Temperature dependent CO$_2$ behavior in microporous 1-D channels of a metal-organic framework with multiple interaction sites

Dongwook Kim$^1$, Jaehun Park$^2$, Yung Sam Kim$^2$ & Myoung Soo Lah$^1$

The MOF with the encapsulated CO$_2$ molecule shows that the CO$_2$ molecule is ligated to the unsaturated Cu(II) sites in the cage using its Lewis basic oxygen atom via an angular $\eta^1$-(O$_A$) coordination mode and also interacts with Lewis basic nitrogen atoms of the tetrazole ligands using its Lewis acidic carbon atom. Temperature dependent structure analyses indicate the simultaneous weakening of both interactions as temperature increases. Infrared spectroscopy of the MOF confirmed that the CO$_2$ interaction with the framework is temperature dependent. The strength of the interaction is correlated to the separation of the two bending peaks of the bound CO$_2$ rather than the frequency shift of the asymmetric stretching peak from that of free CO$_2$. The encapsulated CO$_2$ in the cage is weakly interacting with the framework at around ambient temperatures and can have proper orientation for wiggling out of the cage through the narrow portals so that the reversible uptake can take place. On the other hand, the CO$_2$ in the cage is restrained at a specific orientation at 195 K since it interacts with the framework strong enough using the multiple interaction sites so that adsorption process is slightly restricted and desorption process is almost clogged.

Recent global warming is closely related to the increased CO$_2$ concentration in the atmosphere$^1$. One way to tackle the problem is CO$_2$ capture-storage and utilization$^2$–$^5$. Microporous metal-organic frameworks (MOFs) received great attentions for CO$_2$ capture and utilization$^6$–$^{12}$. The properties of a MOF can be modulated to meet necessities for CO$_2$ sequestration agents using not only its organic residues but also metal ions in pore surface.

Though there are many structural reports on the CO$_2$ bound species, most of them are of molecular complexes that contain metal sites interacting with CO$_2$ in strong coordination bond such as $\eta^2$-(C, O)$^5$, $\eta^1$-(C)$^6$ and linear $\eta^1$-(O$_C$) coordination modes$^{13}$ (Fig. 1). There are also several reports on the structural characteristics of the CO$_2$ encapsulated in micropores of MOFs$^{13}$–$^{21}$, however, the interactions are weak non-covalent interactions, such as hydrogen bonding, dipole-quadrupole interaction and van der Waals interactions (C⋯H⋯O and O⋯H⋯O interactions), only with organic residues in the pore surface.

The structural information of the encapsulated CO$_2$ interacting with the framework using multiple binding sites is scarce$^2$. On the other hand, while there are several reports on the vibrational mode analysis of CO$_2$ strongly bound to metal centers$^{13,19,20}$, only a few investigations on the correlation between the vibrational modes of the encapsulated CO$_2$ weakly interacting with the framework and its structural data are reported$^{16}$–$^{19,22}$.

Here, we report the temperature dependent single crystal structure analyses of a microporous MOF with bound CO$_2$ in microporous 1-D channels via both a weak angular $\eta^1$-(O$_A$) coordination of CO$_2$ to Cu(II) sites and a dipole-quadrupole interaction between a Lewis basic atom of a framework and the Lewis acidic carbon atom of CO$_2$. Temperature dependent vibrational modes of bound CO$_2$ are investigated using infrared (IR) spectroscopy together with temperature dependent crystal structures with the encapsulated CO$_2$. Reversible CO$_2$ uptakes at...
around ambient temperatures but irreversible CO₂ uptake at 195 K could be explained based on temperature dependent structural information and vibrational band analyses of the bound CO₂.

Results and Discussion

Single crystal structure analyses of the MOF and its CO₂-bound MOF derivatives. A solvothermal reaction of CuCl₂ with tetrazole (tz) in methanol (MeOH)/acetonitrile mixed solvent produced block-shaped blue crystalline product of the formula unit, Cu₃Cl₂(tz)₄(MeOH)₂ ((MeOH)₂@1a). A single crystal structure analysis revealed that the product is a 3-D MOF of an 8-c topology with linear trinuclear [Cu₃Cl₂] secondary building unit (SBU) as an 8-c node and tridentate tz as a ditopic linker (Fig. 2). A partially occupied methanol molecule is weakly ligated at the Jahn–Teller elongated sixth coordination site of the terminal copper atom (Cu1–O1M: 2.617(4) Å) of the linear [Cu₃Cl₂] SBU (Fig. 2e).

Figure 1. Coordination modes of CO₂ to metal center.

Figure 2. (a) Ball-and-stick diagrams of 1 viewed along the crystallographic c-axis (left) and b-axis (right), respectively, where [Cu₃Cl₂] SBU as 8-c nodes are represented by pink spheres and the linkers between the 8-c nodes are represented by the green sticks. (b) A [Cu₃Cl₂(tz)₈] cluster (top) and the [Cu₃Cl₂] SBU as an 8-c node (bottom). (c) The space-filling diagrams of 1 drawn as the same view as the ball-and-stick diagrams in (a). (d) A 1-D channel viewed along the crystallographic b-axis, where a CO₂ is ligated to Cu(II) site. Ball-and-stick diagrams of the single crystal structures of (e) MeOH-bound 1a at 173 K, (f) CO₂-bound 1a at 195 K, and (g) CO₂-bound 1a at 296 K.
sites in the cage (Figs 2e and S2a) and the remaining lattice methanol molecules are statistically disordered in the vicinity of the ligated methanol molecules (Figs 2e and S2b). The properly activated sample, 1a, was prepared by vacuuming (~10⁻² torr) (MeOH)₉@1a at 160 °C for 7 d (Figs S3–S7). Single crystals of the CO₂-bound 1a ((CO₂)₀.₈@1a-195K) were obtained by keeping single crystals of 1a in CO₂ at ~1.5 bar and at ambient temperature for an hour, and then cooling down to 195 K. The structure analysis revealed that CO₂ was ligated to unsaturated metal site using its Lewis basic oxygen atom in an angular η¹-(O₂) coordination mode with a 0.40(1) site occupancy factor (0.8 CO₂ molecule per cage) (Figs 2f and S8). The Cu – O₁CO₂ distance, 2.94(2) Å, is slightly longer than the Cu – O₀MeOH distance, 2.61(7)(Å), of the bound MeOH in the as-synthesized crystal but still significantly shorter than the sum of their van der Waals radii sum (3.55 Å)¹⁴. The Cu – O₁CO₂ distance is, much longer than the corresponding Ni – O₁CO₂ bond distances in Ni-MOF-74 structure obtained from the PXRD experiment, 2.29(2) Å²²; the corresponding Mg – O₁CO₂ bond distances in Mg-MOF-74 structures obtained from the NPD experiments, from 2.24(3) Å to 2.39(6) Å³¹, but is comparable to that of Cu-MOF-74, 2.86(3) Å³². It is well-known that the Lewis acidic carbon atom of the CO₂ molecule can interact with the Lewis basic nitrogen atom in the pore environment¹⁶–¹⁹,²⁹. Similar Lewis acid-base interaction is also observed in the CO₂-bound 1a. The Lewis acidic carbon atom (C1C) of the bound CO₂ molecule is interacting with the two Lewis basic nitrogen atoms (N3) of two symmetry-related t₇ ligands. The distance between C1C and N3 (3.21(2) Å) is slightly shorter than their van der Waals sum (3.30 Å). The Cu₂–O₁C–C₁C bond angle (124(2)°) is in the range those in Ni-MOF-74, Mg-MOF-74, and Cu-MOF-74 structures (from 117(2) to 144(2)°). The observed site occupancy factor of the ligated CO₂ molecule, 0.40(1), is due to steric repulsion between the CO₂ molecules bound to the symmetry-related Cu(II) sites in a cage. The interatomic distance between the symmetry-related ligated oxygen atoms of the bound CO₂ molecules (2.64(7) Å) is much closer than their van der Waals radii sum (3.1 Å) (Fig 2f). The steric repulsion between the CO₂ molecules hinders simultaneous ligation of two CO₂ molecules in the same cage. The geometry of the bound CO₂ at 195 K is similar to that of a reported free CO₂ structure. The C – O bond distances, 1.12(3) Å and 1.13(3) Å, in the bound CO₂ structure are not distinguishable to that in the free CO₂ structure, 1.155(1) Å³³. The little variation in the C – O bond distance in the bound CO₂ is due to the combined effect of two different interactions between the CO₂ molecule and the framework. The CO₂ molecule donates its σ electron to the metal center through the ligating oxygen atom but simultaneously accepts σ electron from the two Lewis basic nitrogen atoms of the t₇ ligands through the Lewis acidic carbon atom of the bound CO₂. The O – C – O bond angle in the bound CO₂ structure, 176(3)°, is also nearly linear as that in the free CO₂ structure.

The single crystal of (CO₂)₀.₈(H₂O)₀.₁₅@1a-296K-5h (the single crystal of CO₂@1a-195K stood at ambient condition for five hours) showed the bound CO₂ molecule in the cage with reduced site occupancy, 0.13(2) (Fig 2g). Though the CO₂ molecule is still ligated to the metal center, the Cu – O bond distance of (CO₂)₀.₈(H₂O)₀.₁₅@1a-296K-5h at 296 K, 3.09(5) Å, is elongated by ~0.15 Å than that of CO₂@1a-195K at 195 K but is still slightly shorter than their van der Waals sum of Cu and O atoms. However, interestingly, the Lewis acidic carbon atom (C1C) of the bound CO₂ molecule is no longer interacting with the Lewis basic nitrogen atom (N3) of the ligand. The distance between C1C and N3 (3.34(4) Å) is slightly longer than their van der Waals radii sum, 3.30 Å. The complete replacement of the bound CO₂ molecules in the pore by the water molecules in air took 10 days at ambient condition (Figs S8 and S9).

The vibrational mode analysis of the CO₂ molecule encapsulated in a cage. Figure 3 shows temperature-dependent infra-red (IR) spectra of the MOF, 1a, filled with ¹³C CO₂ molecules in the bending and asymmetric stretch regions (see Fig S10 for whole-range spectra). In the IR spectra of the bending region given in Fig 3a, two separate peaks are observed, which is distinctly different from the case of a free CO₂ molecule where only a single bending peak appears because its two bending modes are degenerate. This observation suggests that the interaction between a CO₂ molecule and 1a disturbs its bending motion to generate two separate peaks. The center frequency of the two peaks is also red-shifted by 11.7 cm⁻¹ compared to the degenerate bending frequency of a free CO₂ of 667.3 cm⁻¹¹³. Similar splitting of the bending modes and red-shift of the center frequency has been reported in the IR spectra of the CO₂ bound in an η¹-(O₂) coordination mode in Ni-MOF-74¹⁰. The splitting of the two bending modes must be related to the binding energy of a CO₂ molecule to the framework and a larger separation indicates stronger interaction between them. The separation between the two peaks decreases as the temperature increases from 5.49 cm⁻¹ at 103 K to 4.92 and 4.67 cm⁻¹, respectively, at 293 and 423 K. Provided that the binding energy is proportional to the separation, the binding energy at 293 and 423 K, respectively, decreases by 11 and 15% compared with that at 103 K. The CO₂ molecule encapsulated in the cage is less strongly bound to the framework at elevated temperatures. These observations are in agreement with the temperature dependent crystal structures of CO₂@1a. Though the bound CO₂ structures themselves from the temperature dependent single crystal structure study are not as sensitive as the bending modes of the bound CO₂ molecule, the temperature dependent interactions between the CO₂ and the framework are in good agreement with the temperature dependent bending modes of the bound CO₂ molecule. While the peak separation of the two symmetric bending modes of the bound CO₂ molecule is temperature dependent, the center frequency is insensitive to temperature. Temperature insensitivity of the center frequency is due to the combined result of simultaneous weakening of both the sigma electron donation to the Cu(II) ion through the Lewis basic oxygen atom of the bound CO₂ and the sigma electron acceptance from the nitrogen atoms of the t₇ ligands via the Lewis acidic carbon atom of the bound CO₂ molecule.

Figure 3b shows that the main asymmetric stretching band at ~2335 cm⁻¹ with a FWHM of ~7 cm⁻¹ is red-shifted by 14 cm⁻¹ compared to the corresponding band of free ¹²C CO₂ at 2349 cm⁻¹⁻¹³. The red-shift indicates that the σ electron donation through the Lewis basic oxygen atom of the bound CO₂ to the Cu(II) center is slightly stronger than the σ electron acceptance through the carbon atom of the bound CO₂ from the nitrogen.
atoms of the tz ligands. An asymmetric stretching band shift is not a good indicator of the binding strength of an encapsulated CO2 to a framework. It is a combined effect of electron donation and acceptance of the encapsulated CO2 to and from the framework. The encapsulated CO2 in the series of isostructural MOFs, M-MOF-74 (where, M = Ni(II), Mg(II), Zn(II), and Co(II)), showed different shifts of the asymmetric CO2 stretching band depending on metal ions even though the frameworks are isostructural. 

The side bands at the high-frequency side are likely to originate from slightly different local structures of CO2 inside the cage. At the lower-frequency side of the main band, two peaks, each of which is enclosed with a rounded rectangle, are observed. The temperature dependent peak at ~2323 cm⁻¹ was assigned as a vibrational hot band coming from the transition, (ν₁, ν₂, ν₃) = (0, 1, 1) → (0, 2, 0). The assignment is further clarified by the hot-band spectra in the bending region shown in Fig. 3c, where the magnitude of the peaks follows the Boltzmann distribution. However, the peak at ~2327 cm⁻¹, which shows similar temperature dependence as the peak at ~2323 cm⁻¹, was assigned to be a peak arising from CO2 double occupancy in a single cage (two CO2 molecules in a single cage interacting to each other). Figure 3d shows the asymmetric stretch region of another sample with low CO2 content in the spectral region of the asymmetric stretch. Note that the spectra in (a, b, and c) were obtained from the same sample (high content of CO2), whereas the spectra in (d) were obtained from a different sample (low content of CO2).

Figure 3. Temperature- and CO2 concentration-dependent FTIR spectra of 1a filled with ¹²CO₂. (a) Spectra in the bending region of ¹²CO₂, Δν₁₀³K(5.49 cm⁻¹), Δν₂₉₃K(4.92 cm⁻¹), and Δν₄₂₃K(4.67 cm⁻¹) represent the separation between the two peak maxima at 103, 293, and 423 K, respectively. Similar spectra are reported for 1a filled with ¹³CO₂ in Fig. S11. (b) Spectra in the asymmetric stretch region of ¹²CO₂. The peaks enclosed with rounded rectangles in the spectral region of ~2323 and ~2327 cm⁻¹ represent a vibrational hot band (starting from the first excited state of the bending mode) and a peak originating from the existence of 2 CO2 molecules in a single cavity, respectively. (c) Spectra in the spectral region of ν₁ = 1 → ν₂ = 2 transition (hot band). (d) Spectra of 1a with low CO2 content in the spectral region of the asymmetric stretch.
sample that contains less amount of CO₂ by a factor of 3. The integrated areas of the peaks in the spectra of the low CO₂ content sample are about 1/3 of those of their corresponding spectra (Fig. S12). The double-occupancy peak is hardly observable at 2327 cm⁻¹ indicating that the integrated area of the peak is not linearly proportional to CO₂ concentration and the peak must be from the double occupancy. In the spectra of the low CO₂ content sample, the peaks are narrower than their corresponding ones in the high-content spectra. This observation suggests that the skeleton of 1a is rather flexible and the structural distribution of the encapsulated CO₂ becomes more inhomogeneous with increase in CO₂ content.

The observation that the magnitude of the double-occupancy peak increases as the temperature increases (Fig. 3b) should also be paid attention to. A single CO₂ molecule in a single cage is preferred at lower temperature under lower concentration of CO₂. The steric repulsion between the CO₂ molecules ligated to the two symmetry-related metal sites in the same cage at low temperature is responsible for this single occupancy. Two CO₂ molecules in a single cage are allowed at high temperature under higher CO₂ content. The reduced restraint on the position of the weakly interacting CO₂ molecule in the cage at high temperature allows two CO₂ molecules in a single cage. The existence of the double-occupancy peak helps us to understand the process of CO₂ filling into the 1-D microporous channels, made of small cages interlinked with small neck-like portal windows. Not only a hopping process of a CO₂ molecule into an empty cage but also a filling process of a second CO₂ molecule into the cage with an encapsulated CO₂ are operating.

CO₂ sorption behaviors. Even though 1a had microporous 1-D channel, it does not show any N₂ (at 77 K and 308 K), H₂ (at 77 K) and CH₄ (at 195 K and 308 K) adsorptions (Fig. S13). However, 1a shows reversible CO₂ uptake at 273 K even though the kinetic diameter of CO₂ (3.3 Å) is much larger than the dimension of the portal window and the isotherms are of a typical type I (Fig. 4a). The large quadruple moment of CO₂ might be responsible for such uptake behavior. The CO₂ molecule can interact with the framework strong enough to induce the portal dimension of the 1-D microporous channels large enough for the reversible CO₂ adsorption and desorption. The total CO₂ uptake amount of 1a at 273 K and 1 bar, 33.9 cm³/g, corresponds to 0.82 CO₂ molecules per cage (or 0.41 CO₂ molecules per unsaturated Cu(II) center). The CO₂ sorption isotherms of 1a at 298 K are also reversible type I isotherms and the total CO₂ uptake amount at ~1 bar is 37.4 cm³/g, which is slightly larger than the CO₂ uptake amount at 273 K and ~1 bar. The increased CO₂ uptake amount at the higher temperature is due to increased double occupancy of CO₂ molecules in a single cage and subsequent pore filling up to the cages located at the inner parts of the microporous 1-D channels. The total CO₂ uptake amount at 313 K and ~1 bar, 31.3 cm³/g, is slightly smaller than that at 298 K. The reduced CO₂ uptake amount is due to decreased CO₂-framework interaction at the higher temperature. The adsorption enthalpy (−ΔH_ads) of CO₂ on 1a calculated using the adsorption isotherms at 298 K and 313 K is −32 − 25 kJ/mol (Fig. S14), which will be the lower bound because the isotherms obtained are not in the true equilibrium condition due to slow adsorption kinetics of CO₂. The adsorption enthalpy (−ΔH_ads) of CO₂ estimated using desorption isotherms at 298 K and 313 K is −41 − 35 kJ/mol, which is comparable to the CO₂ adsorption enthalpy on Mg-MOF-74, ~40 kJ/mol, but is much larger than that on Cu-MOF-74, ~22.1 kJ/mol.

Interestingly, the CO₂ sorption on 1a at 195 K is irreversible (Fig. 4b). The CO₂ adsorption isotherm at 195 K shows steep rise at very low pressure. The adsorption reaches to its maximum amount, 24.3 cm³/g, at 0.022 bar. The amount of adsorbed CO₂ corresponds to ~0.60 CO₂ molecule per cage (or ~0.30 CO₂ molecule per unsaturated metal site), which is much smaller than the expected maximum amount, one CO₂ molecule per cage (or 0.5 CO₂ molecule per unsaturated metal site). Significant portions of the inner cages in the 1-D porous channel may not be accessible for CO₂ molecule because the channels are already blocked by the CO₂ molecules bound strongly to the framework as shown in Fig. 2d. There is no indication of CO₂ desorption even at 0.0015 bar.

At high temperature, not only hopping process of a CO₂ molecule into an empty cage but also filling process of a second CO₂ molecule into the cage with an encapsulated CO₂ molecule are allowed so that even most of the empty inner cages of the 1-D channels could be occupied with CO₂ molecules and the CO₂ molecules in the inner cages the 1-D channels can also be emptied via a reverse process. However, at low temperature, the filling of a second CO₂ molecule into the cage with an encapsulated CO₂ molecule is severely hindered because the CO₂ molecule bound strongly at the open metal center does not allow the second CO₂ molecule into the same cage. At
low temperature, the main pore filling is via hopping process so that the maximum amount of CO₂ captured in the 1-D channels is significantly smaller than the expected maximum amount of CO₂. When multiple consecutive cages in a 1-D channel are filled with CO₂ molecules, desorption of CO₂ is also severely hindered, which is the cause of the irreversible CO₂ sorption behavior of 1a at 195 K.

Conclusions
Although each cage contains two unsaturated Cu(II) sites and the cage size is large enough for two CO₂ molecules, the CO₂ uptake amount is only ~0.9 CO₂ molecule per cage at 298 K and 1 bar, which is due to the steric repulsion between the two bound CO₂ molecules in the cage. Temperature dependent single crystal structure analyses revealed that the interaction between the bound CO₂ molecule and the framework of the MOF is temperature-dependent. As temperature increases, both the CO₂ coordination to the unsaturated Cu(II) center and the Lewis acid—base interaction between the carbon atom of the CO₂ molecule and the nitrogen atom of the ligand are simultaneously weakened.

The red-shifted vibrational peaks of the bound CO₂ molecule indicate that the electron donation to the metal center through the oxygen atom of the CO₂ molecule is slightly larger than the electron acceptance from the nitrogen atom of the ligand through the carbon atom of the CO₂ molecule. The temperature dependent separation of the two bending peaks also supports the temperature dependent interaction between the CO₂ molecule and the framework of the MOF. As temperature increases, the separation of the bending peaks decreases. The observation tells that the CO₂-framework interaction decreases as the temperature increases as observed in the temperature dependent structure analyses of a single crystal of CO₂@1a.

The reversible CO₂ uptakes on 1a were observed at 298 and 313 K, respectively, even though the portal dimension of the cage along the microporous 1-D channel is smaller than the kinetic diameter of CO₂ molecule. The CO₂ uptake amount per unsaturated Cu(II) center at 298 K, 1 bar is 0.45 molecule, which is only slightly smaller than the probable maximum value, 0.5. On the other hand, the irreversible CO₂ uptake was observed at 195 K and the CO₂ uptake amount per unsaturated Cu(II) center is only 0.30 molecule, which is much smaller than the expected value, 0.5. The decreased uptake amount at the lower temperature is due to the strongly interacting CO₂ molecules in the cages aligned along the microporous 1-D channel hindering the other CO₂ molecules to access the available inner cages in the 1-D channel. The different uptake behaviour at different temperature is related to both the small portal dimension of the cages aligned along the 1-D channel and the different strength of CO₂-framework interaction in the cage. The reversible CO₂ uptakes were observed at the higher temperature since the CO₂ molecule interacting weakly with the framework could have proper orientation for the escape from the cavity through the small portal of the cages. However, the irreversible CO₂ uptake was observed at the lower temperature since the CO₂ molecule interacting strongly enough with the metal center in the cage could not have proper orientation for the escape from the cavity through the small portal.

Methods
Synthesis of the MOF, Cu₃Cl₂(tz)₄(MeOH)₂, 1. A 27.3 mg amount of CuCl₂ (0.203 mmol) was dissolved in 4 mL MeOH in a 10 mL vial, and then 0.90 mL 0.45 M tetrazole (tz) acetonitrile solution (0.41 mmol) was added into the solution. The Teflon-sealed vial was heated to 70 °C for 7 days and then slowly cooled down to ambient temperature. The block-shaped blue crystals obtained were washed using 10 mL fresh methanol and filtered.

IR spectroscopy. The samples were in the form of a KBr pellet of a ~200-μm thickness and a 13-mm diameter. The samples were located in a cell composed of two circular KBr windows of a 25-mm diameter. The windows were separated by a 200-μm thick spacer with 25 and 20 mm outer and inner diameters, respectively. FTIR spectra were recorded by a Varian 7000e FTIR spectrometer. The samples with KBr windows were placed in a home-made oxygen-free high thermal conductivity (OFHC) copper cell, and the temperature of the samples was controlled and maintained in a variable temperature cell holder (GS21525, Specac, UK). Details of the temperature-controllable cell assembly have been reported elsewhere.

Gas sorption measurements. All gas sorption isotherms were measured using a BELSORP-max (BEL Japan, Inc.) with a standard volumetric technique.

References
1. Creamer, A. E. & Gao, B. In Carbon dioxide capture: an effective way to combat global warming, Ed. Sharma, S. K. (Springer, Switzerland 2015).
2. Figueroa, J. D. et al. Advances in CO₂ capture technology—The U.S. department of energy’s carbon sequestration program. Int. J. Greenh. Gas Control 2, 9 (2008).
3. Benson, E. E. et al. Electrocatalytic and homogeneous approaches to conversion of CO₂ to liquid fuels. Chem. Soc. Rev. 38, 89 (2009).
4. Cokoja, M. et al. Transformation of carbon dioxide with homogeneous transition-metal catalysts: A Molecular Solution to a Global Challenge? Angew. Chem., Int. Ed. 50, 8510 (2011).
5. Omae, I. Recent developments in carbon dioxide utilization for the production of organic chemicals. Coord. Chem. Rev. 256, 1384 (2012).
6. Schoedel, A. et al. The role of metal–organic frameworks in a carbon-neutral energy cycle. Nature Energy 1, 16034 (2016).
7. Sumida, K. et al. Carbon dioxide capture in metal—organic frameworks. Chem. Rev. 112, 724 (2012).
8. McDonald, T. M. Cooperative insertion of CO₂ in diamine-appended metal—organic frameworks. Nature 519, 303 (2015).
9. Kumar, A. et al. Direct air capture of CO₂ by physiosorbent materials. Angew. Chem. Int. Ed. 54, 14372 (2015).
10. Xu, H.-Q. et al. Visible-light photoreduction of CO₂ in a metal—organic framework: boosting electron—hole separation via electron trap states. J. Am. Chem. Soc. 137, 13440 (2015).
11. Darensbourg, D. J. et al. Sequestering CO₂ for short-term storage in MOFs: copolymer synthesis with oxiranes. ACS Catal. 4, 1511 (2014).
Temperature dependent CO2 behavior in microporous 1-D channels of a metal-organic framework with multiple interaction sites.

© The Author(s) 2017

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/

© The Author(s) 2017