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Chlorination of a Zeolitic-Imidazolate Framework Tunes Packing and van der Waals Interaction of Carbon Dioxide for Optimized Adsorptive Separation

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ABSTRACT: Molecular separation of carbon dioxide (CO₂) and methane (CH₄) is of growing interest for biogas upgrading, carbon capture and utilization, methane synthesis and for purification of natural gas. Here, we report a new zeolitic-imidazolate framework (ZIF), coined COK-17, with exceptionally high affinity for the adsorption of CO₂ by London dispersion forces, mediated by chlorine substituents of the imidazolate linkers. COK-17 is a new type of flexible zeolitic-imidazolate framework Zn(4,5-dichloroimidazolate)₂ with the SOD framework topology. Below 200 K it displays a metastable closed-pore phase next to its stable open-pore phase. At temperatures above 200 K, COK-17 always adopts its open-pore structure, providing unique adsorption sites for selective CO₂ adsorption and packing through van der Waals interactions



with the chlorine groups, lining the walls of the micropores. Localization of the adsorbed CO₂ molecules by Rietveld refinement of X-ray diffraction data and periodic density functional theory calculations revealed the presence and nature of different adsorption sites. In agreement with experimental data, grand canonical Monte Carlo simulations of adsorption isotherms of CO₂ and CH₄ in COK-17 confirmed the role of the chlorine functions of the linkers and demonstrated the superiority of COK-17 compared to other adsorbents such as ZIF-8 and ZIF-71.

■ INTRODUCTION

Adsorption of carbon dioxide on a porous material for its separation from methane in CO₂/CH₄ gas mixtures is a green key technology compared to conventional amine scrubbing. For renewable fuel production, CO₂/CH₄ separation is an essential step for upgrading biogas and sustainable methane production from carbon dioxide via carbon capture and utilization (CCU) strategies.^{2,3} Metal-organic frameworks (MOFs) composed of metal ions/clusters and organic linkers are particularly useful for designing selective adsorbents for CCU because of their exceptional host-guest interactions.^{4,5} Especially, tuning of the chemical functionality of the linker can enhance the interaction with CO2.6 Furthermore, MOFs can easily be structured into macroscopic bodies⁷ or dispersed into membranes, 8,9 desirable for use in direct separation in traditional PSA or TSA operations, 10 or as MOF-based membranes. 11-18

Zeolitic-imidazolate frameworks (ZIFs), a subfamily of MOFs, consist of tetrahedrally coordinated transition-metal cations, mostly Zn²⁺ or Co²⁺, bridged by imidazolates and display zeolite-like topologies (e.g., SOD, LTA, and RHO

zeolite structure types). ^{19,20} They have attracted considerable attention as molecular sieves for gas separation because of their hydrophobic cages, framework flexibility, and considerable chemical and thermal stability. ²¹ Hundreds of ZIF structures have been synthesized to date. ^{19,20} However, with use of the 4,5-dichloroimidazolate (dcim) linker, until now only two ZIF polymorphs have been synthesized directly, microporous RHO-type ZIF-71 and nonporous lcs-type ZIF-72. ²⁰ The corresponding [Zn(dcim)₂]-SOD has been obtained indirectly via solvent-assisted ligand exchange ²² or modulated synthesis. ²³ To the best of our knowledge, a direct one-pot synthesis of Zn(dcim)₂ with SOD topology has not yet been reported.

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RESULTS AND DISCUSSION

Synthesis, Characterization and Gas Adsorption Properties. Phase-pure Zn(dcim)₂, coined COK-17 (COK-17: Centrum voor Oppervlaktechemie en Katalyse No. 17), with SOD topology was obtained by direct one-pot solvothermal synthesis. COK-17 crystallizes from a zinc nitrate and 4,5-dichloroimidazoleDMF/water solvent mixture. Scanning electron microscopy (SEM) shows COK-17 crystals as well-defined, micrometer-sized rhombohedra (Figure S1, Supporting Information). The high crystallinity of the material was confirmed by X-ray diffraction (XRD, Figure 1). The

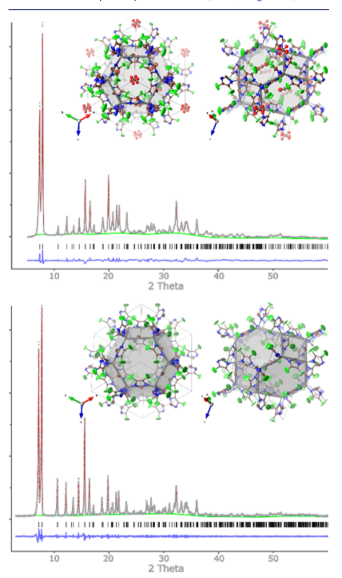


Figure 1. Room temperature XRD pattern and Rietveld refinement of $Zn(dcim)_2$ COK-17. Rietveld refinements of activated COK-17 (bottom) and saturated with CO_2 at a pressure of 1 bar and a temperature of 293 K (top). Framework with highlighted SOD-type cage are shown in the insets.

diffractogram could be excellently described in the rhombohedral space group $R\overline{3}$ with a=b=c=14.10 Å and $\alpha=\beta=\Upsilon=107.23^\circ$. Rietveld refinement^{24,25} revealed the SOD topology with the same arrangement of cations and linkers as found in ZIF-8, but elongated along the space diagonal of the cubic structure, resulting in the decreased angle of 107.23° between

the rhombohedral axes (Figure 1, inset, see Supporting Information for more details). COK-17 is thermally stable up to 673 K in nitrogen, according to thermogravimetric analysis (Figure S2). COK-17 is stable for at least 2 years when stored under ambient humidity. Line broadening of the Bragg reflections indicated the rhombhedral angle varied in the stored sample. After activation at 150 °C in vacuum, the sample returned to its original behavior. Exposure of COK-17 to water at room temperature shows no effect, while upon steaming the unit cell expands to pseudocubic dimensions, though it still is best described in the original rhombohedral space group $R\overline{3}$ (see Figure S3). Modeling the structure using the FOX simulation software, 26 using an unstructured electron density in a large cage, resulted in an excellent fit of the data, demonstrating the connectivity between the linker and Zn ions persisting during steaming.

According to the structure determination by Rietveld refinement, COK-17 at room temperature is always observed in its open-pore state, independent of whether it is fully evacuated or contains adsorbed molecules. The potential occurrence of structural flexibility of the empty COK-17 framework was investigated by force-field-based molecular dynamics simulations at different temperatures and pressures, a method that was shown to provide accurate information on the flexibility of MOFs.²⁷ The free energy of the empty COK-17 framework as a function of its unit cell volume at different temperatures and pressures was estimated, following the procedure explained in the Supporting Information (see computational section "Force field simulations").28 The resulting free energy profiles at various temperatures (Figure S4 and Table S1) clearly reveal that the open-pore phase is the only stable phase at temperatures above 200 K. At lower temperatures, a metastable closed-pore phase appears next to the stable open-pore phase. As a result, host-guest interactions may steer the system from its open-pore phase to the closedpore phase upon adsorption at low temperatures, even though the closed-pore phase of the empty host remains less stable than its open-pore state. The occurrence of such a structural variation was also hinted at by the observation of a stepped Type I N₂ adsorption isotherm at a temperature of 77 K (Figure S5). Two distinctive steps with hystereses are observed in the relative pressure (P/P_0) range of 0.15–0.25 and 0.5–0.8, respectively. In flexible frameworks, such behavior can be due to framework rearrangement and reorganization of the adsorbate. 29-31 The peculiar double-stepped adsorption profile indicates a similar breathing mechanism as recently reported for the isostructural Zn(dcim)₂-SOD material, which also shows two different hysteresis loops in argon physisorption.²³ According to the N₂ adsorption branch, the low-temperature form of COK-17 has a micropore volume of 0.26 cm³ g⁻¹ (plateau I of the isotherm in Figure S5a) and a micropore diameter of 1.2 nm according to nonlinear density functional theory. Brunauer-Emmet-Teller (BET) and Langmuir specific surface areas amount to 500 and 628 m² g⁻¹, respectively.

Adsorption of CO_2 and CH_4 molecules on COK-17 powder at ambient temperature was investigated by determination of adsorption isotherms for CH_4 and CO_2 , by powder XRD and structure refinement of COK-17 saturated at 1 bar pressure with the two gases, and by computational modeling. The experimental adsorption isotherms (Figure 2a) reveal the high affinity of COK-17 for CO_2 and CH_4 , with uptakes at 1 bar and 298 K of 2.5 and 1.24 mmol g^{-1} , respectively. These values are

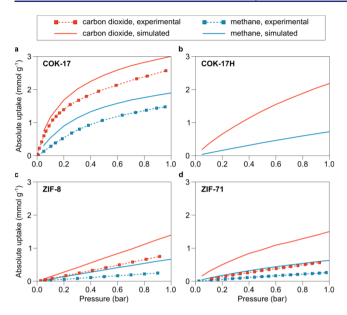


Figure 2. Experimental and simulated single-component adsorption isotherms of CO_2 and CH_4 collected at 298 K. Experimental (dotted lines with symbols) and computationally simulated (full lines) adsorption isotherms of CO_2 and CH_4 on (a) COK-17, (b) a hypothetical isostructural COK-17H framework with CI atoms systematically replaced with CI atoms, and (c) CIF-8 and (d) CIF-71 benchmark materials for which experimental adsorption data were taken from refs33 34, and36

much higher compared to those of related ZIF materials, like ZIF-8, where corresponding CO₂ and CH₄ uptakes are limited to 0.7 and 0.25 mmol g⁻¹, and ZIF-71 with uptakes of 0.6 and 0.27 mmol g⁻¹ only (Figures 2c,d). $^{32-35}$ Breakthrough separation experiments on COK-17, performed at 303 K, revealed its separation performance for binary gas CO₂/CH₄ mixtures with different compositions (CO₂/CH₄ = 20:80 and 50:50 v/v) (Figure S6 in the Supporting Information). The effluent from the bed was monitored by mass spectrometry. The breakthrough curves show pure CH₄ as the first eluent, indicating a high selectivity of COK-17 for CO₂ over CH₄ under flow conditions. The retention time for CH₄ over COK-17 was 100 s in CO₂/CH₄ (20:80) mixed-gas flow versus 50 s in an equimolar CO₂/CH₄ mixture. These values correspond to CO₂ adsorption capacities of 0.40 and 0.50 mmol g⁻¹ at 303 K and partial pressures of 0.2 and 0.5 bar, respectively.

We calculated the isosteric heats of ${\rm CO_2}$ adsorption for COK-17, according to the Clausius—Clapeyron eq (eq 1) from the adsorption isotherms measured at 273 and 298 K:

$$q_{st} = -R[\Delta(\ln P)/\Delta(1/T)]N \tag{1}$$

where $q_{\rm str}$ R, P, T, and N correspond to isosteric heat of adsorption, gas constant, pressure, temperature, and amount of adsorbed ${\rm CO_2}$, respectively. The results are presented in Figure S7 and in agreement with a diminution of the heat of adsorption with increasing surface coverage, which agrees with the proposed physisorption process. The obtained values $(-28.5 \text{ to } -23.5 \text{ kJ mol}^{-1})$ are in agreement with the DFT calculation values for the nondistorted structure.

Host-Guest Interaction. The CO₂ and CH₄ adsorption isotherms on COK-17 at 298 K did not show any steps that would indicate a structure change upon adsorption (Figure 2a, Figures S8 and S9). Given the large uptake, localization of the adsorbed CO₂ molecules by Rietveld refinement of X-ray

diffraction data was feasible, as shown in Figure 1.^{24,25} The diffraction experiments clearly indicated two possible positions for adsorbed CO₂, which are situated in the symmetric and the distorted 6-ring (6R) windows of the SOD cavities, located at the origin and the faces of the rhombohedral unit cell, respectively (Figure 3). While the symmetric 6-ring (site A)

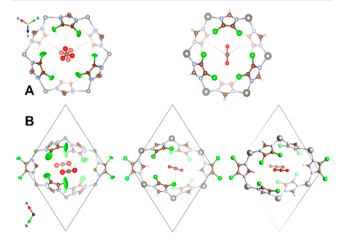


Figure 3. CO_2 sites in COK-17. Top row: adsorption site A for CO_2 in symmetric 6R of COK-17 (left, experiment; right, simulation). Bottom row: adsorption sites(s) B for CO_2 in distorted 6R of COK-17 (left, experiment; middle and right, simulation).

contains only one CO_2 molecule, two crystallographically equivalent sites for CO_2 were found in the distorted rings (site B), which are both occupied simultaneously by a guest molecule at a pressure of 1 bar (Figure 3). Refinement of the structure of COK-17 loaded with CO_2 at 1 bar revealed an overall loading of ca. 0.7 CO_2 molecules per $Zn(dcim)_2$ formula, in agreement with the experimental value of ca. 0.8. The theoretical maximum with all sites A and B occupied corresponds to 1.0 CO_2 per $Zn(dcim)_2$.

Further insight in the adsorption sites was obtained from periodic density functional theory (DFT) calculations performed on the experimentally determined unit cell of the COK-17 lattice. Stable adsorption sites of 1 molecule of CO₂ on sites A and B were found with similar adsorption energies, namely, $E_{\rm ads} = -28 \text{ kJ mol}^{-1}$ on site A and $E_{\rm ads} = -31 \text{ kJ mol}^{-1}$ on site B. This allows them to appear identical in an experimental adsorption process so that both sites are expected to contribute simultaneously to CO₂ adsorption. A similar conclusion for CH₄ can be drawn: at site A an adsorption energy of -25 kJ mol⁻¹ is calculated, while at site B a slightly more stable value of -27 kJ mol^{-1} is found. As the singlemolecule adsorption energies favor CO2 over CH4, it can already be inferred that CO2 will certainly be preferably adsorbed at low pressures, which is discussed in more detail later on (vide infra).

The adsorption of two CO₂ molecules in the distorted 6Rs of site B was simulated next. A distinct energetic minimum upon adsorption of two molecules in position B, closely resembling the refined structure, was found (Figure 3). The total adsorption energy for this configuration amounts to -55 kJ mol⁻¹, which should result in a free energy release of 24 kJ mol⁻¹ upon adsorption of the second molecule on site B, assuming the first molecule contributes -31 kJ mol⁻¹. Interestingly, the large cage of COK-17 appeared empty in the diffraction study and showed significantly lower stabiliza-

tion of CO_2 molecules by DFT as compared to sites A and B. In accordance with simulation and structure determination, the experimental adsorption isotherm for CO_2 on COK-17 could not be fitted with a single-site Langmuir model (Figure S10), but only be described with a dual-site model with different adsorption energies. In contrast, the adsorption of the larger CH_4 molecule on COK-17 could be well described by a single-site Langmuir model.

Molecular Simulations. The exceptional adsorption performance of COK-17 compared to ZIF-8 and ZIF-71 (Figure 2) suggests that the presence of Cl atoms in the imidazole linkers is responsible for the strong adsorption on sites A and B. For an understanding of the exceptional behavior of COK-17 in terms of its structure and Cl content, adsorption capacities were simulated for four different ZIF frameworks, which all contain 6-rings, though with different chemical functionalities and/or topologies (see Table S2). Like COK-17, ZIF-8 with 2-methylimidazole linkers displays the SOD topology but in a cubic undistorted configuration. ZIF-71, on the other hand, shows the same chlorinated linkers as COK-17, but manifests in the RHO rather than in the SOD topology. For specifically the gaining of insight into the effect of the Cl substituents, a purely theoretical framework with the same structure as COK-17 but with the Cl atoms substituted with H atoms was also investigated. Hereafter, this structure will be referred to as COK-17H.

The adsorption isotherms of CO₂ and CH₄ in COK-17, COK-17H, ZIF-8, and ZIF-71 were estimated using grand canonical Monte Carlo (GCMC) simulations with rigid frameworks according to their experimental geometry and using the Dreiding-MBIS/TraPPE model (for more details, see the Supporting Information). The simulated values show the same trends as the experimental isotherms, although they slightly overestimate the experimental observations, especially for ZIF-71. COK-17 outperforms COK-17H, ZIF-8, and ZIF-71 with respect to the affinity for CO₂ and methane. All frameworks systematically show a higher adsorption capacity for CO₂ compared to CH₄.

Minimum energy locations of the guest molecules computed for COK-17, COK-17H, ZIF-8, and ZIF-71 highlight the differences even further. Of all these frameworks only the linker conformation in COK-17 shows inversion centers in the 6R of the structure (Figure 4). This leads to the formation of two-sided cradles formed from six or four linkers on sites A and B, respectively (Figure 4). In ZIF-71 only one-sided cradles are expressed in the 6Rs, while in ZIF-8 the three linkers in the 6Rs are in an almost coplanar arrangement with the ring (Figure 4). Energy minimization by DFT revealed that the low energy configuration of CO2 in both frameworks is not located in the 6Rs, but in the large cavities, close to a 4R in ZIF-71 and at a rather unspecific site for ZIF-8 (Figure 4). ZIF-8 shows no expressed energetic minimum in the structure at large, with the configurations of CO2 in the 6Rs similar in energy compared to the global minimal energy in the large cavities. The adsorption energies for all sites are listed in Table S3.

The contribution of London dispersion to the adsorption energy (Table S3 in the Supporting Information) shows that London dispersion forces dominate the adsorption energies irrespective of the particular framework. Among all sites in the different ZIFs, the distorted 6-rings in COK-17 have the highest $\rm CO_2$ adsorption energy ($\rm -31~kJ~mol^{-1}$) and the highest London dispersion interaction ($\rm -28~kJ~mol^{-1}$). The presence

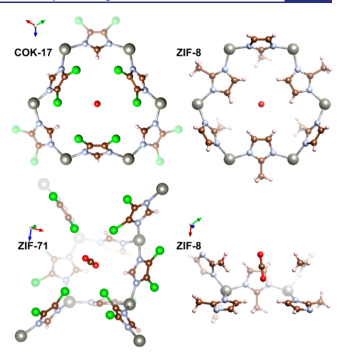


Figure 4. Adsorption sites of CO_2 as determined from simulations. Top left, 6R COK-17; top right, 6R ZIF-8. Bottom left, 4R ZIF-71; bottom right, large cage ZIF-8.

of Cl substituents on the imidazole linker leads to increased van der Waals (vdW) interactions by the enhanced dispersion forces of the electron-rich Cl atom as compared to that of N, C, and H atoms of imidazolate linkers devoid of it. COK-17H has the same structure as COK-17, but does not feature Cl substituents. DFT calculations for molecules on sites A and B in symmetric and distorted 6-rings systematically revealed increased adsorption energies in the presence of Cl-substituted linkers, especially for CO_2 on site B, but even more strongly for CH_4 on both sites, highlighting the impact of chlorination.

Figure 5a shows the selectivities of CO2 versus CH4, extracted from GCMC simulations of CO₂/CH₄ mixtures with different compositions. As a reference, also the experimental selectivities derived from the measurement of singlecomponent adsorption isotherms using the Ideal Adsorbed Solution Theory (IAST) are shown.³⁷ The CO₂ selectivity of ZIF-8 is the lowest, followed by ZIF-71. Both show only slight variations with respect to pressure or composition. COK-17 exhibits varied CO₂/CH₄ selectivities depending on CO₂/CH₄ ratios. The CO₂/CH₄ mixture composition does not affect the CO₂/CH₄ selectivity of ZIF-8 nor of ZIF-71. COK-17 shows the highest selectivity, which, interestingly, increases with pressure and significantly depends on the CO2 content. For instance, the calculated mixed-gas selectivity of COK-17 at 1 bar in a 80:20 binary CO₂/CH₄ mixture composition was 4, while a 50:50 mixture gave a slightly decreased selectivity of 3.5, followed by a selectivity of 3.25 achieved in a 20:80 binary CO₂/CH₄ mixture. The ideal selectivity of CO₂ over CH₄ increases at higher pressures from 1 to 10 bar for different CO₂/CH₄ mixtures according to the results obtained using GCMC simulations. Similarly, the simulation results demonstrate that lower CO₂/CH₄ selectivities were obtained for COK-17 in the mixtures containing an excess of methane. For instance, the calculated ideal selectivity of COK-17 at 10 bar in a 80:20 binary CO₂/CH₄ mixture was 7.8, while a 50:50 mixture gave a slightly decreased selectivity of 7.0 based on

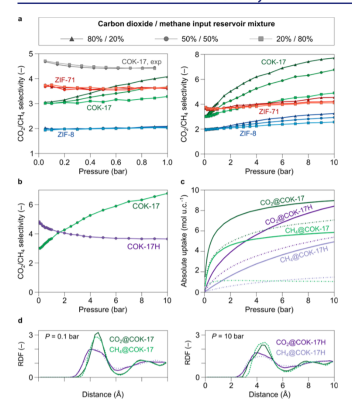


Figure 5. Mixed-gas CO_2/CH_4 selectivity. (a) Static selectivity of CO_2 over CH_4 as modeled via GCMC simulations when the framework is in contact with a reservoir of the CO_2/CH_4 mixture at different volume ratios of 80:20, 50:50, and 20:80 (v/v) and at different pressure ranges from 0 bar to either 1 or 10 bar. (b) Selectivity of CO_2 over CH_4 for a 50:50 mixture obtained using GCMC. (c) Adsorption isotherms of pure components (full lines) and a 50:50 mixture (dotted lines) obtained using GCMC. (d) Radial distribution functions between the center-of-mass of the guest molecules and the chlorine atoms of COK-17 versus the corresponding hydrogen atoms of COK-17H and at pressures of either 0.1 or 10 bar.

GCMC simulations. At very low pressures, the selectivity can be estimated from the Boltzmann weights of the adsorption energies for the most stable adsorption sites:

$$\alpha_{\mathrm{CO_2/CH_4}} \approx \, \mathrm{e}^{-\beta(E_{\mathrm{ads,CO_2}}^{\mathrm{min}} - E_{\mathrm{ads,CH_4}}^{\mathrm{min}})}$$

 $E_{\rm ads,CO_2}^{\rm min}$ and $E_{\rm ads,CH_4}^{\rm min}$ being the adsorption energies of the most stable sites for CO2 and CH4, respectively. For COK-17, the energy difference amounts to -4 kJ mol⁻¹, while for COK-17H an even higher value of $-7 \text{ kJ} \text{ mol}^{-1}$ is obtained. On the basis of this analysis, a higher selectivity in COK-17H compared to that in COK-17 is expected. While the chlorine atoms in COK-17 apparently stabilize the CO₂ guests, the same is true to an even higher extent for CH4. This astonishing result is in apparent contradiction with the hypothesis that increased and preferential vdW interactions of the chlorinated linkers with CO₂ leads to the very high selectivity of COK-17 for CO₂, as observed by experiment as well as simulation. This result together with the selectivity dependence on pressure and particular CO₂/CH₄ composition urged us to investigate the competition and/or synergy of the guest molecules using a force-field methodology. The selectivity, obtained from a GCMC simulation of a 50:50 (v/v) CO₂/CH₄ mixture, as a function of the pressure of the gas mixture is shown in Figure

5b. At low pressures COK-17H indeed shows higher selectivity. Interestingly, the COK-17H selectivity decreases with increasing pressure, while for COK-17 the inverse is true. The curves cross at a pressure of about 2 bar. The behavior can be better understood by analysis of the separate components during simulation, shown by dotted lines in Figure 5c.

For COK-17, Figure 5c reveals that the uptake of CH₄ in the mixture saturates already below a pressure of 1 bar. This means that, above a certain pressure, CO2 molecules win the competition for the remaining adsorption sites. As the CO₂ uptake increases while the CH₄ uptake remains constant, obviously selectivity increases. For COK-17H, both CO2 and CH₄ loadings are still increasing at a pressure of 10 bar. This can be related to the much higher surface area and pore volume of COK-17H compared to those of COK-17 as a consequence of the presence of the large chlorine atoms being replaced by hydrogen in COK-17H. At higher pressures there is still enough free space in COK-17H for both, CO₂ and CH₄ molecules, to adsorb. But in COK-17, the smaller pore volume prevents efficient packing of the almost spherical CH₄ molecules, so saturation already occurs at low loadings. These results are in full agreement with the observation that the favored adsorption site B in COK-17 can be occupied by only one CH₄ molecule, but by two CO₂ molecules, which also led to the necessity to assume a two-site model for the description of the adsorption isotherm (vide supra).

Moreover, the radial distribution functions (RDFs) of Cl around the guest molecules at 0.1 and 10 bar obtained from snapshots of guest-molecule configurations extracted from GCMC simulations of mixtures clearly rules out preferred vdW interactions with CO2 as the sole cause for high selectivities at low pressures (Figure 5d). The similarity of CO₂ and CH₄ RDFs confirms that both molecules compete for the same adsorption sites. In COK-17, the peaks are higher and more defined compared to those of COK-17H, indicating more locally defined adsorption sites. Interestingly, a comparison of RDFs obtained from simulations at 0.1 and 10 bar reveals a shoulder of the CO₂ RDF of COK-17, not present at very low pressures and starting to emerge at 1 bar. At higher pressure and higher total loading, CO2 occupies site B, with two molecules in close contact with the chlorine atoms, while these sites are not accessible for more than a single CH₄. Increased proximity to Cl leads to significant gain in interaction energy and clearly can be related to the more efficient packing of CO₂ with increasing pressure.

CONCLUSIONS

COK-17 zeolitic-imidazolate framework with Zn ions bridged by 4,5-dichloroimidazolate has a permanent microporosity and is highly selective for CO₂ adsorption. CO₂ and CH₄ compete for the same adsorption sites, but the smaller CO₂ molecules win the competition because they can pack more densely in higher numbers, maximizing their van der Waals interaction. CO₂ molecules, neatly fitting into the cavities this way, exclude the uptake of CH₄. Insight from the refined structure and molecular simulations explain the details of the adsorption isotherms of the individual molecules as well as the mixtures. The favored adsorption site can be occupied by only one CH₄ molecule but by two CO₂ molecules, explaining the necessity to assume a two-site model for the description of the CO₂ adsorption isotherm. On the basis of this insight, COK-17 could potentially be used to improve the separation of CO₂

from other molecules where the van der Waals interaction is a dominant mechanism.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c08942.

Synthesis and characterization methods, SEM image, TG profile, free energy and pressure equations of state, OP-to-CP and CP-to-OP transition pressures, gas adsorption—desorption isotherms, DFT and BJH pore size distribution curves, breakthrough curves, single-site and dual-site Langmuir isotherms for CH₄ and CO₂, framework properties, simulated adsorption energies, and contribution of London dispersion (PDF)

Accession Codes

CCDC 2057222–2057223 contain the additional 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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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

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