Carbon Dioxide Capture Chemistry of Amino Acid Functionalized Metal–Organic Frameworks in Humid Flue Gas

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ABSTRACT: Metal–organic framework-808 has been functionalized with 11 amino acids (AA) to produce a series of MOF-808-AA structures. The adsorption of CO₂ under flue gas conditions revealed that glycine- and dl-lysine-functionalized MOF-808 (MOF-808-Gly and -DL-Lys) have the highest uptake capacities. Enhanced CO₂ capture performance in the presence of water was observed and studied by using single-component sorption isotherms, CO₂/H₂O binary isotherm, and dynamic breakthrough measurements. The key to the favorable performance was uncovered by deciphering the mechanism of CO₂ capture in the pores and attributed to the formation of bicarbonate as evidenced by ¹³C and ¹⁵N solid-state nuclear magnetic resonance spectroscopy studies. On the basis of these results, we examined the performance of MOF-808-Gly in simulated coal flue gas conditions and found that it is possible to capture and release CO₂ by vacuum swing adsorption. MOF-808-Gly was cycled at least 80 times with full retention of performance. This study significantly advances our understanding of CO₂ chemistry in MOFs by revealing how strongly bound amine moieties to the MOF backbone create the chemistry and environment within the pores, leading to the binding and release of CO₂ under mild conditions without application of heat.

INTRODUCTION

The carbon dioxide capture problem is of paramount societal importance. Although aqueous amine solutions are currently being used to capture CO₂, they require significant energy input because of the high heat capacity of water and suffer from loss of amines. Porous solid-state materials are being studied for their potential in offering a platform for mounting amines and forego the heat requirement of aqueous solutions. In this regard, materials such as carbon, zeolites, silica, covalent organic frameworks, and metal–organic frameworks (MOFs), are being investigated (as summarized in Table 1; see Table S1 in the Supporting Information for a more detailed comparison). At present, they all fall short of meeting the stringent performance requirements: high uptake capacity, high selectivity, low regeneration energy, fast kinetics, and long cycling lifetime. We believe that to solve the CO₂ problem, significant efforts are required in understanding the CO₂ chemistry in the pores of such materials and in using this knowledge to build viable structures.

Along these lines, we focus on MOFs and report a series of 11 amino acid functionalized MOF-808 structures (MOF-808-AA), in which the amino acid moieties are strongly bound to the backbone. We find that this construct creates an environment inside the pores where CO₂ uptake under humid conditions leads to the formation of bicarbonates and therefore requiring no application of heat to regenerate the material. In this way, application of rough vacuum is sufficient to remove the CO₂ and obviate the need for heating customarily used in aqueous amines and even some solid sorbents.

We and others have appended amines as coordinatively or covalently bound functionalities into MOFs. Specifically, MOF-74 has been used in obtaining high capacity for CO₂ uptake. In Mg-MOF-74 [termed Mg₆(dobdc), dobdc = 2,5-dioxido-1,4-benzenedicarboxylate], the amines are bound directly to the Mg²⁺ to make a weak bonding interaction. Although this interaction allows for high CO₂ uptake mechanism, it is the Achilles’ heel of the material as it results in (1) ease of hydrolysis of the MOF backbone and (2) competition of water to CO₂ for Mg²⁺ sites and therefore the loss of amines. Furthermore, the Mg-MOF-74 system and their derivatives require heating to 140–240 °C for regeneration. Here, we use a robust MOF-808 [Zr₆O₂(OH)₄-(BTC)₂(HCOO)]₆⁶, BTC = 1,3,5-benzenetricarboxylate] sys-

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tem38 and bind the amino acids through their carboxylates to Zr(IV) centers to create Zr(IV)–carboxylate interactions, which are 2–3 times stronger than the Mg(II)–amine interactions.39−41 The fact that the amino residue of the amino acid is dangling into the pores creates a basic environment for CO2 binding in the presence of water. This structural design prevents water or CO2 from interfering with the bond between Zr(IV) and the carboxylate of the amino acids.

In this study, we used MOF-808 as the basis for generating the amino acid functionalized forms. We measured and confirmed their enhanced CO2 capture performance in the presence of water through sorption studies. These involved single-component sorption isotherms, CO2/H2O binary sorption Table 1. Overview of CO2 Capture Properties for State-of-the-Art Sorbents

| sorbent classes | CO2 uptakea dry/humid | binding mechanism | regeneration conditions | operational challengesb |
|----------------|------------------------|-------------------|-------------------------|-------------------------|
| physisorbents  |                        |                   |                         |                         |
| carbon         | L/L (1)                | physisorption     | 25–100 °C/vacuum        | low cyclability         |
| zeolites       | M to H/none (1)        | polar interaction | 120–250 °C/vacuum       | sensitivity to humidity, high regeneration T |
| MOFs           | H/L (1)                | open metal sites  | 100–200 °C              | hydrolysis, low uptake in humidity, high regeneration T |
|                | H/H (1)                | polar interaction | 150–350 °C/vacuum       | hydrolysis, high regeneration T |
| chemisorbents  |                        |                   |                         |                         |
| hydrides       | H                      | carbonate         | 400–1000 °C             | solid management, high regeneration T |
| carbonates     | M                      | bicarbonate       | 150–350 °C              | pore blockage, high regeneration T |
| amine liquids  | M to H/M to H (†)      | carbamate−bicarbonate | 80–150 °C             | amine loss, amine oxidation |
| amino acid salts | M/M (†)   | carbamate−bicarbonate | 120 °C                     | high regeneration T |
| aminosilicones | M to H/M to H (†)      | carbamate−bicarbonate | 110 °C                     | capacity loss (urea formation) |
| grafted amines | M to H/M to H (†)      | carbamate−bicarbonate | 100–180 °C             | hydrolysis, high regeneration T, amine loss |
| MOFs, functionalized | H/H († or 1) | metal carbamate−bicarbonate | 70–140 °C/vacuum | hydrolysis, high regeneration T |
|                | M/M (†)                | carbamate−bicarbonate | 70–140 °C/vacuum | hydrolysis, high regeneration T |

*Qualitative ranges of general uptake values indicated as high (H), medium (M), or low (L) with an arrow to signify whether the uptake increases (†) or decreases (↓) with humidity using flue gas-relevant conditions. See Table S1 and corresponding references for further details. bT, temperature.

Figure 1. (a) Structure of MOF-808-AA and (b) general synthetic scheme for MOF-808-AA from crystallization of MOF-808 to postsynthetic formate removal and amino acid loading, represented with zoom-in views of the Zr SBUs to highlight the changes of coordination in each step. Structural schemes of the coordinatively loaded amino acids are provided in (c) corresponding to abbreviations in Table 2. Atomic models of MOF-808-Gly and its SBU with a loading number N = 6 are used to represent the general structure of MOF-808-AA. Zr atoms are represented as light blue polyhedra, while other atoms are represented as spheres (color code: C, gray; O, red; N, green). H atoms are omitted, and the yellow spheres highlight the small tetrahedral pores for clarity.
and this led us to apply the CO₂ capture and release process in NMR measurements of the MOFs fully hydrolyzed into their hydrochloric acid (HCl) according to reported procedures.38,44 Acid-loaded MOFs was determined through liquid-state 1H spectroscopy. The bicarbonate formation is further corroborated by evidence from solid-state nuclear magnetic resonance (NMR) measurements of MOF-808-AAs with a general formula of [Zr₆O₄(OH)₄(BTC)₄(RS₃-Abu)]ₐ, which has 27 exchangeable Zr-binding sites (excluding the OH-bridging sites). This number of exchangeable sites is nearly matched by the CO₂ uptakes obtained from single-component adsorption measurements, which is consistent with the CO₂ uptake extracted from single-component isotherms at 25 °C, unless specified otherwise. CO₂ uptake in the presence of water extracted from CO₂/H₂O binary adsorption isotherm at RH ~10% (3.6 mmol g⁻¹ loading of water).1 mol L⁻¹ aqueous solutions instead of saturated solutions.

### Results and Discussion

**Synthesis of MOF-808-AA.** MOF-808-AA (Figure 1a) was obtained through the postsynthetic incorporation of amino acids into MOF-808, for which the formate ligands have been removed to give what is termed MOF-808-FR. Typically, MOF-808 was first synthesized, and the formate ligands were removed by treatment with hydrochloric acid (HCl) according to reported procedures.38,44 The product was then immersed in a saturated aqueous or dimethyl sulfoxide (DMSO) solution of an amino acid at a given temperature for 3 days (Figure 1b, Table 2, and section S3), during which exchange of fresh solutions was performed three times per day. The solid was then washed with deionized water three times per day, followed by treatment with pH 10 buffer solutions of the amino acid or a 10% 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) solution in tetrahydrofuran for 1 day, and further washed with acetone for an additional day before activation under dynamic vacuum at 140 °C. In this work, 11 amino acids (Figure 1c) were loaded into MOF-808-FR to yield a series of MOF-808-AAs with a general empirical formula of [Zr₆O₄(OH)₄(BTC)₂-(RS₃-Abu)]₂₋(AA)ₓ(OH)₆-x(NH₃)ₓ(H₂O)y [AA = amino acid anion], where loading number N denotes the molar equivalence of amino acids per Zr₆O₄(OH)₄ cluster (or secondary building unit, SBU).

For easy reference, the names and abbreviations of the amino acids and relevant information about the resulting MOF-808-AAs are summarized in Table 2. The composition of the amino acid-loaded MOFs was determined through liquid-state ¹H NMR measurements of the MOFs fully hydrolyzed into their components with hydrofluoric acid (HF) and deuterium oxide (D₂O) (termed “digest NMR”, section S5). To determine the loading of amino acids, the integrations of amino acid signals were compared to that of the BTC linker, thereby giving the ratio between amino acids and the linker. Fourier-transform infrared spectroscopy (FT-IR, section S6), powder X-ray diffraction (PXRD, section S5), scanning electron microscopy (SEM, section S7), and N₂ as well as CO₂ sorption studies were performed to further characterize the samples.

**Structure of MOF-808-AA.** The structure of MOF-808 is composed of the SBUs Zr₆O₄(OH)₄(−COO⁻)(HCOO)₆, which are connected to each other through 1,3,5-benzene-tricarboxylate linkers, yielding a porous, extended, three-dimensional framework (Figure 1a). Each of the SBUs is composed of six Zr₆ octagonal antiprisms linked by sharing tetragonal caps through μ₃-O atoms. On the periphery of the SBU are a total of 12 exchangeable coordination sites bridging adjacent Zr atoms. In the precursor framework, MOF-808, these sites are occupied by formate (HCOO⁻) ligands in bidentate or monodentate binding mode, while in MOF-808-FR, the sites are most likely occupied by H₂O and OH⁻ for charge balance (section S3). The loading of amino acids involves binding of their carboxylate (−COO⁻) groups to these sites to form the desired derivatives used in this work (Figure 1a). The extended linking of Zr SBUs and BTC linkers form two kinds of interconnected pores: small, tetrahedral pores of 1.2 Å in diameter that are inaccessible to guest molecules and large, adamantane-shaped pores of 18 Å in diameter that are highly interconnected through apertures of 14 Å in diameter.44 The installation of amino acid ligands allows the amino groups (−NH₂ or −NH⁻) on the alkyl chains to point to the inside of the pores, which serve as the primary sites to interact with and thus capture CO₂ entering the pores from the exterior.

To validate this, single-crystal samples of a representative variant, MOF-808-Gly [Zr₆O₄(OH)₄(BTC)₂(Gly)₇(NH₃)ₓ(H₂O)y] [Gly = NH₂CH₂COO⁻,], were prepared by using a similar procedure as for the microcrystalline samples, with the acetone washing steps specifically avoided to eliminate possible assignment difficulties during the structural refinement. The 20−40 μm sized single crystals of MOF-808 were prepared according to the reported procedure,38 which were further treated with 1 mol L⁻¹ HCl solution to remove formate ligands.

| amino acid | abbrev | pKₐ 35,46 | loading (N) | temp (°C) | solvent | amine loading (mmol g⁻¹) | uptake (mmol g⁻¹, 25 °C)² |
|------------|--------|-----------|------------|----------|---------|-----------------|------------------------|
| glycine    | Gly    | 9.60      | 5.89       | 85       | H₂O     | 3.84            | 0.247                  |
| sarcosine  | Sar    | 10.35     | 1.71       | 85       | H₂O     | 1.23            | 0.239                  |
| l-alanine  | l-Ala  | 9.69      | 3.55       | 100      | H₂O     | 2.38            | 0.250                  |
| DL-alanine | DL-Ala | 9.69      | 2.98       | 100      | H₂O     | 2.04            | 0.213                  |
| (R)-3-aminobutanoic acid | R-3-Abu | 10.53 | 2.12 | 85 | H₂O⁴⁻ | 1.47 | 0.132 |
| (RS)-3-aminobutanoic acid | RS-3-Abu | 10.53 | 2.10 | 85 | H₂O⁴⁻ | 1.45 | 0.189 |
| l-isoleucine | l-Ile | 9.60 | 1.44 | 85 | H₂O⁴⁻ | 1.00 | 0.203 |
| l-serine  | l-Ser  | 9.15      | 3.61       | 85       | H₂O     | 2.32            | 0.248                  |
| l-threonine | l-Thr | 9.10      | 4.48       | 100      | H₂O     | 2.67            | 0.192                  |
| l-histidine | l-His | 9.17      | 4.03       | 85       | H₂O     | 2.26            | 0.108                  |
| DL-lysine | DL-Lys | 8.95, 10.53 | 1.87   | 120      | DMSO    | 2.48            | 0.705                  |

CO₂ uptake extracted from single-component isotherm at 25 °C, unless specified otherwise. CO₂ uptake in the presence of water extracted from CO₂/H₂O binary adsorption isotherm at RH ~10% (3.6 mmol g⁻¹ loading of water).
Glycine was incorporated with retention of morphology and crystallinity of the crystals, which were found suitable for single-crystal X-ray diffraction (SXRD) studies. Fully activated crystals were used for collection of SXRD data with synchrotron radiation at Beamline 12.2.1 of the Advanced Light Source (ALS) at Lawrence Berkeley National Laboratory at 100 K by using a nitrogen cold stream. The structure of MOF-808-Gly was solved in the cubic space group $Fd\bar{3}m$, with the unit cell parameter of $a = 35.1413(12)\ \text{Å}$ (section S4). The extended backbone of the MOF was well-resolved to show the same structure as in MOF-808, and relatively small thermal displacement parameters, indicating the retention of crystallinity during the postsynthetic modification (Figure S14).

An average of 2.9 glycinate ligands, as determined by the occupancies of the $\alpha$-C and N atoms in the SXRD result, were found bound to each Zr SBU in a bidentate bridging fashion between two Zr atoms. The amino N atom was found in significant disorder, which was largely attributed to the free rotation of the C–C bond in the glycinate ligand, indicating no significant interaction with other parts of the MOF (Figure 2).

![Figure 2. Single-crystal structure of a segment of MOF-808-Gly focused on one Zr$_6$ SBU, highlighting the bridging binding mode of glycinate to adjacent Zr atoms. Thermal ellipsoids of atoms are shown at the 50% probability level. Color code: C, gray; O, red; N, green; Zr, blue. For clarity, H atoms, disordered $\mu_1$-O atoms, and $\text{OH}^-$/H$_2$O groups are omitted. One randomly selected conformation among the refined rotationally disordered positions of each $=\text{NH}_2$ group is shown to help clarify the spatial arrangement of the amino acid residues in MOF-808-Gly.](https://pubs.acs.org/doi/10.1021/jacs.1c13368)

For the bulk sample, the retention of crystallinity was first established through PXRD measurements (section S5). The composition was then characterized with FT-IR measurements (section S6). Compared to MOF-808, nearly all formate ligands were absent in MOF-808-FR, as was evidenced by the absence of absorbance bands at 1582 and 1337 cm$^{-1}$ which have been assigned as $=\text{COO}$ stretches of the formate ligands. Furthermore, it was determined that >98.5% of the formate ligands were removed based on digest NMR results on MOF-808-FR. The incorporation of glycine molecules into MOF-808-FR was first probed by the appearance of FT-IR absorbance bands at ~2960 cm$^{-1}$ assigned as the C–H stretch of glycinate ligands present in the product MOF-808-Gly. Representative scanning electron micrographs of MOF-808 and MOF-808-Gly showed that the octahedral microcrystals of the MOF remained unaltered before and after the reactions (Figures S20 and S21). Energy dispersive X-ray spectroscopy (EDX) on a set of randomly sampled microcrystals indicated that no remaining chlorine (Cl) was found after the last step of the coordinative functionalization (Figure S22).

**N$_2$ Sorption Isotherm Studies.** N$_2$ sorption isotherms were measured for MOF-808, MOF-808-FR, and MOF-808-AAs at 77 K (Figure 3a and section S8.1) to confirm their permanent porosity. Comparison among MOF-808, MOF-808-FR, and the representative MOF-808-Gly (Figure 3a) showed that the formate removal step increased the porosity, as indicated by the change of the Brunauer–Emmett–Teller (BET) surface area from 1680 to 1971 m$^2$ g$^{-1}$ (Figures S23 and S24). By contrast, the amino acid loading reaction generally reduced the porosity. This was exemplified by MOF-808-Gly, whose BET surface area was reduced to 1427 m$^2$ g$^{-1}$ (Figure S25).

It is likely that besides contributing to the total mass of the MOF, the residues of the amino acids occupy space in the pores, thus considerably reducing the surface area. Evidence from the pore size distribution data (Figures S26–S32) derived from N$_2$ sorption isotherms further indicates that the pore structures were retained in the intermediate and the final products. As expected, the amine ligands were introduced without altering the periodic porous MOF structures. N$_2$ isotherms and BET surface areas of other MOF-808-AAs are given in Figures S33–S41.

**Single-Component CO$_2$ Sorption Isotherms.** Single-component CO$_2$ adsorption and desorption isotherms were measured at 25 °C (~298 K) for all obtained MOF-808-AA variants and their precursor frameworks (Figures 3b,c and section S8.2.1). Comparison of the isotherms of MOF-808, MOF-808-FR, and MOF-808-Gly showed that the latter has increased uptake at carbon capture-relevant pressures (4 and 15 kPa, Table 2). In addition, a significant hysteresis is observed between the adsorption branch and desorption branch, characteristic of CO$_2$ chemisorption, which is not fully reversible at 25 °C. This phenomenon was generally observed for other MOF-808-AA variants, while absent for MOF-808 or MOF-808-FR. This eliminates the possibility of having any unoccupied sites (open metal sites) in the MOF backbone contributing to the observed strong adsorption.

Comparison of CO$_2$ isotherms at 25 °C among the MOF-808-AAs (Figure 3c, Figures S43–S53, and Table 2) yielded a range of CO$_2$ uptake performances, varying between 0.092 and 0.705 mmol g$^{-1}$ at 4 kPa (4% CO$_2$ in 1 atm gas mixture, relevant to natural gas combined cycle flue gas capture) and between 0.277...
We found that multiple factors influence the CO2 uptake: total amine loading, pK_a of the loaded amine species, steric hindrance at the local environment in the pore, and proximity of adjacent functional groups. Most significantly, a larger gravimetric amine loading and a higher pK_a of the loaded amine species are correlated to a higher CO2 uptake. The incorporation of DL-
lysine into MOF-808-FR, despite a relatively low amino acid loading number N of 1.87, yielded a gravimetric amine loading of 2.48 mmol g$^{-1}$, third highest among the series due to the 2-fold amine stoichiometry per amino acid loading. The pK$a$ of the ε-NH$_2$ is 10.53, highest among the amino species, ensuring a high affinity even at a low CO$_2$ pressure. These factors combined resulted in the highest CO$_2$ uptake observed for MOF-808-dl-Lys among all the MOF-808-AA sorbents at both CO$_2$ levels of interest.

To quantitatively evaluate the sorbents’ affinity for CO$_2$, the isosteric heat of adsorption (Q$_s$) was determined by measuring CO$_2$ sorption isotherms at 15 °C ($\sim$288 K), 25 °C, and 35 °C ($\sim$308 K) for selected MOF samples (section S8.2.2). The Q$_s$ value of MOF-808-dl-Lys at zero loading was estimated as $\sim$80 kJ mol$^{-1}$ (Figures S54 and S55), which is significantly higher when compared to $\sim$46 kJ mol$^{-1}$ of MOF-808-Gly (Figures S56 and S57). This is significant for MOF-808-dl-Lys, of which the ε-NH$_2$ were from glycine, having a comparatively lower pK$a$ value of 9.60. This observation further supports the proposed relationship between higher pK$a$ of the amino species and stronger CO$_2$ binding in relatively low CO$_2$ pressure among the MOF-808-AA series.

**Binary CO$_2$/H$_2$O Adsorption Isotherms and Dynamic Breakthrough Experiments.** As moisture is unavoidable toward achieving low-cost, practical flue gas capture, studying the influence of water on the sorbent’s CO$_2$ uptake is important. To probe this, we measured the binary CO$_2$/H$_2$O adsorption isotherm of the representative MOF-808-Gly. The experiment was performed in a custom-built volumetric system (section S8.3.1). The sample was first fully regenerated in situ under vacuum at 150 °C. Water was injected to achieve a headspace relative humidity of $\sim$10% (3.6 mmol g$^{-1}$, Figures 3d,e), and then CO$_2$ was loaded on the water-loaded sorbent. The loadings of both CO$_2$ and H$_2$O were monitored with a gas chromatograph (GC), and the loadings of each species were determined via a mass balance. The Toth model$^52$ was applied to fit the binary isotherm to extract the uptake at the given CO$_2$ pressures (Table 2 and section S8.3.2).

The resulting isotherm revealed a significant increase of CO$_2$ uptake of MOF-808-Gly in the presence of water compared to dry conditions. Specifically, overlay of the binary CO$_2$/H$_2$O and single-component CO$_2$ sorption isotherms of MOF-808-Gly indicated a more than 2-fold increase in CO$_2$ uptake at 4 kPa, which continued to be above the dry isotherm until $\sim$20 kPa (Figure 3d).

The increased uptake in the presence of water was further studied in a dynamic manner with a breakthrough system (section S9): MOF samples were loaded in a packed bed, and a stream of CO$_2$-containing gas mixture was passed through the sorbent and analyzed by GC. Normalized breakthrough curves of MOF-808, MOF-808-FR, and MOF-808-Gly are overlaid to compare the behavior of the sorbents with or without humidity (RH 15%, 25 °C, Figure 3f). Both MOF-808 and MOF-808-FR exhibited no significant change or a slightly earlier breakthrough time upon humidification of the stream. In contrast, MOF-808-Gly exhibited a significant delay of the breakthrough, corresponding to an increased uptake in the presence of humidity. This is in line with the increased uptake observed in the binary adsorption isotherm study.

These observations combined support the sorbent’s characteristics of humidity-enhanced CO$_2$ uptake performance in such scenarios. This is being attributed to the sorption mechanism of MOF-808-AAs, achieved by the introduction of amino acids as presented in the next section.

**Mechanistic Studies of CO$_2$ Chemisorption Using Solid-State NMR.** To understand the mechanism of the CO$_2$ adsorption chemistry of MOF-808-AAs in the presence of water, solid-state cross-polarization magic-angle spinning (CP-MAS) $^{13}$C and $^{15}$N NMR experiments were conducted on the representative variant MOF-808-Gly to probe the change of chemical species before and after loading of $^{13}$CO$_2$, with and without the presence of water (Figure 4 and section S10). $^{15}$N-labeled glycine was used for the preparation of $^{15}$N-labeled MOF-808-Gly following the same synthetic and activation procedure. $^{13}$C and $^{15}$N CP-MAS solid-state NMR (ssNMR) experiments were performed on the resulting sample to establish the composition of the pristine MOF sorbent (Figure 4, bottom). Signals at $\delta$ $^{13}$C = 170.3 and 133.6 ppm were attributed to the carboxylate and aromatic carbons of the BTC linker, respectively. Glycine ligand signals were found with chemical shifts of $\delta$ $^{13}$C = 179.5 and 45.2 ppm, attributed to the carboxylate and α-carbons, respectively. $^{15}$N labeling of the α-amino N of the glycinate ligand allowed tracking any changes in its chemical environment, which is key to understanding the chemical nature of the sorption phenomenon. The signals at $\delta$ $^{15}$N = 34.3 and 27.1 ppm are assigned to the amino groups in the protonated and deprotonated states.

The $^{15}$N-labeled MOF-808-Gly sample was kept in the same rotor and dosed with 98.7 kPa $^{13}$CO$_2$ at 25 °C, followed by...

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*Figure 4. Stacked plots of solid-state $^{13}$C (left) and $^{15}$N (right) CP-MAS ssNMR spectra of $^{15}$N-labeled MOF-808-Gly before and after adsorption of $^{13}$CO$_2$ with or without the presence of water. In dark gray (bottom) are the spectra for MOF-808-Gly fully activated under dynamic vacuum. In green (middle) are the spectra for MOF-808-Gly after exposure to 98.7 kPa $^{13}$CO$_2$ for 24 h. In red (top) are the spectra for MOF-808-Gly after exposure to a N$_2$ atmosphere with 95% relative humidity at 25 °C for 24 h, followed by 90.0 kPa $^{13}$CO$_2$ for 24 h. Signals of interest are marked with their $^{13}$C or $^{15}$N chemical shifts, and the assignments of major species identified are shown in dashed boxes. Spinning sidebands are marked with an asterisk.*
repeating the CP-MAS ssNMR measurement. In the resulting spectra, signals appearing at δ $^{13}$C = 169.4 and 162.5 ppm indicate the formation of carbamic acid (−NHCOOH) and carbamate (−NHCOO−) species, along with free CO$_2$ at δ $^{13}$C = 121.4 ppm. This is further confirmed by the emerging signal at δ $^{13}$N = 79.2 ppm in the $^{15}$N ssNMR spectrum, while the presence of signals at δ $^{15}$N = 34.4 and 27.1 ppm indicates the presence of ammonium species as the counterions of the carbamate as well as unreacted α-amines.

For comparison, a fully activated, $^{15}$N-labeled MOF-808-Gly sample was humidified through exposure to 95% relative humidity in a N$_2$ atmosphere at 1 atm at 25 °C for 24 h. This was followed by exposure to 90.0 kPa $^{13}$CO$_2$ for 24 h prior to CP-MAS ssNMR measurements under the same conditions as the dry sample. The resulting $^{13}$C NMR spectrum showed that the predominant species exhibited a new signal at δ $^{13}$C = 167.3 ppm, which was assigned to ammonium bicarbonate (−NH$_3^{+}$−CO$_2$H) species. This was accompanied by a shoulder peak observed at δ $^{13}$C = 163.1 ppm, likely indicating small amounts of carboxamides and carbamic acids. However, their $^{15}$N ssNMR signals expected around δ $^{15}$N = 79.2 ppm were not observed, possibly due to their low population. A single peak was found at δ $^{15}$N = 31.1 ppm predominant in the spectrum, which can be attributed to protonated α-ammonium in the glycinate ligands. We thus conclude that the predominant chemisorption product is ammonium bicarbonate.

To further probe the speciation of the sorption process, activation treatment at ambient and elevated temperature was applied to the samples dosed with $^{13}$CO$_2$ under dry and humid conditions. $^{14}$N-labeled MOF-808-Gly dosed with $^{13}$CO$_2$ in a dry condition was kept in the same rotor and evacuated under dynamic vacuum at room temperature (−25 °C) for 48 h, followed by measurements of CP-MAS ssNMR under the same conditions (Figure S62). The absence of signals around δ $^{13}$C = 121.4 ppm, where free or physisorbed $^{13}$CO$_2$ was expected, indicated the removal of $^{13}$CO$_2$ in the rotor headspace being physisorbed in the MOF. Additionally, we observed relative attenuation of the signals at δ $^{13}$C = 169.4 and 162.5 ppm which we attributed to carbamic acid or carbamate species. Similarly, the signal at δ $^{13}$N = 79.2 ppm in the $^{15}$N NMR spectrum was still present, indicating the retention of carbamic acid and carbamate species. This observation was in accordance with the nonzero uptake at very low pressures in the desorption branch of the single-component CO$_2$ isotherm of MOF-808-Gly and similar phenomena in other MOF-808-AA variants. In a further step, the same sample ($^{14}$N-labeled MOF-808-Gly, $^{13}$CO$_2$ dosed, activated at room temperature) was reactivated under dynamic vacuum while heated to 120 °C for 24 h (Figure S62). The resulting $^{13}$C and $^{15}$N NMR spectra largely resembled the ones measured on pristine $^{15}$N-labeled MOF-808-Gly prior to exposure to CO$_2$, confirming the requirement of heating to fully regenerate MOF-808-Gly from carbamate or carbamic acid sorption products.

By contrast, vacuum regeneration of $^{15}$N-labeled MOF-808-Gly that was loaded with $^{13}$CO$_2$ under humid conditions exhibited considerable bicarbonate removal at ambient temperature (Figure S63), as evidenced by significant attenuation of signals in ssNMR spectra from around δ $^{13}$C = 167.3 ppm (bicarbonate) to a set of peaks at δ $^{13}$C = 168.0, 165.6, and 162.0 ppm. In addition, emerging signals at δ $^{15}$N = 82.8 ppm further supported that part of the bicarbonate species transformed into carbonate and carbamic acid species during the vacuum regeneration process.

On the basis of these observations, we propose the overall mechanistic process of CO$_2$ chemisorption in MOF-808-AAs as summarized in Scheme 1: (1) in dry conditions, CO$_2$ reacts with amino groups to form carbamates or carbamic acids (1–2 amines to 1 CO$_2$) which necessarily require heat for regeneration; (2) in humid conditions, CO$_2$ reacts with amino groups to form carbamates or carbamic acids (1–2 amines to 1 CO$_2$) with a higher affinity at lower CO$_2$ pressures; and (3) in humid conditions, where abundant CO$_2$ is present, CO$_2$ reacts with amino groups to form mostly bicarbonates (net ~1 amine to 1 CO$_2$) with a likely equilibrium with carbamic acids and carbanisates, which can be reversed through vacuum regeneration at ambient temperature. We thus hypothesized that the increased stoichiometry of the bicarbonate species formation in abundant CO$_2$ and humid conditions partly accounts for the enhanced capture performance of MOF-808-AA sorbents. More importantly, we envisioned that their facile vacuum regeneratability should enable capturing CO$_2$ through vacuum swing adsorption under ambient conditions.

### Vacuum Swing Adsorption Capture of CO$_2$ from Simulated Coal Flue Gas

In light of the discoveries above, we were encouraged to probe the practicality of applying MOF-808-AA sorbents in VSA cyclic capture scenarios. Particularly, we expected to utilize the formation of bicarbonate in humid coal flue gas conditions to allow for efficient CO$_2$ capture while regenerating by using vacuum, a low-energy driving force, at ambient temperature. To examine this, the dynamic capture of CO$_2$ was implemented in an orthogonal packed-bed vacuum swing adsorption setup (detailed in section S11). Cylindrical pellets (6 mm diameter × 5 mm height) were produced through pelletization of pure microcrystalline MOF-808-Gly with a packing density of 0.45 g cm$^{-3}$. Such pellets, weighing 9 g in total, were activated and loaded into a cylindrical sorption bed (35 mm diameter). A simulated coal flue gas composed of 15% CO$_2$ (RH 20%) balanced with N$_2$ was passed through at 1 atm, 20–23 °C, and a fixed flow rate of 30 mL min$^{-1}$. 

![Scheme 1 Proposed Chemisorption Mechanism of CO$_2$ Uptake in MOF-808-AAs with (Humid) and without (Dry) the Presence of Water](https://doi.org/10.1021/jacs.1c0368)
The adsorption–regeneration process was measured in a cyclic manner. The CO₂ concentration was monitored at the effluent of the sorption bed, and a threshold of 2% was set to switch to vacuum regeneration, which was set for a fixed time. For a typical cycle, the adsorption step lasted 5.9 min, when 6.1 min of vacuum regeneration (∼10 Torr) was applied to remove the adsorbed CO₂ and H₂O, which were collected at the exhaust. The cyclic dynamic capture capacity was measured by numerical integration of the effluent CO₂ pressure as 0.42 mmol g⁻¹ h⁻¹, and a continuous monitoring of 80 cycles showed no decay in uptake capacity indicated by the downstream CO₂ concentration profile (Figure S67) and calculated CO₂ uptake capacity per cycle (Figure S68).

**CONCLUSIONS**

We showed in this work that MOF-808, when functionalized with amino acids where their carboxylate is bound directly to the Zr(IV), produce pores capable of selectively binding CO₂ from flue gas. The amine residues of the amino acid units point to the interior of the pores of the MOF, making them of sufficient affinity for CO₂ to allow the formation of bicarbonate along with carbamate and carbamic acid. This CO₂ chemistry also allows for the CO₂ to be desorbed without heating. On the basis of this work, it is clear that creating the right environment in the pores of MOFs for maximization of CO₂ in the form of bicarbonate would contribute greatly to making energy-efficient carbon dioxide adsorption–desorption cycle.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c13368.

Detailed experimental procedures and data for reported compounds, supplementary figures and tables for characterization, and description of instrumental setups (PDF)

**Accession Codes**

CCDC 2118566 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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**Notes**

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