Increased CO$_2$ Affinity and Adsorption Selectivity in MOF-801 Fluorinated Analogues

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ABSTRACT: The novel Zr$^{IV}$-based perfluorinated metal–organic framework (PF-MOF) [$\text{Zr}_6\text{O}_4\text{F}_3\text{O}_2\text{FUM}$]$_{16}$ (ZrTFS) was prepared under solvent-free conditions using the commercially available tetrafluorosuccinic acid (H$_2$TFS) as a bridging ditopic linker. Since H$_2$TFS can be seen as the fully aliphatic and perfluorinated C$_3$ analogue of fumaric acid, ZrTFS was found to be isoreticular to zirconium fumarate (MOF-801). The structure of ZrTFS was solved and refined from X-ray powder diffraction data. Despite this analogy, the gas adsorption capacity of ZrTFS is much lower than that of MOF-801; in the former, the presence of bulky fluorine atoms causes a considerable window size reduction. To have PF-MOFs with more accessible porosity, postsynthetic exchange (PSE) reactions on (defective) MOF-801 suspended in H$_2$TFS aqueous solutions were carried out. Despite the different H$_2$TFS concentrations used in the PSE process, the exchanges yielded two mixed-linker materials of similar minimal formulae [$\text{Zr}_6\text{O}_4\text{F}_3\text{O}_2\text{FUM}_2\text{FUM}_2$]$_{16}$ (PF-MOF1) and [$\text{Zr}_6\text{O}_4\text{F}_3\text{O}_2\text{FUM}_2\text{FUM}_2\text{HTFS}$]$_{16}$ (PF-MOF2). This confirms the beneficial effect coming from the introduction of fluorinated linkers in MOFs on their CO$_2$ adsorption ability. Finally, solid-state density functional theory calculations were carried out to cast light on the structural features and on the thermodynamics of CO$_2$ adsorption in MOF-801 and ZrTFS. Due to the difficulties in modeling a defective MOF, an intermediate structure containing both linkers in the framework was also designed. In this structure, the preferential CO$_2$ adsorption site is the tetrahedral pore in the “UIO-66-like” structure. The extra energy stabilization stems from a hydrogen bond interaction between CO$_2$ and a hydroxyl group on the inorganic cluster.

KEYWORDS: metal–organic frameworks (MOFs), carbon dioxide capture, zirconium, fluorinated linkers, DFT calculations

INTRODUCTION

The urgent need to reduce CO$_2$ emissions is pushing scientists and engineers toward the development of technologies that should prevent further buildup of this greenhouse gas in the atmosphere. Among the technologies under scrutiny, carbon capture and storage (CCS)\(^4\) from large point sources and direct air capture (DAC)\(^5\) are seen as viable options for rapid large-scale deployment. The state-of-the-art sorbents for CCS and DAC are typically amine-based: aqueous methanolamine solutions for CCS\(^3\) and solid-supported amine adsorbents for DAC.\(^4\) In both cases, the working principle exploits the formation of covalent bonds between the amine groups and CO$_2$ to afford carabamates (chemical adsorption or chemisorption). The formation of strong covalent bonds is associated with a high heat of absorption or adsorption, which is convenient to achieve high capture loading at low partial pressures but poses challenges in terms of regeneration of the sorbent and achievable working capacity.\(^5,6\) While DAC demands a sorbent with a high enthalpy of ab/adsorption, due to the large entropic loss associated with CO$_2$ capture from a feed that contains only ~415 ppm of CO$_2$, CCS deals with CO$_2$ concentrations in the range between 4 and 30% and could greatly benefit from a sorbent with a mild heat of adsorption, i.e., a physisorbtent.\(^7\) To this end, adsorbents such as porous...
starting from pure zirconium fumarate MOF-801. The relationship between CO₂ adsorption capacity/selectivity and the extent of fluorination in solid adsorbents has been systematically studied.

**EXPERIMENTAL SECTION**

**Materials and Methods.** All chemicals were commercially available and used as received from the specified vendors without further purification. Zirconium chloride (ZrCl₄) was purchased from Alfa Aesar. Tetrafluoro succinic acid (H₂TFS) was purchased from Fluorchem. Fumaric acid (H₂FUM) was purchased from Merck. Acetic acid (AA) was purchased from Carlo Erba. Powder X-ray diffraction (PXRD) patterns were collected in reflection geometry in the 4°–40° 2θ range, with a 40 s step⁻¹ counting time and with a step size of 0.016° on a PANalytical XPERT PRO diffractometer, a PW3050 goniometer, equipped with an X'Celerator detector and using a Cu Kα radiation source. The long fine focus (LFF) ceramic tube operated at 40 kV and 40 mA. Quantitative ¹H NMR and ¹⁹F NMR analyses were performed at an ambient temperature on either a Bruker Ascend 400 MHz spectrometer (MOF-801) or a Jeol JNM-ECZ500 instrument equipped with a RoyalProbe Broadband probe (PF-MOF1 and PF-MOF2). ¹H acquisition time 4.56 s, relaxation delay 3 s, 4 scans; ¹⁹F: acquisition time 0.27 s, relaxation delay 4 s, 16 scans. The solids (ca. 20 mg) were held in an oven at 120 °C for 2 h before being digested for 24 h in 1.5 mL of 1 M NaOH in D₂O (MOF-801) or in 1.0 mL of 1 M NaOH in D₂O (PF-MOF1 and PF-MOF2). The NMR tubes were then loaded with the solution, which was filtered through cotton wool to avoid the presence of solid particles in suspension. 3-Fluorobenzoic acid (0.029 M) was used as the ¹H internal standard for the fumarate and acetate quantification in MOF-801. 2,6-Difluorobenzoic acid (0.1 M) was used as both ¹H and ¹⁹F internal standards for the fumarate and tetrafluoro succinate quantification in the mixed-linker MOFs (see the Supporting Information for details on the calculations). Thermogravimetric analysis (TGA) was performed using a Netzsch STA490C thermoanalyzer under a 20 mL min⁻¹ air flux with a heating rate of 10 °C min⁻¹. Transmission Fourier-transform infrared spectroscopy (FT-IR) spectra (KBr pellets) were recorded on a Perkin-Elmer Spectrum BX Series FT-IR spectrometer, in the 4000–400 cm⁻¹ range, with a 2 cm⁻¹ resolution. Inductively coupled plasma optical emission spectroscopy (ICP-OES) analysis was carried out using a Varian 700-ES series. A calibration curve was obtained with four standard zirconium solutions (0.5, 1, 2.5, and 5 mg L⁻¹, respectively). The analyses were performed on the supernatant after dissolving the solids in an HNO₃ (2%) aqueous solution. Scanning electron microscopy (SEM) images were acquired with an FEI Quanta 450 ESEM FEG, working at a 15.00 kV acceleration voltage. The samples were sputtered with graphite prior to the analysis.

**Synthesis of [Zr₂O₂(OH)₄(TFS)₄] (Zirconium Tetrafluoro succinate, ZrTFS), ZrCl₄ (1 mmol, 233 mg), H₂TFS (4 mmol, 773 mg), and 1 mL of deionized water were put in a ball-mill vessel. After 15 min of milling, the mixture was put in a 5 mL hydrothermal bomb and was heated to 120 °C for 72 h. The obtained white solid was recovered by centrifugation and washed with ethanol (2 × 5 mL), water (1 × 5 mL), and acetone (2 × 5 mL) and finally dried in an oven at 80 °C. Yield: 267 mg (88%, based on zirconium). IR (KBr, cm⁻¹; Figure S1): 1663 (s) [ν(COO)], 1145/1128 (m) [ν(C–F)]

**Synthesis of [Zr₄O₆(µ₃-OH)₄(µ₁-OH)₂C₄H₄(COO)₂]₄(AA)₄ (Zirconium Fumarate, MOF-801), ZrCl₄ (1 mmol, 233 mg), H₂FUM (3 mmol, 348 mg), and acetic acid (100 mmol, 5.7 mL) were placed together in a hydrothermal bomb with 20 mL of deionized water, and the solution was sonicated until complete reagent dissolution. Then, the reactor was held in a thermostatic oven at 120 °C for 24 h. The obtained white solid was recovered by centrifugation, washed with water (2 × 10 mL) and acetone (1 × 10 mL) and finally dried in an oven at 80 °C. Yield: 200 mg (87%, based on zirconium). The phase identity and purity were checked through PXRD. The amount of AA present in the material was estimated through ¹H NMR (see the Supporting Information). IR (KBr, cm⁻¹; Figure S1): 1576 (s) [ν(C=O)FUM]. The [ν(C=O)FUM] and [ν(C=O)F] bands are not visible, falling underneath of ν(C=C)FUM.

**Postsynthetic Exchange on MOF-801: Preparation of [Zr₄O₆(µ₃-OH)₄(µ₁-OH)₂C₄H₄(COO)₂]₄(AA)₄-HTFS₄]₁₄ (PF-MOF-801) and [Zr₄O₆(µ₃-OH)₄(µ₁-OH)₂C₄H₄(COO)₂]₄(AA)₄-HTFS₄]₁₄ (PF-MOF2). MOF-801 (200 mg, 0.14 mmol) and H₂TFS (0.28 mmol, 68 mg for the synthesis of PF-MOF1 and 0.84 mmol, 160 mg for the synthesis of PF-MOF2,
respectively) were placed together in a reaction flask with 2.5 mL of deionized water. The mixture was left under stirring at 60 °C for 4 h. The obtained solids were recovered by centrifugation, washed with ethanol (2 × 5 mL), water (1 × 5 mL), and acetone (1 × 5 mL), and finally dried in an oven at 80 °C. Yield: 224 mg and 237 mg (99.99 and 99.93%, based on zirconium present in the pristine MOF-801 and using the results from ICP measurements for quantifying the metal leaching after the exchange; Table S1) for PF-MOF1 and PF-MOF2, respectively. The proposed minimal formulae reported above come from quantitative 1H and 19F NMR spectroscopies after digestion and TGA carried out to quantify the number of defects and the amount of exchanged TFS⁻ ligands. PF-MOF1. IR (KBr, cm⁻¹; Figure S1): 1645 (vs) [ν(COO⁻)]TFS⁻, 1576 (vs) [ν(C≡C)]PFUM, 1135/1118 (s) [ν(C≡F)]PFUM/sym]. PF-MOF2. IR (KBr, cm⁻¹; Figure S1): 1653 (vs) [ν(COO⁻)]TFS⁻, 1576 (vs) [ν(C≡C)]PFUM, 1136/1118 (s) [ν(C≡F)]PFUM/sym]. As found in MOF-801, the [ν(COO⁻)]PFUM bands are not visible, falling underneath that of [ν(C≡C)]PFUM.

N₂ and CO₂ Adsorptions. All of the samples were activated at 120 °C under a high vacuum (10⁻³ Torr) for 12 h before any measurement. The Brunauer–Emmett–Teller (BET) specific surface area (SSA) and porosity were estimated by volumetric adsorption with an ASAP 2020 Micromeritics instrument, using N₂ as the adsorbate at −196 °C and an equilibration time of 30 s (an optimal value for surface area measurements when using N₂ as a probe with ultramicroporous materials). A typical measurement used 40 mg of sample. For the BET specific surface area calculation, the 0.01–0.1 p/p₀ pressure range was used for fitting the data. Within this range, all of the Rouquerol consistency criteria were satisfied. The material (micro)porosity was determined from the CO₂ adsorption isotherms collected at T = 0 °C, using a 2D nonlocal density functional theory (2D-NLDFT) method successfully employed for carbonaceous materials with heterogeneous surfaces. The same kind of pore size distribution (PSD) can also be obtained from N₂ isotherms at T = −196 °C. CO₂ adsorption isotherms were measured at 0 and 25 °C up to the maximum pressure of 1.2 bar. The isobaric heat of adsorption (Qₛ) was calculated working on the CO₂ adsorption measured at 0 and 25 °C, by applying a variant of the Clausius–Clapeyron equation (eq 1):35,54

\[
\ln \left( \frac{p_1}{p_2} \right) = Q_s \times \frac{T_2 - T_1}{R \times T_1 \times T_2}
\]

where pₙ (n = 1 or 2) denotes the pressure value for the nth isotherm, Tₙ (n = 1 or 2) denotes the temperature value for the nth isotherm, and R is the gas constant expressed in the appropriate units (8.314 J K⁻¹ mol⁻¹). In order to validate the calculated Qₛ values on two temperatures, an additional isotherm was also collected at T = −20 °C for PF-MOF1 and ZrTFS and the calculation was repeated on three temperature points using the differential form of the Clausius–Clapeyron equation (see the Supporting Information). The isotherms were also fitted with the Virial equation following the guidelines of a recent paper:55

\[
\ln p = \ln n + \frac{1}{T} \sum_{n=0}^{m} a_n p^n + \sum_{n=0}^{m} b_n p^n
\]

where p is the pressure at which loading n is reached, aₙ (m = 7) is −Qₛ/R, and bₙ (m' = 2) is a constant. Qₙ was then derived using the following equation

\[
Q_n = -R \sum_{n=0}^{m} a_n p^n
\]

To evaluate the CO₂/N₂ selectivity at 25 °C, the N₂ adsorption isotherms were acquired at 25 °C and up to 1.2 bar. The CO₂/N₂ Henry selectivity was calculated as the ratio of the initial slopes of the two adsorption isotherms in the 0.01 ≤ p ≤ 0.1 bar range. The CO₂/N₂ ideal adsorbed solution theory (IAST) selectivity of binary mixtures at a total pressure of 1 bar and at T = 25 °C was determined as the ratio of the adsorbed molar fractions of the two gases divided by the ratio of the gas-phase initial molar fractions:56

\[
S_{CO_2/N_2} = \frac{\frac{c_{CO_2}}{c_{N_2}}}{\frac{c_{CO_2}}{c_{N_2}}}_{gas}
\]

The (p)ₙ values were derived from the application of the free software pyiAST (https://github.com/CorySimon/pyiAST) to the experimental single-component isotherms collected at the chosen temperature. The initial composition (%) for the calculation was [15:85] for the [CO₂/N₂] pair. This relative ratio was selected to mimic the general feed composition of a coal-fired power station.8 A Henry or a BET model was employed for the isotherm fitting. For a detailed explanation of these models and the related parameters, see the pyiAST webpage and documentation.

Computational Details. All calculations were carried out with the CP2K code. Atom-centered Gaussian-type basis functions are used to describe the orbitals. The MOLOPT basis set for Zr, O, C, F, and H was employed, with a cut-off of 500 Ha for the plane waves along with the PBE functional. Core–valence interactions are described by Goedecker–Teter–Hutter pseudopotentials. In this simulation, the experimental structures derived from PXRD data were taken as the initial geometry guess and a joint atomic position and cell parameter optimization was carried out, keeping the three axes orthogonal and symmetric. The Brillouin zone was sampled at the Γ point. Mulliken atomic charges on a reference model (SiF₆²⁻) and on the H₄TFS organic linker at comparison were calculated by performing a Gaussian 09 geometry optimization using the
B3LYP exchange–correlation functional and a 6-31++g** basis set on all atoms.

## RESULTS AND DISCUSSION

### Synthesis and Structural Characterization of ZrTFS.

ZrTFS was prepared with a “Shake ‘n Bake” procedure that we have previously employed for the synthesis of several UiO-66 MOF analogues. In this procedure, a dense slurry of reagents in a small amount of a liquid is obtained instead of a clear solution upon a preliminary treatment using a ball mill. This probably induces a partial MOF crystallization before the heating stage, which provides a crystalline material at the end of the synthesis. If the milling stage is bypassed, a much less crystalline product is obtained (Figure S2) and the reaction yield decreases from 88 to 69%. No solid product was obtained when larger volumes of water were used, suggesting that a high concentration of reagents is necessary to induce the crystallization of ZrTFS. This is likely due to the high solubility of H$_2$TFS in water, a result of its highly acidic character ($pK_a$ values of 1.64 and 3.68, determined using Chemicalize, https://chemicalize.com/, developed by ChemAxon). Notably, to the best of our knowledge, ZrTFS is only the second material with an extended three-dimensional framework built with tetrafluorosuccinic acid reported in the Cambridge Structural Database, the other one being a lithium derivative.

The structure of ZrTFS was solved ab initio from PXRD data using the parallel tempering algorithm implemented in the FOX program and refined with the Rietveld method using TOPAS. Details of structure solution and refinement are reported in the Supporting Information (Figure S3). A polyhedral representation of the structure as obtained from the Rietveld refinement is reported in Figure 1. The structure was solved in the same cubic space group as MOF-801 (Pn3), and it is fully consistent with the optimized DFT structure (see below). The lattice parameter is 18.0690 Å, slightly larger than that reported for MOF-801 (17.8469 Å). The framework topology is fcu with 12-connected $[\text{Zr}_6\text{O}_{12}\text{(OH)}_4]^{12}$ hexanuclear clusters bridged by the TFS$^{2-}$ linkers. The geometry of one crystallographically independent linker recalls that of fumaric acid in MOF-801 with the O–C–C–O carbon–oxygen chain torsion angle close to 180° (i.e., O11–C11–C12–C12#1 176.9° #1 = –x, y + 1/2, z + 1/2), whereas the second TFS$^{2-}$ displays a bent geometry between the carboxylic plane and the C–F chain (O21–C21–C22–C22#1 146.5° #1 = –x, y + 1/2, z + 1/2). Fluorine atoms linked to internal carbon atoms (C–F distance 1.37 Å) are pointing toward the cavities, thus reducing the window size with respect to those of MOF-801 from 4.8 to 3.5 Å (average distances measured from atomic centers). These small window sizes strongly limit the gas diffusion into the cavities, as observed in its gas adsorption isotherms (vide infra). SEM analysis (Figure S4) reveals a peculiar truncated octahedral morphology, with the crystallite size in the range of 500 nm.

### Synthesis of Mixed-Linker MOFs via Postsynthetic Exchange.

The lack of accessible porosity in the framework of ZrTFS limits its ability to adsorb CO$_2$. This prompted us to prepare alternative materials where the (flexible) fully aliphatic TFS$^{2-}$ linker and its (more rigid) unsaturated analogue FUM$^{2-}$ could coexist in the same framework. Mixed-linker tetrafluorosuccinate/fumarate MOFs should feature both satisfactory accessible surface area and high fluorine content. To achieve this result, we carried out postsynthetic exchange (PSE) reactions starting from pure MOF-801. MOF-801 was synthesized from fumaric acid and Zr$^{IV}$ chloride according to the literature procedure of Zahn et al., using acetic acid as the crystallinity modulator. Its PXRD pattern is reported in Figure 2, confirming the phase purity and the high crystallinity degree. The use of AA as a modulator leads to the obtainment of a defective material where the bridging fumarate linkers are partially replaced by terminal acetates and water/hydroxide couples. The $^1$H NMR signal integration of the digested sample using 3-fluorobenzoic acid as the internal standard (Figure S5) allowed us to quantify the ratio of fumarate (FUM) and acetate (AA) in the structure (calculation details are provided in the Supporting Information). The general formula found is $[\text{Zr}_6\text{O}_{12}(\mu_3-\text{OH})_6(\mu_2-\text{OH})_{112}(\text{H}_2\text{O})_{312}(\text{FUM})_{210}(\text{AA})_{90}]$. From MOF-801, fumarate/tetrafluorosuccinate partial exchange performed in water with a (1:2) or (1:6) $[\text{Zr}_6]\text{H}_2\text{TFS}$ stoichiometric ratio gave the two mixed-linker MOFs labeled as PF-MOF1 and PF-MOF2. Their PXRD patterns are also shown in Figure 2, whereas their thermogravimetric (TG) profiles are shown in Figure 3. PSE did not seem to affect the structural integrity of the pristine fumarate; the diffractograms are nearly identical, and they clearly show that both PF-MOFs are isostructural to MOF-801. The only difference is in the intensity of a small peak at 2θ of ~13° (already present in MOF-801) that slightly increases with the increasing amount of $\text{H}_2\text{TFS}$ used for the exchange. To gain additional information on the structural features of the presented PF-MOFs, an ab initio indexing using the N-TREOR suite was carried out. In all cases, cubic cells were found: $a = 17.8469$ Å, $V = 5684$ Å$^3$ (MOF-801); $a = 17.8498$ Å, $V = 5687$ Å$^3$ (PF-MOF1); $a = 17.8621$ Å, $V = 5699$ Å$^3$ (PF-MOF2); $a = 18.0690$ Å, $V = 5899$ Å$^3$ (ZrTFS). The results indicate that the unit cell size remains almost unchanged when passing from MOF-801 to PF-MOF1 to PF-MOF2, thus suggesting that very small differences exist between these three materials, whereas ZrTFS displays a larger unit cell. Integration of $^{19}$F and $^1$H NMR peaks of the digested samples allowed us to quantify the relative amounts of FUM$^{2-}$, AA, and TFS$^{2-}$ present in the lattice. In the framework structure of MOF-801, the externally added

![Figure 2. PXRD patterns of MOF-801 (black), PF-MOF1 (red), PF-MOF2 (green), and ZrTFS (blue) in comparison.](https://doi.org/10.1021/acsami.2c07640)
TFS$^-$ linkers may be incorporated through the replacement of either the pristine fumarates or the capping ligands (either acetates or water/hydroxide) placed in its defective sites, giving the hypothesis that AA induced missing cluster defects. In the exchanged samples, after treatment with H$_2$TFS, the signal of AA disappears. This suggests that the tetrafluorosuccinate replaces AA at defective sites in its monoprotonated form (HTFS$^-$). Based on the combination of NMR analysis (Figures S6 and S7) and TGA (Figure 3), the proposed formulae for PF-MOF1 and PF-MOF2 are $[\text{Zr}_2\text{O}_4(\mu_3-\text{OH})_2(\mu_1-\text{OH})_2\text{H}_2\text{O}]_2\text{FUM}_2\text{HTFS}_2$ and $[\text{Zr}_2\text{O}_4(\mu_3-\text{OH})_2(\mu_1-\text{OH})\text{H}_2\text{O}]_2\text{FUM}_2\text{HTFS}_2$, respectively. Based on these formulae, we can conclude that no TFS$^-$/FUM$^-$ exchange has occurred and that HTFS$^-$ has been grafted at defective sites, affording compounds with similar stoichiometry. Apparently, the fluorination degree is not appreciably increased by the use of more concentrated H$_2$TFS solutions (1:2–1:6 MOF-801:H$_2$TFS stoichiometric ratios) to foster a larger fluorinated linker uptake. TGA analysis of the four MOFs (Figure 3) shows a different thermal behavior depending on their fluorination extent. MOF-801 contains almost 30 wt % of water, and it starts to decompose at temperatures higher than 380 ºC. ZrTFS contains a smaller amount of solvent than MOF-801 (about 7%), indicative of its lower porosity. In addition, the thermal stability is much lower ($T_{\text{dec}} \approx 260$ ºC), as expected for the lower Zr–O coordination bond strength of the (less basic) TFS$^-$ compared to that of FUM$^-$. The two PF-MOFs have a thermal behavior, which is intermediate between ZrTFS and MOF-801. The only difference is the water content, higher for PF-MOF2 (23%, with a higher tetrfluorosuccinate loading) with respect to PF-MOF1 (15%); the thermal stabilities though are comparable ($T_{\text{dec}} \approx 280$ ºC).

Gas Adsorption Properties. $N_2$ Adsorption. The porous nature of the four samples was investigated through $N_2$ adsorption at −196 ºC on desolvated samples. As shown in Figure 4, all compounds display type I isotherm, typical of microporous materials. MOF-801 has the highest BET specific surface area (SSA) value (948 m$^2$ g$^{-1}$), and this result is in line with that obtained by other groups in the literature working with the same compound (833 m$^2$ g$^{-1}$ found by Serre and co-workers; $^{99}$ 990 m$^2$ g$^{-1}$ reported by Taghi and co-workers). $^{49}$ The accessible surface area then decreases with an increasing content of the aliphatic fluorinated linker TFS$^-$, passing from 649 and 626 m$^2$ g$^{-1}$ for PF-MOF1 and PF-MOF2, respectively, to only 46 m$^2$ g$^{-1}$ for ZrTFS. This drastic reduction of the surface area was also found in the hydrogenated counterpart of ZrTFS, zirconium succinate MIP-203-S.$^{58}$ In there, it is claimed that this MOF does not show any accessible porosity to $N_2$ at 77 K, presumably because after thermal activation under vacuum to remove the guest molecules, it tends to stay in a closely packed form that is not accessible to nitrogen. Even in the presence of bipyridyl-based auxiliary ligands, the framework empty volume is not accessible to $N_2$, as observed in a family of cadmium succinate mixed-linker MOFs.$^{59}$ Despite the different crystal topologies, it is reasonable to draw the same kind of conclusion for ZrTFS as well. This is further confirmed by the fact that the surface area of ZrTFS does not change when the equilibration time during the isotherm collection is doubled from 30 to 60 s. To gain additional insights into the textural properties of ZrTFS, the theoretical MOF surface area was estimated using a Monte Carlo procedure, which randomly places spheres with a given diameter in the free space ($N_2$ to match experiments—sphere diameter = 3.681 Å)$^{60}$ and calculates the interface between the spheres and the atoms (for further details and explanations, see the webpage: https://mausdin.github.io/MOFsite/mofPage.html). This is a commonly used procedure, and it normally gives reasonable results.$^{61}$ By applying this procedure on ZrTFS, we have obtained a surface area of 226.8 m$^2$ g$^{-1}$. This value is reasonable if the pore size is taken into consideration, but it is much higher than the experimentally measured surface area. This discrepancy may be justified by a window size limitation for $N_2$ free diffusion through the MOF pores. The micropore size distribution of the four samples retrieved from the NLDF analysis (Figure S8) does not show a significant change when passing from the homolinker to the mixed-linker materials. The main contribution to the total pore volume comes from pores in the 10 Å range, in line with the Zr–Zr distance found in the crystal structure between adjacent $[\text{Zr}_6]$ nodes (∼13 Å).$^{59}$ The percentage of the ultramicropore volume decreases with an increasing extent of fluorinated linker insertion, passing from 77% in MOF-801 to 72 and 62% in PF-MOF1 and PF-MOF2, respectively. Degradation coming from PSE cannot be considered the main cause of
Adsorbents at volume derived from the N are preserved upon PSE (Figures S9 supported by the SEM analysis, which shows that the very small degree of degradation (Table S2). This is also of zirconium found in solution after the exchange indicates a the observed microporosity reduction since the small amount performance in CO₂ adsorption depends on two factors: BET area and fluorinated linker content. The former is predominant in MOF-801, where the absence of F atoms is compensated by the high SSA value. Consequently, MOF-801 shows the best performance among the MOFs considered in this study in terms of CO₂ loading on a gravimetric basis: 2.42 (10.6 wt %) and 3.51 (15.4 wt %) mmol g⁻¹ at T = 25 and 0 °C, respectively. On the other hand, the introduction of TFS partially reduces the accessible surface area (and the related CO₂ loading) but considerably improves the thermodynamic affinity of the material for CO₂ and its CO₂/N₂ selectivity. The latter is particularly important in the purification of postcombustion industrial flue gases, where the amount of CO₂ is very low (4–30%). The high Henry and IAST selectivity achieved by PF-MOF2 and ZrTFS, in particular, may be ascribed to the presence of fluorine-decorated ultramicropores that hamper N₂ diffusion (because of its large kinetic diameter) but favor CO₂ adsorption through the beneficial gas–fluorine interaction. On this basis, the best compromise can be found in the PF-MOF2, with a good CO₂ loading of 2.1 mmol g⁻¹ (9.3 wt %) at an ambient temperature and 1 bar, a Qₒ value of almost 30 kJ mol⁻¹, and the good CO₂/N₂ selectivity of 95 (Henry)/41 (IAST). The same conclusion can be drawn from the isotherms reported in Figure 5a, where PF-MOF2 shows the highest CO₂ uptake: 3.7 mmol CO₂/mmol ZrMOF⁻¹ at p = 1 bar. In comparison with other PF-MOFs reported in the literature (Table S7), at ambient (T,p) conditions (25 °C, 1 bar), PF-MOF2 outperforms F4-Uio-66(Ce) (1.5 mmol g⁻¹ at 293 K), but it is less efficient than F4-MIL-140A(Ce) (2.4 mmol g⁻¹) and AlIFIVE-1-Ni/ NbOFFIVE-1-Ni (2.7/2.2 mmol g⁻¹, respectively) published by the team of Eddaoudi in 2016.23,24,62

DFT Study of the CO₂–Framework Interaction. To gain insight into the effect of the fluorinated ligand on the
MOF structural properties and \( \text{CO}_2 \) adsorption ability, DFT simulations in the gas phase at the \([\text{PBE/MOLOPT}]\) level of theory were carried out. As a first step in the computational analysis, the structures of \text{MOF-801} and \text{ZrTFS} were optimized. For \text{MOF-801}, the starting geometry was the defect-free crystal lattice with formula \([\text{Zr}_6\text{O}_{4}(\text{OH})_4\text{(FUM)}_6]\), while for \text{ZrTFS} the initial guess \([\text{Zr}_6\text{O}_{4}(\text{OH})_4\text{(TFS)}_6]\) was considered. To evaluate the effect of the presence of both linkers in the same solid phase on the \( \text{CO}_2 \) adsorption performance from a theoretical viewpoint, a hypothetical (simplified) structure of minimal formula \([\text{Zr}_6\text{O}_{4}(\text{OH})_4\text{(FUM)}_4\text{(TFS)}_2]\) (PF-MOF) was also built in silico and subsequently optimized. Albeit this computational model cannot be considered fully representative of the real exchanged materials PF-MOF1 and PF-MOF2, it is a useful reference for a comparison between a mixed-linker MOF and its homolinker counterparts. Indeed, the design of computationally representative defective structures is too challenging, as the linker is placed onto the defects randomly. On these models, a joint variable-cell and atomic position optimization was performed (see the Computational Details section). As shown in Table 2, the optimized lattice parameters are in perfect agreement with the experimentally determined values, with the reproduction of the slight unit cell size increment moving from pure fumarate to the fluorinated systems. These MOFs feature tetrahedral and octahedral pores interconnected by triangular windows, with a lattice structure similar to that of \text{UiO-66}. The windows are smaller than the pores (Figure 6 and Table 2), and they could be responsible for the percolation of the gas inside the MOF. The gas adsorption process is driven by two main factors: the host–guest chemical affinity and the (high) accessible SSA. Another interesting factor that

| Structural parameters | MOF-801 | PF-MOF | ZrTFS |
|-----------------------|---------|--------|--------|
| cell (Å)              | 18.00   | 18.04  | 18.17  |
| window area (Å²)      | 9.3     | 9.1    | 5.0    |
| pore (Å)              | 9.6/8.6 | 10.0/6.3−9.0 | 10.1/6.6 |
| \( \Delta E_{\text{ads}} \) (eV) | -0.19  | -0.18  | -0.18  |
| pore Oct. center (1)  | -0.29   | -0.27  | -0.31  |
| pore Zr (2)           | -0.34   | -0.47  | -0.49  |
| pore Tetr. center (3) | -0.29   | -0.29  | -0.29  |
| window (4)            | -0.29   | -0.29  | -0.32  |

*a* We report both octahedral and tetrahedral pore sizes, respectively.

Table 2. Main Calculated Structural Parameters and \( \text{CO}_2 \) Adsorption Enthalpies for MOF-801, PF-MOF, and ZrTFS

Figure 6. Top: optimized DFT structure of \text{ZrTFS}. The blue and the orange spheres represent the octahedral and tetragonal pores, respectively; the yellow curved line represents the window between the two pores; the numbers indicate the different sites taken into account for the evaluation of \( \text{CO}_2 \) adsorption enthalpies (\( \Delta E_{\text{ads}} \)) in reference to Table 2. Bottom: detail of the hydrogen bonds between \( \text{CO}_2 \) and the OH group for (a) \text{MOF-801}, (b) PF-MOF, and (c) \text{ZrTFS}.
deeply influences the adsorption performance is the gas percolation and diffusion inside the material. Literature evidence suggests that MOFs featuring high CO₂ chemical affinity but with very small pore windows are poor adsorbents. To compare the window and pores of the different models, the surface of the window is defined as the area of the triangle defined by the three hydrogen (or fluorine) atoms, as indicated in Figure S21a, while the pore dimension is evaluated as the distance between opposite hydrogen or fluorine atoms in Figure S21b. The average dimensions of the cavities and window size are summarized in Table 2. Moving from MOF-801 to ZrTFS, the octahedral pore size does not change significantly, while the window area and the tetrahedral pore size are considerably reduced. Moreover, the mixed-linker PF-MOF shows a variable pore width (between 6 and 9 Å; Table 2) that depends on the presence of either TFS⁻⁻ or FUM⁻⁻. To gain deeper insights into the preferential CO₂ adsorption sites in these materials, CO₂ was introduced in the computational model and located in four different lattice positions (Figure 6): in the center of the octahedral cavity (1), close to the zirconium ions (2), in the center of the tetrahedral pore (3), and in the middle of the pore windows (4). Afterward, the ensemble was reoptimized. The CO₂ adsorption energy (ΔE_ads) is then calculated as the energy difference between the optimized [MOF + CO₂] ensemble and the separated components; a negative value indicates a favorable interaction, while the opposite holds for positive ΔE_ads values. The most favorable adsorption site is the tetrahedral pore, even if it is slightly smaller than the octahedral one. This energy stabilization comes from a hydrogen bond interaction between CO₂ and a hydroxyl group on the inorganic cluster (Figure 6), which is known to be the most favorable adsorption site for polar and quadrupolar species in UiO-66. Apparently, there is no simple correlation between F-functionalization and MOF–CO₂ interaction. We can only state that the interaction of CO₂ with the linker F atoms is not particularly strong, at least at the computational level used here. To better understand the role of fluorine atoms in increasing the affinity with CO₂, the Mulliken partial charges on F species were evaluated as already reported in the literature for the SIFSIX family MOFs. In particular, the SiF₂⁻⁻ anion was taken as the benchmark model, obtaining a partial F charge of −0.76e, in agreement with the values found for similar systems. Subsequently, the partial charge on the fluorine atoms of the H₃TFS linker was calculated at the same level of theory, obtaining an average value of −0.25e. This clearly indicates a reduced C–F bond polarization in H₃TFS if compared to the Si–F bond polarization in SiF₂⁻⁻. Consequently, these results suggest a lower CO₂ affinity of ZrTFS with respect to a general SIFSIX system. If the (ΔE_ads) values found for the different materials are compared, an increase in the interaction energy is found when moving from MOF-801 to ZrTFS, in perfect agreement with the Qₐ trend reported in Table 1. It is to be noted that the window site in ZrTFS is associated with a positive (ΔE_ads) value, which suggests that CO₂ diffusion through the windows is unfavorable, explaining the low adsorption capacity of this MOF.

CONCLUSIONS

The novel perfluorinated MOF ZrTFS containing tetrafluorosuccinic acid and isostructural to MOF-801 has been synthesized and its crystal structure was solved and refined from PXRD data. The presence of fluorine atoms on the alkyl chains hinders gas adsorption; consequently, the measured surface area of this MOF is very low. To obtain a fluorinated material with an acceptable surface area, MOF-801 was taken as the starting point to carry out PSE in aqueous solutions with variable amounts of H₃TFS. In the two exchanged materials, the amount of a fluorinated linker is about the same despite the different concentrations used in the synthesis, and it is most likely placed in the precursor defective sites. This suggests that the postsynthetic exchange involves only acetate and not fumarate. The as-obtained PF-MOFs show lower CO₂ adsorption capacity with respect to MOF-801 (most likely because of the lower surface area) but higher CO₂/N₂ selectivity and CO₂ heat of adsorption. These parameters are directly proportional to the extent of a fluorinated linker present in the lattice, proving the beneficial effect of the presence of fluorinated groups on the CO₂ affinity of the resulting material. DFT-optimized structures of ZrTFS account for the low diffusion of gases into the framework due to the reduction of the window size. The theoretical model of [CO₂@PF-MOF] ensembles shows the formation of strong hydrogen-bonding interactions between CO₂ and hydroxyl groups of the [Zr₆] clusters dangling in the tetrahedral cavities and favored by the exposed fluorine atoms. Although a real linker exchange did not occur in the present work, being a defect functionalization, the PSE methodology is very efficient for the preparation of mixed-linker MOFs with tunable properties and opens new horizons for selected applications such as gas mixture separation and purification. Current work is undergoing in our laboratories in this direction.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.2c07640.

IR spectra, PXRD patterns of ZrTFS obtained through different synthetic methods, PXRD structure determination of ZrTFS, its Rietveld refinement plot and crystallographic details, ¹H NMR and ¹³C NMR spectra of the digested samples of MOF-801, PF-MOF1, and PF-MOF2 and related linker quantification, ICP-OES analysis of the reaction supernatant after PSE on MOF-801, MOF pore size distribution, calculation of the CO₂/N₂ Henry selectivity, CO₂ adsorption isotherm plot in [mmol g⁻¹] units at 25 °C, further Qₐt calculations using three temperatures and geometrical considerations on the DFT-optimized structures (PDF)

Accession Codes

CCDC-2160377 contains the crystallographic data for ZrTFS that can be obtained free of charge from the Cambridge Crystallographic Data Centre (http://www.ccdc.cam.ac.uk/data_request/cif).

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Notes

The authors declare no competing financial interest.

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