Electron-mediated control of nanoporosity for targeted molecular separation in carbon membranes

Carbon molecular sieve (CMS) membranes are considered game-changers to overcome the challenges that conventional polymeric membranes face. However, CMS membranes also confront a challenge in successfully separating extremely similar-sized molecules. In this article, high-precision tuning of the microstructure of CMS membranes is proposed by controlled electron irradiation for the separation of molecules with size differences less than 0.05 nm. Fitting CMS membranes for targeted molecular separation can be accomplished by irradiation dosage control, resulting in highly-efficient C2H4/C2H6 separation for low dosages (~250kGy, with selectivity ~14) and ultra-selective H2/CO2 separation for high dosages (1000~2000kGy with selectivity ~80). The electron irradiated CMS also exhibits highly stabilized permeability and selectivity for long-term operation than the pristine CMS, which suffers from significant performance degradation due to physical aging. This study successfully demonstrates electron irradiation as a possible way to construct “designer” nanoporous carbon membranes out of the standard components mostly confined to pyrolysis conditions.

Energy-efficiency improvements in the industrial separation sectors will accelerate the global decarbonization efforts, as these processes account for more than 10% of global energy consumption and carbon emissions1,3. For example, the current industrial separation of ethylene from ethane performed with energy-intensive cryogenic distillation solely consumes 0.1% of global energy use due to their extremely similar size and chemical properties1. This challenging separation could benefit from the low-energy approach of membranes. Nanoporous carbon membranes (i.e., carbon molecular sieve, CMS), which are derived from the thermal decomposition of polymeric materials, have demonstrated the fine separation of molecules with sub-0.1 nm size differences: gas separations for light hydrocarbons (C2 or C3 compounds), natural gases (N2/CH4), and liquid hydrocarbon separations for xylene isomers or hexane isomers5–10. The CMS provides slit-like transport pathways for the molecules by its disordered sp2-hybridized graphenic layers. The unique bimodal pores of the CMS are composed of supermicropores (0.7–2 nm) and ultramicropores (<0.7 nm)11. The connectivity between these two types of pores in the CMS allows high permeability, which is derived from the pore gallery in supermicropores, and high selectivity, which is due to the limiting pore diameter with sufficient structural rigidity to enhance the diffusion selectivities12,13–14.

However, CMS structures have limited separation performance for gas pairs with extremely small size differences (i.e., Δsize < 0.05 nm; for example, C2H4/C2H6 or H2/CO2). This is primarily due to insufficient pore confinement and limited populations of sub-0.5 nm pores. As an example, CMS membranes prepared from Matrimid® and 6FDA(4,4’-(hexafluoroisopropylidene)diphthalic anhydride)-based polyimide were tested for C2H4/C2H6 separation (Δsize = 0.01nm) with different heating protocols during the fabrication of CMS membranes15. Although those CMS membranes exhibited moderate C2H4 permeability (~100 Barrer) in a relatively low pyrolysis temperature range of 500–550 °C, the C2H4/C2H6 selectivity remained at around 3–5, indicating the minor possibility of sharp C2H6 differentiation by the CMS.
window size. When the pyrolysis temperature was raised to 800 °C, the selectivity was improved to 9 while the C2H4 permeability dropped an order of magnitude. As another challenging example, H2/CO2 (Δsize = 0.041 nm) selectivity of less than 10 is usually reported in CMS membranes regardless of the final pyrolysis temperature or the types of precursors. In addition to the strong adsorption between CO2 and carbon surfaces, which enhances intrinsic CO2 permeability, the incomplete development of sub-0.5 nm ultramicropores in the CMS matrix suppresses the effective discrimination between H2 and CO2.

Several post-synthetic ultramicropore tuning mechanisms for CMS to align the specific pore size have been recently proposed. Hyperaging with mild post-pyrolysis heat treatment allows pore-size tuning to enhance the selectivity of H2 vs. larger hydrocarbon gases (C2H4/C2H6 and H2/CO2, which are key energy-intensive separation processes). The effective discrimination between H2 and CO2 reduces the number of accessible transport channels for large gas molecules, thus facilitating the selective discrimination of the closely-related gas pairs— which pushes pore size distribution of the pristine CMS to a smaller ultramicropore regime (Δsize < 0.5 nm). On the other hand, when a trace amount of O2 is present in an inert gas environment during pyrolysis (known as “O2-doping”), chemically adsorbed O2 molecules on the carbon edge sites induce the formation of a closed ultramicropore window and result in enhanced permeation performance such as in CO2/CH4. Likewise, low concentration hydrogen gas dosage during the pyrolysis created mid-sized ultramicropores (Δsize > 0.5 nm), which are suitable for separating vapor-phase xylene isomers (i.e., p-xylene and o-xylene; Δsize = 0.58 nm). In addition, cellulose hollow fiber membranes which had not been reported before were fabricated and pyrolyzed into CMS membranes without severe structural collapse. The cellulose-originated CMS membranes achieved distinguished H2/CO2 separating capability (e.g., 101 Barrer of H2 permeability with H2/CO2 selectivity ~28), indicating the importance of suitable polymeric precursor selection and pyrolysis temperature.

However, the major handles for tuning ultramicropores so far are limited to optimizing the pyrolysis temperature, pyrolysis environment, and precursors. Therefore, it is crucial to seek advanced and adjustable tuning factors for the fine calibration of both ultramicropores and supermicropores of the CMS microstructure to achieve molecular separations for challenging gas pairs with Δsize < 0.05 nm.

We present highly-controlled electron beam irradiation as an effective method to precisely tailor the ultramicropore dimensions in CMS membranes (Fig. 1a). It is well known that high-energy electron irradiation causes structural changes in carbon-based materials, including carbon nanotubes (CNTs) and fullerenes (C60). Such high-energy interaction can induce point defects in carbon sheets and strands, making vacancies and interstitial junctions on sp2-structures, thereby creating unique carbon landscapes. We successfully tuned ultramicropore dimensions by interacting carbon strands with different electron dosages (50–2000 kGy, 1 kGy is equivalent to 1 kJ/kg of energy transfer). We found that electron irradiation drives mid-size ultramicropores to be concentrated below 0.4 nm, which intensifies the size-selective permeation of similarly-sized molecules, such as C2H4/C2H6 and H2/CO2, which are key energy-intensive separation model pairs. Notably, the permeability of the membranes sharply increased 3 to 4-fold when the controlled amount of electrons was irradiated on the carbon membrane, crossing the upper bound limits. We showed that both pore tightening and oxygen-rich surface formation by electron irradiation were responsible for performance enhancements. High-energy interaction between electrons and carbon surfaces further allowed the retention of original membrane performance over time with greatly improved aging resistance.

**Results**

**Electron beam induced CMS tuning**

To investigate the effect of electron irradiation on a model CMS membrane, we selected 6FDA:BDPA(DI):DAM [4,4’-(hexafluoropropylidene)diphthalic anhydride (6FDA), 4,4’-biphalic anhydride (BDPA), 2,4,6-trimethyl-1,3-phenylenediamine (DAM), 1:1 indicates equimolar copolymer] as a precursor (Supplementary Fig. 1). 6FDA-based polyimides have shown high gas separation performances and resistance to swelling due to structural rigidity and inefficient chain packing from trifluoromethyl group (-CF3) on the backbone. Analogous to the precursor properties, the inefficient packing and high free volume of 6FDA-based polyimide induced an open, selective, and highly porous CMS membrane structure in the pyrolysis process. Therefore, Carbon molecular sieve membranes fabricated with 6FDA-based polyimides indicated superior permeability for gas separation, especially in light hydrocarbons, than other commercialized or studied precursor derived CMS. Given the coexistence of bulky 6FDA and planar BPDA monomers, 6FDA:BDPA(DI):DAM derived CMS provides a more selective pore with moderate permeability than other 6FDA-based polyimide such as 6FDA-DAK. CMS membranes were fabricated with thickness of 75–80 μm by pyrolyzing the symmetric dense film of 6FDABDPA-DAM under an ultra-high-purity (UHP) argon environment (Fig. 1a). The CMS were prepared as self-standing and symmetric films then irradiated with a range of 50–250 kGy by a high-energy electron-beam accelerator in ambient air. From our extensive experience with electron-irradiated CMS membranes, we believe that the macro-mechanical properties were not altered upon electron irradiation of CMS membranes. For example, mechanical property test on a bundle of carbon fibers (not membranes, diameter <10 μm) revealed stable tensile stress and elongation after electron irradiation, therefore, mechanical property of electron-irradiated CMS would be insignificantly changed. While scanning electron microscope (SEM) cross-section images did not catch apparent differences between the pristine and electron irradiated CMS membranes, energy-dispersive X-ray spectrometer (EDS) scan revealed an asymmetric chemical composition with an oxygen-rich surface of the CMS film after electron irradiation (Fig. 1b–d and Supplementary Fig. 2). High-resolution transmission electron microscopy (HR-TEM) images of the CMS films illustrated the maze-like microstructure of amorphous carbon membranes (Fig. 1e, g). The corresponding Fast-Fourier Transformed (FFT) patterns (Fig. 1f, h) show continuous ring shapes distinguished by vivid contrast differences in the TEM images, indicating the representative pattern of disordered carbon strands. When the pristine CMS was exposed to 250 kGy (250kGy-CMS, hereafter) of electron irradiation, the radial intensity of the FFT patterns changed with different the number of rings, indicating the different packing degrees of the carbon strands. The breadth of radial intensity for 250kGy-CMS was from 5.65 to 3.27 nm−1 (corresponding to 0.177 to 0.306 nm of micropore spacing, respectively, Fig. 1h) showing that denser, compact ultramicropore dimensions were obtained when compared to those of pristine CMS from 5.23 to 2.72 nm−1 (0.191 to 0.367 nm of micropore spacing, Fig. 1f). Furthermore, the disappearance of radial intensity at 3.94 nm−1 (0.254 nm of micropore spacing, Fig. 1f) of the pristine CMS after electron irradiation indicates the formation of condensed ultramicropore dimensions. The formation of tighter ultramicropores reduces the number of accessible transport channels for large gas molecules, thus facilitating the selective discrimination of the closely-sized gas pairs.

To precisely analyze the orientation of the amorphous carbon strands in the electron irradiated CMS, synchrotron grazing-incidence wide-angle X-ray scattering (synchrotron GI-WAXS) was used. GI-WAXS utilizes a very small incidence angle to confine the penetration depth to the nanometer level, probing the molecular length-scale structures in the amorphous CMS membranes. 2D scattering images show the ring patterns of both the precursor and the irradiated CMS films (Fig. 2a, b and Supplementary Fig. 3). The anisotropic packing of the CMS structures is observed by the more dominant intensity development of qyy than that of qxx, and this tendency becomes more pronounced as the pyrolysis temperature increases (500 °C to 600 °C, Fig. 2b and Supplementary Fig. 3). The ‘out-of-plane’ (qyy, qzz) and ‘in-plane’ (qxx, qyy) scattering vectors indicate that the aromatic strands constituting the CMS micropores are anisotropic...
with face-on orientation, not edge-on packing perpendicular to the substrate. Notably, 2D scattering image of 250kGy-CMS shows that the “out-of-plane” vector \( q_z \) intensity in the range of 0.5 to 1.0 Å\(^{-1} \) was mitigated, while the peak around 1.4 Å\(^{-1} \) became strong and narrow (Fig. 2b, c). The vertical line-cut as a function of \( q_z \) shows a broad peak along \( q_{\text{out}} \approx 1.365 \text{ Å}^{-1} \) involved with imperfect lamellar-like ultramicropores with approximately 0.46 nm \( d \)-spacing for 250kGy-CMS and 0.514 nm \( d \)-spacing for the pristine CMS (Fig. 2c). In addition, the GI-WAXS peak analysis of 250kGy-CMS (Fig. 2d) revealed that the assumed pore size distribution started from 1.635 Å\(^{-1} \) (i.e., 0.385 nm), thus confirming that the sharp molecular sieving of the ethane (0.4 nm) is possible.

We further found that the electron irradiation not only successfully tuned the ultramicropore dimensions but also modified the surface of the CMS via forming oxygen-rich carbon strands. The survey scan of X-ray photoelectron spectroscopy (XPS) compares the atomic composition of the surface on the pristine CMS and 250kGy-CMS, showing the intensified OIs peak (∼530 eV) and relatively decreased Cls peak (∼285 eV) after electron irradiation (Fig. 2e and Supplementary Fig. 4). It is well known that the high-energy electron interaction generates atomic vacancies, typical carbon defects that are vulnerable to oxidation or reaction with other gas molecules\(^{25,39} \). Furthermore, the formation of interstitial defects is responsible for locally reduced carbon dimensions, liable to rearrangement of bonds and topological transformation of the carbon structures. The defective strands exposed to ambient air would combine with \( \text{O}_2 \) and develop an oxidized surface, creating a narrow ultramicropore window on the CMS surface that can reject large penetrants. However, electron irradiation can be distinguished from \( \text{O}_2 \) doping treatment on CMS membranes. The \( \text{O}_2 \) doping process uses trace amounts of oxygen.
the disordered carbon networks in the CMS membrane, the development implies defects in crystalline carbon, but in the case of activity of 4 when pyrolyzed at 550 °C in a UHP Ar environment (<1 ppm)

250kGy-CMS along the out-of-plane axis.

250kGy-CMS (ID/IG = 0.621) is lower than that of the pristine CMS DAM showed C2H4 permeability of 110 Barrer with C2H4/C2H6 selectivities for example, previously reported CMS membranes produced from 6FDA-BPDA(L1)-DAM showed C2H4 permeability of 110 Barrer with C2H4/C2H6 selectivity of 4 when pyrolyzed at 550 °C in a UHP Ar environment (<1 ppm O2). When the O2 concentration in the pyrolysis chamber was increased to 30 and 50 ppm, the CMS membranes showed C2H4 permeability of 85 and 68 Barrer, respectively, with slight improvements in the C2H4/C2H6 selectivities (5.2–5.5, respectively). While the high-energy electron irradiation directly combines the advantages of a defect-mediated oxidized carbon surface with reduced ultramicropore dimensions, a higher degree of molecular specificity of the resultant membrane is expected. As shown in the depth profiles of XPS (Fig. 3f, g and Supplementary Fig. 5), the pristine CMS shows a consistent atomic composition along with an etched depth down to 3.5 μm from the surface. However, 50 kGy-CMS has an oxidized surface and the oxygen composition continuously decreases with the etched depth. In addition, all the 100kGy-CMS, 250kGy-CMS and 500kGy-CMS show a 3 or 4-fold increase in oxygen composition on the surface, indicating the formation of oxygen-rich carbon surfaces. When the electron irradiation increased from 50kGy to 500kGy, the oxygen composition and the thickness of the oxidized carbon layer also increased (from 1.61 to 4.13 μm Supplementary Fig. 6). Raman spectroscopy was also used to measure the defectiveness of the carbon structure after electron irradiation (Fig. 2h and Supplementary Fig. 7). D peak (100–1350 cm−1) development implies defects in crystalline carbon, but in the case of the disordered carbon networks in the CMS membrane, the D peak represents the proportion of the aromatic rings in the carbon clusters. G peak (1500–1580 cm−1) involves in-plane bond-stretching of sp2-carbon atoms and occurs at all sp2 sites, including aromatic and olefinic carbons. Therefore, the defectiveness in carbon materials observed during XPS depth proﬁles of CMS membranes, CMS membranes, mixed matrix membranes (MMMs), and metal-organic framework (MOF) membranes (Fig. 3c and Supplementary Tables 1–4), and found that separation performances of the permeation of gas molecules in these CMS membranes to understand the structure-property relationship in the electron-irradiated CMS membranes.

**Gas permeation and diffusion.** The electron-induced ultramicropore tuning can be directly verified through gas permeability measurements for gas pairs. We first measured the unary gas permeabilities of pristine CMS (pyrolyzed at 500 °C) film and electron-irradiated CMS films (50kGy-CMS, 100kGy-CMS, and 250kGy-CMS) via a constant-volume gas permeation apparatus (Fig. 3b and Supplementary Figs. 8 and 9). When a low electron dosage (50 kGy) was applied to the CMS films, the unary gas permeabilities of relatively small gases (from He to C2H4) did not dramatically change (Fig. 3b) but sharp molecular cut-off appeared between C1 hydrocarbon gases (i.e. C2H4/C2H6 selectivity increased from 13.28 to 47.13) when the electron dosage exceeded 100kGy, the permeability of the larger gases significantly decreased (e.g., 55% reduction for CH4 and 84% reduction for C2H6) compared to smaller gases—indicating the formation of tight but size-selective ultramicropores. In the highly irradiated CMS (250kGy-CMS) showed dramatically reduced permeability for most gases (e.g., 95% and 98% reduction for CH4 and C2H6, respectively, even 90% reduction in N2 except for helium, which means that the ultramicropore distribution of the electron-irradiated CMS had shifted toward the sub-0.4 nm region, with the highest pore populations concentrated at 0.33 to 0.40 nm (Fig. 3a and Supplementary Fig. 10). Especially in the case of C2 hydrocarbons (C2H2/C2H4), the permeability of the ethane drastically decreased to 19.9 Barrer compared to that of ethylene (180.6 Barrer) in 100kGy-CMS, with a significantly increased ideal C2H2/C2H4 selectivity of 9.06 (Supplementary Fig. 11). However, when a very high electron dosage was applied (500 kGy) to the CMS membranes, the formation of the tightest structures with negligible ethylene permeabilities were observed (Supplementary Fig. 12).

We compared the single-component separation performance of the electron-irradiated CMS with other state-of-the-art polymeric membranes, CMS membranes, mixed matrix membranes (MMMs), and metal-organic framework (MOF) membranes (Fig. 3c and Supplementary Tables 1–4), and found that separation performances of the
Electron irradiation led to similar C$_2$H$_4$/C$_2$H$_6$ permeabilities but with much-improved C$_2$H$_4$/C$_2$H$_6$ selectivity up to 14 (following the red colored trajectory in Fig. 3d). Electron irradiation (up to 250 kGy) led to similar C$_2$H$_4$ permeabilities, but with much-improved C$_2$H$_4$/C$_2$H$_6$ selectivity up to 14 (following the red colored trajectory in Fig. 3d).

To elucidate the possible reason for the excellent C$_2$H$_4$/C$_2$H$_6$ separation via electron-irradiation, diffusion analysis was conducted via a custom-built pressure-decay sorption setup (Supplementary Fig. 14). C$_2$H$_4$ and C$_2$H$_6$ uptake curves for CMS films at different fugacities were fitted by the Fickian diffusion model to calculate their diffusivities. For all CMSs tested, there was a relatively delayed uptake of C$_2$H$_6$ over C$_2$H$_4$ (Supplementary Figs. 15–17), and the gaps between relative half-time diffusivities (C$_2$H$_4$/C$_2$H$_6$) increased as the electron dosage increased. Figure 3e shows the Fickian diffusivities of C$_2$H$_4$ and C$_2$H$_6$ for all CMS films revealing the enhanced diffusion selectivity of C$_2$H$_4$/C$_2$H$_6$ at higher electron dosages (up to 250 kGy). While the C$_2$H$_4$ and C$_2$H$_6$ sorption isotherms at 308.15 K (Supplementary Fig. 18) imply
negligible sorption selectivity influence on the improved separation capability, regardless of C2H6 and C2H4, almost the same isotherm curve and final uptake amount were observed in pristine and electron-irradiated CMS, respectively.

To explain the fast and selective ethylene transport in the electron-irradiated CMS membranes, the sp1/sp2-hybridized carbon ratio for pristine and electron-irradiated CMS (50, 100, 250 kGy) samples were determined via XPS depth profiles (Fig. 3f). By deconvoluting the XPS C1s spectrum to three Gaussian peaks, the sp2-hybridized carbon peaks were obtained, and the ratio was estimated by their peak area (Supplementary Figs. 19-23). It is generally known that the sp2-hybridized carbon obstructs the packing of the strands due to the three-dimensional atomic configuration, which makes faster penetration of smaller gas molecules. In contrast, the two-dimensional sp2-hybridized carbon is concerned with closely packed carbon strands, thus a more selective CMS structure for specific gas pairs. As shown in Fig. 3f, the sp2/sp2 ratio is consistent (≈0.6) for all depths in pristine CMS, whereas electron-irradiated CMS shows the sp2/sp2 ratio gradient depending on the depth. At the surface, 50kGy-CMS, 100kGy-CMS and 250kGy-CMS had a low sp2/sp2 ratio (0.40, 0.35 and 0.28, respectively), and the ratio gradually increased to arrive at a similar value of the pristine CMS (≈0.6) below 2.75 μm, which indicates that the penetrant molecules could undergo significant transport resistance on the electron-irradiated CMS surface. Therefore, the substantial portion of accessible ultramicropores on the surface are both oxygen-functionalized and tightly packed, rejecting the permeation of C2H6 over C2H4.

We extended our finding in the unary permeation of electron-irradiated CMS to binary gas mixture permeation, revealing the possible effects of multicomponent interaction within the CMS membranes. For this, the Wicke-Kallenbach mode was used to test equimolar C2H4/C2H6 mixtures (Supplementary Fig. 23). For the 50kGy-CMS, the permeability of C2H6 was 368.3 Barrer in the unary permeation, while that of mixed gas was 352.6 Barrer. Roughly a 4% loss in permeability was observed in this case, and the selectivity of the mixture permeation slightly decreased from 5.4 to 4.7 (Fig. 3g). The 100kGy-CMS and 250kGy-CMS showed comparable selectivity between unary (9.1 and 14.0 for 100kGy-CMS and 250kGy-CMS, respectively) and binary permeation (8.0 and 13.8 for 100kGy-CMS and 250kGy-CMS, respectively). For all CMS membranes tested, including pristine and electron-irradiated ones, a slight reduction (about 2-15%) in the mixture selectivity was noticed, which was mainly due to the coupling effect among different gas species within the ultramicropores of the CMS membrane41,42. The equimolar binary permeation tests revealed more realistic performances of electron-irradiated CMS membranes under industrially relevant gas mixtures.

In the previous unary permeation tests, we noticed that when the electron irradiation dosage increased to the extreme level, exceeding 1000 kGy, almost complete blocking of the N2 permeation (0.364 nm) while maintaining a moderate permeability of He (0.26 nm, Supplementary Fig. 12) was achieved. Motivated by this dramatic change in permeabilities at high dosages, we further broadened our sight to another key energy-intensive separation pair of H2/CO2. The 100kGy-CMS membrane at permeation temperatures of 35 °C, 45 °C, and 55 °C and predictions at 100 °C and 130 °C. Source data are provided as a Source Data file.

Fig. 4 | Gas separation performances of the highly irradiated CMS membranes in H2/CO2 pair. a H2/CO2 permeation data of pristine and highly electron-irradiated CMS membranes for two different temperatures (500 °C and 675 °C) including other state-of-the-art CMS membranes. b H2/CO2 permeation data of 500 °C 1000kGy- and 675 °C 1500kGy-CMS including other state-of-the art membranes. c H2/CO2 separation performances of 675 °C CMS membranes in unary and binary (equimolar mixture) systems. d H2/CO2 experimental data of 675 °C 1500kGy-CMS membrane at permeation temperatures of 35 °C, 45 °C, and 55 °C and predictions at 100 °C and 130 °C. Source data are provided as a Source Data file.
the penetration of CO₂ as well as the strong sorption of CO₂ onto the micropores of the CMS. With high-dosage irradiation (1000–2000 kGy) of those CMS membranes, notable performance enhancements were observed toward making highly H₂ selective CMS membranes. In the case of the CMS membrane pyrolyzed at 500 °C, with 1000 kGy irradiation, ideal H₂/CO₂ selectivity was increased by a factor of 40. Remarkably H₂-selective membranes were formed when CMS membranes pyrolyzed at 675 °C were irradiated with 1500 kGy, showing outstanding urinary H₂ permeability over 350 Barrer and ideal H₂/CO₂ selectivity over 80. Figure 4b compares the separation performances of 1000kGy-CMS (pyrolysis at 500 °C) and 1500kGy-CMS (pyrolysis at 675 °C) membranes in the Robeson plot, revealing the superior molecular sieving properties of the high-dosage electron-irradiated CMS membranes (Supplementary Tables 5–8).

Binary gas permeation tests were also completed for a H₂/CO₂ pair to elucidate the multicomponent permeation effects (Fig. 4c). The binary permeation of the pristine CMS membrane (pyrolysis at 675 °C) showed quite a large deviation from unary permeation, especially with a noticeable decrease in both H₂ permeability and H₂/CO₂ selectivity. Favorable and competitive CO₂ adsorption over H₂ of the micropores of the CMS membrane is a plausible cause of this result45. On the other hand, both 1000kGy-CMS and 1500kGy-CMS showed largely unchanged separation performances for binary permeation, which might have been caused by the decrease of the population of ultramicropores that could favorably accommodate CO₂ molecules in those heavily irradiated membranes, which reduced the impact of competitive adsorption for binary permeation. Notably, binary permeation in 2000kGy-CMS showed a marginal permeability loss among other irradiated samples, with rather increased selectivity than ideal selectivity. This powerful molecular sieving would have resulted from the further removal of transport pathways favorable to larger CO₂ molecules in massively irradiated CMS membranes. We further confirmed the superior H₂/CO₂ separation performances of electron-irradiated CMS membranes at high temperatures covering from 100 °C to 130 °C, the realistic feed condition directly coupled with industrial water-gas shift reactors. Experimental data (1500kGy-CMS) from permeation tests at 35 °C, 45 °C, and 55 °C were used to calculate the apparent activation energy of permeation using Arrhenius regression (Fig. 4d).

We expected that the permeability of the electron-irradiated CMS membranes would significantly increase at such high temperatures, keeping slightly reduced H₂/CO₂ selectivities, which is still a remarkable separation performance.

Aging resistance of electron-irradiated CMS

Physical aging generally occurs in non-equilibrium state glassy polymers. It entails a segmental rearrangement process that leads the polymer with excess free volume to densification and then further approaches the thermodynamically stable state over time31,44. This self-retarding process was also reported in CMS membranes and significantly degraded the separation performance for long-term operation44,45. To investigate the effect of aging on electron-irradiated CMS membranes, both pristine CMS and 100kGy-CMS were tested for a C₂H₄/C₂H₆ pair. We monitored the change of C₂H₄ permeability and C₂H₄/C₂H₆ selectivity for up to 36 days (Fig. 5). The pristine CMS membranes immediately lost 60% of the initial permeability within 6 days. They converged to only 30% remaining permeability after 36 days, whereas the selectivity gradually increased to 120%, which is a typical aging behavior for CMS membranes. In contrast, 100kGy-CMS preserved more than 80% of C₂H₄ permeability during the same period (36 days), and unlike the pristine sample, a slight decrease (~20%) in selectivity was observed, showing excellent aging resistance. 87 K Ar physisorption experiments (Supplementary Figs. 25 and 26) demonstrated the relatively reduced pore volume (pristine: 0.286 cm³/g; 100kGy-CMS: 0.269 cm³/g) and BET surface area (pristine: 733.9 m²/g and 100kGy-CMS: 691.9 m²/g) in 100kGy-CMS, implying the relaxation of the excess free volumes in CMS. Moreover, the previously discussed structure characterization, such as TEM and GI-WAXS (Figs. 1 and 2), also indicated that the electron irradiation tuning reduced the ultrimicropore dimensions. The reductions in pore volume can illustrate relatively small C₂H₄ permeability loss in 100kGy-CMS since the already tightened CMS structure less induce pore shrinkage. Additionally, elemental analysis (EA) was performed to figure out the aging characteristic of the CMS membranes (Supplementary Table 9). The aged pristine CMS (6.92%) revealed a relatively higher oxygen fraction than that of fresh pristine CMS (6.68%) because reactive carbon edges slowly chemisorb oxygen in air condition47,48. The oxygen fraction in 100kGy-CMS decreases from 9.95% to 9.06% in 30 days. This result is comparable to the loss of the oxygen-containing group in aged graphene oxide (GO) and can be considered as a process where over-oxidized CMS surface rebound back to equilibrium structure67. The unusual aging behavior (selectivity decrease) of the electron-irradiated CMS is conceivable with the hypothetical scheme based on the changes in oxygen fraction of the 100kGy-CMS after aging process (Supplementary Fig. 27). The D₀ and D₁ are supermicropore dimensions and D₂, D₃, D₄, and D₅ are ultramicropore of the fresh and aged electron-irradiated CMS, respectively, according to the IUPAC definition. When aging process proceeds, the supermicropore dimension D₀ shrinks to D₁ and entails overall permeability reduction in both C₂H₄ and C₂H₆. Oxygen adsorbed on the carbon edge sites is less developed after aging and it makes ultramicropore dimensions (D₄ and D₅) increased than the fresh sample, resulting in slightly lessened selectivity of the aged CMS. That is, contrast structural evolution in supermicropore and ultramicropore explains the unique permeation behavior of 100kGy-CMS after aging process.
Discussion

In summary, we successfully demonstrated that post-pyrolysis electron irradiation tuned the microstructure of 6FDA:BPDA(1:1)-DAM derived CMS membranes by precisely adjusting the irradiation dosage. During electron irradiation, the generated carbon defects induced ultramicropore tightening and oxidized surface formation on the CMS. The electron-tailored ultramicropore size distribution was shifted to be more concentrated in the sub-0.4 nm range, which fits well for discriminating small hydrocarbon gases. As the electron dosage increased, the size-sieving effect of the membrane enormously increased, which in turn enhanced aging resistance. The binary gas permeation test on the studied gas pairs (C2H2,C2H6, and H2/CO2) showed comparable separation capabilities to that of unary gas test and more realistic performances of electron-irradiated CMS membranes. H2/CO2 performances at high temperature (100 and 130 °C) were also calculated to confirm separation behavior and viability of the electron-irradiated CMS, indicating remarkable performance preservation in industrially relevant conditions. In the perspective of post-treatment of nanoporous carbon membranes, electron-irradiation shares similarities with the previously reported hyperaging process. Both methods intentionally reduce the distance between carbon strands or plates, decreasing the free volume of the membrane film and thus changing the gas permeation. However, electron irradiation is not a time-dependent thermal treatment and, more importantly, it combines the advantages of both surface oxidation and pore tightening. Compared to other surface treatment methods, ozone treatment to Matrimid® derived CMS was reported by purging ozone to CMS at room temperature. As with the case of O2 doping treatment, ozone-modified CMS shows similar performance behavior, namely, a selectivity increase and a permeability decrease. However, an insignificant tuning degree of the modified CMS does not create an attractive separation performance for challenging gas pairs such as H2/CO2. In this regard, it can be acknowledged that electron irradiation opens a stage for an attractive tuning factor distinguished from the conventional CMS tuning, which is mainly limited to thermal treatment.

Methods

Materials

4,4-(Hexafluoropropylidene)diphenic Anhydride(6FDA, 98.0%, TCI chemicals), 2,4,6-trimethyl-1,3-phenylenediamine (DAM, 98.5%, TCI chemicals) and 4,4'-Biphenylthalic Anhydride (BPDA, 98%, TCI chemicals) were sublimated in vacuum state before use. Acetic anhydride (AcAn, >99%, Sigma-Aldrich) and 3-methylpyridine (beta picoline, 99%, Sigma-Aldrich) were dried by soaking 3 Å zeolite sieve before use. N-methyl-2-pyrrolidone (NMP, 99%, Alfa Aesar), methacrylic acid for 1 day. Then chemical imidization was proceeded to convert polyamic acid to cyclic polyimide at ambient temperature, using beta picoline as a catalyst and AcAn as a dehydrating agent respectively. The polymer precursor film fabrication

To remove the moisture, the polymer powder was prepared in vacuum oven at 120 °C for 12 h before use. 23 wt% polymer solution was formed by dissolving the dried powder in NMP and placed on a roller for homogeneous quality. The polymer solution, film applicator and a glass plate and beak containing NMP were prepared in glove bag. Then the bag was purged with N2 two or three times, sealed and saturated with NMP for slow solvent evaporation in solution. Next, the solution was cast on a glass plate with film applicator and evaporated thoroughly for 1 day. The polymer film was removed from the plate and placed in DI water, MeOH and n-hexane sequentially for solvent exchange. Afterwards, the polymer film was dried in 120 °C vacuum oven for overnight.

Spin coated film fabrication for GIWAXS analysis

5–8 wt% polymer-NMP solution was prepared and spin-coated on the surface of silicon wafer at 6000 rpm for 60 s. After the coating process, the coated wafer samples were moved immediately into the NMP saturated glove bag for thin and dense film formation. The resultant films had thickness approximately below 1 μm.

Grazing incidence wide angle X-ray scattering (GIWAXS) analysis

Grazing incidence wide angle X-ray scattering (GIWAXS) measurement was performed on beamline 9-A(U-SAXS) in Pohang Accelerator Laboratory (Pohang, Korea) with \( E_r = 11.07 \text{ keV} \) and a wavelength of 1.101 Å. The beam incidence angle was set in the range 0.12–0.15° which is between the critical angle of the sample (polymer or CMS) film and the silicon substrate. The scattered beams from thin film were recorded by a two-dimensional CCD detector, a sample to detector distance of 0.22 m.

CMS membrane fabrication (pyrolysis protocol)

The polymer films were placed on a corrugated quartz plate and the plate was loaded in three-zone tube furnace (SH scientific, SH-80LTG-0M). Providing ultra-high purity (UHP, 99.999%) argon through mass flow meter (Omega Engineering), oxygen level inside the quartz tube was maintained around 30ppm prior to pyrolysis. Then the polymer films were soaked at 500 °C, 600 °C and 675 °C for 2 h (Ramping protocol: RT–250 °C: 10 °C/min, 250–X °C: 3 °C/min, X–20 °C: 0.2 °C/min, 2 hr soaking at X °C, X=500 °C, 600 °C and 675 °C).

Electron irradiation

Electron beam was irradiated on the fabricated CMS membranes using conveyer type electron accelerator No.4 at EB-Tech Co., Ltd. (Daejeon, Korea). The samples were placed on thin aluminum foils and the foils were taped to steel plate stage to hold the sample. Each turn was irradiated by 25kGy, and the number of turns was adjusted according to the final irradiation dosage (For 1 turn, beam energy: 1 MeV, current: 9.3 mA, conveyer speed: 10 m/min).

Single gas permeation

First, the prepared CMS membrane was loaded on permeation cell. Permeation test was carried out with pure single gas, He, H2, CO2, N2, CH4, C2H6, C6H14, and CH3OH at 35 °C under 2 bar upstream pressure using constant volume method. The downstream was kept under vacuum state before measurement start. When the feed is provided from the upstream and passes the membrane, the pressure transducer detects the pressure rise in downstream volume. The pressure rise rate was utilized to calculate the permeability of each gas (Supplementary Fig. 8).

Mixed gas permeation test

Schematic diagram of mixed-gas permeation test is shown in Supplementary Fig. 23. The prepared CMS membranes was mounted on the membrane cell and temperature was maintained at 35 °C. By mass flow
controller (5850E, BROOKS), the equimolar C2H4/C2H6 mixture feed and Helium sweep gas were provided to the membrane cell with a flow rate of 15scm respectively. The rejected feed was vented to retentate side and the permeate penetrants were swept by He to gas chromatography (YL6500 GC, Young Lin Instruments) equipped with flame ionization detector and thermal conductivity detector. In the case of H2/CO2 separation, the equimolar H2/CO2 mixed gas was used as feed gas and Ar was flown as sweep gas to push the permeated gases to GC. For GC quantitative evaluation, additional calibrations using standard samples with various known concentration were carried out.

Pressure decay sorption test
For custom-built pressure decaying system, the fabricated CMS film was smashed into small pieces and loaded into a sample tube. Then the tube was wrapped with heating wire to evacuate the system inside and degas the sample prior to test. After the evacuating process, the system was placed in a water bath with a thermostat to preserve uniform temperature (35°C). The gas reservoir was filled with target gas, equilibrating the temperature for about 10 min, then the target gas was introduced into the sample tube by controlling the valves. The pressure of the reservoir and the sample tube was measured via a high-resolution pressure transducer. (Supplementary Fig. 14).

Data availability
All data supporting the results of this study are available within the article and its supplementary information file. Source data are provided with this paper.

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Author contributions
B.O., H.S. and D.-Y.K. conceived the concept of the research and implemented the experiment. B.O. carried out the unary and binary gas permeation test, pressure decay experiment, TEM, XPS, 87 K Ar physirsorption and elemental analysis. B.O. and H.S. performed 6FDA-BPDA(1:1)-DAM polyimide synthesis and Raman analysis. B.O. and J.C. performed SEM, EDS and GIWAXS analysis. B.O. and D.-Y.K wrote the manuscript. H.S., J.C. and S.L. helped to revise the manuscript. All authors contributed development of research discussion and approved the final manuscript.

Competing interests
The authors declare no competing interests.

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