

Understanding How Ligand Functionalization Influences CO₂ and N₂ Adsorption in a Sodalite Metal–Organic Framework

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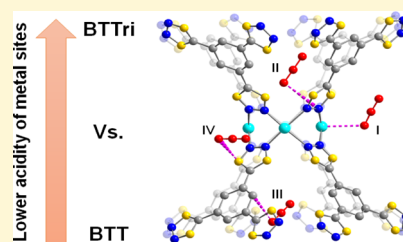


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Supporting Information

ABSTRACT: In this work, a detailed study is conducted to understand how ligand substitution influences the CO₂ and N₂ adsorption properties of two highly crystalline sodalite metal–organic frameworks (MOFs) known as Cu–BTT (BTT^{3−} = 1,3,5-benzenetristetrazolate) and Cu–BTTri (BTTri^{3−} = 1,3,5-benzenetristriazolate). The enthalpy of adsorption and observed adsorption capacities at a given pressure are significantly lower for Cu–BTTri compared to its tetrazole counterpart, Cu–BTT. In situ X-ray and neutron diffraction, which allow visualization of the CO₂ and N₂ binding sites on the internal surface of Cu–BTTri, provide insights into understanding the subtle differences. As expected, slightly elongated distances between the open Cu²⁺ sites and surface-bound CO₂ in Cu–BTTri can be explained by the fact that the triazolate ligand is a better electron donor than the tetrazolate. The more pronounced Jahn–Teller effect in Cu–BTTri leads to weaker guest binding. The results of the aforementioned structural analysis were complemented by the prediction of the binding energies at each CO₂ and N₂ adsorption site by density functional theory calculations. In addition, variable temperature in situ diffraction measurements shed light on the fine structural changes of the framework and CO₂ occupancies at different adsorption sites as a function of temperature. Finally, simulated breakthrough curves obtained for both sodalite MOFs demonstrate the materials' potential performance in dry postcombustion CO₂ capture. The simulation, which considers both framework uptake capacity and selectivity, predicts better separation performance for Cu–BTT. The information obtained in this work highlights how ligand substitution can influence adsorption properties and hence provides further insights into the material optimization for important separations.



INTRODUCTION

Since the industrial revolution, the combustion of carbon-based fuels has led to a significant rise in atmospheric CO₂ levels, one of the main causes of global warming.¹ During the last few years, the concentration of atmospheric CO₂ has surpassed a record value of 400 ppm.² This, combined with a slow transition to clean, renewable energy sources, has brought a sense of urgency toward the design of new porous solid adsorbents^{3–6} able to capture CO₂ from a variety of gas mixtures, such as postcombustion flue gas.⁷

One possible solution to this problem is a class of porous crystalline materials, known as metal–organic frameworks (MOFs), which consist of metal-ions or metal-ion clusters that are interlinked by organic ligands. These unique materials offer unprecedented internal surface area and easy chemical tunability, allowing chemists to readily adjust their adsorption properties.^{4,8–10} The presence of metal ions and multiple types of chemical moieties on the organic building blocks create a potential energy landscape with multiple minima for an incoming guest species; this corresponds to well-defined adsorption sites with varying binding energies.¹¹ Considering this, these materials offer a unique opportunity to gain direct insight into their structure-derived function using X-ray and neutron diffraction techniques. Diffraction data provide

binding mechanisms, reveal the relative differences in binding energies between neighboring adsorption sites, and deliver understanding of how the local framework structure changes as a function of external stimuli, such as temperature or pressure. Such insight makes it possible to pinpoint the structural features that give rise to enhanced or diminished adsorption properties. Further, diffraction data can be employed to validate computational methods meant to predict the structure and adsorption properties of MOFs.⁶ This validation process is necessary for the eventual deployment of simulation tools that can accurately predict the properties of hypothetical MOF structures, a process that might allow for the rapid identification of target materials having optimal properties for important gas separations, such as CO₂ capture.

MOFs bearing azole-containing ligands are becoming increasingly prevalent in the literature.^{12,13} Their popularity stems from strong metal–nitrogen bonds, a phenomenon that often leads to higher chemical and thermal stability when

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compared to their carboxylate counterparts.^{14,15} One prevalent example of a family of azole frameworks is M-BTT ($M = \text{Mg, Ca, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, and Zn}$; $\text{BTT}^{3-} = 1,3,5\text{-benzenetristetrazolate}$), which has the following general formula: $[(\text{M}_4\text{Cl}_x)_3(\text{BTT})_8]^{y-}$ ($x = 0 \text{ or } 1$; $y = 0 \text{ or } -3$).¹⁶ The MOF features truncated, octahedral cages constructed by six $[\text{M}_4\text{Cl}_x]^{(8-x)+}$ units and eight $[\text{BTT}]^{3-}$ ligands that are interlinked to form a porous, three-dimensional sodalite-based network. In addition to high stability, this family of MOFs possesses a high density of open metal coordination sites (OMCs), which are capable of forming strong interactions with small molecules such as CO_2 ; thus, the adsorption properties of the framework are readily modified by altering the metal identity. Given this, in previous work, we employed in situ diffraction techniques, combined with density functional theory (DFT) to rationalize how metal-substitution influences the CO_2 adsorption properties of several isostructural M-BTT ($M = \text{Fe, Mn, Cu, Cr}$) frameworks.¹⁶ In addition to varying metals, the tetrazole ligand can also be replaced with a triazole or pyrazole (BTT^{3-} vs $\text{BTTri}^{3-} = 1,3,5\text{-benzenetristriazole}$ or $\text{BTP}^{3-} = 1,3,5\text{-benzenetrispyrazolate}$) forming similar sodalite architectures;^{15–19} however, the resulting materials show dissimilar adsorption properties because of the distinct structural features that are induced by the ligand selection. Given that the aforementioned sodalite MOFs can undergo extensive chemical substitution and often exhibit exceptional crystallinity, they are of interest for in situ diffraction studies.¹⁵ We have previously shown that the substitution of the BTT^{3-} ligand with BTTri^{3-} causes a change in the structure of the metal cluster and the framework charge, and eliminates the existence of charge-balancing cations and anions in the channel of Cu-BTTri.²⁰ In the current work, we extend this study to unveil how ligand substitution influences the CO_2 and N_2 adsorption properties of Cu-BTTri compared to Cu-BTT. For this, both single-component N_2 and CO_2 adsorption isotherms and in situ diffraction techniques, coupled with DFT calculations are used. The diffraction results unveil the structures of the CO_2 and N_2 adsorbed Cu-BTTri framework, providing molecular level insight into how the presence of the triazole influences the CO_2 adsorption isotherms. The characterization of the structure and adsorption properties of the different sites are compared to DFT calculations used to predict the structure and binding energies of Cu-BTTri containing CO_2 . Finally, transient breakthrough simulations are used to provide insight into the performance of the Cu-BTT and Cu-BTTri for CO_2/N_2 separation under dry, postcombustion flue gas conditions. This work links the structural parameters of two Cu-containing frameworks to the CO_2 and N_2 adsorption properties, which influence the materials' overall performance for postcombustion flue gas separations. The combined experimental and computational effort allows us to rationalize the structure–property relationships in these two sodalite MOFs, work that can hopefully help inform the design of MOFs optimized for such separations.

■ EXPERIMENTAL SECTION

Synthesis of Cu-BTTri. The H_3BTTri ligand was first synthesized following a procedure that was reported previously in the literature.¹⁵ Subsequently, Cu-BTTri was synthesized, solvent exchanged and activated following a procedure reported previously by us.²⁰

Standard Adsorption Isotherm Measurements. For gas adsorption isotherms, UHP-grade (99.999% purity) helium, nitrogen,

and carbon dioxide were used for all measurements at pressures that ranged from 0 to 1 bar using a commercial BELSORP-max instrument. The sample was transferred to preweighed analysis tubes that were capped with a TRANSEAL. The sample was first evacuated on the activation station based on the program described in the literature.¹⁵ The evacuated analysis tube containing the sample was then carefully transferred to an electronic balance and reweighed to determine the mass of the activated sample (typically ranging 50–200 mg). Then, the tube was transferred to the BELSORP-max. N_2 adsorption isotherms were collected at 77 K and Brunauer–Emmett–Teller (BET) surface areas and pore volumes were calculated assuming a value of $16.2 \text{ \AA}^2$ for the molecular cross-sectional area of N_2 . In addition, N_2 adsorption isotherms were measured at temperatures ranging from 278 K to 298 K. For CO_2 , standard adsorption isotherms were measured using a commercial water bath dewar vessel connected to a Julabo F32-MC isothermal bath at temperatures ranging from 278 to 318 K.

In Situ Neutron Diffraction. High-resolution neutron powder diffraction (NPD) experiments were carried out on the Cu-BTTri sample using BT1 at the National Institute of Standards and Technology (NIST) Center for Neutron Research (NCNR). All measurements were carried out on activated samples of $\approx 0.8 \text{ g}$. At NIST, the sample was first activated via heating under dynamic vacuum at 450 K and then transferred into a He purged glove-box, loaded into a vanadium can equipped with a gas loading valve, and sealed using an indium O-ring. NPD data were collected using a Ge(311) monochromator with an in-pile 60 collimator corresponding to a wavelength of $2.0728 \text{ \AA}$. The sample was loaded onto a closed cycle refrigerator and then data were collected at 10 K. After data collection on the activated framework, CO_2 and N_2 with different dosing levels were then loaded into the pores. The dosing levels used for CO_2 include 0.40, 1.25, and 1.60 CO_2 per Cu^{2+} , whereas for N_2 the loading levels were 0.33 and 2.19 N_2 per Cu^{2+} . For data collection, the samples were first exposed to a predetermined amount of gas at room temperature. Upon reaching an equilibrium pressure, the sample was then slowly cooled (1 K per min) to ensure complete adsorption of the CO_2 and N_2 , and then data were recollected at 10 K. The resulting diffraction patterns were assessed via Le Bail and subsequent Rietveld analysis. To locate the CO_2 or N_2 molecules, Fourier difference maps were used to unveil excess scattering density in the MOF channels.

In Situ Synchrotron X-ray Diffraction. Bare and CO_2 -loaded Cu-BTTri samples were measured at the Swiss-Norwegian Beamline (SNBL, BM01) at the European Synchrotron Radiation Facility (ESRF) in Grenoble, France. A custom-built in situ diffraction powder cell was designed to mount directly on the goniometer head. Before the measurements, a sample of Cu-BTTri was activated at $180 \text{ }^\circ\text{C}$ for 24 h. Then, both the sample and the in situ powder cell were placed into an Ar-filled glove box. Afterwards, a capillary was loaded with 2 mg of activated sample that was attached directly to the in situ cell, and then sealed using a valve. The in situ cell was subsequently removed from the glove box and mounted on the goniometer head for data collection ranging from 190 to 290 K. An Oxford Cryostream 700 with working temperature of 80–500 K was used to maintain the desired temperature. Data were collected using a DECTRIS PILATUS 2M detector and the wavelength was adjusted at $0.67522 \text{ \AA}$. The sample to detector distance was set at 339 mm and the measurement time was varied between 20 and 40 s. After data collection of the bare framework from 190 to 290 K, the in situ cell was attached to a custom-built gas dosing manifold that is equipped with a turbo pump and pressure gauges. The Ar was evacuated from the cell, and next, the sample was dosed with 12.5 Torr of CO_2 , pressurized with He to the ambient pressure. The cooling rates in all experiments were 2 K per min . The azimuthal integration of raw images was performed with the Bubble software.²¹ Further, Le Bail analysis and Rietveld refinements of the acquired powder diffraction patterns were performed using TOPAS 5.²² The CIF files for the structure of the guest loaded frameworks based on the Rietveld analysis have been produced by Topas and uploaded to the CCDC database. The table of the atomic parameters corresponding to the

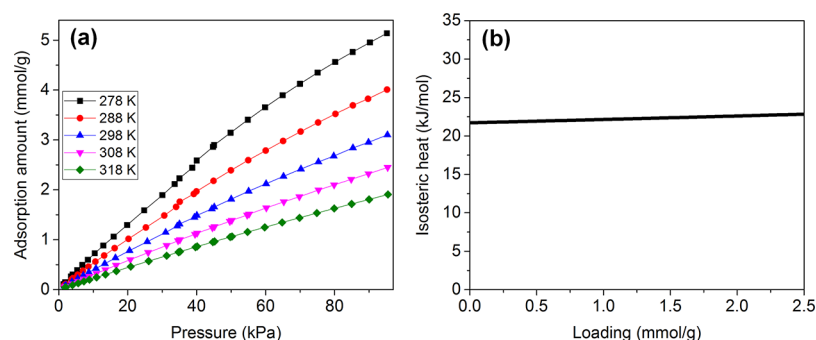


Figure 1. (a) Measured CO₂ adsorption isotherms for Cu–BTtri at different temperatures. (b) Calculated isosteric heat of CO₂ adsorption for Cu–BTtri.

bare and loaded structures has been also attached to the [Supporting Information](#). It should be noted that, for the gas-loaded structures, the standard deviation errors related to the occupancy of guest species have been produced based on the least squares matrix of Topas. Although this is the standard procedure for the calculation of the errors associated with the occupancies in Rietveld analysis, the actual errors can deviate from the reported values.

Sequential Le Bail fitting and Rietveld analysis was carried out on the variable temperature X-ray diffraction data that ranged from 190 to 290 K for the activated sample, and the samples dosed with CO₂ or N₂. After initial analysis on the data set collected at the lowest temperature (190 K), the crystal structure obtained from Rietveld refined was used as the starting model for subsequent, sequential Le Bail/Rietveld analysis.

Electronic Structure Simulation Details. Binding energies, structural details, and charge distribution were obtained from DFT calculations, under the generalized gradient approximation by Perdew, Burke, and Ernzerhof (PBE functional).²³ All calculations were performed by using the plane-wave self-consistent field package from the Quantum Espresso²⁴ suite of codes. We used the following ultrasoft pseudopotentials²⁵ extracted from <http://materialscloud.org/sssp>: Cu_pbe_v1.2.uspp.F.UPF, O_pbe_v1.2.uspp.F.UPF, C_pbe_v1.2.uspp.F.UPF, H_pbe-rrkjus_psl.0.1.UPF, and N_pbe.theo-s.UPF. The former three were generated using the Vanderbilt code²⁶ and the latter two using the “atomic” code by Dal Corso.²⁷ We employed kinetic energy cutoffs for wave functions and charge density and potential of 55 and 660 Ry, respectively. All calculations were performed for the Γ point of the Brillouin zone because of the large size of the MOF (228 atoms in the unit cell). Spin-polarized calculations were performed to take into account the most stable spin state for Cu–BTtri, which was found to be antiferromagnetic. Following previous work²⁸ where other chemically similar MOFs including Cu–BTT were studied, dispersion corrections were considered under the Grimme-D2 scheme.²⁹ We also tested the Tkatchenko–Scheffler³⁰ dispersion correction scheme, and the results were found to be in good agreement with those obtained by applying the Grimme-D2 method (see the [Supporting Information](#)).

We took as a starting point the experimental configuration for Cu–BTtri, and then allowed the nuclei to relax while keeping the cell parameters fixed to experimental values (tests allowing the cell parameters to change were also carried out and have shown very small changes of 1–3% in volume). Subsequently, several calculations were set, each of them containing one gas molecule adsorbed in a different site of the MOF. We took as initial configurations those found in the experiment and allowed atomic positions to relax, again without changing the cell parameters. Binding energies were computed as the difference between the energy of the MOF/gas binary complex and the sum of the energies of MOF and gas molecule.

RESULTS AND DISCUSSION

Characterization of the Cu–BTtri Framework. The Cu–BTtri framework was first synthesized using previously published procedures^{15,20} and subsequently characterized via

powder X-ray diffraction to confirm the sample purity. Synchrotron X-ray diffraction data ([Figure S1](#)) reveals Bragg scattering up to a of $42^\circ 2\theta$ (wavelength 0.67522 Å, ≈ 0.94 Å d -spacing), confirming that the sample has exceptional crystallinity, a prerequisite for in situ diffraction studies. The BET surface area and the pore volume for Cu–BTtri were calculated based on the nitrogen adsorption isotherm collected at 77 K ([Figure S2](#)) to be 1850 m²/g and 0.88 cm³/g, respectively, which are consistent with the originally reported values for this material.¹⁵

We recently determined that Cu–BTtri,²⁰ which crystallizes in a cubic $Fm\bar{3}c$ space group (no. 226), has the following molecular formula: Cu₃(BTtri)₂. The structure features square [Cu₄]⁸⁺ clusters that are interlinked by triangular [BTtri]^{3−} ligands to form a neutral sodalite-type framework. The structure does not fully replicate that of its ligand-substituted counterpart, Cu–BTT,³¹ or the other metal-substituted triazole analogs, such as Co–BTtri¹⁷ and Fe–BTtri.¹⁸ Unlike these other sodalite MOFs, the Cu–BTtri structure has no chlorine at the center of the cluster. As such, the framework is neutral and hence is devoid of any extra-framework charge-balancing species inside the channel. When compared to Co- or Fe–BTtri, the difference can be justified by the stronger Jahn–Teller distortion for octahedral copper(II) complexes, which results in the elongation of the bonds in the z -direction. Whereas in some cases this leads to the formation of distorted octahedral coordination environments,³² in others it leads to a loss of z -bound ligands^{33,34} as is the case of Cu–BTTri. When compared to Cu–BTT, the lower basicity of the tetrazolate compared to the triazolate likely helps to stabilize the presence of the Cl[−] ligand. The pK_a s of the triazole and tetrazole rings are approximately 13.93 and 8.23, respectively.³⁵ The higher basicity of BTtri^{3−} implies that it is also a stronger electron donor. Consequently, the Jahn–Teller effect is more pronounced for Cu–BTtri^{3−},³⁶ likely promoting the elimination of the chlorine in the center of square-like metal cluster. The higher ligand basicity not only influences the structural properties of the framework, but it also enhances the chemical and thermal stability of Cu–BTtri over Cu–BTT, a direct result of the higher bond strength found between the Cu and the triazolate.¹⁵

Standard Adsorption Isotherms and Isosteric Heat of Adsorption for CO₂ and N₂. To assess the adsorption properties of Cu–BTtri, single-component CO₂ adsorption isotherms were collected up to 1 bar and at temperatures ranging from 278 to 318 K, [Figure 1a](#). Interestingly, the isotherm does not exhibit a steep slope in the low-pressure regime, in contrast to what is normally observed for other

MOFs having OMCs, such as M–BTT frameworks (Figure S3).¹⁶ This is likely indicative of a low zero-coverage isosteric heat of CO₂ adsorption, which was extracted from the variable temperature data using a dual site Langmuir model, followed by fitting with the Clausius–Clapeyron equation (Figures 1b, S4, and S5). The zero-coverage isosteric heat was determined to be 21.7 kJ/mol for Cu–BTTri, which is significantly lower than the value previously reported for Cu–BTT, 30.7 kJ/mol.¹⁶ This observation is not surprising, considering that BTTri^{3−} is a stronger electron donor, which likely decreases the framework's affinity for CO₂ because of the lower Lewis acidity of the Cu²⁺. It is noted that, as the CO₂ loading increases in Cu–BTTri, the isosteric heat is relatively flat. This suggests that there are other, secondary adsorption sites, which have similar isosteric heats. Further, as the pressure is increased, the interactions between neighboring CO₂ could further promote an increase in the average binding energy at constant coverage.

Nitrogen adsorption isotherms were also collected for Cu–BTTri (Figure S6) at temperatures ranging from 278 to 298 K. As expected, the amount of N₂ adsorbed in the framework is significantly less than that of CO₂, because of the lower quadrupole moment and polarizability of the N₂ compared to CO₂.⁸ The weaker interaction of N₂ with the framework is further supported by the low zero-coverage isosteric heat of N₂ adsorption, which is 12 kJ/mol (Figure S7). Further, when comparing the N₂ adsorption properties of Cu–BTTri and Cu–BTT at 298 K (Figure S8), the triazole offers a lower total N₂ uptake. As such, the selectivity factors were calculated from isotherms collected at 298 K to be 27.8 and 20.8 for Cu–BTTri and Cu–BTT, respectively. The selectivity factor is the ratio of the quantity of CO₂ adsorbed at 0.15 bar/the quantity of N₂ adsorbed at 0.85 bar, normalized by the concentrations of both adsorbents. The chosen pressure points (0.15 bar for CO₂ and 0.85 bar for N₂) are those relevant to postcombustion flue gas capture. Whereas Cu–BTTri offers a higher CO₂/N₂ selectivity, Cu–BTT offers an overall higher CO₂ capacity, leaving questions as to which of these frameworks would effectively perform better in a postcombustion capture process.

Location of Adsorption Sites for CO₂ and N₂. To obtain molecular level insight into CO₂ and N₂ adsorption, neutron and synchrotron X-ray powder diffraction data were collected on the activated Cu–BTTri and then again on the sample dosed with various amounts of CO₂—0.40, 1.25, and 1.60 CO₂ per Cu²⁺. The location and orientation of CO₂ and N₂ molecules were determined via Rietveld analysis followed by subsequent Fourier difference analysis. A total of four CO₂ adsorption sites were observed inside Cu–BTTri, which are outlined in Figure 2. Fractional atomic coordinates, occupancies, and isotropic displacement parameters for all adsorption sites are reported in Tables S6–S8, and then a comparison of experimental and computationally obtained binding energies and shortest framework distances are shown in Table S3.

The first identified CO₂ adsorption site, I, is found as expected at the OMC, Cu²⁺. The occupancies of this adsorption site are 0.21(2), 0.74(2), and 0.82(2) for loadings of 0.40, 1.25, and 1.60 CO₂ per Cu²⁺, respectively. (CO₂)O⋯Cu²⁺ distances range from 2.7(1) to 2.96(8) Å, depending on the CO₂ loading level (Figure 2). This results in a distance of 3.63(7) Å between the nitrogen atom of the triazole ring and the carbon atom of the CO₂. For Cu–BTT, the distance of CO₂ to the OMC was previously reported by us to be 2.60(3) Å at a loading level of 1.56 CO₂ per Cu²⁺;¹⁶ this value is

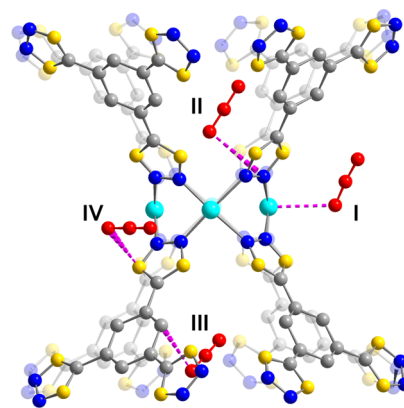


Figure 2. Ball and stick model of the Cu–BTTri framework dosed with 1.60 CO₂ per Cu²⁺. The adsorbed CO₂ molecules are shown in red. The Cu, C, N, are denoted as cyan, gray, and blue spheres, respectively. Hydrogen atoms are excluded for the sake of clarity. Yellow spheres represent mixed sites containing both C and N. The pink dotted lines represent nearest neighbor interactions ranging from ~2.8 to 3.3 Å.

considerably shorter than the corresponding distance in Cu–BTTri, a direct result of the weaker electrostatic interaction between the metal site and the adsorbed CO₂ molecules (Figure S9). This weaker interaction is due to the reduced bonding affinity of the Cu²⁺ sites in the axial direction as a result of the more pronounced Jahn–Teller effect in Cu–BTTri compared to Cu–BTT.^{28,37}

In order to obtain further insight into the mechanism of host–guest interactions in this framework, we took advantage of the theoretical support provided by DFT calculations. First, the bare Cu–BTTri and the CO₂ molecules were allowed to relax, whereas the cell parameters were kept constant (see the section on Electronic Structure Simulation Details and the Supporting Information). The distance between the OMC and CO₂ for the computationally simulated site I is 2.96 Å, which matches well with the experimentally obtained value (Table S3). Moreover, the energy of the resulting relaxed structures was computed by a single point calculation, and the binding energies were then extracted as follows (Table S3)

$$E_{\text{binding}} = E_{\text{Cu-BTTri+CO}_2} - E_{\text{Cu-BTTri}} - E_{\text{CO}_2}$$

The calculated binding energy of site I is estimated to be −25.0 kJ/mol, which matches well with the aforementioned measured zero-coverage isosteric heat of adsorption, 22 kJ/mol. To unveil the main interactions responsible for the adsorption of CO₂, the charge density difference induced by CO₂ binding was also computed. For this, the sum of the self-consistent densities for the empty Cu–BTTri framework and an isolated CO₂ molecule were first determined and then subsequently subtracted from the self-consistent density for the Cu–BTTri + CO₂ binary complex. Plots for two isosurfaces representing the charge density differences are shown in Figure S10. Based on the plot (Figure S10a) for site I, the main interactions responsible for the CO₂ binding predominantly originate from the OMC, whereas secondary interactions take place with the nitrogen atoms in the triazole ring. The charge density differences provide a qualitative indication of the perturbations induced by the CO₂ molecule on the framework, but they do not necessarily indicate the sign and magnitude of the interactions. Therefore, to better understand the mechanism of host–guest interactions at different adsorption

sites, decomposition of the binding energies into different contributions was carried out using the PBE functional; the results include electrostatics and short-range exchange–correlation effects. Further, the contributions dictated by van der Waals interactions were additionally computed using the Grimme-D2 method, Table S4. For the primary adsorption site, the binding energy at the DFT–PBE level is calculated to be -7.4 kJ/mol, whereas van der Waals (vdW) forces contribute a further -17.6 kJ/mol to the binding energy. Based on these results, it can be concluded that the van der Waals contribution is dominating. Most probably, the combination of the strong electron donating ligands to the Cu^{2+} OMC lends to a considerably reduced electrostatic affinity toward the guest species when compared to Cu–BTT.^{16,28} For other MOFs with OMCs, like HKUST-1 and MOF-74, contributions to CO_2 binding are mainly attributed to strong electrostatic interactions.³⁸ For Cu–BTTri, the highly electron donating ligand can reduce the charge, and hence electrostatic contribution at the OMC.²⁰ It is noted that whereas vdW interactions are the dominant force found in Cu–BTTri, electrostatic interactions, and other terms that are well-described at the DFT–PBE level, also play an important role in the overall binding energy.

The second CO_2 adsorption site, II, located just above the metal cluster (Figure 2), is disordered around the fourfold rotation axis that passes through the center of the square-like metal cluster; this static disorder is dictated by vdW interactions between this CO_2 and the ligand. The shortest distance is $3.27(4)$ Å, which is found between the oxygen atom of the CO_2 molecule and the nitrogen atom on the triazole ring. Within the error of the experimental data, the DFT-based simulations provide a secondary adsorption site that is nearly identical to site II. The shortest framework– CO_2 distance is approximately 3.16 Å. Like site I, site II can be observed in all refined structures regardless of the CO_2 loading. The occupation of this adsorption site ranges from $0.101(6)$ at the loading of 0.40 CO_2 per Cu^{2+} and reaches a value of $0.67(3)$ at the highest loading of 1.60 CO_2 per Cu^{2+} . The fact that the occupancies for site I and II are similar indicates that the enthalpy of adsorption for these sites are also comparable, a hypothesis that is further supported by the linear isosteric heat of CO_2 adsorption obtained from Cu–BTTri (Figure 1b). This is also confirmed by DFT calculations that yield a value of -26.5 kJ/mol for the isosteric heat of adsorption of site II, which is only 1.5 kJ/mol higher than that of site I (Table S3). The isosurface plot for site II confirms the interaction between the electron-rich triazole groups with the CO_2 molecule. Moreover, DFT calculations show that binding in this site is entirely due to strong vdW interactions ($E_{\text{dispersion-D2}} = -30.3$ kJ/mol), with a repulsive interaction between the host framework and the adsorbed CO_2 at the DFT–PBE level ($E_{\text{PBE}} = +3.8$ kJ/mol, see Table S4). This can probably be explained by electrostatics, because both moieties, including the triazole rings and oxygen atom of the carbon dioxide molecule, have a surplus of partial negative charge.

The third CO_2 adsorption site, site III, is located nearest to the benzene rings on the ligand (see Figure 2).¹⁶ The main mechanism for the stabilization of this adsorption site is through vdW interactions between the oxygen atom of the CO_2 with the carbon atoms of the benzene ring. The shortest framework distance for site III is $2.8(1)$ Å, which is similar to other vdW interactions found between CO_2 and other MOFs.^{39–41} DFT calculations have also successfully modeled

this adsorption site and predicted its binding energy to be -17.0 kJ/mol, which is lower than those calculated for site II (Table S4). This observed drop in binding energy is consistent with the observation that the relative occupancy of site III is significantly lower than sites I and II at all CO_2 loadings evaluated. For example, at the highest loading, 1.60 CO_2 per Cu^{2+} , the occupancy of site III is $0.124(8)$, whereas sites I and II are $0.82(2)$ and $0.67(3)$, respectively. Despite the lower predicted binding energy of site III, the simulated site III binding energy might not be precise, because the DFT calculations do not take intermolecular CO_2 – CO_2 interactions into account. The shortest contact distance for site III is approximately 3 Å from site II CO_2 and hence, this likely implies that the adsorption is further stabilized by intermolecular $\text{CO}_2(\text{II})$ – $\text{CO}_2(\text{III})$ interactions. The importance of these intermolecular interactions also explains the very low occupancy of site III at lower CO_2 loadings. It should be noted that, unlike sites I and II, there are more pronounced differences between the experimental and DFT-determined location of site III CO_2 (see Table S3), which can be attributed to the fact that only a single adsorbed CO_2 molecule is considered in the DFT calculations.

The fourth adsorption site, IV, was found inside the large pore of Cu–BTTri, at the same location where the extra cations and anions were previously observed in Cu–BTT (Figure 2).¹⁶ Site IV is in close proximity to the triazole ring, with the shortest framework– CO_2 distance equal to $3.02(9)$ Å. This adsorption site has additional interactions with other atoms of the triazole ring with distances that range from ~ 3 to 3.5 Å. The DFT-determined binding energy for site IV is -15.8 kJ/mol, almost entirely due to vdW interactions ($E_{\text{dispersion-D2}} = -15.8$ kJ/mol and $E_{\text{PBE}} = +0.05$ kJ/mol). The latter could result from a minimal interaction with the Cu^{2+} , consisted with a distance of approximately 4.6 Å between site IV and the OMC. This is no surprise given the long distance combined with the low Lewis acidity of Cu^{2+} in this structure.

In order to further understand the performance of the material for CO_2/N_2 separations in Cu–BTTri, the N_2 adsorption sites were also determined via Rietveld analysis of neutron diffraction data dosed with the 0.33 and 2.19 N_2 per Cu^{2+} . This analysis has led to the elucidation of six adsorption sites, I–VI, shown in Figure 3.

The first N_2 adsorption site is found at the OMC, with a Cu^{2+} – N_2 distance of $2.74(5)$ Å (Figure 3). The occupancy of site I N_2 is $0.145(7)$ and $0.326(8)$ at loadings of 0.33 and 2.19 N_2 per Cu^{2+} , respectively. The binding energy for this adsorption site is calculated to be -11.9 kJ/mol, which is in excellent agreement with the zero-coverage isosteric heat of adsorption ($Q_{\text{st}} = 11.9$ kJ/mol) obtained from variable temperature N_2 adsorption isotherms collected from 278 to 298 K. The contribution of vdW and DFT–PBE terms have been calculated to be -9.9 and -2.0 kJ/mol, respectively.

The energy decomposition based on DFT + vdW calculations can help understand better the role of the physicochemical properties of the adsorbate on the nature and extent of the host–guest interaction. Whereas the DFT–PBE baseline involves many other terms, for example, Pauli repulsion and exchange–correlation energy, it largely reflects electrostatic interactions between the adsorbate and the framework. For instance, in our previous work,²⁰ we demonstrated that DFT–PBE interactions between the OMC and D_2 molecule are responsible for less than 5% of the total binding energy, whereas in this work the share is

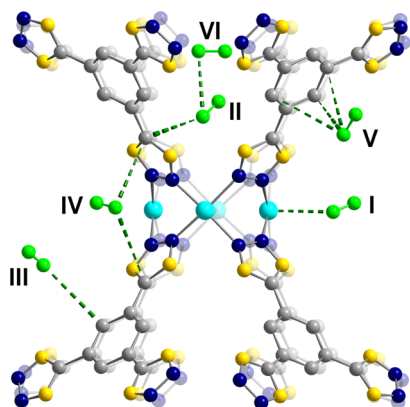


Figure 3. Ball and stick model of the Cu-BTtri framework dosed with 2.19 N₂/Cu²⁺. The adsorbed N₂ molecules are depicted in green. The Cu, C, N, are denoted as cyan, gray, and blue spheres, respectively. Hydrogen atoms are excluded for the sake of clarity. Yellow spheres represent atomic sites that are mixed with both C and N. The green dotted lines represent nearest neighbor interactions, and have distances ranging from 2.74(5) to 3.67(5) Å.

calculated to be approximately 17 and 31% for N₂ and CO₂, respectively (Tables S4 and S5). The electrostatic nature of these interactions is consistent with the relation between their magnitude and that of the adsorbate's permanent multipoles.²⁰ For example, CO₂, N₂, and H₂ have quadrupole moments of 43.0×10^{-27} , 15.2×10^{-27} , and 6.6×10^{-27} esu¹.cm⁻¹, respectively,⁸ that follow the same trend as the DFT-PBE binding energies of CO₂, N₂, and H₂, that is -7.41, -1.98, and -0.49 kJ/mol, respectively. For the gases with stronger permanent multipoles, tuning the charge in the framework wall can have a larger impact on the adsorption properties, whereas for those with weaker or no permanent multipole, the vdW interaction tends to dominate.²⁰ Understanding this relationship can provide insight into how to alter the structural features to optimize the adsorbate's physicochemical properties for different separations.

The second N₂ adsorption site, II, is located just above the Cu₄⁸⁺ cluster, albeit slightly farther away from the cluster than observed for CO₂ (Figure 3). Like I, site II N₂ can be detected in the data obtained from 0.33 and 2.19 N₂ per Cu²⁺ loadings with occupancies of 0.020(4) and 0.149(8), respectively. The shortest distance, 3.21(9) Å, is found between a framework carbon atom and site II N₂ (Figure 3). This interaction is predominantly vdW in nature, and this is further corroborated by the calculated shares of vdW and DFT-PBE interactions as well as the charge difference isosurface plot (Table S5 and Figure S11). The other four N₂ adsorption sites, III to VI, only appear in the diffraction data collected at the highest N₂ loading, 2.19 N₂ per Cu²⁺. These sites have occupancies ranging from 0.296(9) to 0.020(5) and all exhibit N₂-framework distances greater than 3 Å with DFT calculated binding energies that range from -4.5 to -7.1 kJ/mol. Whereas site VI is not found in close proximity to the framework wall, there are N₂-N₂ intermolecular interactions with site II, which likely help to stabilize the last adsorption site.

When comparing Cu-BTtri to Cu-BTT, the total N₂ uptake at room temperature of the former is significantly lower at pressures below 1 bar (Figure S8). Considering the low-pressure regime is typically dominated by the strongest adsorption sites, we surmise that the low N₂ adsorption of

Cu-BTtri stems from sites I and II. Obviously, the OMC in Cu-BTtri has a lower Lewis acidity reducing N₂ adsorption at low pressures. Moreover, for the second adsorption site, the triazole rings have a high partial negative charge that destabilizes N₂ adsorption (check the electrostatic contribution at Table S5). Given the higher basicity of the triazole ligand, it is hypothesized that such a destabilizing interaction would be sufficiently smaller for the tetrazole, making N₂ binding in site II stronger. Therefore, the presence of weaker primary and secondary binding sites in the triazole framework is an explanation for the material's significantly lower N₂ adsorption when compared to Cu-BTT.

Evolution of Lattice Parameters and Guest Occupancies as a Function of Temperature. In addition to CO₂ and N₂ binding, the bulk properties of the material, such as unit cell parameters and cell volume, were investigated as a function of temperature and gas loading level. This allows us to follow adsorption/desorption processes and better understand how the material's structure changes as a function of external stimuli. In this work, in situ synchrotron X-ray diffraction measurements were employed rather than neutron diffraction, in order to obtain rapid measurements using small sample sizes. First, data were collected at 100 K on a Cu-BTtri sample dosed with 12.5 Torr of CO₂ (1.09 CO₂/Cu²⁺) as it was necessary to determine if the position of CO₂ could be elucidated via synchrotron X-ray diffraction. Indeed, Rietveld analysis permitted the elucidation of the same four adsorption sites (Table S11) as observed via NPD. Afterward, variable temperature measurements were performed on the activated sample and then again on the aforementioned sample dosed with 12.5 Torr of CO₂ gas. For both samples, data were collected during cooling from 290 to 190 K (rate of 2 K per min). For the bare Cu-BTtri, Figure 4, the lattice parameter *a*

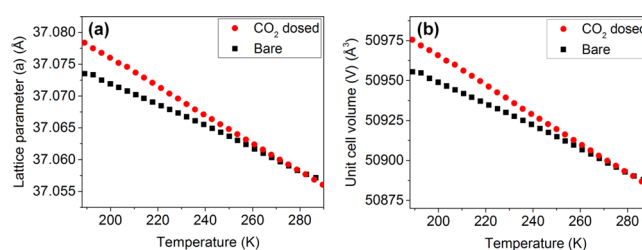


Figure 4. Change of (a) lattice parameter (a) (b) unit cell volume (V) of activated Cu-BTtri with temperature.

increases from 37.0571(1) to 37.0735(1) Å upon cooling, which leads to an increase in cell volume by 0.13%, representative of negative thermal expansion (NTE). The average thermal expansion coefficient, α , was calculated for the lattice parameter *a*, to be approximately -4.5 (MK)⁻¹. Albeit small, NTE was previously reported for only a few MOF structures to date.^{42–44} This magnitude of NTE behavior for this framework is similar to what was observed for HKUST-1, a well-known cubic MOF structure that has an average thermal expansion coefficient of -4.1 (MK)⁻¹.⁴⁴ Understanding the extent to which a material contracts or expands as a function of temperature is important to better understand how it might change throughout the course of a separation, a factor that can significantly impact process engineering. Interestingly, the NTE becomes even more pronounced when the sample is under 12.5 Torr of CO₂. Upon decreasing the temperature from 290 to 190 K, the lattice parameter, *a*, increases from

37.0560(1) to 37.0783(1) Å and results in a 0.18% change in the unit cell volume, this leads to a slightly larger thermal expansion coefficient, α , of -6.0 (MK)^{-1} (Figure 4). The more intense NTE in the presence of CO_2 gas can be attributed to the development of host–guest interactions inside the framework as the amount of loaded adsorbate increases upon decreasing temperature. Similar behavior was previously reported for $\text{Fe}_2(\text{dobdc})$ and $\text{Mg}_2(\text{dobdc})$,⁴⁵ where an increase in CO_2 dosing led to the increase in the degree of the lattice expansion for the framework.³⁷

Subsequent to the unit cell analysis, carried out via LeBail fitting, sequential Rietveld analysis was performed on the aforementioned variable temperature diffraction data obtained from Cu–BTTri dosed with 1.09 CO_2 per Cu^{2+} . The occupancies of sites I–IV are plotted as a function of temperature in Figure 5. Whereas the occupancies of site III

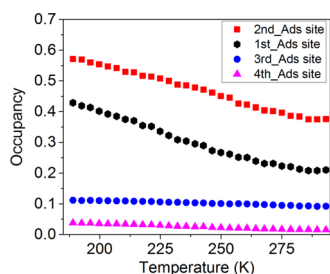


Figure 5. Occupation of different CO_2 adsorption sites in Cu–BTTri as a function of temperature.

and site IV are minimal over the entire temperature regime, the occupancies of sites I and II increase significantly as the temperature is decreased, which is due to the higher isosteric heats of CO_2 adsorption, as predicted by the van't Hoff equation.⁴⁶ Further, it is noted that at these temperatures, the second adsorption, site II, has a higher occupancy than site I, which is counterintuitive to the observations seen via NPD data collected at 10 K, which shows a higher occupancy of site I over site II. The occupancies are 0.21(1) and 0.38(1) at the highest temperature, 290 K, whereas the occupancies increase to 0.43(1) and 0.57(1), for sites I and II respectively, at 190 K. Whereas an energy difference as small as $\approx 3 \text{ kJ/mol}$ is sufficient to cause a sequential occupation of adsorption sites at 10 K, at higher temperatures, where adsorption isotherms are collected, it is possible that the distribution of the occupancies is also influenced by entropic effects.⁴⁷ Therefore, despite the higher occupancy of site II at room temperature, it is still assumed that site I is energetically favorable, and hence has a higher occupancy at temperatures approaching 0 K where entropy effects are highly attenuated.

Transient Breakthrough Simulations in Fixed Bed Adsorbers. Given that the single-component adsorption isotherms imply that Cu–BTTri has a higher selectivity for CO_2 over N_2 , whereas Cu–BTT has a higher CO_2 capacity, it was of interest to evaluate the potential of these frameworks in a dry postcombustion capture process (see scheme in Figure 6). For a clear comparison, Figure 7a shows the adsorption selectivities of Cu–BTT and Cu–BTTri over the whole pressure range and Figure 7b compares the volumetric CO_2 uptake capacities of both Cu–BTT and Cu–BTTri. We note that Cu–BTT has an uptake capacity that is significantly higher than Cu–BTTri by a factor of 2, despite the higher selectivity for the latter.

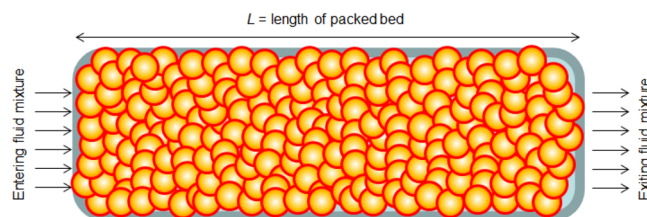


Figure 6. Schematic figure of a fixed bed adsorption bed for the postcombustion capture process.

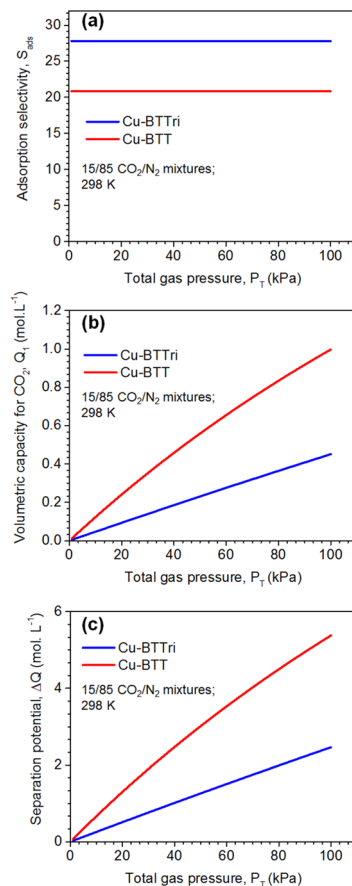


Figure 7. (a) Adsorption selectivity (S_{ads}), (b) volumetric capacity for CO_2 , Q_1 , and (c) separation potential, ΔQ , as a function of total pressure.

The typical composition of the flue gas feed is approximately 15/85 v/v CO_2/N_2 and the separations are normally carried out in fixed bed adsorbers (see schematic in Figure 6); such adsorbers are commonly operated in transient mode, and the compositions of the gas phase, as well as component loadings within the crystals, vary with position and time.^{48–50} During the initial stages of the transience, the pores are gradually loaded, and only toward the end of the adsorption cycle is pore saturation attained. For a given separation task, transient breakthroughs provide a more realistic evaluation of the efficacy of a material, as they reflect the combined influence of CO_2/N_2 adsorption selectivity and CO_2 uptake capacity.^{49,50}

The adsorption selectivity is defined by

$$S_{\text{ads}} = \frac{q_1/q_2}{y_1/y_2} \quad (1)$$

where the q_1 and q_2 represent the molar loadings of $\text{CO}_2(1)$ and $\text{N}_2(2)$ within the MOF that is in equilibrium with a bulk fluid mixture with mole fractions y_1 and $y_2 = 1 - y_1$. The molar loadings, also called gravimetric uptake capacities, are usually expressed with the units mol kg^{-1} . The component loadings are commonly calculated on the basis of unary isotherm data fits, along with the ideal adsorbed solution theory (IAST) of Myers and Prausnitz⁵¹ for mixture adsorption equilibrium (see the Supporting Information). The volumetric uptake capacities are

$$Q_1 = \rho q_1; \quad Q_2 = \rho q_2 \quad (2)$$

where ρ is the crystal framework density of the MOF, expressed in units of kg m^{-3} , or kg L^{-1} .

High uptake capacities are desirable because these result in longer breakthrough times and reduced frequency of bed regeneration. Higher values of S_{ads} are also desired because they lead to sharper breakthrough fronts and larger differences between the breakthrough times of individual constituents. If high product purities are desired, then this also demands $S_{\text{ads}} \gg 1$. Most commonly, however, high uptake capacities do not go hand-in-hand with high selectivities.^{30,52}

In order to overcome the selectivity/capacity trade-off for these two MOFs, we adopted an approach previously developed by Krishna,⁵² which defines a combined selectivity/capacity metric that is derived using the “shock wave” model for fixed bed adsorbers. The maximum achievable productivity of purified N_2 is realized when both intracrystalline diffusion and axial dispersion effects are completely absent and the concentrations “fronts” of the fluid mixture traverse the fixed bed in the form of shock waves.⁵² For the separation of $\text{CO}_2(1)/\text{N}_2(2)$ mixtures, the maximum achievable productivity, ΔQ , called the separation potential, can be calculated using the shock-wave model; the result is⁵²

$$\Delta Q = Q_1 \frac{y_2}{1 - y_2} - Q_2 \quad (3)$$

The quantity ΔQ can be described as an appropriate combination of selectivity and uptake capacity that is reflective of the separation potential for a specific adsorbent in a fixed bed.⁵² The ΔQ , expressed in moles of N_2 per L of adsorbent, represents the maximum amount of pure N_2 that can be recovered during the adsorption phase of the fixed bed separation and hence, is also indicative of the amount of CO_2 that is adsorbed on the bed until the point of breakthrough. Of course, it should be noted that if the purity of the CO_2 gas, which is released in the desorption stage, is of utmost importance, then the evaluation of the materials may be done based on the selectivity rather than separation potential.

A comparison of the separation potential, ΔQ , of Cu-BTT and Cu-BTTri for 15/85 $\text{CO}_2(1)/\text{N}_2(2)$ mixtures at 298 K, is plotted as a function of the total pressure, p_t , in Figure 7c. At 100 kPa total pressure, the ΔQ is 5.4 mol L^{-1} for Cu-BTT and 2.7 mol L^{-1} for Cu-BTTri. These values imply that Cu-BTT has the potential to produce twice the amount of pure N_2 as Cu-BTTri. In order to confirm the superior performance of Cu-BTT, transient breakthrough curves were also simulated for 15/85 CO_2/N_2 gas mixtures at an operating pressure of 100 kPa and 298 K using a methodology reported previously.^{49,50,52,53} For the breakthrough simulations, the following parameter values were used: length of packed bed, L

= 0.3 m; voidage of packed bed, $\varepsilon = 0.4$; superficial gas velocity at inlet, $u = 0.04 \text{ m/s}$.

The results for the transient breakthrough simulations obtained from a fixed bed packed with (a) Cu-BTT and (b) Cu-BTTri are shown in Figure 8. The transient

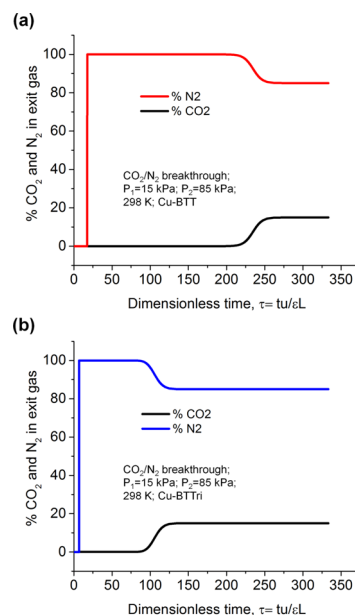


Figure 8. Transient CO_2 and N_2 breakthrough curves for (a) Cu-BTT and (b) Cu-BTTri frameworks.

breakthrough simulation results are presented in terms of a dimensionless time as the x -axis, τ , defined by dividing the actual time, t , by the characteristic time, $L\varepsilon/u$. The y -axis represents the % CO_2 and % N_2 in the gas mixture exiting the adsorber. The time at which CO_2 “breaks through” is significantly higher for Cu-BTT, as compared to Cu-BTTri; this can be primarily ascribed to the higher CO_2 capacity of Cu-BTT, as shown in Figure 7b. During the time interval indicated by $\Delta\tau$ in Figure 8, nearly pure N_2 can be produced. Note that the time interval $\Delta\tau$ is higher for Cu-BTT than that of Cu-BTTri by nearly a factor of 2. Arbitrarily setting the minimum purity requirements for N_2 as 99.95%, we can determine the productivity of N_2 from a material balance on the adsorber. The calculated values for Cu-BTT and Cu-BTTri are, respectively, 4.3 and 1.8 mol L^{-1} , confirming that the separations with Cu-BTT yield a significantly higher productivity of pure N_2 and thus a higher amount of CO_2 can be captured. The actual values of the productivities for Cu-BTT and Cu-BTTri are lower than the corresponding values of ΔQ , calculated using the IAST (see Figure 7c), because of the distended nature of the transient breakthroughs in Figure 8.

Overall, neither the adsorption capacity of the single gases nor the selectivity, S_{ads} , is a proper metric for judging the effectiveness of a specific MOF for separations in fixed bed adsorbers. Separation potential can give a better estimation of how effective a material’s performance is. However, depending on the particular requirements for each case, such as CO_2 recovery, CO_2 purity, or the effective amount of gas captured per mass unit of the material, the final choice might be different. For example, in the present case, for optimizing CO_2 recovery, Cu-BTT is a better option because it yields higher

amount of cyclic CO₂ adsorption because of its higher separation potential. However, if the CO₂ purity is the main target, Cu-BTTr, which shows better CO₂/N₂ selectivity, would be a better choice for this separation.

CONCLUSIONS

This work presents a comprehensive study of CO₂ and N₂ adsorption in a sodalite MOF known as Cu-BTTr; this was carried out using a series of characterization tools including single-component adsorption measurements, in situ neutron and X-ray diffraction, DFT calculations, and finally breakthrough simulations. Whereas CO₂ and N₂ isotherms are used to assess selectivity and adsorption capacity, in situ diffraction is used to provide molecular level insight into the location and orientation of these two small molecules throughout the framework. As such, the diffraction technique allows further rationalization of the observed isotherms, provides a means to monitor adsorption/desorption processes, and allows one to assess changes in the framework as a function of temperature. Albeit small, both the activated and CO₂-loaded Cu-BTTr are found to exhibit a rare NTE phenomenon, with average thermal expansion coefficients of -4.5 and -6.0 (MK)⁻¹, respectively. In addition, DFT calculations are used to predict the binding energies and geometries of potential adsorption sites, results which are found to be in good agreement with those determined experimentally. To better understand the host–guest interaction at each adsorption site, the calculated binding energies were further decomposed into separate contributions from DFT–PBE (predominantly electrostatic) and van der Waals interactions.

Finally, the performance of Cu-BTTr is compared to a ligand-substituted counterpart, Cu-BTT. While the sodalite structures are similar, the ligand exchange has an extreme effect on the performance of the material with regard to CO₂ and N₂ adsorption. Despite that Cu-BTTr exhibits a lower overall adsorption capacity and weaker interactions with both CO₂ and N₂, the material surprisingly offers a higher selectivity for CO₂ over N₂ compared to that of its tetrazole counterpart. These results stem from the fact that the triazolate offers a stronger Cu–ligand bond compared to the tetrazolate, and hence, significantly weakens the interactions along the *z*-axis, where small molecules bind. This assumption is validated by neutron diffraction and adsorption measurements, which show elongated Cu–CO₂ distances and lower initial isosteric heats of CO₂ adsorption, respectively. We finally compared these materials' performance in a dry, postcombustion flue gas separation. For this, the separation potential, ΔQ , of Cu-BTT and Cu-BTTr was determined, and transient breakthrough curves were simulated; it was concluded that, despite its higher selectivity for CO₂ over N₂, Cu-BTTr does not outperform Cu-BTT for the separation of interest. It was demonstrated that adsorption capacity and selectivity alone cannot be appropriate metrics to evaluate the efficiency of an adsorbent to be employed in a chosen separation.

It should be reiterated that this study is based on the thermodynamic and hydrodynamic properties of the frameworks of interest in dry postcombustion capture conditions. For the actual application of a material in such a large-scale separation, other techno-economic factors like price, long-term stability/cyclability, ease of regeneration, temperature spikes during adsorption, and the influence of water and minor impurities, like SO_x and NO_x, must also be considered.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.chemmater.9b04631>.

Experimental neutron and synchrotron powder X-ray diffraction patterns, calculated diffraction patterns obtained by Rietveld analysis and Fourier difference patterns, nitrogen adsorption measurements, BET surface area calculations, carbon dioxide measurements, fit of experimental gas adsorption data by Langmuir and dual-site Langmuir model, experimental and computational CO₂ and N₂ gas adsorption measurements, experimental and computational shortest framework distance for different individual adsorption sites, computational decomposition of electrostatic and vdW interactions, and isosurfaces of the charge density plots (PDF)

CuBTTr_NPD_0_5_CO2_10K crystallographic data (CIF)

CuBTTr_NPD_0_5_N2_10K crystallographic data (CIF)

CuBTTr_NPD_1_5_CO2_10K crystallographic data (CIF)

CuBTTr_NPD_2_5_CO2_10K crystallographic data (CIF)

CuBTTr_NPD_1_5_N2_10K crystallographic data (CIF)

CuBTTr_1_CO2_Cu_synchrotron crystallographic data (CIF)

CO2_SITE1_DFT data (XYZ)

CO2_SITE2_DFT data (XYZ)

CO2_SITE3_DFT data (XYZ)

CO2_SITE4_DFT data (XYZ)

N2_SITE1_DFT data (XYZ)

N2_SITE2_DFT data (XYZ)

N2_SITE3_DFT data (XYZ)

N2_SITE4_DFT data (XYZ)

N2_SITE5_DFT data (XYZ)

N2_SITE6_DFT data (XYZ)

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Notes

The authors declare no competing financial interest.

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