Journal of Materials Chemistry A



PAPER

View Article Online
View Journal | View Issue



Cite this: *J. Mater. Chem. A*, 2023, **11**, 5568

Cooperative CO₂ adsorption mechanism in a perfluorinated Ce^{IV}-based metal organic framework†

Margherita Cavallo, ^a Cesare Atzori, ^{bab} Matteo Signorile, ^{ba}
Ferdinando Costantino, ^{bc} Diletta Morelli Venturi, ^c Athanasios Koutsianos, ^{‡d}
Kirill A. Lomachenko, ^{bb} Lucia Calucci, ^{bef} Francesca Martini, ^{bfg}
Andrea Giovanelli, ^{bg} Marco Geppi, ^{bfg} Valentina Crocellà ^{b*a}
and Marco Taddei ^{b*fgh}

Adsorbents able to uptake large amounts of gases within a narrow range of pressure, *i.e.*, phase-change adsorbents, are emerging as highly interesting systems to achieve excellent gas separation performances with little energy input for regeneration. A recently discovered phase-change metal-organic framework (MOF) adsorbent is F4_MIL-140A(Ce), based on Ce^{IV} and tetrafluoroterephthalate. This MOF displays a non-hysteretic step-shaped CO_2 adsorption isotherm, reaching saturation in conditions of temperature and pressure compatible with real life application in post-combustion carbon capture, biogas upgrading and acetylene purification. Such peculiar behaviour is responsible for the exceptional CO_2/N_2 selectivity and reverse CO_2/C_2H_2 selectivity of F4_MIL-140A(Ce). Here, we combine data obtained from a wide pool of characterisation techniques – namely gas sorption analysis, *in situ* infrared spectroscopy, *in situ* powder X-ray diffraction, *in situ* X-ray absorption spectroscopy, multinuclear solid state nuclear magnetic resonance spectroscopy and adsorption microcalorimetry – with periodic density functional theory simulations to provide evidence for the existence of a unique cooperative CO_2 adsorption mechanism in F4_MIL-140A(Ce). Such mechanism involves the concerted rotation of perfluorinated aromatic rings when a threshold partial pressure of CO_2 is reached, opening the gate towards an adsorption site where CO_2 interacts with both open metal sites and the fluorine atoms of the linker.

Received 14th December 2022 Accepted 9th February 2023

DOI: 10.1039/d2ta09746j

rsc.li/materials-a

Introduction

Solid adsorbents for gas separations are the object of intensive research, due to their potential advantages, in terms of energy intensity and recyclability, over state-of-the-art technologies based on either cryogenic distillation or absorption/scrubbing.¹⁻⁵ Metal-organic frameworks (MOFs) are among the candidates investigated to this end, alongside zeolites, activated carbons and amine-functionalised porous silicas.^{3,4,6,7} If

compared to the other classes of adsorbents, MOFs offer superior structural control at the atomic level and virtually unlimited possibilities for functionalisation and tuning of the physicochemical properties. One of the most intriguing aspects of MOF adsorbents is that they can display unique adsorption phenomena, associated with either highly specific interaction of the adsorbate with the surface, structural flexibility, or combinations thereof. 8-16

[&]quot;Dipartimento di Chimica, Centro di Riferimento NIS e INSTM, Università di Torino, Via G. Quarello 15, I-10135 and Via P. Giuria 7, I-10125 Torino, Italy. E-mail: valentina.crocella@unito.it

^bEuropean Synchrotron Radiation Facility, 71 Avenue des Martyrs, CS 40220, 38043 Grenoble Cedex 9, France

Dipartimento di Chimica, Biologia e Biotecnologie, Unità di Ricerca INSTM, Università di Perugia, Via Elce di Sotto 8, 06123 Perugia, Italy

^aCentre for Research & Technology Hellas/Chemical Process and Energy Resources Institute, 6th km. Charilaou-Thermis, 57001, Greece

[&]quot;Istituto di Chimica dei Composti Organo Metallici, Unità di Ricerca INSTM, Consiglio Nazionale delle Ricerche, Via Giuseppe Moruzzi 1, 56124 Pisa, Italy ^fCentro per l'Integrazione della Strumentazione Scientifica dell'Università di Pisa (CISUP), 56126, Pisa, Italy

^{*}Dipartimento di Chimica e Chimica Industriale, Unità di Ricerca INSTM, Università di Pisa, Via Giuseppe Moruzzi 13, 56124 Pisa, Italy. E-mail: marco.taddei@unipi.it hEnergy Safety Research Institute, Swansea University, Fabian Way, Swansea, SA1 8EN, UK

[†] Electronic supplementary information (ESI) available. CCDC 2212834 and 2212835. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d2ta09746j

[‡] Present address: Anorganische Materialchemie, Fakultät für Chemie & Chemische Biologie, Technische Universität Dortmund, Otto-Hahn-Straße 6, 44227 Dortmund, Germany.

So-called "phase-change" MOFs are of particular interest for gas separations because they display steep gas uptake when a threshold partial pressure of a given adsorbate is reached, with potential benefits in terms of achievable working capacity, selectivity and energy efficiency. 17,18 This behaviour is determined by an adsorption-induced structural rearrangement, which can be either (i) a breathing/swelling effect, due to the flexible nature of the framework, which leads to significant variations in the volume of the unit cell;9,19-22 (ii) a cooperative adsorption mechanism, such as the CO2 insertion occurring in so-called amine-appended MOFs, which involves changes in the coordination environment of the metal and highly specific hydrogen bonding interactions; 18,23,24 or (iii) the concerted rotation of organic rings within the framework, which results in ring configuration that maximises the host-guest interaction.25-27

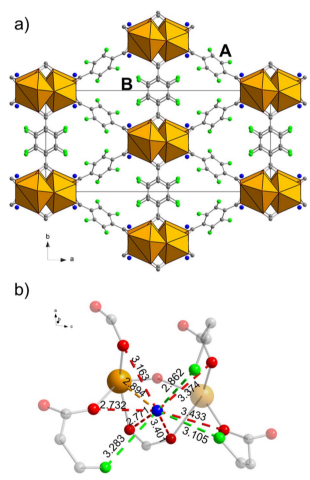


Fig. 1 Crystal structure of as-synthesised F4_MIL-140A(Ce) viewed along the crystallographic c axis (a) and local environment around the water molecule (b). In (a) the two crystallographically independent linkers sitting on an inversion centre and on a two-fold axis are identified with A and B, respectively. In (b) a threshold distance of 3.5 Å between the O atom of water and surrounding atoms was chosen to identify possible interactions, represented as dashed lines having the same colour of the interacting atom. Non-interacting atoms in (b) are shaded. Colour code: Ce, orange; F, green; C, grey; O, red; H₂O, blue. H atoms are not shown because the structure was determined from PXRD data.

Some of us recently reported the discovery of F4_MIL-140A(Ce), an ultramicroporous MOF based on Ce^{IV} and tetrafluoroterephthalate (Fig. 1).28 F4_MIL-140A(Ce) is built from the connection of one-dimensional inorganic building units, made up of octacoordinated Ce^{IV} ions, carboxylate groups belonging to the linker and μ_3 -O species, *via* the perfluorinated aromatic rings of the linker. This structure features narrow triangular channel-like pores lined with the fluorine atoms belonging to the linkers (Fig. 1a). Different from its ZrIV-based MIL-140 analogues,29 F4_MIL-140A(Ce) contains one water molecule per Ce atom, located in the proximity of the metal atom and at distances compatible with a number of non-covalent interactions with the organic linker (Fig. 1b).

F4_MIL-140A(Ce) displays a non-hysteretic step-shaped CO₂ adsorption isotherm, with steep uptake increase at pressure <0.2 bar at 298 K, leading to reach saturation within a narrow range of pressure.28 Such behaviour is not observed when the MOF is exposed to other gases, e.g., N2 or C2H2, endowing this material with high CO₂/N₂ selectivity and with reverse CO₂/C₂H₂ selectivity, a highly sought-after feature.30 Furthermore, this MOF can be synthesised from commercially available reagents in mild conditions and in aqueous medium, an attractive method for prospective bulk production.31 Considering such a unique combination of features, we believe that understanding the adsorption mechanism in F4_MIL-140A(Ce) becomes an issue of primary concern. Besides the fundamental interest in describing a potentially novel phenomenon, the knowledge deriving from such an endeavour might help to engineer new materials with the same working principle and improved separation performance.

A mechanism was in fact recently proposed based on powder X-ray diffraction (PXRD) data, infrared (IR) spectroscopy data, and density functional theory (DFT) simulations, claiming that CO2 adsorbs in a site located among three fluorinated rings, where the Lewis acidic C atom interacts with four electron-rich F atoms,30 akin to what observed in SIFSIX-3 materials32,33 and in fluorinated bis(pyrazolyl)-based MOFs.34 In the present work, we propose an alternative mechanism, deduced from the combination of data obtained from a wide pool of characterisation techniques, including gas sorption analysis, in situ IR spectroscopy, in situ PXRD, in situ X-ray absorption spectroscopy (XAS) at the Ce K-edge, ¹H, ¹³C and ¹⁹F solid state nuclear magnetic resonance (SSNMR) spectroscopy and adsorption microcalorimetry, with periodic DFT simulations.

Results and discussion

Adsorption properties

Gas adsorption volumetry. CO2 adsorption/desorption volumetric isotherms were collected at 298, 313, 328 and 343 K in the 0-5 bar pressure range to study the pure CO_2 adsorption capacity of the material (Fig. 2a). All the isotherms exhibit a peculiar step, which moves to progressively higher pressure by increasing the temperature and the pressure range over which saturation occurs becomes broader. The maximum amount of adsorbed CO₂ on F4_MIL-140A(Ce) is around 2.6 mmol g⁻¹ at 298 K and 5 bar and shows the usual decrease by increasing the

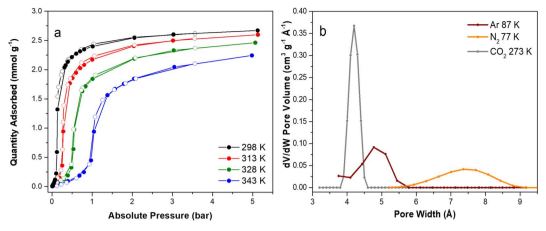


Fig. 2 CO₂ adsorption (filled circles) and desorption (empty circles) isotherms collected on F4_MIL-140A(Ce) at 298 K (black), 313 K (red), 328 K (green) and 343 K (blue) up to 5 bar (a) and pore size distributions of F4_MIL-140A(Ce) computed by NL-DFT of: Ar at 87 K (dark red), N2 at 77 K (orange) and CO₂ at 273 K (grey) (b).

temperature (2.2 mmol g⁻¹ at 343 K and 5 bar). The isotherms were fitted with a dual site Langmuir-Freundlich model (Fig. S1-S4†) and the pressure at which they display the maximum slope (P_{max}) was determined from the first derivative curve. Plotting $\log P_{\text{max}}$ versus temperature, a linear trend emerges between 298 and 343 K, in principle allowing to predict which pressure the step will take place at within the investigated temperature range (Fig. S5†). The isosteric heat of adsorption (Qst), extracted from the fitted isotherms using the linear version of the Clausius-Clapeyron equation (Fig. S6†), ranges between 35 and 45 kJ mol⁻¹.

The sharp uptake of gas (i.e. the isotherm step) moves to higher pressures by increasing the temperature. This peculiar adsorption feature allows easily adapting the phase transition mechanism of this MOF to the partial pressure of common postcombustion streams. Additionally, this characteristic isotherm is known to be particularly suitable for adsorption processes thanks to the large working capacity that can be achieved upon small swings in temperature and/or pressure. Such step-shaped isotherms have previously been observed in other MOFs exhibiting higher CO2 adsorption capacities at saturation, as in the amine-appended $Mg_2(dobpdc)^{18}$ (3.4 mmol g^{-1} at 298 K and 5 bar) and in ELM-11 (ref. 9) (3.6 mmol g^{-1} at 273 K and 0.3 bar). However, in the former case, the material displays a much higher isosteric heat of CO_2 adsorption (around -70 kJ mol^{-1}), whereas in the latter the phase transition occurs in a higher pressure range (300 mbar at 273 K). This leads, on the one hand, to stronger interaction energies and, consequently, to higher regeneration energies, on the other hand, to a low adsorption of CO₂ at low pressures. To further investigate the peculiar adsorption mechanism of F4_MIL-140A(Ce), the textural properties were accurately studied by collecting low pressure adsorption/desorption volumetric isotherms using three different adsorptives: N2 at 77 K, Ar at 87 K and CO2 at 273 K (Fig. S7† in the ESI†). The isotherms were measured consecutively using the same MOF powder. Therefore, the differences observed with the selected probe molecules can be attributed to the nature of the probe molecule at the analysis temperature. Ar

and N2 show the typical type I(a) isotherm characteristic of microporous materials, while CO2 exhibits the less common step-shaped isotherm specific of the so-called "phase change" adsorbents (Fig. S7a† in the $0 < p/p^0 < 0.03$ range and Fig. S7b† in the whole p/p^0 range), in agreement with the previous results reported by D'Amato et al.28 The semi-logarithmic isotherms reported in the inset of Fig. S7a† show that the pore filling with Ar occurs at higher relative pressure regions compared to N₂. The difference in relative pressures becomes more significant if the MOF possesses open metal sites, highlighting the advantage of using monoatomic inert Ar for the characterization of ultramicroporous MOFs. The BET area values (Table S2 and Fig. S8-S10†) display an incremental trend, suggesting a different interaction of the molecules with the material (Table S2†). The BET area obtained by Ar isotherm (213.8 m^2 g^{-1}) is lower compared to N₂ (244.8 m² g⁻¹). The spherical Ar molecules display a more ideal packing on the surface compared to diatomic N₂ molecules, whose orientation on the surface can be affected by the possible presence of open metal sites able to interact with N2 itself. As a consequence, an uncertainty in the evaluation of the cross-sectional area of nitrogen gas is very likely, resulting in an overestimation of the computed BET area with this probe. 35,36 The highest BET area value is the one derived from the CO_2 isotherm at 273 K (262 m² g⁻¹), suggesting that a structural rearrangement might occur when F4_MIL-140A(Ce) interacts with CO₂, as highlighted by the peculiar step-shaped isotherm collected by using this molecule as adsorptive (Fig. S7†).

The pore size distribution (PSD) was also evaluated by applying the non-local density functional theory (NL-DFT) to the adsorption isotherms. It is well-known that the PSD computed by NL-DFT methods depends on the selected kernel of isotherms. However, despite the high number of MOF structures reported to date, proper DFT kernels to study the unique pore size and shapes of these microporous materials are still missing.36 For this reason, the PSD data for MOFs should be considered as merely indicative. Moreover, the goodness of the PSD obtained by DFT depends on the goodness of the fitting of

the experimental isotherm (see Fig. S11, S12 and S13†). In all the investigated cases, the fit between the calculated and the experimental isotherms was imperfect (the best fit is the one obtained with the N2 isotherm). Again, this suggests that PSD data should be merely considered as a possible (and not unquestionable) description of the intrinsic porosity of the investigated MOF. The PSDs computed from the three molecular probes are compared in Fig. 2b and Table S1.† N2, a nonideal molecule for the analysis of microporous materials, gives the highest value (\sim 7.4 Å), while Ar and CO₂ PSDs show the presence of a family of ultramicropores of around 4.8 and 4.2 Å diameter, respectively, values in better agreement with that estimated on the basis of the crystal structure of assynthesised F4_MIL-140A(Ce) (between 4.8 and 5.8 Å, measured from the atomic centres).28 The result obtained with N₂ does not surprise, indeed it is known that this probe can significantly overestimate the PSD of MOFs with very small micropores, especially if the pore dimension is close to the kinetic diameter of the molecule (3.64 Å for N₂).36 The micropore size obtained by both Ar and CO₂ is more reliable and the PSD derived from CO₂ at 273 K is very narrow. This could be a further proof of a possible structural rearrangement of the material in the presence of CO2, which allows the detection of the actual micropores size just with an adsorptive inducing a "phase change" adsorption. Still, it is worth noting that N2 and Ar show a compatible micropore volume (Fig. S14†) while CO₂ displays a higher value, which might be seen as a further proof of a structural transition caused by this molecule.

In situ IR spectroscopy. As mentioned in the Introduction, a notable feature of as-synthesised F4_MIL-140A(Ce) is that it contains one H2O molecule per formula unit, located in the proximity of a Ce atom.28 Hydrogen bonding-like interactions are also likely to exist with F atoms and carboxylate groups from the surrounding organic linkers (Fig. 1b). According to the thermogravimetric curve (Fig. S15†), removal of H₂O is completed at a temperature as high as 423 K, suggesting that it is strongly adsorbed. The evacuation process of F4_MIL-140A(Ce) was further investigated using in situ IR spectroscopy (Fig. 3). Activation at 393 K for 12 h under vacuum was effective to remove the adsorbed H₂O before the spectroscopic experiments (see black spectrum in Fig. 3). The evacuated F4_MIL-140A(Ce) was then exposed to increasing pressures of H₂O, observing significant changes in the IR spectrum, especially in the 3700–3500 cm⁻¹ and 800–770 cm⁻¹ spectral ranges (Fig. 3b and c). In the former spectral region, two well-defined bands at 3643 and 3556 cm⁻¹ appear upon sending increasing H₂O doses. The presence of those peculiar narrow bands in the O-H stretching region is ascribable to the interaction of H₂O with specific acid sites.³⁷ In the 800-770 cm⁻¹ region, two initial bands at 790 and 779 cm⁻¹ are observed in the evacuated sample (see black spectrum in Fig. 3c). After sending 0.3 mbar of H₂O (corresponding to about 0.3-0.4% relative humidity, assuming a local temperature of around 313-318 K under the IR beam; vapour pressure of H₂O at 313 K is 74 mbar, at 318 K is 96 mbar) the 779 cm⁻¹ band shifts to 776 cm⁻¹, before going back to the original position upon further increasing the H2O pressure. A new band at 771 cm⁻¹ also appears at high H₂O

pressure. The 790 cm⁻¹ signal decreases until 0.3 mbar of H₂O, then it slowly shifts to 793 cm⁻¹, changing again its intensity. The spectral modifications observed upon H₂O contact in this region are probably due to changes occurring on the fluorinated rings. A more specific assignment of the bands present in those two spectral regions is provided by DFT calculations: in the high frequency region, the signals at 3643 and 3556 cm⁻¹ can be disambiguated as the O-H stretching mode of water forming distinct H-bonds with F atoms on aromatic rings and O atoms from carboxylates, respectively. The assignment of bands in the low frequency region is less trivial, due to the coupling of different vibrations generating complex vibrational modes. In a simplified view, the observed signals can be attributed to the out-of-plane bending modes of carboxylate groups with respect to the aromatic rings. The coexistence of multiple features is explained by the presence of two crystallographically independent linkers (Fig. 1), as well as by the perturbation induced by adsorbates on the linkers themselves. It is worth noting that the main changes in the 800-770 cm⁻¹ spectral region were observed with a H₂O dose of 0.3 mbar (green curve in Fig. 3). This suggests that the exposure of the sample to a specific H₂O pressure is responsible for a sudden change in the structure, hinting at an adsorption process similar to the one displayed by CO2 and possibly involving the same adsorption sites. This is supported by the very similar saturation loading of about 2.5 mmol g⁻¹ for both water and CO₂, corresponding to one adsorbate molecule per formula unit (see Fig. 2a and S15†).

In situ IR analysis using CO and N2 as molecular probes at liquid nitrogen temperature was performed to further study the presence of Lewis acidic adsorption sites in the evacuated MOF (Fig. S16†). Both the CO and N₂ IR measurements highlight that, after H₂O removal, F4_MIL-140A(Ce) features accessible Ce^{IV} sites available for interaction with Lewis bases, even as weak as N2. Detailed discussion of these results is available in Section S4 of the ESI.†

Next, we investigated the vibrational behaviour of evacuated F4_MIL-140A(Ce) when exposed to an increasing CO₂ pressure in the range between 110 and 550 mbar, at beam temperature (i.e., around 313 K). The CO2 pressure range was accurately selected by considering the position of the step of the CO₂ isotherm collected at 313 K (Fig. 4a). The incremental CO2 doses are responsible for significant changes in the CO₂ asymmetric stretching region between 2400 and 2250 cm⁻¹ (Fig. 4b) and in the 800–770 cm^{-1} spectral range (Fig. 4c). Upon sending CO_2 doses below the isotherm step (i.e., until around 110 mbar), a new, quite broad band at 2344 cm⁻¹ appears, slightly shifted to lower frequencies compared to the antisymmetric CO2 stretching vibrational mode (at 2349 cm⁻¹). At the same time, no significant changes are detected in the low frequency region, showing only a slight decrease in intensity of the 790 and 779 cm⁻¹ bands. These spectral modifications suggest that, below the isotherm step, CO₂ weakly interacts with the sorbent, likely because access to the open metal sites is prevented by the structural arrangement within channels. By increasing the CO2 pressure and approaching the isotherm step, a new band at 2361 cm⁻¹, probably due to the interaction of CO₂ with Ce^{IV}, appears, while the 2344 cm⁻¹ band increases giving rise to an

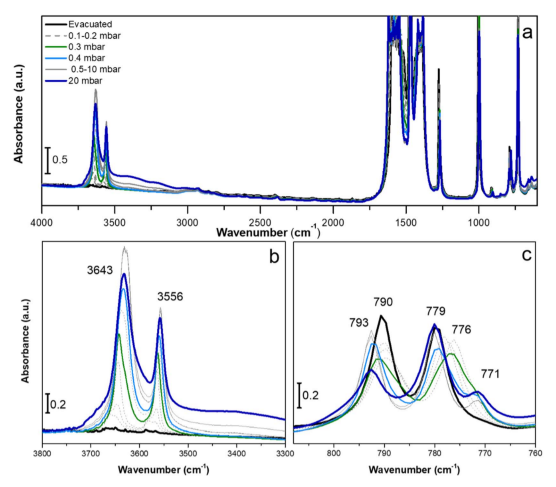


Fig. 3 IR spectra of F4_MIL-140A(Ce) collected dosing 0.1-20 mbar of H_2O on the evacuated sample, reported in the full spectral range (a), and in the $3800-3300 \text{ cm}^{-1}$ (b) and $810-760 \text{ cm}^{-1}$ (c) regions.

intense band with apparent maxima at 2344 and 2330 cm⁻¹. In general, the assignment of the above-mentioned bands is not straightforward because the rotovibrational profile of gaseous CO2, which falls in the same spectral region and prevails at high CO₂ coverages, superimposes on those components. Concurrently, the bands at 790 and 779 cm⁻¹ are shifted to 788 and 782 cm⁻¹, respectively, suggesting that changes are occurring on the fluorinated aromatic rings. The CO2 doses corresponding to the final part of the adsorption step and the beginning of the isotherm plateau cause no further changes in the low frequency spectral range, while both the 2361 and the 2344 and 2330 cm⁻¹ bands go out of scale. The predominance of the band at 2361 cm⁻¹, possibly related to the direct interaction of CO₂ with Ce^{IV} open metal sites, at CO₂ pressures that correspond to the isotherm step could be a further proof of the occurrence of a phase-change adsorption process that enables the establishment of a strong adsorbate/surface interaction.

DFT optimisation of molecular adducts. IR experiments provided good evidence of the presence of open metal sites able to interact with various adsorbates. The energetics of interaction of F4_MIL-140A(Ce) with H_2O , CO, N_2 and CO_2 were then investigated by means of periodic DFT calculations. Initially, the experimental structure of the MOF from ref. 28 was

optimized in the absence of any adsorbed molecule, i.e. in its evacuated form. The adsorption of H2O, CO, N2 and CO2 was then simulated by manually placing the adsorbates at opportune positions within the MOF (i.e., in the vicinity of the inorganic building unit) and by reoptimising the obtained structure. To fully exploit the framework symmetry, only full coverage (i.e., 1:1 adsorbate/Ce ratio) was studied. For H₂O, CO and N₂, the molecule was positioned in direct interaction with the Ce open metal site through the O atom, the C atom and one of the N atoms, respectively. In the case of CO₂, two possible adsorption sites were considered: (i) as for the other adsorbates, with an O atom in direct interaction with Ce (referred as "Ce" site); and (ii) positioned in the pore and interacting with the surrounding perfluorobenzene rings, as recently proposed by Zhao and coworkers30 (labelled here as "channel" site). The graphical representation of these adducts is presented Section 5 of the ESI,† whereas the main structural parameters are listed in Table S3.† The adsorption energetics for these adducts were also computed (details in Section 5 of the ESI†), as reported in Table 1.

The energetic description of the adsorption phenomena suggests a strong preference of F4_MIL-140A(Ce) for H2O over the other investigated adsorbates. Interestingly, in the case of

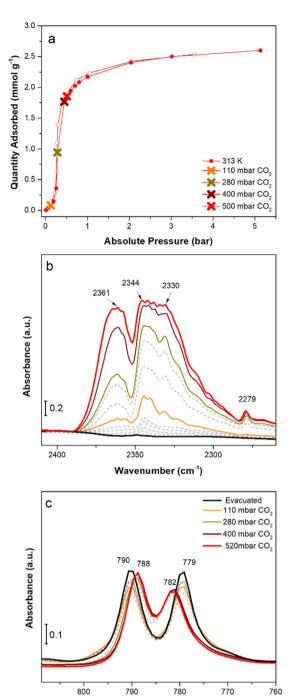


Fig. 4 CO₂ adsorption (filled circles) and desorption (empty circles) isotherms collected on F4_MIL-140A(Ce) at 313 K (40 °C) up to 5 bar (a). IR spectra of CO₂ adsorption on evacuated F4_MIL-140A(Ce) reported in the 2410-2260 cm⁻¹ (b) and 810-760 cm⁻¹ (c) spectral ranges. The same incremental pressure doses of CO2 reported with coloured crosses in the volumetric isotherm of panel (a) were employed during the IR experiment. Dashed lines represent intermediate pressures.

Wavenumber (cm⁻¹)

CO₂, distinct interaction energies are computed depending on the adsorption site. In particular, the "channel" site, analogous to what previously proposed by Zhao and coworkers as the

Table 1 Adsorption electronic energy (ΔE) enthalpy (ΔH) and Gibbs free energy (ΔG) for F4_MIL-140A(Ce) in interaction with H₂O, CO₂ (for the latter, at both the "Ce" and the "channel" sites), CO and N_2 . All values are corrected for the basis set superposition error and given per adsorbed molecule. ΔH and ΔG values are computed at a temperature of 298.15 K and a pressure of 1013 mbar

Model	$\frac{\Delta E}{\text{(kJ mol}^{-1})}$	ΔH (kJ mol ⁻¹)	ΔG (kJ mol ⁻¹)
+H ₂ O	-86.0	-77.8	-35.8
+CO ₂ (Ce)	-55.2	-46.1	-7.5
+CO ₂ (channel)	-35.2	-26.1	10.5
+CO	-40.5	-36.1	1.0
+N ₂	-32.2	-28.0	6.8

preferred one for CO2 adsorption, shows a lower interaction energy compared to that of CO₂ in direct interaction with the open metal site. Furthermore, the ΔG values estimated for the adsorption occurring at the two sites point out that the interaction is thermodynamically favoured only for the "Ce" site, whereas the "channel" site is not providing a stable adduct at ambient conditions. In fact, the computed ΔG value for CO₂ adsorption at the "channel" site is the highest observed in Table 1, suggesting that the involvement of the Ce^{IV} site is key for a favourable interaction with the adsorbates. A close inspection of the optimised crystal structures reveals that both H₂O and CO₂ are stabilised by interactions with both the metal centre (a Lewis acid), through their negatively polarised oxygen atoms, and the fluorine atoms belonging to the organic linker (Lewis bases), through the H atoms and C atom, respectively (Fig. S18-S24†). Thus, the ability of H₂O and CO₂ to behave as both Lewis acids and bases appears to be crucial to maximise their affinity for the adsorbent surface. We also simulated the interaction of CO₂ with a fully hydrogenated MIL-140A(Ce) (Fig. S25†), observing a significant reduction in binding energy to the "Ce" site down to $-39.4 \text{ kJ mol}^{-1}$. This value is comparable to the strength of the interaction computed in the case of CO with F4_MIL-140A(Ce). Notably, the Gibbs free energy becomes positive in MIL-140A(Ce) (5.8 kJ mol⁻¹), suggesting that the interactions with the linker also play a key role in stabilising the MOF-CO2 adduct.

Structural characterisation

Powder X-ray crystallography: long range structure. To investigate the structural response of F4 MIL-140A(Ce) upon evacuation and CO2 adsorption, we carried out an in situ synchrotron PXRD study. The as-synthesised MOF was first kept under dynamic vacuum, progressively heating up to 403 K. A phase transition occurs already below 373 K, corresponding with the removal of the adsorbed H₂O molecules, and the new phase is stable when cooled down to 298 K (Fig. S26†). The structure retains the monoclinic C2/c space group during the phase transition. The main change in the unit cell is the increase of the β angle from 96.35(5)° in the as-synthesised MOF to 103.99(2)° in the evacuated one, which leads to a 2.7% decrease in volume from 2402(2) Å³ to 2336.0(5) Å³, respectively

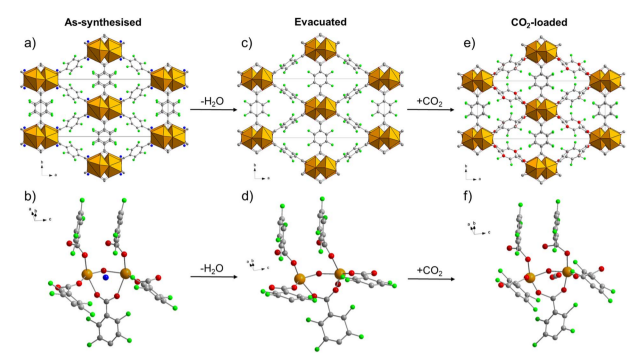


Fig. 5 Comparison of the crystal structure viewed along the c axis and the local environment around the adsorption site of the as-synthesised (a and b, respectively), evacuated (c and d, respectively) and CO_2 -loaded (e and f, respectively) forms of F4_MIL-140A(Ce). Colour code: Ce, orange; F, green; C, grey; O, red; H_2O , blue. H atoms not shown because their positions cannot be determined from PXRD data.

(Table S4†). The trend is in agreement with what suggested by DFT optimisation (Table S3†). Rietveld refinement of the PXRD pattern for the evacuated MOF was carried out starting from the DFT-optimised model, finding that H₂O removal leads to rotation of the aromatic rings belonging to linker A (Table S5, Fig. 5a–d, S28–S29 and S44†). This structural rearrangement occurs in response to the loss of several interactions between the framework and the water molecules: H₂O interacts with one Ce atom *via* its electron rich O atom and with at least three coordinated carboxylic O atoms and three F atoms (belonging to both linkers A and B) through hydrogen bonding (Fig. 1 and S44†).

The evacuated F4_MIL-140A(Ce) was then held at a constant temperature of 328 K and exposed to increasing CO₂ pressures in the 0.055-1.485 bar range. As displayed in Fig. S30,† a phase transition occurs between 0.367 and 0.745 bar, corresponding to the pressure range where the CO2 adsorption isotherm collected at the same temperature displays the step (Fig. S31†). Upon reduction of the pressure from 1.485 down to 0.322 bar, the reverse transition takes place, regenerating the evacuated form of the MOF (Fig. S32†). An analogous, reversible phase transition is observed between 0.222 and 0.527 bar at 313 K, and between 0.833 and 1.490 bar at 343 K, in both cases in excellent agreement with the step position in the corresponding CO₂ adsorption isotherms (Fig. S33-S36†). The PXRD pattern of the CO₂-loaded MOF is consistent at the three temperatures, suggesting that the equilibrium structure is the same in the investigated temperature range (Fig. S37†). We also performed an isobaric experiment exposing the MOF to a constant pressure of 0.498 bar and decreasing the temperature from 343 to 313 K in 5 K steps (Fig. S38†). The MOF switches from the evacuated to the CO₂-loaded form between 328 and 323 K, again in excellent agreement with what expected from the adsorption isotherms measured at different temperatures (Fig. S39†). Again, the unit cell parameters are only slightly affected by the phase transition, with the β angle decreasing from 103.99(2)° in the evacuated MOF to $97.68(1)^{\circ}$ in the CO₂-loaded one (Table S4†). This trend is also in agreement with what suggested by DFT optimisation (Table S3†). For the sake of Rietveld refinement of the CO₂-loaded MOF, a structural model with CO₂ adsorbed at the "Ce" site was adopted as the most probable guess, based on the DFT results previously discussed. Rietveld refinement was carried out on the PXRD pattern for the CO₂-loaded MOF at 313 K, finding that the insertion of CO₂ triggers rotation of linker A to a position similar to that observed for the as-synthesised MOF (Table S5,† Fig. 5e-f and S41). The CO₂ molecule displays coordination-like interaction with two Ce atoms, via one of its electron-rich O atoms, besides additional interactions with three F atoms (belonging to both linkers A and B) and with two coordinated carboxylic O atoms, via the electron-poor C atom (Fig. S42†). Thus, CO₂ appears to be adsorbed in a pocket where it is stabilised by a wealth of non-covalent interactions, akin to what seen for H₂O in the as-synthesised MOF. The step observed in the CO₂ adsorption isotherm most likely arises from the concerted rotation of perfluorinated aromatic rings upon the insertion of CO₂ in such an adsorption site, once the activation barrier for ring rotation is overcome and the site is made accessible. Further support to this interpretation is provided herein by means of other characterisation techniques.

Comparing the three forms of F4_MIL-140A(Ce), it is evident that the as-synthesised and the CO2-loaded ones have common features, ascribable to the presence of the adsorbate, which are not shared by the evacuated form. By examining the torsional angles between the carboxylate groups and the aromatic rings for each linker in the different forms of the MOF (Table S5†), it can be seen that O2-C1-C2-C3 (linker A) and O5-C10-C9-C8 (linker B) torsional angles significantly change when adsorbates are present, by bringing the aromatic rings more out-of-plane respect to the carboxylates, as a result of interactions with the adsorbate. The third torsional angle, O4-C5-C6-C7 belonging to linker B, is instead only slightly affected, because of the lack of interactions with the adsorbates on that side of the linker (Fig. S43†).

XAS (Ce K-edge): local structure of the metal ion. To complement the long-range structural analysis carried out by PXRD, Ce K-edge XAS spectra were collected on the assynthesised, evacuated and CO₂-loaded F4_MIL-140A(Ce) samples, respectively (Fig. S45†). This allowed us probing the local structure of Ce centres and gathering sensible information by fitting the extended X-ray absorption fine structure (EXAFS) signal as already done for several MOFs. 38-42 The experiments were carried out at 328 K in dynamic conditions, i.e., under a He flow for the evacuation step and under a controlled mixing of He and CO₂ to follow the adsorption/desorption of CO₂ within the porous system. The changes displayed by the EXAFS signal (Fig. S46a†) collected on F4_MIL-140A(Ce) during the adsorption process agree with what expected on the basis of the CO₂ adsorption isotherm measured at 328 K (Fig. S9†). Moreover, the Fourier transform of the EXAFS signal (Fig. S46b†) shows evident changes in the 2.5-4 Å range, i.e., the second shell of ligands from Ce centres. The signals observed in this region can be assigned, by looking at the crystallographic structure, to Ce-Ce, Ce-H2O and Ce-CO2 distances which are indeed being modified by adsorption/desorption processes. The scattering length of these three paths was parametrised in order to achieve a satisfactory data/parameter ratio allowing us to refine one Ce-O_w and two Ce-C_{CO_o} distances in addition to the Ce-Ce ones. By fitting the EXAFS signal using paths whose starting position is calculated from the crystallographic data coming from the Rietveld refinement, results in Fig. 6 and Table S6† are obtained.

Ce-Ce paths, whose refinement was compulsory because their distances fall very close to the Ce-adsorbates ones, gave values in excellent agreement with the ones observed by PXRD. The Ce-adsorbate distance for the as-synthesised material (Ce- $O_{\rm w}$) was refined at 3.368(3) Å, whereas the Ce- $C_{\rm CO}$ distances for the CO2-loaded one were fitted respectively to 4.06(3) Å and 4.44(4) Å. Such values are in agreement with those refined from PXRD data through Rietveld method (2.89 and 3.62 for Ce-Ow; 4.04 and 4.44 for Ce-C_{CO₂}) and from DFT optimisation (2.63 and 3.13 for Ce-O_w; 3.93 and 4.20 for Ce-C_{CO_s}), thus strengthening their reliability and consistency.

SSNMR: local structure of the adsorbate and the organic linker. The local structure of as-synthesised, evacuated, and CO₂-loaded F4_MIL-140A(Ce) was further characterized by multinuclear SSNMR spectroscopy exploiting the ¹⁹F and ¹³C nuclei of the linkers. Moreover, for the as-synthesised and the CO₂-loaded samples, ¹H and ¹³C nuclei of water and CO₂ were respectively investigated to unravel the state of the adsorbates in the MOF.

The ¹H direct excitation (DE) static spectrum of assynthesised F4_MIL-140A(Ce) indicates the presence of both weakly physisorbed and structural water, which is completely removed after heating under vacuum, as shown by the absence of any signal in the spectrum of evacuated F4_MIL-140A(Ce) (Fig. S47†). The ¹³C DE spectrum of CO₂-loaded F4_MIL-140A(Ce) (Fig. 7a), recorded under magic angle spinning

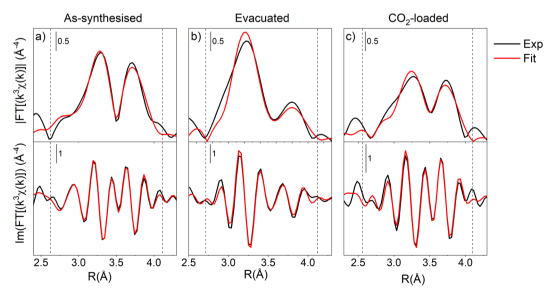


Fig. 6 Comparison between experimental (black lines) and fitted values (red lines) of magnitude (top) and imaginary part (bottom) of phaseuncorrected Fourier transform of K^3 -weighted Ce K-edge EXAFS spectra for the as-synthesised (a), evacuated (b) and CO_2 -loaded (c) states. Dashed lines delimit the fitting range.

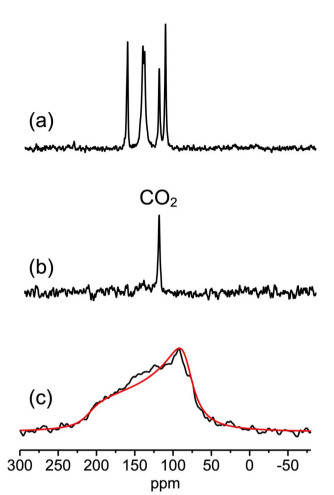


Fig. 7 13 C DE spectra of CO₂-loaded F4_MIL-140A(Ce) recorded (a) under MAS with a recycle delay of 150 s, (b) under MAS with a recycle delay of 2 s, and (c) in static conditions with a recycle delay of 2 s.

(MAS) at a rate of 15 kHz and with a recycle delay between subsequent scans (150 s) sufficient to quantitatively observe the signals of MOF and CO₂ carbons, indicates that almost one CO₂ molecule (signal at 124.7 ppm) per Ce atom was adsorbed in the MOF when loaded at a CO₂ pressure of 1 bar. When the ¹³C DE-MAS spectrum is acquired with a short recycle delay of 2 s, the sole signal of CO₂ is selectively observed (Fig. 7b), since the ¹³C longitudinal relaxation time of CO2 carbons is much shorter than those of the linker carbons, in agreement with ref. 43. The ¹³C DE static spectrum recorded with the short recycle delay shows a powder pattern with a line shape arising from the anisotropic ¹³C chemical shift (CS) interaction averaged by CO₂ dynamics (Fig. 7c), as found in other MOFs.44 The observed residual chemical shift anisotropy (CSA) indicates that CO2 reorientation in F4_MIL-140A(Ce) is not isotropic, the CO2 mobility being restricted by the interactions with the MOF linkers and Ce atoms and/or by steric hindrance in the pores. Generally, the CS interaction can be described with a second rank tensor with three principal components δ_{11} , δ_{22} , and δ_{33} , with $\delta_{11} \ge \delta_{22} \ge \delta_{33}$ ("Mehring notation"). Three parameters are sufficient to describe the tensor: following the "Maryland

notation", they are the isotropic chemical shift $[\delta_{iso} = (\delta_{11} + \delta_{22} +$ δ_{33})/3], the span ($\Omega = \delta_{11} - \delta_{33}$), and the skew [$\kappa = 3(\delta_{22} - \delta_{iso})$ / Ω]. The span measures the breadth of the observed powder pattern and is used to indicate the extent of the CSA, while the skew, ranging from -1 to 1, gives an indication of the symmetry of the CS tensor, the two limit values of κ corresponding to an axial symmetry. For CO₂ in the solid state, an axial CS tensor is reported in the literature, with $\Omega = 315-335$ ppm and $\kappa = 1.46,47$ The signal observed for CO2 adsorbed on F4_MIL-140A(Ce) is instead characterised by $\Omega \simeq 127$ ppm and $\kappa = -0.98$. The span reduction with respect to fully immobilised CO₂ indicates that a motion with a rate higher than the spectral broadness in frequency units occurs for CO2 in F4_MIL-140A(Ce) at room temperature. Wobbling on an individual adsorption site and hopping among symmetry equivalent adsorption sites have been found for CO2 in other MOFs by 13C SSNMR spectroscopy. 48-53 In our case, the negative value of κ suggests that a wobbling or hopping motion occurs with an angle (β) between the C_{∞} symmetry axis of CO₂ and the motion axis larger than the magic angle (54.74°),⁵³ while the axial line shape indicates a 3-fold or higher symmetry for the motion, which should occur in the fast regime. A satisfactory simulation of the spectral line shape (Fig. 7c) could indeed be obtained with a wobbling in a cone motion among 3 (or more) equivalent sites in the fast regime with $\beta = 75^{\circ}$. It must be pointed out that this motion could be associated either to a reorientation of CO₂ about a local axis or to the hopping of CO₂ molecules among equivalent adsorption sites, also in combination with translation. The occurrence of both wobbling and hopping motions cannot be excluded. Further variable temperature 13C SSNMR static experiments on F4_MIL-140A(Ce) loaded with 13C isotopically enriched CO2 and line shape analyses are necessary to characterise CO2 dynamics in more detail.

¹⁹F DE and ¹⁹F-¹³C cross polarization (CP) experiments were also recorded under MAS on all samples to highlight possible effects of H₂O or CO₂ adsorption on linkers. The spectra, showing signals from the linker fluorine and carbon atoms, respectively, are reported in Fig. S48† and 8.

¹⁹F DE-MAS spectra show broad spinning sideband patterns (Fig. S48†) due to the large chemical shift anisotropy typical of ¹⁹F nuclei.⁵⁴ The ¹⁹F isotropic signals (Fig. 8a) lie in the spectral region between -150 and -135 ppm. A spectral deconvolution of the whole spinning sideband profiles showed that inequivalent fluorine atoms are present in all samples: two in the assynthesised MOF, with isotropic chemical shifts of -143.1 and -144.8 ppm and intensities in a 3:1 ratio; three in the evacuated MOF, with chemical shifts of -141.0, -142.0, and -143.0 ppm and intensities in a 1:2:1 proportion; two in the CO_2 -loaded MOF with isotropic chemical shifts of -138.9 and -142.1 ppm and 1:3 relative intensities. In the $^{19}F^{-13}C$ CP-MAS spectra (Fig. 8b), the signals of fluorinated (CF), quaternary (Cq), and carboxylic (COO⁻) carbons of the linkers can be recognised in three distinct spectral regions. For each kind of carbon, several peaks are observed, as better appreciated after spectral deconvolution (Fig. S49†). The observed ¹⁹F and ¹³C multiple signals are ascribable to the two crystallographically inequivalent linkers in the structure of F4 MIL140 (Ce), namely,

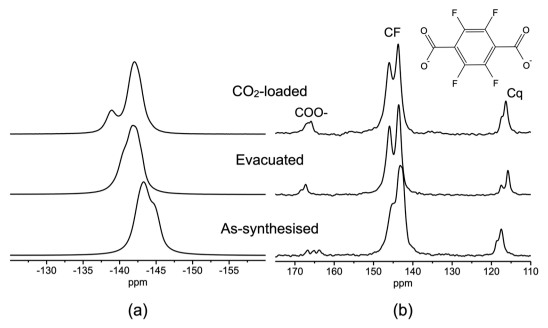


Fig. 8 (a) 19 F DE-MAS spectra and (b) 19 F $^{-13}$ C CP-MAS spectra (ct = 1 ms) recorded on (from bottom to top) as-synthesised, evacuated, and CO $_2$ -loaded F4 $_{\rm MIL}$ -140A(Ce).

linker A and linker B. Analogous results were reported for MIL-140A(Zr). To distinguish signals arising from linkers A and B, the correlations occurring in 2D $^{19}\mathrm{F}^{-13}\mathrm{C}$ HETCOR spectra (Fig. S50–S52†) between signals of $^{13}\mathrm{C}$ and $^{19}\mathrm{F}$ nuclei in spatial proximity were also used. Therefore, signal assignment was possible combining the analysis of 1D $^{19}\mathrm{F}$ and $^{19}\mathrm{F}^{-13}\mathrm{C}$ CP-MAS spectra (recorded at different contact time values) with that of 2D $^{19}\mathrm{F}^{-13}\mathrm{C}$ HETCOR spectra. The results, shown in Table 2 for as-synthesised, evacuated, and CO₂-loaded F4_MIL-140A(Ce), clearly indicate the presence of two different linkers with symmetry compatible with that found by the PXRD structural analysis (see Section 8 in the ESI† for a detailed description of signal assignment).

Interestingly, shifts of 13 C and 19 F signals are observed among the as-synthesised, evacuated, and 19 F observed MOF

(Fig. 8 and Table 2), ascribable to the change of local chemical environment caused by adsorption of water or CO₂. For instance, the shielding of ¹⁹F nuclei could be related to the interaction between the linker fluorine atoms and the adsorbate molecules, ^{56,57} although the rotation of the phenyl rings out of the plane of carboxylates could also play a role. For assynthesised F4_MIL-140A(Ce), ¹³C signals from COO⁻ and Cq carbons spatially close to water ¹H nuclei are highlighted in the ¹H-¹³C CP-MAS spectrum shown in Fig. S53,† since only for these carbons ¹H-¹³C magnetisation transfer can effectively take place through dipolar coupling.

On the other hand, the signal of CO_2 carbons was not observed in $^{19}F^{-13}C$ CP MAS spectra of CO_2 -loaded F4_MIL-140A(Ce) (Fig. 8b), even at contact time values as long as 10 ms. The distances between linker fluorines and CO_2 in the

Table 2 Assignment of signals in 1D 19 F DE-MAS and 19 F $^{-13}$ C CP-MAS spectra (Fig. 8) and in 2D 19 F $^{-13}$ C HETCOR spectra (Fig. S50–S52) of assynthesised, evacuated, and CO₂-loaded F4_MIL-140A(Ce)

Linker	Position	Atom	As-synthesised		Evacuated		CO ₂ -loaded	
			$\delta(^{13}\mathrm{C})\mathrm{(ppm)}$	$\delta(^{19}F)$ (ppm)	$\delta(^{13}\text{C}) \text{ (ppm)}$	$\delta(^{19}\mathrm{F})\mathrm{(ppm)}$	$\delta(^{13}\text{C}) \text{ (ppm)}$	δ(¹⁹ F) (ppm)
A	CF	C4	145.3	-143.1	146.0	-142.0	143.6	-142.1
		C3	145.3	-143.1	146.0	-142.0	145.7	-142.1
	Cq	C2	117.6		115.8		116.3	
	COO^-	C1	163.7		168.4		165.8	
В	\mathbf{CF}	C8	143.5	-143.1	143.6	-141.0	145.7	-142.1
		C7	142.6	-144.8	143.6	-143.0	143.6	-138.9
	Cq	C9	117.6		117.4		117.2	
	•	C6	118.7		115.8		116.3	
	COO^-	C10	166.9		167.3		167.2	
		C5	165.2		167.3		166.5	

Rietveld-refined structure of CO_2 -loaded F4_MIL-140A(Ce) (Fig. S42 and S43†) would allow sufficiently strong dipolar interactions to occur for an efficient magnetization transfer between ¹⁹F and ¹³C nuclei, as observed by other authors.⁵⁸ The scarce CP efficiency found in our case should therefore be ascribed to a reduction of the dipolar interactions and/or of the residence time of CO_2 in the adsorption site due to reorientational and translational motions.⁵⁹⁻⁶¹

Energetics of the cooperative CO₂ adsorption mechanism. The foregoing discussion on the adsorption properties and structural response of F4 MIL-140A(Ce) to adsorption and desorption of H2O and CO2 suggests that a concerted mechanism takes place when CO2 is involved, due to the ease of rotation of the perfluorobenzene rings. The peculiar CO2 adsorption mechanism was further investigated by performing adsorption microcalorimetry at 303 K to directly measure the CO₂ molar adsorption heat at different coverages (see the ESI† for further details). Fig. 9 reports the volumetric CO₂ isotherms (a) and the differential molar adsorption heats (b) of F4_MIL-140A(Ce). The volumetric isotherm collected in the 0-0.6 bar pressure range with the microcalorimetric apparatus (dark grey curve in Fig. 9a) is perfectly in agreement with the isotherm obtained by the standard automatic volumetric instrument (light blue curve in Fig. 9a and S54† shows a comparison with the isotherms collected at 298 and 313 K). The differential molar adsorption heat curve $(-q_{\text{diff}}, \text{ Fig. 9b})$ unusually starts from low values, close to the molar heat of liquefaction of CO2 (17 kJ mol⁻¹). The differential heat then increases until 25 kJ mol⁻¹ and rapidly decreases to 22 kJ mol⁻¹. After this intermediate step, $-q_{\text{diff}}$ reaches a plateau at \sim 34 kJ mol⁻¹ corresponding to a coverage above 0.2 mmol g⁻¹, i.e., exactly where the step in the isotherm is taking place. These unusual values of differential heat clearly suggest that some endothermic modification of the adsorbent is occurring concurrently with the exothermic adsorption of CO₂. Indeed, microcalorimetry provides the "average" heat of adsorption which could

derive from different contributions generated by the interaction between the porous adsorbent and the adsorptive. If the endothermic event is particularly heat-demanding, the resulting net differential heat displays this peculiar trend.62 After reaching the value of 34 kJ mol⁻¹, $-q_{\text{diff}}$ remains constant until a coverage of 2 mmol g^{-1} (a value corresponding to the decrease in the slope of the adsorption isotherm immediately after the step). The relatively high value of $-q_{\text{diff}}$ over a wide range of coverages suggests that CO2 strongly interacts with a homogeneous set of adsorption sites distributed on the internal surface of F4 MIL-140A(Ce).38 When all the available adsorption sites are covered by CO2, the differential heat curve steeply decreases, again approaching the heat of liquefaction of CO₂. At the step, the differential heat curve is in agreement with the isosteric heat of adsorption plot, indirectly extracted from the CO₂ isotherms by applying the Clausius-Clapeyron equation (Fig. S6†). However, the Clausius-Clapeyron model fails in identifying a region of low heat of adsorption before the step takes place.

DFT also provides further insights in the description of the adsorption phenomena: the initial q_{diff} is closely approached by the ΔH value computed for CO₂ adsorbed at the "channel" site (see Table 1). At plateau, the q_{diff} value is instead comparable to that computed for CO2 at the "Ce" site, though slightly lower. The peculiar energy profile revealed by adsorption microcalorimetry could be explained by a scarce initial affinity of CO₂ for the surface of the adsorbent. As a matter of fact, even though the "channel" site is not the most favourable adsorption site for CO2, the adsorptive must necessarily pass through the porous system to reach the most favourable "Ce" site. Furthermore, the displacement of CO₂ implies the rearrangement of the material, i.e., the concerted rotation of the perfluorobenzene rings, to enable access to the most energetic adsorption site. The occurrence of this endothermic phenomenon is clearly proved by the differential heat trend observed at the onset of the isotherm step.

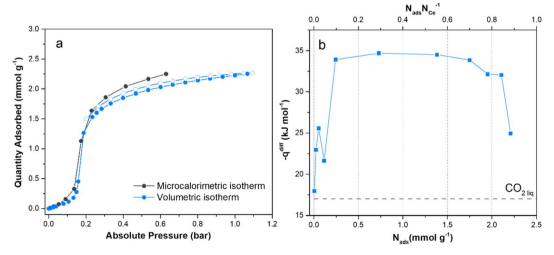


Fig. 9 Comparison of volumetric isotherms obtained by performing CO_2 adsorption at 303 K with the automatic volumetric instrument (blue) and with the volumetric line coupled with the microcalorimeter (black) (a). Differential molar adsorption heats curve related to the adsorption of CO_2 at 303 K (b). The dotted horizontal line represents the standard molar heat of liquefaction of CO_2 at 303 K: 17 kJ mol⁻¹.

To give a quantitative description of the phenomenon, a relaxed potential energy surface scan was performed by DFT. In brief, CO2 initially modelled in the "channel" site, was stepwise approached to the "Ce" site. At each step, the whole structure was reoptimized, keeping the distance between the centre of mass of CO₂ (i.e., its C atom) and Ce constant. The

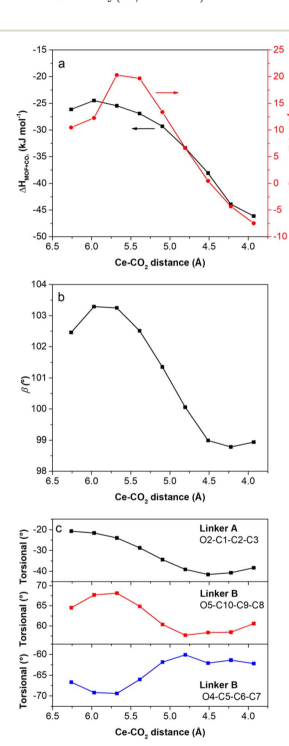


Fig. 10 Trends of: (a) ΔH and ΔG for CO₂ adsorption; (b) β angle of the monoclinic unit cell; and (c) perfluorobenzene ring-carboxylate torsional angles as a function of the Ce-CO₂ distance. ΔH and ΔG values are computed at a temperature of 298.15 K and a pressure of 1013 mbar.

trends of the energetic (ΔH and ΔG) and selected structural (β cell angle, O2-C1-C2-C3 torsional angle in linker A, O5-C10-C9-C8 and O4-C5-C6-C7 torsional angles in linker B) descriptors are reported as a function of the Ce-CO₂ distance in Fig. 10.

The energy trend clearly shows the existence of an activation barrier (ΔG barrier of ~ 10 kJ mol⁻¹) that the CO₂ molecule must overcome to reach the most favourable "Ce" site. As the barrier is overtaken, a drastic modification in the structural features of the MOF takes place, accompanied by a progressive stabilisation of the MOF-CO₂ adduct. This rearrangement implies the distortion of the unit cell through a decrease of the β cell angle, as well as the rotation of the perfluorobenzene rings, as testified by the concomitant variation of their torsional angles compared to the carboxylates. The trend from DFT simulation is in full agreement with that observed by PXRD. The key role played by the Ce^{IV} open metal site in F4 MIL-140A(Ce) also explains the lack of step observed in the CO2 isotherm of its Zr-based analogue, recently reported by Zhao and co-workers.30 Indeed, Zr^{IV} is seven-coordinate in the MIL-140 topology, with no H₂O coordinated in its as-synthesised form and, therefore, no available open metal sites for adsorbates to interact upon evacuation.

Conclusions

Through a combination of experimental and computational methods, we have produced a coherent set of evidence supporting the existence of a cooperative mechanism of CO2 adsorption in F4_MIL-140A(Ce), which involves the concerted rotation of perfluorinated aromatic rings. This phenomenon opens the gate for CO2 to reach a highly favourable adsorption site, where it is stabilised by interactions with both coordinatively unsaturated Ce^{IV} sites and the organic linkers. The step-shaped isotherm of F4 MIL-140A(Ce) resulting from this phase-change behaviour allows to reach saturation in a narrow range of partial pressure compatible with real-life postcombustion capture requirements. While several MOFs are known to display either adsorption at coordinatively unsaturated metal sites or concerted ring rotation, to the best of our knowledge the combination of these two features is observed in F4_MIL-140A(Ce) for the first time. Furthermore, the phase transition has a low activation barrier and does not involve major restructuring of the framework, resulting in a complete reversibility in desorption and leading to a non-hysteretic behaviour. Based on the knowledge acquired with this work, we expect that the CO2 adsorption behaviour of F4_MIL-140A(Ce) can be finely tuned by both varying the fluorination degree of the organic linker and preparing mixed Ce/Zr analogues, paving the way towards the development of a class of adsorbents based on the same working principle. In view of the practical application of F4_MIL-140A(Ce) in postcombustion CO₂ capture, its high affinity for water could be detrimental to the capture performance, as water might preferentially adsorb over CO2. We are currently investigating this aspect using dynamic breakthrough analysis, testing both dry and humid gas mixtures.

Experimental procedures

Gas adsorption volumetry

N₂, Ar and CO₂ isothermal physisorption measurements were performed on a Micromeritics ASAP 2020 analyser at 77 K, 87 K and 273 K, respectively. For CO₂ adsorption the temperature was kept constant thanks to a home-made patented glass coating cell⁶³ in which a coolant or heating fluid, connected to a thermostatic bath (JULABO F25), is recirculating. Prior to the measurement, the sample was heated overnight at 393 K under vacuum. Specific surface areas were determined by using the Brunauer–Emmett–Teller (BET) method, following the Rouquerol consistency criteria.⁶⁴ Cumulative pore volume and pore size distribution were derived from adsorption isotherms by applying the NL-DFT method (see ESI,† Section 3, for further information).

High pressure CO_2 adsorption–desorption isotherms up to 5 bar were measured with a Quantachrome iSorb High Pressure Gas Analyser at 298, 313, 328 and 343 K. About 200 mg of sample was used for the adsorption studies. The sample was degassed at 393 K under dynamic vacuum for 12 h prior to analysis and at 393 K for 1 h in between subsequent measurements.

In situ IR spectroscopy

IR spectroscopy was performed on a Bruker Vertex 70 instrument equipped with a MCT (mercury cadmium tellurium) cryodetector. All the spectra were recorded in the 4500-600 cm⁻¹ spectral range with a resolution of 2 cm⁻¹ and an average of 32 scans. Before the analysis, F4_MIL-140A(Ce), as self-supported pellet, was held in vacuum at 393 K for 12 h. Both CO and N2 interactions with the sample were studied by dosing a proper amount of gas (~60 mbar) into the cell and cooling down to 77 K with liquid nitrogen. The spectra reported for both the experiments were collected in outgassing through subsequent expansions until complete desorption. H2O adsorption was carried out by sending incremental doses of H₂O (from 0.1 to 20 mbar) on the evacuated F4_MIL-140A(Ce). The CO2 interaction with F4_MIL-140A(Ce) was studied by sending increasing CO₂ doses on the sample after applying the same evacuation procedure previously described. The increasing doses of CO₂ were chosen to be compatible with the equilibrium pressures involved in the adsorption/desorption isotherm at 313 K (40 °C) which is approximately the IR beam temperature.

Periodic DFT simulations

Vibrational and energetic features of F4_MIL-140A(Ce), also in the presence of relevant adsorbates, were simulated at the DFT level of theory with the CRYSTAL17 code.^{65,66} In detail, the B3LYP hybrid GGA functional^{67,68} was exploited in conjunction with a double-ζ quality basis set (C and O from ref. 39; F from ref. 69; and Ce from ref. 70). A larger Ahlrichs TZV2p basis³⁹ described atoms belonging to adsorbed molecules (H₂O, CO₂, CO and N₂). Dispersive interactions were empirically included in the Hamiltonian according to the Becke-Johnson dumped version of the Grimme's D3 scheme.^{71,72} Such computational

setup provided a satisfactory structural and electronic description of Ce^{IV}-MOFs.⁷³

In situ PXRD

The in situ PXRD study here described took place at Beamline P02.1 (PETRA III, DESY, Hamburg).74 PXRD patterns were collected using 60.0 keV (0.207124 Å) radiation. The sample was held in a 0.5 mm internal diameter glass capillary, which was mounted on a custom-built spinner device that allows for rotating a gas-filled capillary mounted on a commercial Huber goniometer head by 360° forth and back. The temperature of the sample was controlled between 298 and 413 K with an Oxford Hot-Air Blower. The beamline was equipped with a PerkinElmer XRD1621 CN3 - EHS detector (200 \times 200 μ m² pixel size, 2048×2048 pixel area). The detector was positioned at 1000 mm from the sample stage. Each PXRD pattern was collected in the 0.0071–16.3041 $^{\circ}2\theta$ range, with a 0.0067 $^{\circ}2\theta$ step size and a total acquisition time of 30 s. The resulting 2D images were azimuthally integrated to 1D diffraction patterns using the software Fit2D.75

In situ XAS

Ce K-edge XAS data were collected at the BM23 (ref. 76) beamline of the European Synchrotron Radiation Facility (ESRF) in Grenoble, France. The storage ring was operating in the 32-bunch mode with a target current of 150 mA. The measurements were conducted in transmission mode using a Si(311) double-crystal monochromator. The intensity of the X-ray beam was measured by means of three ion chambers (30 cm, 1 kV) placed before the sample (I_0 , filling 0.28 bar Kr + 1.72 bar He), after the sample (I_1 , filling 1.35 bar Kr + 0.65 bar He) and after the CeO₂ reference sample (I_2 , filling 1.35 bar Kr + 0.65 bar He). This experimental setup allowed us to reference the energy to the edge of the reference sample and thus calibrate the energy for each spectrum.

SSNMR spectroscopy

SSNMR spectroscopy measurements were carried out on a Bruker Avance Neo spectrometer working at Larmor frequencies of 500.13, 470.59, and 125.77 MHz for ¹H, ¹⁹F, and ¹³C nuclei, respectively, equipped with a double-resonance cross polarization – magic angle spinning (CP-MAS) probe head accommodating rotors with an external diameter of 4 mm. MAS experiments were performed at a spinning frequency of 15 kHz. All spectra were acquired at room temperature using air as spinning gas. The chemical shift for all nuclei was referenced to the ¹³C signal of adamantane at 38.46 ppm. Further details on experimental settings for the acquisition of specific spectra are given in Section 8 of ESI.†

Evacuated and CO₂-loaded F4_MIL140A(Ce) samples for SSNMR measurements were prepared using a home-made cell provided with a mechanical lever operated from outside enabling the capping of the rotor without disturbing the cell atmosphere. For preparing the evacuated sample, the unsealed rotor containing the as-synthesised packed sample was heated overnight under vacuum (0.1 mbar) at the temperature of 423 K

within the cell and then sealed under N₂ atmosphere. For CO₂ loading, the home-built cell containing the evacuated sample packed into the rotor was loaded with CO₂ at 1 bar pressure and the rotor was sealed under the gas atmosphere.

Adsorption microcalorimetry

The molar differential heat of CO₂ adsorption was evaluated on F4_MIL-140A(Ce) by means of a heat flow microcalorimeter (Calvet C80 by Setaram) connected to a high-vacuum ($\approx 10^{-4}$ mbar) glass line equipped with a Varian Ceramicell 0-100 mbar gauge and a Leybold Ceramicell 0-1000 mbar gauge. The sample was held in vacuum at 393 K for 12 h before being placed into the calorimeter under isothermal conditions. The measurement was performed at 303 K by following a wellestablished step-by-step procedure described elsewhere.⁷⁷

Author contributions

M. C., V. C. and M. T. carried out the gas sorption experiments and data analysis. M. C., C. A. and V. C. carried out the IR and adsorption microcalorimetry experiments and data analysis. C. A., D. M. V., K. A. L. and M. T. carried out the EXAFS experiments. C. A. carried out EXAFS data analysis. M. S. performed the DFT calculations and data analysis. F. C., A. K. and M. T. carried out the PXRD experiments. M. T. carried out PXRD data analysis. L. C., F. M., A. G. and M. G. carried out the SSNMR experiments and data analysis. M. C., C. A., M. S., L. C., F. M., V. C. and M. T. drafted the manuscript. V. C. and M. T. coordinated the work. M. T. conceived the study.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

The in situ PXRD experiments were carried out at P02.1/ PETRAIII at DESY, a member of the Helmholtz Association (HGF) (Proposal I-20190224 EC). The research leading to this result has been supported by the project CALIPSOplus under the Grant Agreement 730872 from the EU Framework Programme for Research and Innovation HORIZON2020. The in situ XAS experiments were performed on beamline BM23 at the European Synchrotron Radiation Facility (ESRF), Grenoble, France. D. M. V and M. T. acknowledge the ESRF for covering the travel expenses for this beamtime. The authors acknowledge the European Union's Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie grant agreement no 663830 (M.T.). The authors acknowledge the University of Pisa for funding through the Progetti di Ricerca di Ateneo scheme (PRA2020_39) (M.T.). The authors thank the Italian MUR through the Project PRIN 2020 doMino (ref. 2020P9KBKZ) (M. S., F. C., L. C., V. C, M. T.). The authors acknowledge the CINECA award under the ISCRA initiative, for the availability of high performance computing resources and support, through the IsC92 project "ULTRADS" (M.S., M. C., C. A., V. C., M. T.).

The authors acknowledge Dr Alexander Schökel and Dr Sergej Wenz (Beamline P02.1 at PETRAIII, DESY) for technical assistance during the beamtime. The authors thank Prof. Silvia Bordiga (University of Torino) for facilitating this work in the early stages. The authors thank Dr Susanna Monti (ICCOM-CNR) and Dr Giovanni Barcaro (IPCF-CNR) for helpful discussion. Centro per l'Integrazione della Strumentazione Scientifica dell'Università di Pisa (CISUP) is acknowledged for the use of the Bruker Avance Neo 500 Solid State NMR Spectrometer.

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