on difference in kinetic diameters of water or water splitting where external water management systems are re-quired (38). Preliminarily, practical feasibility in H₂O/H₂ for c-TR and c-PIM-1 hollow fiber membranes was assessed and they exhibited a selectivity improvement by ~2.9- and ~3.2-fold, respectively (table S7), also indicating a universal tool for efficient and specific applications.

**DISCUSSION**

We demonstrated here that tailoring the membrane surface architecture with elemental fluorine provides a clear approach to address performance trade-off relationships between productivity and separation efficiency, especially in natural gas processing and other important gas separation applications as well. Experimental pore size distributions and gas separation experiments were successfully combined with conceptual MD simulations as well as the simplified series resistance models for heterogeneous membrane surfaces. This combination demonstrated the impact of the shrunken throat-like pores localized to the membrane subsurface on the conspicuous enhancement in gas separation performances. Toward practical applications, our approach can additionally provide two more benefits in the fabrication steps: (i) scalable production with membrane uniformity due to precise control of the reactive domain and (ii) no need to use expensive perfluoropolymers and to recover expensive perfluorinated solvents after fabrications. It can also be applied to absorptive materials, catalysts, and interfaces to control mass transport. Future efforts using this approach can be extended to microporous materials, sensing materials or catalysts with monolithic or hybrid polycrystalline moieties (such as the zeolitic imidazole framework and covalent organic frameworks or their mixed matrix membranes).

### MATERIALS AND METHODS

#### Materials

2,2’-Bis(3-aminophenyl) hexafluoropropane (bisAPAF) was purchased from Central Glass Co. Ltd. (Tokyo, Japan), and 3,3’-di(p-carboxyphenyl) (HAB) was obtained from Wakayama Seika Kagyo Co. Ltd. (Wakayama, Japan). 2,4,6-Trimesityl-m-phenylenediamine (DAM) was obtained from Chemtec (China; sublimed twice and stored under argon). 3,5-Diaminobenzoic acid (DABA; Aldrich) was recrystallized from oxygen-free water and sublimed before use. All diamines in this study were stored in a vacuum oven at 40°C before preparation. 3,3’-Diaminobenzidine (DAB) was purchased from Aldrich and purified by vacuum sublimation at 85° to 90°C. Isophthaloyl dichloride (IPC) and terephthaloyl dichloride (TPC) were purchased from Aldrich. After sublimation, DAB, IPC, and TPC were stored in a glove box purged with argon. 4,4’-(Hexafluoro- roisopropylidenedi) phthalic anhydride (6FDA) and 2,2-bis(4-3,4-dicarboxyphenoxono) phenyl]propane dianhydride (BPADA) were purchased from Daikin Industries Ltd. (Osaka, Japan). 4,4’-Oxydiphthalic anhydride (ODPA) was purchased from Chemtrec. 6FDA, BPADA, and ODPA were dried in a vacuum oven at 100°C for 24 hours to remove the absorbed water. 5,5’,6,6’-Tetrahydroxy-3,3,3’3’-tetramethyl-1,1’-spiropiobisindane (TTTSBI; Alfa Aesar) was purified by crystallization from methanol. 2,3,5,6-Tetrafluoroterephalonitrile (TFTPN; Aldrich) was purified by vacuum sublimation at 150°C under an inert atmosphere before use. For synthesis of PIM-1 for hollow fiber spinning, TTTSBI was purified by crystallization from methanol and TFTPNN (from Sung Young Chemical Technology) was purifed by vacuum sublimation at 140°C. N-methyl-2-pyrrolidinone (NMP), chloroform, o-xylene, ethylene glycol (EG), propionic acid (PA), tetrahydrofuran (THF), dimethylformamide (DMF), dimethylacetamide (DMAc), methanol, ethanol, and butanol were purchased from Aldrich. The reactants and solvents used in this study were of reagent-grade quality and were used without further purification. Commercially available P84 PIs were purchased from Lenzing.

#### Synthesis of PHAs for the TR-A series

Synthesis of three poly(p-hydroxyamid) (PHA) precursors was performed for this study: HAB-IPC (TR-A0) (37), APAF-IPC (TR-A1) (28, 38), and APAF-TPC (TR-A2) (28, 38). TR-A1 is used as a representative example for the TR-A series. TR-A1 was synthesized by dissolving 3.66 g (10 mmol) of bisAPAF in 20 ml of NMP in a three-necked round-bottom flask under a nitrogen atmosphere. Then, 2.03 g (10 mmol) of IPC was poured into the solution and stirred vigorously in an ice bath for 4 hours, resulting in a viscous and pale yellowish solution. After termination of the reaction, the solution was precipitated in distilled water, filtered repeatedly, and dried at 100°C under vacuum.

#### Synthesis of poly(hydroxyl amic acid)-co-(amine-containing amic acid)s for the TR-P series

TR-P denotes equimolar TR poly(benzoxazole-co-pyrrolone) (TR-PBO-co-PPL) polymers from copolymers containing hydroxyl poly(amic acid) and amine-containing poly(amic acid) moieties. In this study, HAB, DAB, and 6FDA were used, and the solutionimidization method (39) was adopted for the preparation instead of thermal imidization as in the previous study (29).
Fabrication of flat sheet dense membranes

Each TR precursor film was formed by slow evaporation from a 15 wt % polymer solution in NMP. For the case of PIM-1 and P84 polymers, the concentration was 5 wt % in chloroform and NMP, respectively. The solution was filtered with a 1.0-µm polytetrafluoroethylene (PTFE) filter cartridge, degassed, and cast onto a well-cleaned glass plate. All TR precursors and P84 solutions were kept at 80°C overnight in a vacuum oven and then steadily heated up to 100°, 150°, 200°, and 250°C in the oven. The resulting film was washed in distilled water and dried again at 120°C. For PIM-1, the film was formed by slow solvent evaporation at room temperature under ambient atmosphere over a period of 4 days, followed by soaking in methanol for perfect removal of trapped solvents (43). All films in this study showed reproducible and controllable membrane thicknesses of 50 to 60 µm except for PIM-1 (~120 µm). For thermal rearrangement, each film was further heated to their corresponding target temperatures under a N₂-purged atmosphere in a muffle furnace (Lenton, London, UK).

Fabrication of asymmetric hollow fiber membranes

For selected TR-X0 and TR-X1 polymers, their precursor hollow fiber membranes were fabricated using a induced phase separation method, i.e., the so-called dry-jet wet spinning process. The degassed dope solution and bore solution were delivered to the spinneret (0.44-0.2-0.12 from Kasen, Osaka, Japan) using a gear pump (GM-S series from Mitsubishi Company, Tokyo, Japan) and a 307 HPLC pump (Gilson Company, Middleton, WI, USA), respectively, and the nascent fibers were extruded and passed through the air-gap distance. Subsequently, the hollow fiber membranes were immersed in a coagulation bath and moved to a godet bath and a take-up roll. Relevant parameters for fabricating asymmetric hollow fiber membranes are summarized in a previous study (32, 41). The fabricated hollow fiber membranes were washed and dried for 3 days. The resulting hollow fiber membranes underwent the same thermal rearrangement as the flat sheet dense membranes.

Fabrication of PIM-1 hollow fibers was also conducted using a dry-jet wet-quench method. Under room temperature of 22°C and relative humidity of 10%, the degassed PIM-1 dope was extruded with a [45 wt % THF, 46.75 wt % DMAC, and 8.25 wt % ethanol] bore fluid, a [15 wt % PIM-1, 69.5 wt % THF, 13.25 wt % DMAC, and 2.25 wt % ethanol] dope fluid, and a [82.5 wt % butanol and 17.5 wt % THF] sheath layer surrounding the dope and bore fluid. This sheath layer slows the evaporation of THF, reducing skin layer thickness and defects in the final fiber. All dopes and fluids were extruded using syringe pumps (1000D for sheath and core, 500D bore fluid, Teledyne Isco). Air gap was set at 1.5 cm, and the flow rates for bore, core, and sheath layers were 90/120/65 ml/hour, respectively. Dope temperature from pump to spinneret was kept at 22°C, while the water quench bath was kept at 50°C. Drum take-up rate was 1.5 m/min. After phase inversion in the water quench bath, the fibers were collected on a rolling drum in a separate water bath. The fibers remained on this drum for at least 45 min to ensure full-phase inversion before removal to a separate bath of deionized water. This separate deionized water bath was replaced every day for 3 days to ensure full removal of solvent molecules within the fiber. After these water exchanges, the fibers underwent three methanol washes for 20 min each, followed by three more 20-min washes in hexane. After the final hexane wash, the fibers were dried under ambient conditions for an hour before being placed in a 60°C vacuum oven (29 in Hg) overnight.

Hollow fiber membrane modules

Fifty TR hollow fibers with a length of 25 cm were bundled, and both ends were blocked using a paste resin. Each bundle was placed into a 20-cm-long cylindrical aluminum housing and sealed with sealing caps. Epoxy resin was injected into the interior of the housing at a steady rate and was cured for 3 hours at 60°C. The caps were then removed to cut excess fiber ends on both sides. Afterward, the oxygen/nitrogen permeselectivity for all modules was investigated to ensure validity before other evaluations. The detailed information is as follows: outer diameter (OD) = 426 ± 22 µm, inner diameter (ID) = 289 ± 31 µm, effective length = 18.5 cm, packing density = 22 ± 2%, and membrane area = 52 cm². For PIM-1 hollow fiber modules, one single PIM hollow fiber with a total length of ~30 cm was used in each module. One-fourth-inch stainless steel tubing was swaged with one-fourth-inch stainless steel union tee on both ends, followed by national pipe thread (NPT) female tube adapter that results in an effective length of 14 cm (see fig. S8B). After placement of the fiber into the tubing and before sealing, PTFE tape was packed into the female tube adapters, preventing uncured epoxy from entering the module interior. 3M DP100 Clear epoxy filled the tube adapters, sealing the module interior. Before the epoxy hardened, NPT male tube adapters were screwed into the epoxy-filled NPT female tube adapter on both ends, which results in a total module length of 23 cm. Epoxy was cured under room temperature for 48 hours before gas separation performance evaluation works, and excess fiber length was removed. Detailed fiber information is as follows: OD ~ 900 µm, ID ~ 580 µm, effective length = 14 cm, and membrane area = 3.96 cm².

Direct fluorination

All flat sheet membranes in this study were treated with dilute fluorine gas (500 ± 15 parts per million F₂/N₂ at atmospheric pressure) at 25°C for 30, 60, 90, 120, 150, and 300 min in a homemade reaction chamber, as shown in fig. S1A (44). Before the actual treatment, the interior of the reaction chamber was kept inert by alternately applying a vacuum and nitrogen gas in the chamber approximately five times. After fluorination, unreacted fluorine gas was removed using a scrubber filled with activated carbon. Direct fluorination was conducted for 1, 5, and 10 min for the modular hollow fiber configurations. Dense membranes were directly fluorinated on both sides.
of the membrane to realize efficient and uniform performance. For TR hollow fiber membrane modules, the lumen side was exposed to fluorine gas, while shell side was exposed for PIM-1 hollow fiber membrane modules, depending on the position of the effective skin layer acting as a role in regulating separation performances. We denoted a direct fluorinated polymer membrane as Sample-Fyy. Here, Sample indicates polymer species for direct fluorination and yyy is the direct fluorination time in minutes. Particularly in the case of TR polymers, Sample is further coded as TR-Xxx, where X represents the precursor types and x is the number of species (refer to fig. S1B).

Surface properties
Electron spectroscopy for chemical analysis (Sigma Probe ThermoVG, UK) was used to investigate surface chemical substitution. The survey mode and high-resolution mode were used with monochromatic AlKα radiation at a general pressure below 7 × 10⁻⁹ torr. In the case of survey mode, 10 sweep cycles were conducted at electron binding energy ranges of 0 to 1000 eV. Approximately 100 scans for C 1s peaks were also performed. Solid-state nuclear magnetic resonance measurements were conducted.

Microscopic image analysis
Field-emission scanning electron microscopy (SUPRA 55VP, Carl Zeiss, Germany) was performed to investigate the substitution of fluorine in the near surface by measuring the depth profile of the membrane. Specific atom sources, such as fluorine, carbon, nitrogen, and platinum, were identified using the focused ion beam technique and energy-dispersive x-ray spectroscopy. The flat sheet membrane samples were coated with platinum (Pt) to prevent surface charging effects. Electron probe x-ray microanalysis (EPMA) was also used to confirm cascaded structures in hollow fiber modular configurations.

Investigation of the pore size distribution with depth profile
A variable monoenergy slow positron beam (VMSPB) system coupled to sb-PALS and DBES (20, 45) (R&D Center for Membrane Technology, Chung Yuan University, Taiwan) was used for investigating the pore size and distribution with depth profiles of pristine and fluorinated membranes. A 50 mCi ²²Na positron source was used. The sb-PALS system used the secondary electrons emitted from the membrane surface as the start signal and the γ-rays induced by positron annihilation in the pore volume as the stop signal. The sb-PALS data were measured using a BaF₂ lifetime detector at a counting rate of approximately 100 to 200 counts per second (cps). Each PALS spectrum contained 2.0 million counts. All the collected sb-PALS data were fitted using the PATFIT program. The analyzed results of positron lifetimes (τ₁, τ₂, and τ₃) and intensities (I₁, I₂, and I₃) from PALS spectra were attributed to positron and positronium annihilation in membrane materials. The shortest τ₁ of 0.125 ns is from p-Ps annihilation, the τ₂ of 0.45 ns is from positron annihilation, and τ₃ is due to o-Ps annihilation. Because Ps is known to preferentially localize in defect sites (particularly in the free volume before annihilation occurs), the parameters from o-Ps annihilation have been successfully used to obtain the electron properties and depth profiles of free volumes in membranes. The o-Ps lifetime τ₃ is on the order of 1 to 5 ns in polymeric materials, corresponding to the so-called pickoff annihilation with electrons in molecules, and is used to calculate the mean free-volume radius R (angstroms to nanometers) based on an established semi-empirical correlation equation from a spherical-cavity model as follows

$$\frac{1}{\tau_3} = 2 \left[ 1 - \frac{R}{R + \Delta R} + \frac{1}{2\pi} \sin \left( \frac{2\pi R}{R + \Delta R} \right) \right]$$

Here, τ₃ is the positron lifetime (in nanoseconds), R is the pore radius (in angstroms), and AR is the electron layer thickness, semi-empirically determined to be 3.823 Å. The pore volume was calculated from the pore radius, assuming a spherical shape. Detailed descriptions of the VMSPB and the data analysis can be found elsewhere (20).

DBES spectra were measured using an HP Ge detector (EG&G Ortec, with a 35% efficiency and an energy resolution of 1.5 at the 511-keV peak) at a counting rate of approximately 1000 cps. The total number of counts for each DBES spectrum was 1 million. DBES spectra were characterized by the S parameter, which was defined as the ratio of the integrated counts between energies of 510.3 and 511.7 keV (S width) to the total counts. PALS was performed by varying the monoenergeticity of the positron beam over a range from 0 to 30 keV. When the positron and Ps are localized in a hole or free volume with a finite size, the observed S parameter is a measure of the momentum broadening based on the uncertainty principle; a larger hole results in a larger S parameter value and an increase in the amount of para positronium (singlet state). The S parameter has been successfully used in detecting the free-volume depth profile in polymeric systems (20). The cascaded structure of fluorinated membranes was obtained from an analysis of the S parameter data using the VEPFIT program. The χ² of the analysis was less than 1.0, indicating a reasonable error in the data curve fitting.

Physicochemical properties
Small-angle x-ray scattering (SAXS) measurements were conducted with the synchrotron radiation source of the 4C SAXS beamline at the Pohang Light Source II of the Pohang Accelerator Laboratory (PAL) (46, 47). Scattering data were adjusted for sample transmission and background scattering. The two-dimensional scattering data were analyzed using a software package provided by PAL to obtain radially integrated SAXS intensities versus the scattering vector q. Here, q is a function of angle (θ) according to the following equation

$$q = \frac{4\pi}{\lambda} \sin(\theta)$$

Here, λ is the radiation wavelength, and θ is half of the scattering angle (2θ). The Bragg spacing d was calculated according to the following equation

$$d = \frac{2\pi}{q}$$

Thermogravimetric analysis (TGA; TGA Q50, TA Instruments, New Castle, DE, USA) was performed to observe the thermal stability of the membranes. The thermal stability was analyzed on the basis of the decomposition temperature as a function of the fluorination time. The TGA measurements were carried out under a nitrogen atmosphere at a heating rate of 5°C min⁻¹ from 60° to 800°C. No changes were observed in the thermal stability up to 450°C. The mechanical properties of the tensile strength and elongation were characterized to evaluate the stress-strain behavior of the polymer samples using a universal testing machine (Autograph AGS-J, Shimadzu, Kyoto, Japan) and at least five film specimens of each sample (1 1/4″ High ASTM D-638 Type). Optical property changes were also monitored.
MD simulations
The Materials Studio program package (BIOVIA Software Inc., CA, USA) was used for MD simulations in this study. Because gas transport through polymeric membranes is subject to various factors, such as intermolecular interactions and complex pore morphologies (in addition to the effect of bottlenecks), we developed ideal and conceptual pore models using carbon nanotube (CNT) structures (Fig. 1, E and F). First, we built a single-walled CNT with a 0.964-nm ID, corresponding to that of TR-A2-F0. Then, the length of the CNT models was scaled to one-half to obtain a tighter carbon atom wall. In addition, according to the fluorine treatment time, the pore structures of the CNT models were modified to a rugby ball shape, as shown in Fig. 1E. The models were geometrically optimized after helium and methane molecules were inserted into the CNT models using the sorption module in the Materials Studio program package. Last, a constant number, volume, and temperature (NVT) simulation was performed under 298 K and 1 atm conditions for 100 ns with an Andersen thermostat and the Ewald nonbond sumation method to obtain the diffusivity coefficients calculated from the mean square displacement of gas molecules in the CNT models simulated by MD. The COMPASS II (Condensed-phase Optimized Molecular Potentials for Atomistic Simulation Studies II) force field was used (48–51). Three different types of TR microstructures, with 0-, 60-, and 300-min exposures, were considered to investigate the efficacy of the squeezed hourglass-shaped microstructure in He/CH4 applications, and their pore size distributions were obtained from the sb-PALs spectra (Fig. 1D and fig. S3).

Gas permeation properties
The constant volume method with varying pressure (the so-called time-lag method) was introduced to measure the gas permeability of P84, TR-PBO, and PIM-1 membranes (table S3). The membrane films were 4 cm² in size and were masked with impermeable aluminum tape on both sides before being placed in the permeation cell apparatus. The constant volume method is based on the following equation

\[
P = \frac{\partial p}{\partial t} \left( \frac{V T_0 l}{p_0 T A \Delta P} \right)
\]

where \( P \) (Barrer, 1 Barrer = \( 3.3 \times 10^{-16} \) mol m/m² s Pa) is the permeability and \( \partial p/\partial t \) is the experimental value of the rate of pressure increase under a thermodynamic steady state. \( V \) (cm³) indicates the volume downstream, and \( l \) (cm) is the thickness of the membrane. \( A \) (cm²) represents the effective area of the membrane, and \( T \) (K) is the absolute temperature of the operating system. \( \Delta P \) (cm Hg) indicates the pressure difference across the membrane and is the driving force of diffusion through the membrane. \( T_0 \) and \( p_0 \) indicate standard temperature and pressure, respectively. The gas permeabilities of six representative single gases of He, H₂, CO₂, O₂, N₂, and CH₄ were measured at 35°C and 760 torr.

Constant pressure methods were applied in the case of pristine and c-TR and c-PIM-1 hollow fiber membrane modules. The gas permeability was calculated using the following equation

\[
P = \frac{Q \times l \times T_0}{A \times \Delta P \times T}
\]

where \( Q \) (cm³ min⁻¹) is the gas flow rate downstream measured using a bubble flow meter, \( \Delta P \) (cm Hg) indicates the pressure difference across the membrane (5 bar for TR and 7 bar for PIM-1, respectively), and \( T \) is the temperature measured at the bubble flow meter (35°C). The ideal selectivity of a single gas (\( \alpha_{i,j} \)) is determined as the ratio of the two gas permeances. The values reported here are an average of at least three measurements. The uncertainty in the data is 5% or less.

Evaluation of the practical feasibility in helium recovery
The mixed-gas permeation tests with 2000-hour-aged TR-X1-F0 and TR-X1-F1 hollow fiber membrane modules were performed using a constant pressure method following the previous report (52). Twenty bar of He/CH₄ (1/99 mol %) gas mixture was fed from bore to shell side of the module at ambient temperature, while permeate pressure was constant at 0.5 bar by a diaphragm vacuum pump (Laboport N86 K18, KNF Neuberger Inc., Trenton, NJ, USA). The stage cut (\( \theta \)) was defined as the ratio of permeate flow rate to feed flow rate and controlled by regulating feed flow rate using mass flow controller (M3030VA, Line Tech, Daejeon, Korea). Gas composition at permeate side was determined by gas chromatography (Micro GC 490, Agilent Technologies Inc. Santa Clara, CA). Helium recovery was determined by multiplying stage cut with the ratio of helium concentration at permeate to that at feed.

A simple single-stage membrane process model that describes the helium concentration in the permeate and retentate stream as a function of stage cut (\( \theta \)) was developed to predict the He purity and He recovery in the permeate stream. The goal of this study was to compare the experimental results and precisely predict the exact condition to achieve 90% helium recovery for the case of TR-X1-F1 hollow fiber membrane modules. This model was conducted using a numerical method, which divides the membrane process into a finite number of stages with the same stage repeated in series (see fig. S10, 1000 in this study). In this model, the retentate stream of each stage becomes the feed stream of the next stage. Therefore, the overall permeation is a summation of each permeate stream, and the overall retentate stream is the final retentate stream. For simplicity, the effect of pressure drop on the helium separation performance was neglected, meaning that the operating pressure along the flow path is constant. The concentration of He in the permeate stream at the pressure ratio \( \varphi \) is as follows

\[
n_{He,\varphi} = \frac{\varphi}{2} \left( \frac{n_{He}}{n_{He,0} + \frac{1}{\varphi} + \frac{1}{\alpha - 1} - \left( \frac{n_{He}}{n_{He,0} + \frac{1}{\varphi} + \frac{1}{\alpha - 1}} - \frac{4\alpha n_{He}}{\alpha - 1} \right)} {\alpha} \right)
\]

Therefore, the He recovery and purity were calculated as follows

\[
\text{Recovery} = \frac{\sum_{k=1}^{n} (P_k \times n_k \varphi)}{F_1 \times n_1}
\]

\[
\text{Purity} = \frac{\sum_{k=1}^{n} (P_k \times n_k \varphi)}{\sum_{k=1}^{n} P_k}
\]

The simulated result with commercial perfluoropolymers (marked in blue pentagons in Fig. 3, D and E) (53) was also presented for comparison.

Preliminary study on assessing feasibility in energy-related applications
To investigate capability of hydrogen dehydration (H₂O/H₂) for c-TR and c-PIM hollow fiber membranes, H₂ and water vapor permeabilities were collected using a homemade gas permeability testing system...
connected with a gas chromatograph (GC, 490 Micro GC, Agilent Technologies, USA) and two mass flow controllers (MFC, M3030V, Line Tech, Korea) operating at 80% relative humidity (RH) at 50°C under a 2.2-bar unilateral back pressure (54). The gas permeability (P) can be obtained by the following equation

\[ P = \frac{V M_{\text{gas}} d}{P_{\text{feed}} RT d_{\text{f}}} \]

Here, A (cm²) and d (µm) represent the effective membrane area and thickness of membrane samples, respectively. \( P_{\text{feed}} \) and \( M_{\text{gas}} \) (g mol⁻¹) are the pressures of each gas (760 mmHg) and molecular weight of the permeating gas, respectively. \( V \) (cm³) is the volume of the measuring device at the bottom of the membrane samples. \( p \) (g cm⁻³) and \( R \) (liter mmHg K⁻¹ mol⁻¹) are the densities of the permeating gas and the gas constant, respectively. \( \frac{d}{d_f} \) is the slope that can be plotted from a change in permeated gas pressure as a function of time. The unit of \( P \) is Barrer, where 1 Barrer = 10⁻¹⁰ cm³ (standard temperature and pressure, STP) cm⁻² s⁻¹ cm Hg⁻¹. Ar [100 standard cubic centimeters per minute (scm)] was used as a sweep gas, and 1000 sccm of feed was injected to the system.

**SUPPLEMENTAL MATERIALS**

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