Chemical Science

EDGE ARTICLE

Cite this: Chem. Sci., 2024, 15, 9641

C All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 5th March 2024 Accepted 15th May 2024 DOI: 10.1039/d4sc01525h

rsc.li/chemical-science

Introduction

Excessive emissions of greenhouse gases, primarily carbon dioxide $(CO₂)$, can lead to the intensification of the greenhouse effect, resulting in global temperature imbalance.¹⁻⁴ It is imperative to develop appropriate and effective methods to address the $CO₂$ emitted from production and daily activities. Until now, carbon capture and sequestration (CCS) and direct air capture (DAC) technologies, which can efficiently capture $CO₂$ from emission sources, are acknowledged as effective approaches for CO_2 treatment.⁵⁻⁸ Exploration of efficient CO_2 capture materials that meet the needs of above-mentioned scenarios is of utmost importance in reducing the impact of greenhouse effect. Although traditional inorganic adsorbents such as activated carbon and zeolite have shown prominent gas separation selectivities,⁹ their low structural designability has limited their use as $CO₂$ capture adsorbents at low concentration in ambient air and indoor confined spaces.

Metal–organic frameworks (MOFs) have emerged as a new family of highly promising porous materials for $CO₂$ capture because of their structural and chemical tunability.¹⁰–¹² Given

Tuo Di[,](http://orcid.org/0000-0002-7904-5003)^{[a](http://orcid.org/0000-0001-6956-9543)} Yukihiro Yoshida, D^{*a} Ken-ichi Otake, D^b Susumu Kitagawa D^b [a](http://orcid.org/0000-0001-6955-3015)nd Hiroshi Kitagawa **D**^{*a}

Exploration of porous adsorbents with high $CO₂/N₂$ selectivity is of great significance for reducing $CO₂$ content in the atmosphere. In this study, a series of isoreticular ultramicroporous fluorinated metal– organic frameworks (MOFs) were prepared to explore the benefits of fluorinated ultramicropores in improving $CO₂/N₂$ selectivity. Gas adsorption measurements revealed that the increase in the number of fluorine atoms in a ligand molecule leads to the increased $CO₂$ uptakes and $CO₂/N₂$ selectivity. Theoretical calculations indicate that the interaction between the fluorine atoms and adsorbed $CO₂$ molecules enhances the $CO₂$ -philicity, offering useful insight into the improvement of $CO₂/N₂$ selectivity in isoreticular frameworks. **EDGE ARTICLE**
 CO Check for undates **Increased CO₂/N₂ selectivity by stepwise**
 CO Check for undates **fluorination in isoreticular ultramicroporous**
 EQUESTANTS CO CO₂ (M₂ selectivity by stepwise
 EQUEST

the self-assembling construction of metal ions/clusters and organic linkers via strong coordination bonds, MOFs have provided new avenues to accurately regulate both their surface and interior by judiciously selecting metal knots and organic ligands to regulate their CO_2 -philicity.^{13,14} It has been found that the interpenetration of microporous frameworks to produce ultramicropores (pore size: $ca.$ 0.5 nm) leads to higher $CO₂$ affinity, thus higher $CO₂/N₂$ selectivity, than those of the parent microporous MOF.¹⁵ In addition, chemical modifications, such as fluorination of component ligands, can also lead to increased $CO₂$ -philicity, because the favourable interactions between the quadrupole of $CO₂$ and fluorine atoms can greatly enhance $CO₂$ $affinity.¹⁶⁻¹⁹$ Thus, it is possible that the combination of fluorination with the ultramicroporous structure offers an optimal platform to systematically investigate the effect of fluorination on $CO₂/N₂$ selectivity. To this end, it is necessary to choose a series of isostructural MOFs with different degrees of fluorination, because uncertainties caused by different structural characteristics inevitably reduce the rationality of the obtained results. It should be mentioned that the fluorination of ligands greatly affects the framework structure in many cases,²⁰⁻²² which will hamper our research to examine the fluorination effect on the gas sorption properties.

Chun and co-workers reported a family of pillar-layered MOFs; Zn^{II}-paddlewheel dimers coordinate with bis-bidentate terephthalate (BDC^{2−}) linkers to construct two-dimensional (2D) square grids, which are connected by bis-monodentate 1,4-diazabicyclo^[2.2.2]octane (DABCO) linkers.²³ Of particular importance is that the three-dimensional (3D) framework structure renders the ultramicropores (pore size ∼0.6 nm) and even remains unchanged by replacing hydrogen atoms on BDC ligands with other functional groups such as methyl and

ROYAL SOCIETY
OF CHEMISTRY

a Division of Chemistry, Graduate School of Science, Kyoto University, Kitashirakawa-Oiwakecho, Sakyo-ku, Kyoto 606-8502, Japan. E-mail: yoshiday@ssc. kuchem.kyoto-u.ac.jp; kitagawa@kuchem.kyoto-u.ac.jp

b Institute for Integrated Cell-Material Sciences (iCeMS), Kyoto University Institute for Advanced Study, Kyoto University, Yoshida Ushinomiya-cho, Sakyo-ku, Kyoto 606- 8501, Japan

 \dagger Electronic supplementary information (ESI) available: Synthesis of H₂1FBDC and additional experimental results such as ${}^{1}\mathrm{H}$ NMR spectra, PXRD patterns, FT-IR spectra, TG profiles, and breakthrough curves. CCDC 2336743 (DMOF-1F) and 2336744 (DMOF-2F). For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d4sc01525h>

halogen groups,²³⁻²⁶ which is inherently suitable for the present research.

In this study, we selected the ultramicroporous Zn_2 (-BDC)(tmBDC)(DABCO) (tmBDC^{2−}: tetramethylterephthalate) as a parent MOF because of its high stability against water vapour compared with its analogues.²⁷ Crystallographic, optical, and sorption studies confirmed the successful substitution of BDC ligands with 2-fluoro-BDC $(1FBDC^{2−})$ or 2,5-difluoro-BDC $(2FBDC^{2−})$ ligands while maintaining the framework structure. Theoretical calculations based on sorption isotherms demonstrated that increasing the fluorine content leads to a pronounced increase in CO_2/N_2 selectivity under low CO_2 concentration conditions, primarily driven by intermolecular interaction between the fluorine atoms and adsorbed $CO₂$ molecules. The present work would provide deeper insights into the rational design of ultramicroporous MOFs for controlling $CO₂/N₂$ selectivity.

Methods

Materials and chemicals

Zinc(II) nitrate hexahydrate $(Zn(NO₃)₂·6H₂O; 99%)$, hydrochloric acid (HCl; 35–37 wt%), potassium permanganate (KMnO4; 99%), and N,N-dimethylformamide (DMF; super dehydrated) were used as purchased from Fujifilm Wako Pure Chemical Industries, Ltd. Terephthalic acid $(H₂BDC; 99%)$ and 1,4-diazabicyclo[2.2.2]octane (DABCO; 98%) were obtained from Tokyo Chemical Industry Co., Ltd. 2,3,5,6-Tetramethylterephthalic acid $(H_2tmBDC, 95%)$ and 3-fluoro-4methylbenzoic acid (98%) were acquired from Combi-Blocks Inc. 2,5-Difluoroterephthalic acid $(H₂2FBDC, 95%)$ was obtained from Enamine Ltd. All these chemicals were used without purification. 2-Fluororoterephthalic acid (H_21FBDC) was synthesized according to the reported literature²⁸ with modest modifications (ESI and Fig. S1†).

Syntheses

 $Zn_2(BDC)(tmBDC)(DABCO)$ (DMOF-0F). DMOF-0F was synthesized according to the reported literature²³ with minor modifications. Typically, $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (0.95 g, 3.2 mmol), H_2BDC (0.27 g, 1.6 mmol), H_2 tmBDC (0.36 g, 1.6 mmol), and DABCO $(0.18 \text{ g}, 1.6 \text{ mmol})$ were mixed in 48 mL DMF. After sonication for 5 min, the mixture was stirred at room temperature for 3 h before being filtered to yield a transparent solution. Then, the solution was transferred into a Teflon-lined stainlesssteel autoclave and heated at 110 °C for 48 h. After being cooled to room temperature, the transparent precipitate was collected and washed with DMF before vacuum drying at 120 °C overnight for further characterization. Yield: 0.47 g (47%). Elemental analysis calcd $(\%)$ for Zn₂(BDC)(tmBDC)(DABCO): C, 49.78; H, 4.50; N, 4.47. Found (%): C, 49.78; H, 4.43; N, 4.55.

 $Zn₂(1FBDC)(tmBDC)(DABCO)$ (DMOF-1F). DMOF-1F was prepared by the procedure described above for DMOF-0F except that H₂1FBDC was used instead of H₂BDC. Yield: 0.52 g (50%). Elemental analysis calcd $(\%)$ for Zn₂(1FBDC)(tmBDC)(DABCO): C, 48.39; H, 4.22; N, 4.34. Found (%): C, 47.51; H, 4.12; N, 4.32.

 $Zn₂(2FBDC)(tmBDC)(DABCO)$ (DMOF-2F). DMOF-2F was prepared by the procedure described above for DMOF-0F except that H₂2FBDC was used instead of H₂BDC. Yield: 0.54 g (51%) and the activation was performed at 105 °C instead of 120 °C. Elemental analysis calcd (%) for $Zn_2(2FBDC)(tmBDC)(DABCO)$: C, 47.08; H, 3.95; N, 4.22. Found (%): C, 46.84; H, 4.02; N, 4.15.

Characterization

Powder X-ray diffraction (PXRD) measurements were performed with a Rigaku MiniFlex600 with 2θ ranging from 5-50° using Cu K α radiation ($\lambda = 1.5418$ Å). Fourier transform infrared (FT-IR) spectra were measured using a PerkinElmer Spectrum 100 spectrometer in attenuated total reflection (ATR) mode. Thermal stability was characterized by thermogravimetric analysis (TGA) on a Bruker TG-DTA2000SA at a heating rate of 5 K min⁻¹ under N₂ atmosphere. The N₂ (77 K and 273 K) and CO₂ (273 K and 298 K) gas adsorption and desorption measurements were performed using a BELSORP-max volumetric adsorption system. Before each measurement, the activated powder sample was activated at 343 K for 12 h. Ideal Adsorbed Solution Theory (IAST) selectivity simulation was performed based on a graphical user interface software, GraphIAST.²⁹ Single-crystal X-ray diffraction (SCXRD) measurements were performed on a Rigaku XtaLAB PRO diffractometer equipped with a HyPix-6000HE detector with a graphite monochromatized Mo Ka radiation ($\lambda = 0.71069$ Å) at 100 K. A single crystal was cooled by a stream of cooled nitrogen gas. Breakthrough experiments were conducted on a BELCAT II-Cryo-10 apparatus. A mixture of $CO₂/N₂$ (15:85 v/v) was passed through a column packed with powdered samples (ca. 0.3 g) at 273 K. The samples were activated at 70 °C for 20-30 min prior to the experiments. Geometry optimizations of all $CO₂$ -adsorbed MOFs were carried out based on grand canonical Monte Carlo (GCMC) simulations using Materials Studio software (Accelrys) based on the universal force field (UFF).³⁰ Chemical Science

Harbors Top Research in the present $P_{\text{eq}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{SPB}}(p_{\text{$

Results and discussion

Crystal structures of the three MOFs were determined by SCXRD measurements, in which as-synthesized crystals were utilized for DMOF-0F and DMOF-1F, whereas heat-treated crystals (see below) were employed for DMOF-2F. Because the guest solvent molecules in the pore space could not be refined due to their disordered nature, the SQUEEZE function in PLATON software³¹ was utilized to remove the residual electron density. The PLATON/SQUEEZE procedure estimated a residual electron count of 175 e−, corresponding to 1.1 DMF molecules per $Zn_2(1FBDC)(tmBDC)(dabco)$ unit. Here, we elaborate only on the refined structures of two fluorinated MOFs, DMOF-1F and **DMOF-2F**, since the structure of the non-fluorinated MOF $(i.e.,$ DMOF-0F) has already been reported in the literature.²³ In fact, the fluorinated MOFs are isomorphous with the reported DMOF-0F. As shown in Fig. 1a–c, 2D (two-dimensional) square grids were constructed by bridging Zn dimers with bis-bidentate BDC-type ligands in a square-planar geometry in the ab plane. The paddlewheel units are connected by nitrogen atoms of

Fig. 1 Expanded cubic-like structure of (a) DMOF-0F, (b) DMOF-1F, and (c) DMOF-2F, in which only one orientation of the disordered xFBDC ligand is shown. (d) Disorder of 1FBDC and tmBDC in DMOF-1F. Hydrogen atoms are omitted for clarity. Colour code: C: grey; F: orange; N: blue; O: red; Zn: green.

DABCO molecules along the c axis, thereby expanding the planar network into a 3D structure. However, their space groups are not identical; namely, **DMOF-1F** belongs to the *I4/mcm* space group, whereas DMOF-2F crystallizes in the P4/mmm space group as in **DMOF-0F.**²³ In the crystals, x FBDC $(x = 1 \text{ or } 2)$ and tmBDC ligands coexist in a disordered manner, where they have different orientations with a dihedral angle of 73.70° for $x = 1$, and 50.67° for $x = 2$ (Fig. 1d). The equivalent occupancy between xFBDC and tmBDC of 0.5 is consistent with the elemental analysis data. In addition, considering the fact that the xFBDC ligands can coordinate to Zn ions in a random manner (Fig. $S2$ and $S3\dagger$), fluorine atoms in *xFBDC* ligands are disordered over four sites with an equivalent occupancy of 0.25 for DMOF-1F (Fig. 1b) and over two orientations, i.e., 2,5- and 3,6-positions, with an equivalent occupancy of 0.5 for DMOF-2F (Fig. 1c). All three MOFs possess large 3D inter-connected voids with one-dimensional channels running along the c axis with windows through the a and b axes. Although the fluorine atoms slightly protrude into the channels, there are very little difference in the sizes of pores and windows, with an average diameter of ca 0.6 nm in all three MOFs. The procedure to estimate the pore sizes is shown in Fig. S4.† It is noteworthy that these pores are apparently larger than the kinetic diameter of common gas molecules (e.g., N_2 (0.364 nm) and CO_2 (0.330 nm)), which allows their practical application in gas transportation and storage. In addition, it is expected that the surface density of fluorine atoms in the pores has a significant effect on the affinity for various gas molecules,^{32,33} which would be manifested in the adsorption isotherms, as discussed below.

PXRD measurement was utilized to confirm the phase purity of the polycrystalline samples. As shown in Fig. 2a and S5,† the

Fig. 2 (a) PXRD patterns, (b) FT-IR spectra, and (c) TGA profiles of assynthesized DMOF-0F (dark green), DMOF-1F (pale blue), and DMOF-2F (orange). In (b), enlarged spectra in the range of $900-1350$ cm⁻¹ are shown in the panel to the right.

PXRD patterns of the three as-synthesized MOFs match well with those simulated from the SCXRD data without any trace impurity peaks, indicating the high purity of the polycrystalline samples. FT-IR spectroscopy was applied to investigate the chemical structure of the as-synthesized MOFs. In Fig. 2b, the peak at 749 cm^{-1} can be assigned to the stretching vibration of Zn–O bonds originating from the dinuclear paddlewheel units.³⁴ The peaks at 1619 is ascribed to the stretching vibration of carboxylate groups,³⁵ whereas the peak at 1057 cm⁻¹ is due to the asymmetric stretching vibration of $N-C_3$ bonds in pillar DABCO molecules.³⁴ In addition, the C-F stretching band was observed at 1227 cm⁻¹ (DMOF-1F) and 1184 cm⁻¹ (DMOF-2F) in fluorinated MOFs. The observation of conspicuous characteristic peaks of DMF, which was used as the reaction solvent, at 1664 cm⁻¹ (C=O stretching), 1256 cm⁻¹ (C–N stretching), and 660 cm⁻¹ (O=C-N bending) indicates the inclusion of DMF in the as-synthesized MOFs.³⁶ Thermal stability of the assynthesized MOFs was evaluated using TGA in the temperature range from room temperature to 600 $^{\circ}$ C under N₂ atmosphere. As shown in Fig. 2c, all the TG profiles exhibit similar thermal behaviour, showing a huge weight loss below 200 °C. Combining the results from FT-IR, we conclude that the decrease in weight is primarily caused by the removal of solvent DMF molecules adsorbed in the channel and on the surface of particles. Subsequently, there is a plateau between 200 and 300 °C followed by a weight loss possibly due to the removal of terephthalate or DABCO molecules. Based on the TG profiles, it can be concluded that the three MOFs are thermally stable up to approximately 300 °C under N_2 atmosphere, which is comparable to those of some well-known MOFs; e.g., HKUST-1 $(Cu₃(BTC)₂; H₃BTC: 1,3,5-benzentericarboxylic acid; up to ca.$ 240 °C)³⁷ and MOF-801 (Zr₆O₄(OH)₄(fumarate)₆; up to *ca.* 260 °C).³⁸

Because the DMF molecules occupy the pore space throughout the frameworks, appropriate methods should be applied to completely remove the solvent species in the pores before subsequent adsorption measurements. After plenty of attempts, we finally determined that the dynamic vacuum at elevated temperature is the optimal method to achieve the complete removal of DMF in the pores while keeping the framework structure. All the activated MOFs after thermal treatment at 120 °C for DMOF-0F and DMOF-1F and 105 °C for DMOF-2F for 10–12 h exhibit the PXRD patterns similar to those of as-synthesized ones with respect to the peak positions and widths (Fig. S6†). Given that the vibrational bands of DMF molecules (1664, 1256, and 660 $\mathrm{cm}^{-1})$ and weight loss due to DMF evaporation disappear in the FT-IR spectra (Fig. S7†) and TG profiles (Fig. S8†), respectively, it is obvious that DMF molecules are completely removed through the dynamic vacuum, and therefore, the activated MOFs are available for gas adsorption measurements.

Porous properties of the three MOFs were evaluated by N_2 adsorption/desorption measurements at 77 K. As shown in Fig. 3a, the three MOFs exhibit a steep increase in the lowpressure region $(P/P_0 < 0.01)$, suggesting the presence of abundant permanent micropores throughout the frameworks. Subsequently, they exhibit negligible N_2 uptakes in the relative pressure range 0.1–0.8 with a typical type-I isotherm in the IUPAC classification.³⁹ In the high-pressure region ($P/P_0 > 0.9$), all three isotherms show a slight increase, indicating the presence of macropores arising from the inter-particulate voids due to the loose packing of particles. The uptake of N_2 adsorption at $P/P_0 = 0.97$ in **DMOF-0F** (263 cm³ g⁻¹) is lower than those of **DMOF-1F** (308 $\mathrm{cm}^3 \mathrm{~g}^{-1}$) and **DMOF-2F** (313 $\mathrm{cm}^3 \mathrm{~g}^{-1}$). The Brunauer–Emmett–Teller (BET) surface areas of DMOF-0F, 1F, and 2F are estimated to be 949, 1123, and 1225 m² g⁻¹ (Table 1), respectively, which are close to the reported value for DMOF-0F

Fig. 3 (a) N_2 adsorption (closed symbols) and desorption (open symbols) isotherms at 77 K and (b) pore size distribution of DMOF-0F (dark green circles), DMOF-1F (pale blue squares), and DMOF-2F (orange triangles).

(1100 m^2 g⁻¹).²³ The total pore volumes (V_{total}) calculated at a P/P_0 of 0.97 are 0.41, 0.48, and 0.48 $\rm cm^3~g^{-1},$ respectively. Pore size distribution calculated using the micropore analysis (MP) method shows that the three MOFs possess prominent ultramicropores with a size of ca. 0.6 nm (Fig. 3b), which is consistent with that expected from the crystallographic data (ca. 0.6 nm).

Considering the relatively high BET surface areas (S_{BET}) , the present MOFs with comparable ultramicropore sizes with different fluorination environments allow us to investigate the effect of fluorination on CO_2 -philicity, because several fluorinated MOFs have been reported to exhibit Coulomb interactions between the negatively-charged fluorine atoms and positivelycharged carbon atom in $CO₂$ molecules.³³ Fig. 4a and b display the CO₂ sorption isotherms of the three MOFs collected at 298 and 273 K, respectively. At both temperatures, all the MOFs show a gradual increase in $CO₂$ uptake over the whole pressure region with a minimal hysteresis loop, indicating the physisorptiondominant process. They exhibit a relatively high $CO₂$ uptake, possibly as a consequence of their high S_{BET} and V_{total} . The CO₂ uptake at 273 K follows the order DMOF-0F (102.1 cm³ g⁻¹, 4.55 mmol g^{−1}, 20.0 wt%) < **DMOF-1F** (106.1 cm³ g^{−1}, 4.73 mmol $\rm g^{-1}, \ 20.8 \ \ wt\%) <$ <code>DMOF-2F</code> $(107.4 \ \rm cm^3 \ \rm g^{-1}, \ 4.79 \ \rm mmol \ \rm g^{-1},$ 21.1 wt%). Although the lower V_{total} value of DMOF-0F estimated from the N_2 sorption isotherm at 77 K may lead to the lower CO_2 uptake, it is apparent that the higher $CO₂$ uptake in DMOF-2F than that of **DMOF-1F** with comparable V_{total} is attributed to the higher fluorine content. These values are comparable or superior to some well-known MOFs, such as SNU-4 $(Zn_2(abtc)(DMF)_2;$ H_4 abtc: 3,3',5,5'-azobenzenetetracarboxylic acid, 20.6 wt%, surface area 1460 m² g⁻¹),⁴⁰ SNU-5' (Cu₂(abtc)(DMF)₂; 19.2 wt%, surface area 1260 m² g⁻¹),⁴⁰ Zn₂(BTTB)(DMF)₂ (H₄BTTB: 1,2,4,5-tetrakis(4-carboxyphenyl)benzene; 19.7 wt%, surface area 1370 m² g⁻¹),¹⁸ and SNU-21 (Cu₂(TCM)(H₂O)₂, H₄TCM: tetrakis[4-(carboxyphenyl)oxamethyl]methane; 18.4 wt%, surface area 905 m² g^{-1}).⁴¹ The reversible isotherm suggests that the MOFs under study can release the adsorbed $CO₂$ molecules without the need of any additional thermal energy, which is advantageous for the real-world $CO₂$ capture and regeneration of porous materials.⁴² The PXRD patterns after the $CO₂$ adsorption/ desorption measurement (Fig. S9[†]) confirmed the retention of the structure during the sorption process. Chemical Science

Published on the matter on 16 May 2022. The two the state of the state of the state of the state of the matter of the state of the matter of the state of the state of the state of the state of the state

To examine the thermodynamics of the $CO₂$ adsorption, isosteric heat of adsorption (Q_{st}) toward $CO₂$ were calculated from the Clausius–Clapeyron equation:⁴³ $ln(P_2/P_1) = Q_{st} \times (T_2 - T_1)$ T_1 /(RT₁T₂), where P_n and T_n (n = 1 or 2) denote the pressure and

 a BET surface area. b Total pore volume. c Values at 0.3 bar.

Fig. 4 CO₂ adsorption (closed symbols) and desorption (open symbols) isotherms of DMOF-0F (dark green circles), DMOF-1F (pale blue squares), and DMOF-2F (orange triangles) at (a) 298 K and (b) 273 K. (c) Isosteric heats of adsorption (Q_{st}) for CO₂ calculated from the data at 298 and 273 K.

temperature values, respectively, for the *n*th isotherm, and R is the gas constant expressed in the appropriate unit (8.314 J K^{-1}) mol−¹), using the adsorption isotherms at 298 and 273 K. As shown in Fig. 4c, the Q_{st} values are estimated to be 17.2-25.7 kJ mol⁻¹ for DMOF-0F, 19.1–25.5 kJ mol⁻¹ for DMOF-1F, and 20.2–26.7 kJ mol⁻¹ for **DMOF-2F**, depending on the degree of $CO₂$ loading, which lie in the typical range for $CO₂$ adsorption of MOFs.^{9,44} It is noteworthy that the Q_{st} values, which are lower than the energy of the chemical bonds, is again indicative of the physisorption process (<40 kJ mol⁻¹),⁴⁵ which facilitates the controlled release of $CO₂$ gas as mentioned above.⁴²

It is noteworthy that for the removal of $CO₂$ from flue gas in coal power plants (also known as post-combustion $CO₂$ capture), the $CO₂$ concentration must be reduced to a certain level with respect to N_2 . Thus, it is necessary to explore adsorbents that exhibit a high $CO₂/N₂$ selectivity to enable efficient capture of $CO₂$ under such conditions. The $CO₂$ and $N₂$ adsorption isotherms of the MOFs under study were compared

to investigate the adsorption selectivity of $CO₂$ over N₂. As shown in Fig. 5a–c, the adsorption amount of $CO₂$ is remarkably higher than that of N_2 at 273 K and the CO_2/N_2 selectivity is estimated to be 8.4, 11.3, and 14.8 for DMOF-0F, DMOF-1F, and DMOF-2F, respectively, based on the initial slope method (Fig. 5d–f).⁴² Although the selectivity values are less than those of a certain number of MOFs reported so far,¹ the results deliver the message that fluorination has a favourable effect on $CO₂/N₂$ selectivity. Dynamic breakthrough measurements using a gas mixture of CO_2/N_2 (15:85) at 273 K demonstrate the effective separation via a one-step experiment, although no significant differences were found among the three MOFs (Fig. S10†). The difference in CO_2/N_2 selectivity depending on the number of fluorine atoms $(8.4-14.8)$ is more pronounced compared with that of ultramicroporous fluorinated MOF, $Ni(tpt)(xF-$ OPA)(H_2O) (tpt: 2,4,6-tri(4-pyridinyl)-1,3,5-triazine; H_2OPA : ophthalic acid; 13.8–15.8), with 1D channels of ca. 0.62 nm in diameter, in which 4-fluoro-OPA $(x = 1; TKL-105)$, 3,6-difluoro-

Fig. 5 CO₂ and N₂ sorption isotherms of at 273 K for (a) DMOF-0F, (b) DMOF-1F, and (c) DMOF-2F. Initial slope fitting for CO₂ and N₂ adsorption isotherms at 273 K for (d) DMOF-0F, (e) DMOF-1F, and (f) DMOF-2F.

OPA $(x = 2; TKL-106)$, and 3,4,5,6-tetrafluoro-OPA $(x = 4; TKL-106)$ 107) were utilized as a bridging ligand.⁴⁶ Although the factors that determine the selectivity remain unclear in these systems, the CH \cdots F hydrogen bonds between the pyridine moieties of and fluorine atoms of $xF-OPA$ within the pore walls in TKL-106 and 107 would suppress the intermolecular $CF\cdots C(CO_2)$ interactions between the $xF-OPA$ ligands and $CO₂$ molecules. Although DMOF-1F and DMOF-2F also involve $CH\cdots F$ hydrogen bonds, the C-H groups are in the fast-rotating DABCO ligands⁴⁷ which may allow the effective intermolecular $CF\cdots C(CO_2)$ interactions between $xFBDC$ ligands and $CO₂$ molecules. Park and co-workers reported the increased $CO₂/N₂$ selectivity by the partial substitution of 2-amino-BDC (NH₂BDC^{2−}) with 2,3,5,6tetrafluoro-BDC (4FBDC²⁻) in amino-functionalized MIL-101(Cr) with a chemical composition of $Cr_3O(OH)(H_2O)_2(NH_2 BDC$ ₃;⁴⁸ the CO_2/N_2 selectivity in the 4FBDC-introduced MOFs (92–108) is signicantly higher than that of the pristine MOF (51). However, the authors concluded that the increased $CO₂$ adsorption in 4FBDC-introduced MOFs arises mainly from the decreased pore sizes (1.7 to 0.9 cm³ g^{-1}) rather than the increased fluorine content. On the other hand, we succeeded in improving the CO_2/N_2 selectivity mainly by controlling the number of fluorine atoms in a BDC ligand while properly keeping the porous properties (0.41 to 0.48 $\rm cm^3~g^{-1}$). Thus, the present result is the first demonstration of fluorinated ultramicroporous MOFs to control the $CO₂/N₂$ selectivity at a significant level by controlling the fluorine content (Table $S1\dagger$).

To demonstrate the ability of the three MOFs to capture trace $CO₂$, their $CO₂/N₂$ selectivity at atmospheric concentration (500 ppm for $CO₂$) was calculated using IAST.²⁹ As illustrated in Fig. 6a, the CO_2/N_2 selectivity varies depending on the fluorination and increases with increasing pressure. The selectivity in the low-pressure region (0.3 bar) is in the order **DMOF-0F** $(12.4) <$ DMOF-1F $(14.5) <$ DMOF-2F (21.9) , which is consistent with the results obtained from the initial slope method (Fig. 6b). The difference between DMOF-0F and DMOF-1F (2.1) is less pronounced than that between DMOF-1F and DMOF-2F (7.4), which is consistent with the calculated Q_{st} values (Table 1).

GCMC simulations were conducted to elucidate the origin of the $CO₂$ capture behaviour at the microscopic level. The geometric structures of the MOFs were optimized with the assumption that there are four $CO₂$ molecules in each 1D channel in the unit cell at $P/P_0 = 0.3$ at 273 K. In **DMOF-0F**, there are several $CH\cdots O(CO_2)$ hydrogen bonding interactions between the tmBDC or DABCO ligands and $CO₂$ molecules with an H \cdots O distance of 2.88 Å (Site I), 2.89 Å (Site II), 2.94 Å (Site III), and 2.97 and 3.01 Å (Site IV) (vs. the sum of the van der Waals radii: 2.72 Å^{49} as shown in Fig. 6c, d and Table S2.† Such interactions were found in CO_2 -incorporated DMOF-1F framework, with a distance of 3.02 and 3.04 Å (Site I), 2.98 and 2.98 Å (Site II), and 2.90 and 3.00 Å (Site III) (Fig. 6e and f). In addition, **DMOF-0F** shows several $\pi \cdots C(CO_2)$ interactions^{19,50} between $CO₂$ molecules and benzene rings of BDC-based ligands, with the nearest C···C distance of 3.45 Å (Site I), 3.60 Å (Site II), and 3.53 Å (Site III). However, there is no signicant interaction between the fluorine atoms and $CO₂$ molecules in the pore of DMOF-1F, although a Coulomb attractive force is expected

Fig. 6 (a) Simulated IAST selectivity of $CO₂/N₂$ for DMOF-0F (dark green line), DMOF-1F (pale blue line), and DMOF-2F (orange line) under atmospheric CO_2 concentration (i.e., 500 ppm of CO_2 to N_2). (b) CO2/N2 selectivity of three MOFs based on initial slope method (pale blue) and IAST method (orange). Simulated interactions between the framework and adsorbed $CO₂$ molecules in (c and d) DMOF-0F, (e and f) DMOF-1F, (g and h) DMOF-2F, where $CO₂$ molecules with no effective interactions are emitted. CH \cdots O(CO₂), CF \cdots C(CO₂), and π \cdots $C(CO₂)$ interactions are shown in pink, red, and light green dotted lines, respectively.

between the negatively-charged fluorine atoms and positivelycharged carbon atom in $CO₂$ molecules.³³ We note that the increase in the number of $CO₂$ molecules in a channel in the unit cell from 4 (corresponding to $P/P_0 \sim 0.3$) to 6 (corresponding to $P/P_0 \sim 0.4$) results in the formation of short CF \cdots $C(CO₂)$ contacts with a distance of 3.30 Å (vs. the sum of the van der Waals radii: 3.17 Å;⁴⁹ Fig. S11†). In **DMOF-2F** (Fig. 6g and h), $CO₂$ molecules in the pores exhibit a significant interaction with fluorine atoms with an $F \cdots C$ distance of 3.33 Å (Site II in Fig. 6g). Such a favourable $CF\cdots C(CO_2)$ interaction has also been found in fluorinated Ce-MIL-140 (CeO(4FBDC)),⁵¹ in which an $CF\cdots C(CO_2)$ interaction with a distance of 3.3 Å results in a higher affinity for $CO₂$ and boosts the $CO₂$ selectivity. Therefore, it is apparent that the $CF\cdots C(CO_2)$ interaction plays a pivotal role in the higher $CO₂/N₂$ selectivity in DMOF-2F than those of DMOF-0F and DMOF-1F.

Conclusions

In this study, a series of isoreticular MOFs, namely, DMOF-0F, DMOF-1F, and DMOF-2F, possessing ultramicropores with varying degrees of fluorination were synthesized through the rational design of component species and framework structure. Both experimental and theoretical studies demonstrate that the $CO₂/N₂$ selectivity in the MOFs under study is increased by incorporating a higher number of fluorine atoms into the ultramicropore walls. It appears that the strong $CF\cdots C(CO_2)$ interaction between 2FBDC ligands and $CO₂$ molecules in DMOF-2F, predicted based on the Monte Carlo simulation, makes it the best adsorbent for $CO₂$ among the three MOFs. Given that the study of the relationship between the fluorine content and $CO₂/N₂$ selectivity in ultramicroporous MOFs is still in its infancy, this work provides deeper insight into the design and synthesis of MOFs with controllable $CO₂/N₂$ selectivity. Edge Article

application of DMOF-DF,

application Access Articles. The Equilibrican Science

Conclusion Conclusion and Creative in the State Article is likely and the state Article is likely and the state of DMOF-DF.

Co

Data availability

Additional details regarding experimental and calculation data are given in the ESI.†

Author contributions

T. D., Y. Y., and H. K. conceived the research and designed the experiments. T. D. performed all the experiments, measurements, and calculations. Dynamic breakthrough measurements were performed by T. D. with the support of K.-i. O. and S. K. T. D., Y. Y., and H. K. cowrote the manuscript. All the authors discussed and commented on the paper.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We are grateful to Prof. Tokutaro Komatsu (Nihon University) for his useful advice on the GCMC simulations. This work was supported by the ACCEL program (JPMJAC1501) of the Japan Science and Technology Agency (JST), JSPS KAKENHI (Grant No. 20H02708 and 20H05623). T. D. was supported by JST SPRING (Grant No. JPMJSP2110).

References

- 1 K. Sumida, D. L. Rogow, J. A. Mason, T. M. McDonald, E. D. Bloch, Z. R. Herm, T.-H. Bae and J. R. Long, Chem. Rev., 2012, 112, 724–781.
- 2 G. Singh, J. Lee, A. Karakoti, R. Bahadur, J. Yi, D. Zhao, K. AlBahily and A. Vinu, Chem. Soc. Rev., 2020, 49, 4360–4404.
- 3 M. T. Dunstan, F. Donat, A. H. Bork, C. P. Grey and C. R. Müller, Chem. Rev., 2021, 121, 12681–12745.
- 4 G. Cui, J. Wang and S. Zhang, Chem. Soc. Rev., 2016, 45, 4307–4339.
- 5 X. Shi, H. Xiao, H. Azarabadi, J. Song, X. Wu, X. Chem and K. Lackner, Angew. Chem., Int. Ed., 2020, 59, 6984–7006.
- 6 X. Zhu, W. Xie, J. Wu, Y. Miao, C. Xiang, C. Chen, B. Ge, Z. Gan, F. Yang, M. Zhang, D. O'Hare, J. Li, T. Ge and R. Wang, Chem. Soc. Rev., 2022, 51, 6574–6651.
- 7 E. S. Sanz-Pérez, C. R. Murdock, S. A. Didas and C. W. Jones, Chem. Rev., 2016, 116, 11840–11876.
- 8 S. Chowdhury, Y. Kumar, S. Shrivastava, S. K. Patel and J. S. Sangwai, Energy Fuels, 2023, 37, 10733–10757.
- 9 S. Choi, J. H. Drese and C. W. Jones, ChemSusChem, 2009, 2, 796–854.
- 10 J. Yu, L.-H. Xie, J.-R. Li, Y. Ma, J. M. Seminario and P. B. Balbuena, Chem. Rev., 2017, 117, 9674–9754.
- 11 L. Jiang, W. Liu, R. Q. Wang, A. Gonzalez-Diaz, M. F. Rojas-Michaga, S. Michailos, M. Pourkashanian, X. J. Zhang and C. Font-Palma, Prog. Energy Combust. Sci., 2023, 95, 101069.
- 12 C. A. Trickett, A. Helal, B. A. Al-Maythalony, Z. H. Yamani, K. E. Cordova and O. M. Yaghi, Nat. Rev. Mater., 2017, 2, 17045.
- 13 D. M. Venturi, M. S. Notari, R. Bondi, E. Mosconi, W. Kaiser, G. Mercuri, G. Giambastiani, A. Rossin, M. Taddei and F. Costantino, ACS Appl. Mater. Interfaces, 2022, 14, 40801– 40811.
- 14 A. S. Palakkal and R. S. Pillai, J. Phys. Chem. C, 2020, 124, 16975–16989.
- 15 P. Nugent, Y. Belmabkhout, S. D. Burd, A. J. Cairns, R. Luebke, K. Forrest, T. Pham, S. Ma, B. Space, L. Wojtas, M. Eddaoudi and M. J. Zaworotko, Nature, 2013, 495, 80–84.
- 16 J. R. Fried and N. Hu, Polymer, 2003, 44, 4363–4372.
- 17 P. Deria, J. E. Mondloch, E. Tylianakis, P. Ghosh, W. Bury, R. Q. Snurr, J. T. Hupp and O. K. Farha, J. Am. Chem. Soc., 2013, 135, 16801–16804.
- 18 Y.-S. Bae, O. K. Farha, J. T. Hupp and R. Q. Snurr, J. Mater. Chem., 2009, 15, 2131–2134.
- 19 A. Torrisi, C. Mellot-Draznieks and R. G. Bell, J. Chem. Phys., 2009, 130, 194703.
- 20 D. Smets and U. Ruschewitz, Z. Anorg. Allg. Chem., 2020, 646, 1157–1167.
- 21 Y. Wang, Z. Liu, Y. Li, Z. Bai, W. Liu, Y. Wang, X. Xu, C. Xiao, D. Sheng, J. Diwu, J. Su, Z. Chai, T. E. Albrecht-Schmitt and S. Wang, J. Am. Chem. Soc., 2015, 137, 6144–6147.
- 22 R. Christoffels, C. Breitenbach, J. Weber, L. Körtgen, C. Tobeck, M. Wilhelm, S. Mathur, J. Neudörfl, M. Sadegh, Z. Farid, M. Maslo, E. Strub and U. Ruschewitz, Cryst. Growth Des., 2022, 22, 681–692.
- 23 H. Chun, D. N. Dybtsev, H. Kim and K. Kim, Chem. Eur. J., 2005, 11, 3521–3529.
- 24 L. K. Cadman, J. K. Bristow, N. E. Stubbs, D. Tiana, M. F. Mahon, A. Walsh and A. D. Burrows, Dalton Trans., 2016, 45, 4316–4326.
- 25 K. Uemura, Y. Yamasaki, F. Onishi, H. Kita and M. Ebihara, Inorg. Chem., 2010, 49, 10133–10143.
- 26 N. C. Burtch, H. Jasuja, D. Dubbeldam and K. S. Walton, J. Am. Chem. Soc., 2013, 135, 7172–7180.
- 27 H. Jasuja, N. C. Burtch, Y.-G. Huang, Y. Cai and K. S. Walton, Langmuir, 2013, 29, 633–642.
- 28 S. T. Meek, J. J. Perry, S. L. Teich-McGoldrick, J. A. Greathouse and M. D. Allendorf, Cryst. Growth Des., 2011, 11, 4309–4312.
- 29 E. Dautzenberg, S. van Hurne, M. M. J. Smulders and L. C. P. M. de Smet, Comput. Phys. Commun., 2022, 280, 108494.
- 30 A. K. Rappé, C. J. Casewit, K. S. Colwell, W. A. Goddard and W. M. Skiff, J. Am. Chem. Soc., 1992, 114, 10024–10035.
- 31 A. L. Spek, Acta Crystallogr., Sect. C: Struct. Chem., 2015, 71, 9–18.
- 32 D. Balestri, I. Bassanetti, S. Canossa, C. Gazzurelli, A. Bacchi, S. Bracco, A. Comotti and P. Pelagatti, Cryst. Growth Des., 2018, 18, 6824–6832.
- 33 S. Kumar, B. Mohan, C. Fu, V. Gupta and P. Ren, Coord. Chem. Rev., 2023, 476, 214876.
- 34 L. Peng, S. Wu, X. Yang, J. Hu, X. Fu, Q. Huo and J. Guan, RSC Adv., 2016, 6, 72433–72438.
- 35 M. Nara, H. Torii and M. Tasumi, J. Phys. Chem., 1996, 100, 19812–19817.
- 36 G. Durgaprasad, D. N. Sathyanarayana and C. C. Patel, Bull. Chem. Soc. Jpn., 1971, 44, 316–322.
- 37 S. S.-Y. Chui, S. M.-F. Lo, J. P. H. Charmant, A. G. Orpen and I. D. Williams, Science, 1999, 283, 1148–1150.
- 38 G. Wißmann, A. Schaate, S. Lilienthal, I. Bremer, A. M. Schneider and P. Behrens, Microporous Mesoporous Mater., 2012, 152, 64–70.
- 39 M. Thommes, K. Kaneko, A. V. Neimark, J. P. Olivier, F. Rodriguez-Reinoso, J. Rouquerol and K. S. W. Sing, Pure Appl. Chem., 2015, 87, 1051–1069. Chemical Science

27 H. Jacques Nr. G. Turks, V. G. Lang, V. G. Lang, V. G. Lang, V. R. Chemical Cypte, V. R. Chemical J. P. Chinese

2024. A. Carolinear art d. V. N. Science. Published Cypte, Coreel Dec. W. N. Science. A.
	- 40 Y.-G. Lee, H. R. Moon, Y. E. Cheon and M. P. Suh, Angew. Chem., Int. Ed., 2008, 47, 7741–7745.
	- 41 T. K. Kim and M. P. Suh, Chem. Commun., 2011, 47, 4258– 4260.
	- 42 H. Gao, L. Ding, H. Bai, A. Liu, S. Li and L. Li, J. Mater. Chem. A, 2016, 4, 16490–16498.
	- 43 H. Pan, J. A. Ritter and P. B. Balbuena, Langmuir, 1998, 14, 6323–6327.
	- 44 J. M. Simmons, H. Wu, W. Zhou and T. Yildirim, Energy Environ. Sci., 2011, 4, 2177–2185.
	- 45 B. von Oepen, W. Kördel and W. Klein, Chemosphere, 1991, 22, 285–304.
	- 46 D.-S. Zhang, Z. Chang, Y.-F. Li, Z.-Y. Jiang, Z.-H. Xuan, Y.-H. Zhang, J.-R. Li, Q. Chen, T.-L. Hu and X.-H. Bu, Sci. Rep., 2013, 3, 3312.
	- 47 N. C. Burtch, A. Torres-Knoop, G. S. Foo, J. Leisen, C. Sievers, B. Ensing, D. Dubbeldam and K. S. Walton, *J. Phys. Chem.* Lett., 2015, 6, 812-816.
	- 48 J. M. Park, G.-Y. Cha, D. Jo, K. H. Cho, J. W. Yoon, Y. K. Hwang, S.-K. Lee and U.-H. Lee, Chem. Eng. J., 2022, 444, 136476.
	- 49 A. Bondi, J. Phys. Chem., 1964, 68, 441–451.
	- 50 A. Torrisi, C. Mellot-Draznieks and R. G. Bell, J. Chem. Phys., 2010, 132, 044705.
	- 51 Z. Zhang, S. B. Peh, R. Krishna, C. Kang, K. Chai, Y. Wang, D. Shi and D. Zhao, Angew. Chem., Int. Ed., 2021, 60, 17198–17204.