A scalable solid-state nanoporous network with atomic-level interaction design for carbon dioxide capture

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Carbon capture and sequestration reduces carbon dioxide emissions and is critical in accomplishing carbon neutrality targets. Here, we demonstrate new sustainable, solid-state, polyamine-appended, cyanuric acid–stabilized melamine nanoporous networks (MNNs) via dynamic combinatorial chemistry (DCC) at the kilogram scale toward effective and high-capacity carbon dioxide capture. Polyamine-appended MNNs reaction mechanisms with carbon dioxide were elucidated with double-level DCC where two-dimensional heteronuclear chemical shift correlation nuclear magnetic resonance spectroscopy was performed to demonstrate the interatomic interactions. We distinguished ammonium carbamate pairs and a mix of ammonium carbamate and carboxylic acid during carbon dioxide chemisorption. The coordination of polyamine and cyanuric acid modification endows MNNs with high adsorption capacity (1.82 millimoles per gram at 1 bar), fast adsorption time (less than 1 minute), low price, and extraordinary stability to cycling by flue gas. This work creates a general industrialization method toward carbon dioxide capture via DCC atomic-level design strategies.

INTRODUCTION

The increasing concentration of greenhouse gases in the atmosphere is a critical environmental concern (1, 2). In 2021, the U.S. government pledged to reach carbon neutrality before 2050, with carbon neutrality referring to the achievement of net zero carbon dioxide emissions (3). In 2018, carbon dioxide emissions from the combustion of fossil fuels were about 75% of the total U.S. anthropogenic greenhouse emissions (4). Carbon capture, utilization, and sequestration (CCUS) comprises essential requirements to achieve carbon neutrality (5, 6). However, there are many challenges toward making CCUS a reality, notably capturing costs and energy consumption (5). The U.S. Department of Energy (DOE) has previously announced projects with a total value of $3.18 billion to boost advanced and commercially scalable technologies with CCUS to reach an ambitious flue gas emission CO2 capture efficiency target of 90% (7, 8). Therefore, extensive efforts have been made to develop advanced materials that simultaneously reduce CO2 capture costs and energy consumption while being greener, cheaper, and more scalable than previous CCUS technologies (5).

Aqueous polyamines generate dynamic libraries of ligands for spontaneous CO2 fixation with the advantage of rapid and reversible reaction with CO2, as well as low solvent cost (8). Aqueous polyamines have significant limitations, however, including decomposition of volatile amines, high energy consumption during solvent regeneration, equipment corrosion, and emissions that can be problematic for both environmental and human health (9). To overcome these limitations, amine-functionalized adsorbents have been proposed as an alternative to aqueous polyamines, having noncorrosive properties while still maintaining the high chemical selectivity and stability of the amine reaction with CO2. Adsorbents such as amine-functionalized metal-organic frameworks (MOFs) could usher in a new era for CCUS due to their high adsorption capacity and cooperative CO2 adsorption (9, 10). Nevertheless, boosting performance in capacities, chemical stability, and scalability of MOFs requires atomic-level understanding and further exploration of material options. Compared to MOFs, polymer-based porous adsorbents can be synthesized to be metal-free, sustainable, and derived from cheap raw materials, all of which are essential for mass production (11, 12).

At the kilogram scale, we demonstrate the new sustainable solid-state, polyamine-appended, cyanuric acid–stabilized melamine nanoporous networks (MNNs) via dynamic combinatorial chemistry (DCC) to achieve effective, scalable, recyclable, and high-capacity CO2 capture. MNNs, synthesized from commercial melamine and paraformaldehyde, are promising for reversible CO2 capture owing to the intriguing advantages of their robust flake-like structures, high surface areas, tunable surface chemistries, and industrial-scale capture capabilities. The nature of the chemisorption mechanism at the atomic level is essential for the design of polyamine-appended networks with high CO2 capacity. A quantitative assessment of the local bonding environment and the mechanism at the atomic level for the adsorption of CO2 with these networks is required to further the design of MNNs with high CO2 adsorption capacity for CCUS.

Solid-state nuclear magnetic resonance (NMR) provides information about an atom-by-atom basis to reveal the chemical structure and dynamics of solid porous materials (13). Because NMR signals are derived from the magnetism of atomic nuclei (especially the magnetic moment and angular momentum of nuclear spins), NMR is a spectroscopic technique in which a specific signal can be detected from each atom in a molecule (13). The frequency peaks are assigned to individual atoms, utterly different from other frequency

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peaks (e.g., infrared and Raman spectra). As a result, multidimensional NMR spectroscopy can be tailored according to a given chemical system to provide precise information about the interaction between atoms that cannot be gained in other structural technologies. For example, in the two-dimensional (2D) heteronuclear correlation (HETCOR) measurement, the correlation is mediated by either through-bond coupling or through-space dipolar coupling (13). In addition, NMR spectroscopy has identified local interatomic interactions in hybrid composites using quantitative measurement techniques (14). High-resolution methods, including 13C cross-polarization magic-angle spinning (CP-MAS) NMR and 2D HETCOR spectroscopy, can provide critical mechanistic and dynamic insights into the local environment, short-range structure, and interatomic interaction mechanism between CO2 molecules and MOFs (14, 15).

DCC is one candidate for creating such a library wherein all constituent atoms are in equilibrium under thermodynamic control (16), and the library members interconvert through reversible chemical processes involving covalent bonds or noncovalent interactions. Research on using DCC to craft binding sites for CO2 on a neutral target is currently scarce (8). MNMs present a diverse array of materials, and thus, a targeted, systematic search for a library of materials for carbon capture is important. In this study, we develop new long-term, polyamine-appended, cyanuric acid–stabilized MNMs via DCC at the kilogram scale for effective, scalable, recyclable, and high-performance CO2 capture.

Here, we use DCC to design interconverting species of amine [e.g., diethylenetriamine (DETA)]–appended MNMs for CO2 capture. We interrogate the cyanuric acid–modified DETA-appended MNMs (MNMsCya ⊃ DETA) structure by multinuclear NMR spectroscopy and density functional theory (DFT) calculations, uncovering the origin of the stability of chemisorbed CO2 during adsorption–desorption cycles. To further demonstrate the applicability of MNMsCya ⊃ DETA for CCUS, we measured adsorption time scales, cycling stability, and regeneration energy of the networks. Our work establishes that these systems exhibit scalable and affordable carbon capture properties and affirms the strategy of DCC-induced solid-state porous network design as a sustainable method for long-term gas storage.

**RESULTS**

**MNN modification and characterization**

To enhance the CO2 chemisorption and its adsorption–desorption stability, we synthesized MNMsCya ⊃ DETA (Fig. 1, A to D) at a kilogram scale (Fig. S1) using a designed scalable setup (Fig. 1E). In this modified network, cyanuric acid was introduced as an MNN dopant during polymerization and used as noncovalent anchor sites for alkylamines to ultimately maintain the CO2 adsorption capacity and improve chemical stability. The addition of DETA enabled an increase in CO2 chemisorption capacity. Figure 1F shows typical transmission electron microscopy (TEM) micrographs of the microtomed particles imaged under different magnifications, indicating that micro- and mesopores were developed (17). The pore size distribution of the networks suggests a mesopore-dominated structure, with a small fraction of micropores, as shown in Fig. S2. The TEM result demonstrates an amorphous phase in the microporous and mesoporous structure, which is consistent with the pore size distribution results (Fig. S2).

We first sought to determine the structures of modified MNMs before the adsorption of CO2. We examined the cyanuric acid–modified amine-appended MNMs in detail as a benchmark material via solid-state NMR. Our investigation began with the structure of MNMsCya (Fig. 1C), which was determined using quantitative 13C MAS NMR spectra by direct excitation (fig. S3). The MNM spectrum presents a sharp peak at 166.5 parts per million (ppm) emanating from the triazine ring of melamine, while the two overlapping broad peaks at around 50.0 ppm arise from the =NH—CH2—NH— aminal group and the =NH—CH2—OH terminal hemiaminal group, respectively (fig. S3A). After cyanuric acid dopant incorporation, the signals observed at 150.0 and 40.3 ppm are attributed to triazine ring C=O bonds and —NH—CH2—O—, respectively (18).

We subsequently measured the 13C CP-MAS NMR spectra of DETA, MNMs ⊃ DETA, and MNMsCya ⊃ DETA via CP from 1H to 13C nuclei with heteronuclear decoupling (fig. S3B). The DETA spectrum exhibits two peaks at 53.6 and 42.7 ppm, attributed to the CH2 groups from =CH2—NH—CH2 and =CH2—NH2, respectively. These DETA signals persist in the MNMs ⊃ DETA and MNMsCya ⊃ DETA spectra, at 53.0 and 42.0 ppm, respectively. Thus, DETA molecules are successfully appended to the networks, presumably by hydrogen bonding during post-synthetic modification.

We performed direct detection of 13C MAS NMR spectra (i.e., without CP) of the MNM series to provide a quantitative spectrum that would assist in identifying the structures of modified MNMs. All spectra were deconvoluted and integrated using the dmfit software (19) (fig. S4) with the spectral simulation parameters (table S1). This analysis provides information about the relative proportion of triazine rings within MNMs and of the triazine rings of cyanurate within MNMsCya. Through this quantitative 13C NMR spectroscopy, we were able to confirm the structures of MNMsCya, MNMs ⊃ DETA, and MNMsCya ⊃ DETA, consistent with the DFT-calculated structures at the TPSS-D3(BJ)/6-31G* level in Q-Chem 5.4 (Fig. 1, A to D) (20). Cyanuric acid molecules were inserted into the triazine rings within MNMs (N2,N4,N6-trimethyl-1,3,5-triazine-2,4,6-triamine) with a ratio of 1:6, based on the results from the quantitative 13C NMR spectra (fig. S4 and table S1). The positions of the inserted DETA molecules fell at the edges of MNMsCya networks and were bound through two amine groups to the framework —NH— sites by hydrogen bonds with an average N...H distance of 2.12 Å (Fig. 1D). The 13C MAS NMR spectra of MNMsCya ⊃ DETA as a function of pressure are shown in fig. S5.

Last, 15N CP-MAS NMR experiments were conducted for further structure determination. The 15N NMR spectrum of cyanuric acid–modified MNMsCya shows a weak resonance at 147.0 ppm (fig. S3C), assigned to the nitrogen within the triazine ring of cyanurate (2,4,6-trimethoxy-1,3,5-triazine), while the 15N signal of the cyanuric acid nitrogen is located at 136.0 ppm. The weak signal within the 15N spectrum of MNMsCya is due to the low concentration of cyanuric acid (7.5 mole percent (mol %)) in MNMsCya. The chemical shift difference (11 ppm) between the aforementioned two spectra is probably due to triazine ring currents from MNMs (N2,N4,N6-trimethyl-1,3,5-triazine-2,4,6-triamine) within MNMsCya, as discussed previously (19). The 15N spectra of MNMs ⊃ DETA and MNMsCya ⊃ DETA also confirm that DETA was successfully appended into the networks (fig. S3C). The two distinct peaks of the 15N spectra of MNMs ⊃ DETA and MNMsCya ⊃ DETA, which appear after DETA postmodification, are located at 30.3 and 20.1 ppm, corresponding to —NH— and NH2— within the appended DETA, respectively. The emergence of these two peaks from DETA plays an essential role in reaction with CO2.
CO$_2$ adsorption properties

We conducted CO$_2$ adsorption isotherms of MNNs and MNNs$_{Cya \supset DETA}$ at room temperature (298 K), shown in Fig. 2A. Both samples have typical Langmuir-type isotherms commonly observed for microporous materials that favor gas adsorption (21). Notably, an increase in CO$_2$ uptake for MNNs$_{Cya \supset DETA}$ is observed, relative to the unmodified material, from 0.91 to 1.82 mmol/g at 1 bar, presumably due to the successful incorporation of cyanuric acid and DETA into MNNs. Our results indicate that the DETA and cyanuric acid-modified MNNs have a high affinity and adsorption capacity of CO$_2$, making them suitable for direct CO$_2$ capture.

Further solid-state $^{13}$C and $^{15}$N NMR measurements enabled us to examine the underlying mechanisms for the formation of ammonium carbamate and carbamic acid during CO$_2$ adsorption. We used a custom-built gas dosing setup to achieve $^{13}$CO$_2$ adsorption onto network materials with a precise gas flow in a sealed system (Fig. S6) (19). This system affords the use of commercial MAS rotors while enabling controlled gas dosing of activated network samples. Here, $^{13}$CO$_2$ gas dosing was performed at various pressures up to ~1 bar (298 K) using the system pressure gauge that precisely monitored the equilibration of $^{13}$CO$_2$ adsorption before the acquisition of the $^{13}$C MAS NMR spectra. In all measurements, $^{13}$C-labeled $^{13}$CO$_2$ gas could be readily distinguished from unlabeled carbon within the networks owing to its high $^{13}$C enrichment level. Starting from bare MNNs, the direct $^{13}$C NMR spectra (Fig. 2B) for two scans of $^{13}$CO$_2$-dosed MNNs display a prominent resonance that corresponds to physisorbed CO$_2$ at 125.4 ppm. As expected, we also observe a peak at 125.4 ppm, assignable to physisorption of CO$_2$ in the $^{13}$C NMR spectra of $^{13}$CO$_2$-dosed MNNs$_{Cya}$, with further signal averaging as compared to $^{13}$CO$_2$-dosed MNNs. Under these conditions, a second peak at 166.4 ppm is also observed and assigned to the resonance from the triazine ring of the networks (Fig. 2B). The chemical shift of physisorbed CO$_2$ within MNNs is shifted by ~2.3 ppm relative to free gaseous CO$_2$ at 127.7 ppm at 1 bar (22), likely due to the
shielding effect of the triazine ring currents in the frameworks. A similar result was also previously reported for MOF-274 because of aromatic ring currents arising from the linkers (22). Figure 2C and fig. S7 depict the $^{13}$C NMR spectra of $^{13}$CO$_2$-dosed MNNs and MNNs$_{Cy}$ obtained via direct excitation and CP experiments.

The latter experiment allows for the much faster detection of chemisorbed CO$_2$ by proton CP from the $^1$H-rich networks. The direct detection spectra exhibit a peak at 125.4 ppm caused by CO$_2$ physisorption in the networks.

The distinct peak observed at ~165 ppm, assigned to chemisorbed CO$_2$, consists of a sharper (164.7 ppm) and a broader (161.1 ppm) component (Fig. 2C). We identify the former chemisorbed species as ammonium carbamate, with the latter being a minor chemisorption product, carbamic acid. The chemical shifts are similar to those previously assigned to ammonium carbamate and carbamic acid (22). Quantitative $^{13}$C NMR spectra reveal that the overwhelming majority (92%) of adsorbed CO$_2$ is chemisorbed at 1 bar, and only a small amount (~8%) is physisorbed at this pressure (table S1). Of the chemisorbed species, carbamic acid comprises a minor percentage (20%) of the reaction products, while ammonium (80%) carbamate dominates the chemisorption products (table S1). The spectra of CO$_2$-adsorbed MNNs$_{Cy}$ ⊃ DETA and MNNs ⊃ DETA are very similar, suggesting similar chemisorbed products (ammonium carbamate and carbamic acid).

**Chemisorption mechanism in the context of DCC**

DCC allows individual species to interconvert through a reversible bond exchange process that involves either covalent bonds or noncovalent interactions (23). Whether covalent or noncovalent, dynamic bonds are crucial to access distinct self-assembled structures that produce the most thermodynamically stable species (24). Here, CO$_2$-derived building blocks are generated via a strategy of combining two reversible covalent bonds (imines and carbamates) and therefore hold enormous potential for CO$_2$ capture (Fig. 3A). Figure 3 (B and C) detail the five hypothetical structures that may be generated from the N–CO$_2$ reactions in the MNNs$_{Cy}$ ⊃ DETA, using a double-level dynamic combinatorial system (25, 26). The system comprises first-level carbamation (Fig. 3B) and second-level ion-pairing reactions (Fig. 3C). At the first level, the exposure of CO$_2$ to MNNs$_{Cy}$ ⊃ DETA leads to the formation of ammonium carbamate and carbamic acid. This occurs through reversible reactions, exchange, and generation of a dynamic system of species (Fig. 3B, eqs. 1 and 2). It has been demonstrated that structural diversity within a dynamic combinatorial system can be expanded by maintaining two communicating exchange processes simultaneously (27, 28). Consequently, in the presence of excess CO$_2$, a system driven by DCC may display a sequential reaction between or within system members, giving rise to second-level paired structures (Fig. 3C). Depending on the degree of protonation and carbamation, these paired structures may potentially encompass ammonium carbamate chains (Fig. 3C, eq. 3), carbamic acid pairs (Fig. 3C, eq. 4), and mixed carbamate and carbamic acid pairs via Lewis acid sites (Fig. 3C, eq. 5). In particular, the above reversible reactions can eventually revert to the initial reactants under thermodynamic control. From this perspective, if the combinatorial species can bind intermolecularly to induce self-pairing, then this process will lead to self-assembly and strongly contribute to favorable CO$_2$ adsorption isotherms.

**Demonstration of the dynamic combinatorial species by solid-state NMR at the atomic level**

DETA was appended to solid-state networks by hydrogen bonds to provide a noncovalent anchoring site for CO$_2$ (vide supra). Compared to liquid DETA, the structure of DETA-appended MNNs has a rigid and porous nature, resulting in a complex milieu of CO$_2$...
reactions and allowing for the potential formation of monocarbamates and dicarbamates (Fig. 4A). A simplified network of dynamic combinatorial species, where protonation is only partially expressed, is shown in Fig. 4A. Specifically, this process consists of two dynamic combinatorial levels. We start from two pristine structures of MNNsCya ⊃ DETA, which are denoted d0. With low CO2 loading, N–CO2 covalent bonds are generated by the CO2 addition reactions to the lone pair electrons of the nitrogen, producing protons that...
generate neighboring ammonium ions. There are two types of amine sites that CO$_2$ molecules might occupy: primary amine (d1a) and secondary amine (d1s), which form via the reversible reactions, eqs. 6 and 7, respectively (Fig. 4A). In our nomenclature for adducts generated in the dynamic combinatorial systems, s and a denote symmetric and asymmetric carbamate patterns on the networks, respectively. With an excess of CO$_2$, both primary amines might react with CO$_2$ molecules, denoted d2s.

Solid-state NMR methods specify the populations within the dynamic combinatorial networks. In particular, natural abundance $^{15}$N CP-MAS NMR spectra corroborate CO$_2$ reaction sites (Fig. 4B and fig. S8). For both MNNs$^{Cya}$ $\supset$ DETA-CO$_2$ and MNNs $\supset$ DETA-CO$_2$, we observe that the resonances at 20.1 ppm, assignable to nitrogen from primary amines (marked with purple circles), disappeared after loading CO$_2$ at ~1 bar. This result indicates a conformational carbamate species generated through a primary amine N$\rightleftharpoons$CO$_2$ reaction, in turn implying that CO$_2$ molecules favor interaction with primary amine nitrogens. This result is also consistent with an analogous study in a diamine-appended MOF (22).

Furthermore, the peak at 30.3 ppm arising from secondary amine nitrogen remains in the spectrum regardless of CO$_2$ loading, allowing us to conclude that the secondary amine nitrogens have not reacted with CO$_2$. In contrast, in the liquid DETA-CO$_2$ system, CO$_2$ readily reacts with secondary amines and forms carbamate (8). We theorize that hydrogen bonding interactions between the secondary amine nitrogen (DETA) and MNNs’ backbone occupies the lone pair of the secondary amine nitrogen, effectively preventing reaction with CO$_2$. Correspondingly, we can exclude eq. 7 in our dynamic combinatorial system (Fig. 4A).

To monitor the further reaction of additional CO$_2$ at high CO$_2$ loadings, we calculate the mole fraction of chemisorbed CO$_2$ (mole) for each DETA (mole) in the network. On the basis of the prior structure of DETA-appended MNNs determined via DFT calculations and NMR lineshape analysis, we identify a stoichiometry of 1 CO$_2$ captured per 1 DETA (table S1). These quantitative calculations reveal that no additional CO$_2$ molecules react with both primary

![Diagram](https://www.science.org/)

**Fig. 4. Demonstration of the dynamic combinatorial species in MNNs$^{Cya}$ $\supset$ DETA by solid-state NMR.** (A) Depiction of the network of reversible carbamate N$\rightleftharpoons$CO$_2$ reaction sites. (B) $^{15}$N CP-MAS NMR spectra of MNNs$^{Cya}$ $\supset$ DETA with and without CO$_2$ adsorption. (C) $^1$H-$^{13}$C HETCOR NMR spectrum, with the assignment of observed correlations, for MNNs$^{Cya}$ $\supset$ DETA loaded with CO$_2$ at 1 bar. The contact time was 100 μs. (D) The two-level dynamic combinatorial system of CO$_2$ chemisorption species as confirmed by NMR. First-level species: ammonium carbamate and carbamic acid. Second-level species: ammonium carbamate/carbamic acid pairs.
Kinetics of CO₂ adsorption and adsorption-desorption cycles

To further elucidate the second level (ion-pairing interactions) of the dynamic combinatorial system (Fig. 3C), we perform additional 2D ¹H-¹³C HETCOR experiments (Fig. 59) on MNNs⁵⁷Ca ⊃ DETA-CO₂ at 1013 mbar (Fig. 4C). This experiment relies on through-space dipole-dipole interactions to selectively detect ¹H-¹³C correlations resulting from spatial proximity. A short contact time (100 μs) enables the detection of only those hydrogen atoms closest to the ¹³C nuclei of the ion-paired species. The HETCOR spectra reveal a dominant correlation at 164.7 ppm and a minor correlation at 161.1 ppm in the ¹³C dimension, corresponding to ammonium carbamate and carbamic acid, respectively (18). For this dominant ¹³C resonance at 164.7 ppm, we assign the correlations to the ¹H peak of the secondary amine NHCOO⁻ with a chemical shift of 6.1 ppm (Fig. 4C, peak labeled A). A weaker correlation to the 11.2 ppm ¹H chemical shift is associated with protons from ammonium carbamate, RNH₂⁺ (Fig. 4C, peak labeled B). As a result, we can see the second-level dynamic combinatorial species (Fig. 3C, eq. 3) in the form of the dominant ion pairing between ammonium and the oxygen atom of the carbamate (Fig. 4D). This conformation is unique among the amine-appended polymer networks, and its formation is likely due to the irregular self-folding of amorphous polymer networks.

The carbamic acid species associated with the ¹³C resonance at 161.1 ppm present an intense cross-peak (5.5 ppm, peak C) and two less-intense ¹H correlations (8.5 ppm, peak D; 12.5 ppm, peak E; Fig. 4C). On the basis of our previous hypothesis, these species could be carbamic acid pairs or mixed carbamate and carbamic acid pairs (Fig. 3C, eqs. 4 and 5). The presence of peak C, however, leads us to conclude that some ammonium ions are strongly correlated to the reacted ¹³CO₂, consistent only with eq. 5—the mixed carbamate/carbamic acid pairs. The intensities of the cross-peaks A and C suggest similar ¹H-¹³C correlations for both moieties, implying that the peaks at 5.5 and 8.5 ppm correspond to protons of NHCOOH (acid) and RNH₂⁺ (carbamate), respectively. These two major (peaks A and C) and minor (peaks B and D) correlations at these short contact times are anticipated for the mixed carbamic acid/carbamate pair via hydrogen bonds, forming important components of the second level of the DCC system (Fig. 3C, eq. 5), with the relatively short C...H distances predicted from DFT calculations (Fig. 5F; 1.95 Å for C...H₃N⁺ and 2.01 Å for C...H₃NHCOOH). Last, a weak proton signal at 12.5 ppm is tentatively assigned to the acid proton NHRCOO⁻ owing to the extreme chemical shift associated with strong O...H hydrogen bonds. Together, the NMR data paint a picture of ion-pairing and hydrogen-bonding interactions that promote the stabilization of the CO₂-inserted phase.

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These data illustrate that amine-appended network materials not only have low energy consumption for CO₂ but also may exhibit cycling times that can be matched to industrial process equipment.

Carbon dioxide uptake during multiple adsorption-desorption cycles, CO₂ isotherms were measured at 298 K for 10 adsorption-desorption cycles to reveal the effects of cyanuric acid modification on CO₂ adsorption performance. Figure 5E shows the recorded CO₂ uptakes of MNNs⁵⁷Ca ⊃ DETA and MNNs ⊃ DETA over 10 cycles. After 10 cycles, we observe a substantial decrease from 99 to 78% CO₂ capacity at 1 atm for the MNNs ⊃ DETA, yet the CO₂ capacities for the MNNs⁵⁷Ca ⊃ DETA sample did not change significantly. We hypothesized that the DETA of MNNs ⊃ DETA sample loses amine functionality over the adsorption-desorption cycles and tested this hypothesis by conducting ¹³C MAS NMR experiments before and after 10 CO₂ adsorption-desorption cycles. The ¹³C NMR spectra of MNNs ⊃ DETA following the CO₂ adsorption-desorption cycles are shown in Fig. 5G (marked in orange), while MNNs⁵⁷Ca ⊃ DETA is shown in Fig. 5G (marked in blue). In the absence of cyanuric acid, the network DETA peaks at 53 and 42 ppm diminish, suggesting that DETA loading decreases after 10 cycles. We surmise that in the adsorption-desorption process, hydrogen bonds break as CO₂ is inserted, significantly hindering subsequent adsorption performance upon cycling. Unexpectedly, we noted that the two DETA resonances in the spectrum of MNNs⁵⁷Ca ⊃ DETA remain unchanged after 10 cycles, from which we conclude that cyanuric acid aids in tethering the DETA molecules to the networks. N...H...O hydrogen bonds between cyanuric acid and networks are formed, providing additional stability. The electron negativity of oxygen in N...H...O is higher than that of nitrogen in N...H...N, and we surmise that this results in greater stabilization of DETA by the cyanuric acid groups. These data illustrate that cyanuric acid- and amine-appended networks, for which the CO₂ chemisorption byproducts can be designed and synthesized via DCC, have extensive and stable CO₂ capture capacities at atmospheric pressures.
DISCUSSION

We have synthesized an amine-appended melamine porous network at a kilogram scale that acts as a CO₂ capture material with efficient adsorption, extraordinary cycling stability, and low regeneration energy. From a dynamic combinatorial framework, we reveal the chemisorption within this porous network as a series of interconnected reactions, elucidated by solid-state $^{13}$C and $^{15}$N NMR spectroscopy at the atomic level. We have demonstrated a double-level DCC system that includes...

Fig. 5. DFT-calculated structures, kinetics of adsorption, and stability of MNNs$^{\text{CYA}}$ ⊃ DETA during CO₂ cycles. (A) Schematic diagram of adsorption column system. (B) Adsorption (filled shapes) and desorption (open shapes) isotherms for CO₂ uptake in MNNs$^{\text{CYA}}$ ⊃ DETA at 273, 298, and 313 K. (C) The $-\Delta h_{\text{ads}}$ (enthalpy) and $-\Delta s_{\text{ads}}$ (entropy) for MNNs$^{\text{CYA}}$ ⊃ DETA determined using the Clausius-Clapeyron equation as a function of CO₂ loading. (D) Adsorption breakthrough and rate for adsorption column of MNNs$^{\text{CYA}}$ ⊃ DETA at 313 K (15% CO₂, 83% N₂, and 2% H₂O). (E) Recorded CO₂ uptakes during 10 adsorption-desorption cycles using the homemade dosing setup. (F) Proposed mixed chemisorption structure with ammonium carbamate pair. The structure was obtained by DFT modeling at the TPSS-D3(BJ)/6-31G* level. (G) $^{13}$C NMR (16.4 T) spectra acquired by CP for MNNs ⊃ DETA and MNNs$^{\text{CYA}}$ ⊃ DETA, before and after 10 CO₂ adsorption-desorption cycles.
reversible carbamate N–CO$_2$ reactions and interlocked formation of ion pairs. In the first level, we identify the formation of two chemisorbed species consisting of ammonium carbamate (dominant) and carbamic acid (minor), quantified via direct detection $^{13}$C NMR spectra. The 2D $^1$H–$^{13}$C HETCOR NMR measurements reveal the second-level dynamic combinatorial species comprising ammonium carbamate pairs and mixed carbamate and carbamic acid pairs. Through quantitative $^{13}$C NMR, we determine the local atomic structure of amine-appended MNNs with cyanuric acid, which were confirmed with DFT calculations, and we discerned the equimolar reaction of chemisorbed CO$_2$ with each DETA molecule. This structure shows conformational carbamate species derived from the reaction of CO$_2$ with primary amine functionalities. In particular, our results highlight the importance of cyanuric acid modification to enhance the CO$_2$ chemisorption cycling stability of MNNs-based materials. Solid-state NMR spectroscopy as a nondestructive and atomic-level tool is proven to be an effective way to supplement standard laboratory methods for atomic structure determination in the sustainable energy field. Industrialization of CO$_2$ capture using networks (e.g., polymers, covalent organic frameworks (COFs), and MOFs) is made possible by this work, which uses atomic-level and DCC design methodologies with accurate mechanism analyses. Integrating scalable nanoporous materials synthesis, chemical engineering, and advanced characterization techniques will accelerate our world’s transition to a negative carbon emissions pathway.

MATERIALS AND METHODS

Synthesis of melamine porous network series

Commercially available melamine and paraformaldehyde (Sigma-Aldrich) were used as the starting materials. The scaffold was synthesized through a condensation polymerization at 150° to 170°C in dimethyl sulfoxide (DMSO) for 3 to 7 days (fig. S10) (29). During the polymerization reaction of MNNs, cyanuric acid (7.5 mol %) was mixed with the hot melamine solution (with DMSO as solvent) before the addition of formaldehyde to tether DETA (denoted as MNNs$_{\text{Cys}}$). DETA was mixed with MNNs/MNNs$_{\text{Cys}}$ (with hexane as solvent) (denoted as MNNs ⊃ DETA/MNNs$_{\text{Cys}}$ ⊃ DETA). The DETA was mixed with MNNs/MNNs$_{\text{Cys}}$ in a sonication bath at 50°C for 3 hours. Afterward, the mixture was filtered and washed using tetrahydrofuran, a polar solvent, to remove the residual surface DETA. The collected solid was activated under a vacuum oven at 85°C for 1 hour. All samples were stored in an Ar-filled glovebox for future use.

NMR sample preparation

The activated sample was packed into 4- or 3.2-mm rotors inside an Ar-filled glovebox. At room temperature, $^{13}$CO$_2$ gas dosing was performed in a custom-built setup shown in fig. S6, featuring an evacuated system in which rotors can be closed with caps using a moveable plunger with an O ring. Gas dosing was carried out at various pressures (10 to 1000 mbar) via a capacitance manometer (model 722B, MKS Instruments). Before gas dosing, the rotor containing the sample was evacuated for at least 10 min inside the glass tube of the custom-built setup. VCR (Swagelok) metal gasket with face seal fittings enabled a metal-to-metal seal, providing a leak-tight system over a pressure range from vacuum to positive pressure. $^{13}$CO$_2$ gas (99 atomic % of $^{13}$C, Sigma-Aldrich) was dosed into the rotor overnight to ensure saturation of the sample. Once equilibrium was reached, the pressure at this condition was recorded and related to $^{13}$CO$_2$ adsorption capacity.

NMR measurement

All 1D NMR experiments were performed at 500.12 MHz for $^1$H (11.7 T) on a Bruker Advance spectrometer with a Bruker narrow bore H/C/N MAS probe (fig. S9A). The $^{13}$C/$^{15}$N MAS NMR spectra were acquired using CP from $^1$H to $^{13}$C/$^{15}$N nuclei with heteronuclear decoupling at a sample spinning rate of 10 kHz (fig. S9B). All CP experiments were conducted at a recycle delay of 2 s with a radio frequency field strength of ~25 to 70 kHz, with a contact time of 2 ms. The quantitative $^{13}$C direct excitation experiments were carried out with heteronuclear decoupling during detection and sufficiently long recycle delays, 200 s, allowing the nuclei to fully relax between scans. The 2D $^1$H–$^{13}$C HETCOR experiments were conducted at 16.4 T (700 MHz for $^1$H) using a Bruker 3.2-mm MAS probe at a spinning rate of 15 kHz with a radio frequency field strength of ~80 kHz. The CP ($^1$H to $^{13}$C) pulse sequence was 90° $^1$H – t$_1$ – CP – t$_2$, with a contact time of 100 μs. A $^1$H 90° one-pulse sequence was used for $^1$H NMR experiments. $^1$H, $^{13}$C, and $^{15}$N chemical shifts were calibrated using $^{13}$C-labeled adamantane and $^{15}$N-labeled glycine, referenced to 1.85 ppm (adamantane; $^1$H), 38.5 ppm (adamantane, tertiary carbon; $^{13}$C), and 33.4 ppm (glycine; $^{15}$N), respectively.

DFT calculations

To determine the DETA and cyanuric acid–modified MNN structure along with CO$_2$-adsorbed local structure, DFT calculations were carried out on the Massachusetts Green High Performance Computing Center (30, 31), using the Q-Chem 5.4 package. The cluster structures of CO$_2$, DETA, MNNs, MNNs ⊃ DETA, MNNs ⊃ DETA–CO$_2$, MNNs$_{\text{Cys}}$ ⊃ DETA, and MNNs$_{\text{Cys}}$ ⊃ DETA–CO$_2$ were optimized at the TPSS-D3(BJ)/6-31G* level (Fig. 5F and figs. S14 to S21).

CO$_2$ adsorption isotherms and breakthrough measurement

The CO$_2$ adsorption isotherms were measured gravimetrically using a sorption analyzer (model VTI-SA, TA Instruments) at 298 K with CO$_2$ as the carrier gas. Samples were first degassed at 85°C under vacuum for 1 hour. The evacuated tube recorded the equilibrium weight of the degassed sample in response to a step change in the concentration of the CO$_2$ (relative pressure range of 0.01 to 1). Approximately 80 to 90 mg of sample was weighed and placed into the tube of the analyzer. Equilibrium was defined to be reached when the weight changed by less than 0.001% over an interval of 30 s. A custom-built setup for breakthrough measurement, shown schematically in Fig. 5A, was used to perform breakthrough adsorption measurements. N$_2$ was used as the carrier gas. During the adsorption process, the mixed N$_2$ and CO$_2$ were obtained from Airgas Inc. The mass flow controller (I-Series IP66, MFC) was used to precisely control the flow rate. The effluent gas concentration was measured via a capacitance manometer (model 722B, MKS Instruments). The outlet concentration was recorded every minute until the outlet gas concentration was stabilized at 99% of the inlet concentration.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abo6849

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Acknowledgments: Z.L. acknowledges the start-up support from the University of Massachusetts Amherst (UMass) and the computational resource from the Massachusetts High Performance Computing Center. We also thank H. Celik, N. Jarenwattananon, and the College of Chemistry NMR Facility, University of California, Berkeley. We appreciate technical discussions with A. C. Forse and T. M. Osborn Popp. Funding: This work was partly supported by the DOE, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering (contract no. DE-AC02-76SF00515). J.A.R. acknowledges partial support from the ACT-Pv5Ma project, which has received joint funding from BEIS, NERC, and EPSRC (UK), funding from the Division of CCS R&D, DOE, and funding from the Office Fédérale de l’Energie (Switzerland). D.M.H. acknowledges support from the Joint Center for Energy Storage Research, an Energy Innovation Hub funded by the DOE, Office of Science, Basic Energy Science. H.-C.Z. and G.S.D. acknowledge the financial support of the DOE Office of Fossil Energy, National Energy Technology Laboratory (DE-FE0006742). Author contributions: Y.C., J.A.R., J.T., and H.M. conceived the idea, planned and conducted all the experiments. H.-C.Z. and G.S.D. synthesized materials and performed CO2 adsorption isotherms. Z.L. calculated molecular and complex structures. All the authors reviewed and commented on the manuscript. Competing interests: The authors declare that they have no competing interests. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials.

Submitted 18 February 2022
Accepted 17 June 2022
Published 3 August 2022
10.1126/sciadv.abd6849
A scalable solid-state nanoporous network with atomic-level interaction design for carbon dioxide capture

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Sci. Adv., 8 (31), eabo6849. • DOI: 10.1126/sciadv.abo6849

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https://www.science.org/doi/10.1126/sciadv.abo6849

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