Synthesis of large-pore stabilized MIL-53(Al) compounds with increased CO2 adsorption and decreased water adsorption

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breathing MOFs, solvent effects

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Synthesis of Large-Pore Stabilized MIL-53(Al) Compounds with Increased CO₂ Adsorption and Decreased Water Adsorption

William P. Mounfield, III* and Krista S. Walton*

This work reports two large-pore stabilized MIL-53(Al) metal-organic frameworks prepared via solvothermal synthesis strategies that possess no or only slight breathing behaviour. Powder X-ray diffraction confirmed one material remains in the large-pore form under all conditions, while the other material undergoes a more gradual breathing transition than is observed for MIL-53(Al) prepared by traditional methods. Solid-state NMR was employed to elucidate additional structural information and gain insight into the role synthesis solvent plays on breathing behaviour. The CO₂ and water adsorption of these large-pore stabilized materials were studied, and the improvement in CO₂ loadings over MIL-53(Al) prepared by traditional methods was discussed.

Introduction

Metal-organic frameworks (MOFs) are a promising class of materials that have garnered much research attention for their potential applications. These inorganic-organic hybrid materials are characterized by metal clusters and organic linkers, large surface areas, and tunable chemical properties that make them excellent candidates for gas separations.¹⁻⁸ Certain MOFs offer unique separation applications due to their ability to ‘breathe’, i.e., undergoing a breathing transition in the presence of certain guest molecules.⁹⁻¹³ MIL-53 is one of the most well-known MOFs that exhibits this breathing transition during specific guest molecule adsorption and has been studied extensively since its discovery in 2002.¹⁴⁻²² Traditionally synthesized in water, MIL-53(Al) is comprised of AlO₄(OH)₂ corner-sharing octahedral chains connected by terephthalate groups to form a three-dimensional structure with 8.5 Å pores.¹⁵ Despite the wide research into its breathing behaviours, there has been much less attention given to understanding the effect of solvent on the synthesis and breathing behaviour of MIL-53(Al). DMF (n,n-dimethylformamide) is commonly used in MIL series MOF synthesis such as the amine-functionalized MIL-53(Al)²³, as well as its polymorph MIL-101(Al)-NH₂.²⁴ In addition, a study on the synthesis of the non-porous MIL-69²⁵ found that the use of DMF as solvent in place of water allowed the formation of a new porous material, DUT-4.²⁶ However, despite its wide use in this and many other MOF syntheses, to the authors’ knowledge, there has not been a complete study on the effect of DMF on the synthesis and breathing behaviour of MIL-53(Al). Recently, a computational and experimental study of the amine-functionalized MIL-53(Al), MIL-101(Al) system investigated the effect of replacing water with DMF during synthesis and found DMF was essential for synthesis of the amine functionalized MIL-101(Al) material.²⁷ In this study, we present two solvothermal synthesis methods for MIL-53(Al) with DMF as the solvent. The synthesis at lower temperature results in a material that is stabilized in the large-pore (lp) form, similar to the non-breathing vanadium analogue, MIL-47(V),¹⁶,²⁸ and exhibits higher CO₂ adsorption in the 1-20 bar pressure range and lower water adsorption at humidity levels less than 50 %RH in comparison to that of the breathing material synthesized by traditional routes. The synthesis at higher temperature results in a material that is stabilized in the lp form, but exhibits a swift breathing transition with higher CO₂ adsorption across the entire pressure range and much lower water adsorption at all humidity levels in comparison to that of the breathing material synthesized by traditional routes as well as the lower temperature, lp stabilized material presented in this study.

Experimental section

Synthesis of MIL-53(Al) materials

Synthesis of MIL-53(Al)₁₁₂₀. The synthesis of MIL-53(Al)₁₁₂₀ was performed by mixing aluminium nitrate nonahydrate (Al(NO₃)₃·9H₂O) and benzene-1,4-dicarboxylic acid (C₆H₄-L-1,4-(CO₂H)₂ (referred to as BDC hereafter) in a 1:2.25 molar ratio. The reagents were added to 30 mL DMF (n,n-dimethylformamide) and stirred for 30 minutes until all solids were dissolved. The mixture was placed in a 60 mL glass vial and reacted at isothermal conditions of
120°C for 72 hours. The resulting milky-white solid was gravity-filtered, washed with DMF three times and washed finally with methanol once. The final product was dried in air overnight.

**Synthesis of MIL-53(Al)_{DMF,220}**

The synthesis of MIL-53(Al)_{DMF,220} (sDMF_{220}) was performed by mixing aluminum nitrate nonahydrate (Al(NO₃)₃·9H₂O) and benzene-1,4-dicarboxylic acid (C₆H₄-1,4-(CO₂H)₂) (referred to as BDC hereafter) in a 1.225 molar ratio. The reagents were added to 30 mL DMF (n,n-dimethylformamide) and stirred for 30 minutes until all solids were dissolved. The mixture was placed in a sealed Teflon-lined reactor and reacted at isothermal conditions of 220°C for 72 hours. The resulting white solid was gravity-filtered, washed with DMF three times and washed finally with methanol once. The final product was dried in air overnight.

**Synthesis of MIL-53(Al)_{H₂O}**

The synthesis of MIL-53(Al)_{H₂O} (sH₂O) was performed following the procedure found in literature. 130 g Al(NO₃)₃ and 0.288 g BDC were stirred for 30 minutes in 5 mL H₂O and placed in a 23 mL Teflon-lined stainless steel autoclave for 72 hours at 220°C. The resulting white product was gravity filtered and washed with water three times. Powder X-ray diffraction and nitrogen adsorption analysis were performed on the resulting white solid. Unreacted BDC was found in the pores, and an extraction process from literature was adapted to remove this leftover material. The filtered product was placed in 10 mL DMF in a 23 mL Teflon-lined stainless steel autoclave overnight at 150°C. The resulting product was filtered with DMF and dried in air overnight.

**Powder X-ray diffraction**

Powder X-ray diffraction patterns were recorded on an X’Pert X-ray PANalytical diffractometer with an X’celerator module using Cu Ka (λ = 1.5418 Å) radiation at room temperature, with a step size of 0.02° in two theta (2θ). All attempts at obtaining single crystal X-ray diffraction data were unsuccessful.

**N₂ physisorption**

Nitrogen physisorption isotherms were measured at 77 K for samples activated at 473K for 16h before and after water exposure using the Quadrasorb system from Quantachrome instruments (Fig. S2). Nitrogen adsorption analysis was performed on all resulting products, and surface areas were determined by applying BET theory over a range of data points applicable to microporous materials. N₂ isotherms showed typical Type I behaviour as per the IUPAC classification.

**Gas adsorption**

An Intelligent Gravimetric Analyzer (IGA-1 series, Hiden Analytical Ltd.) was used to collect pure gas (CO₂, CH₄, N₂) adsorption isotherms at 293K and pressures up to 20 bar (Figure 3, S3). Samples were activated in situ at 473 K under vacuum until no further weight loss was observed. After activation, the system was maintained under vacuum, and the temperature was adjusted to the desired value. A sample size of approximately 25 mg was used for the measurements, and a maximum equilibration time of 40 min was used for each point in the isotherm.

**Water adsorption**

An Intelligent Gravimetric Analyzer (IGA-3 series, Hiden Analytical Ltd.) was used to obtain water vapour adsorption isotherms at 293K (Figure 4). A sample size of approximately 35 mg was used for collecting the water vapour adsorption isotherm. Dry air was used as the carrier gas, with a percentage being bubbled through a vessel filled with deionized water. Two mass flow controllers were used to vary the ratio of saturated air and dry air so that the relative humidity (%RH) could be controlled. Due to water condensation in the equipment at higher humidities, experiments were conducted up to 80 %RH. The total gas flow rate was set at 200 cm³/min, and each adsorption/desorption step was allowed a maximum time of 24 h to approach equilibrium. Samples were activated in situ at 473 K under vacuum until no further weight loss was observed before starting the experiment. Samples were regenerated by heating at 473 K under vacuum for 16 h after the water adsorption isotherm measurement prior to BET analysis.

**FTIR spectroscopy**

FTIR spectra were recorded on a Bruker Vertex 70, equipped with a DTGS detector in transmission geometry. Samples were prepared in a 1:100 sample:KBr pellet. The spectra were collected with an accumulation of 24 scans and a resolution of 4 cm⁻¹. A flow of N₂ at 20 mL min⁻¹ was maintained during the measurement. The spectra were recorded at room temperature without any pre-treatment of the samples.

**Solid-state NMR spectroscopy**

²⁷Al, ¹³C, and ¹H NMR spectra were recorded on a Bruker Avance III spectrometer with a 9.4T magnetic field operating at 400 MHz using an automated procedure for analysis. A 4mm Bruker MAS probe with a ZrO₂ rotor spinning at 12 kHz was used in the ¹H and ²⁷Al MAS NMR studies. ²⁷Al MAS NMR spectra were recorded over 4096 scans on a simple block decay single-pulse excitation of 0.6 μs pulse duration with a repetition delay of 500 ms. Deconvolution of the ²⁷Al NMR spectra was performed using Bruker TopSpin 3.2.

**Results and discussion**

Due to the larger kinetic diameter in comparison to water, 5.5 Å versus 2.6 Å, and its known ability to induce a large-pore structural transition, DMF offers the ability to stabilize the lp form of MIL-53(Al) much like CO₂. It possesses a kinetic diameter of 3.6 Å. Recently, there has been work on alternative synthesis using an ionic liquid as the solvent in place of water for increasing the hydrophobicity of MIL-53. The ionic liquid possesses a kinetic diameter similar to DMF, illustrated in Fig. S5. Therefore, in accordance with the findings of Liu et al. and Senkovska et al., we hypothesized that the size of the
solvent molecule directs the synthesis such that DMF prevents the formation of the original literature framework, as illustrated in Fig. 1(c), where DMF molecules in orthogonal and parallel orientations are seen to block the pores and occupy a much larger space than the water molecule in Fig. 1(a). As seen by Senkovska et al. for DMF synthesis of MIL-69, the framework must assemble in the lp form in the presence of DMF to accommodate the larger solvent molecule as shown in Fig. 1(b), and when synthesized at lower temperatures (120 °C) is unable to undergo the breathing transition shown in Fig. 1(a) for dehydration/hydration/cooling of MIL-53(Al) materials prepared by published procedures. Interestingly, synthesis at higher temperatures (220 °C) results in a material that is able to undergo a breathing transition similar to that observed for MIL-53(Al) in literature.

![Fig. 1 Schematic of (a) MIL-53(Al)dehydration, hydration and cooling to room temperature after activation, (b) removal of DMF from pores of MIL-53(Al)_DMF and stability of lp form after cooling to room temperature (c) DMF molecules occupying pore space of MIL-53(Al) in orthogonal and parallel orientations.](image)

BET analysis was performed on each of the resulting products and it was determined that the unreacted BDC was removed from the SH2O material by the DMF extraction process and no calcination step was necessary. Further, the extraction step allows for one of the highest reported BET surface areas for MIL-53(Al), 1410 m²/g, much higher than previously reported literature values. In comparison, sDMF120,220 possess activated surface areas of 1472 m²/g, and 1570 m²/g, respectively. In order to determine the degree to which the three materials were activated, the theoretical surface area was calculated and compared to the experimental surface area results. As all surface areas are nearing the theoretical value of 1632.8 m²/g, it was believed that the materials were sufficiently activated. However, analysis of FTIR spectra (Fig. 2) for sDMF220 showed the presence of a spectral peak at 1701 cm⁻¹ attributed to ν(C=O) in DMF molecules that remain trapped in the material; an observation that has been made in other MOFs when using DMF in place of H₂O as the synthesis solvent. Several washing steps with methanol were repeated in an attempt to remove any free, unreacted BDC molecules from the framework, but these repeated washing steps did not eliminate the presence of the spectral peak in the FTIR spectra. In order to confirm the peak was not due to free BDC molecules trapped in sDMF220, the material was again washed six times with methanol and activated at 300 °C for 24 hours. A BET analysis confirmed a complete structural collapse of the material and the continued presence of the spectral peak at 1706 cm⁻¹ indicates a free carboxylic group of BDC still exists within the material and cannot be removed through activation. Furthermore, the continued presence of this group after numerous activation procedures, suggests that this peak is not, in fact, due to trapped BDC molecules, but rather an indication of the defects of the material, i.e., uncoordinated linkers that contribute to the framework’s inability to undergo a structural transition.

Analysis of FTIR spectra for sDMF220 showed the presence of a spectral peak at 1701 cm⁻¹ attributed to ν(C=O) in DMF molecules that remain trapped in the material; an observation that has been made in other MOFs when using DMF in place of H₂O as the synthesis solvent. Several washing steps with methanol were repeated in an attempt to remove any free, unreacted BDC molecules from the framework, but these repeated washing steps did not remove the trapped species, suggesting these molecules are coordinated to the metal center or to the benzene ring of the ligand. Although there are some coordinated DMF molecules in the material, the observed high surface area indicates that these molecules are few in number and should not reduce the material’s gas adsorption capability.

![Fig. 2 FTIR spectra of activated MIL-53(Al)sDMF, SH2O, green), activated MIL-53(Al)_DMF,220 (sDMF220, blue), MIL-53(Al)_DMF,120 (sDMF120, 24h, light blue) activated for 24 hours at 300 °C after six methanol washes, activated MIL-53(Al)_DMF,220 (sDMF220, red).](image)
solvent molecule. This hypothesis is supported by the slightly larger cell parameters observed with LeBail refinement (Figure S8, Table S1). Adsorbed water molecules in sH$_2$O material account for the spectral band observed at 1629 cm$^{-1}$.

However, after activation, it is clear that the MIL-$53$(Al)$_{sH_2O}$ sample is present in the narrow-pore (np) form. The stability of MIL-$53$(Al)$_{sDMF}$ in the lp form is quite unlike the MIL-$53$(Al) sample synthesized with water, designated MIL-$53$(Al)$_{sH_2O}$, as seen when referencing its as-synthesized XRD pattern, Fig. 3(5). After extraction of remaining BDC molecules from the pores of the sample, and subsequent activation, it is clear that the MIL-$53$(Al)$_{sH_2O}$ sample is present in the narrow-pore (np) form.$^{15}$ The material synthesized at higher temperatures in DMF, designated as MIL-$53$(Al)$_{sDMF,220}$, exhibits an as-synthesized pattern similar to MIL-$53$(Al)$_{sDMF}$ after extraction with DMF, where both materials are present in the lp form. However, after activation, unlike the material synthesized in DMF at lower temperatures, MIL-$53$(Al)$_{sDMF,220}$ is present in the np form, similar to MIL-$53$(Al)$_{sH_2O}$. In conjunction with the presence of DMF in the IR spectra from the activated MIL-$53$(Al)$_{sDMF,220}$ sample, we hypothesized that the material would be able to undergo a complete structural transition similar to the literature framework but that the transition would be affected by the coordinated DMF molecules; a hypothesis that is tested with gas adsorption experiments later in the text. The LeBail refinement performed on the PXRD pattern for MIL-$53$(Al)$_{sDMF,220}$ reveals the material exists in a structural form that is slightly larger than the traditional material’s np form. After intermittent exposure to laboratory atmosphere, ~50 %RH, there was no observed change in the samples’ XRD pattern, as seen in Fig. S4.

![Fig. 3 XRD patterns of (1) large-pore form, (2) narrow-pore form, (3) as-synthesized MIL-$53$(Al)$_{sDMF,120}$, (4) activated MIL-$53$(Al)$_{sDMF,120}$, (5) as-synthesized MIL-$53$(Al)$_{sH_2O}$, (6) extracted MIL-$53$(Al)$_{sH_2O}$, (7) activated MIL-$53$(Al)$_{sH_2O}$, (8) as-synthesized MIL-$53$(Al)$_{sDMF,220}$, (9) activated MIL-$53$(Al)$_{sDMF,220}$.](image)

![Fig. 4 (top) $^1$H, and (bottom) $^{27}$Al MAS NMR spectra for sDMF$_{120}$ (blue), sDMF$_{220}$ (red), and sH$_2$O (green).](image)
determination of the relative hydrophobicity of the materials. The signal at $\delta = 3$ ppm in the $^1$H MAS NMR spectra in Fig. 4 for sDMF$_{120}$, sDMF$_{220}$, and sH$_2$O is indicative of the formation of weak hydrogen bonds between water molecules and neighbouring oxygen atoms of the organic linkers. The increased intensity of this signal in sH$_2$O clearly illustrates the traditionally synthesized sample is hydrated to a greater degree when exposed to laboratory air than the samples synthesized in DMF at high temperature. In addition, the presence of the water signal at 6.1 ppm in the sDMF$_{120}$ spectrum indicates a very low degree of hydration due to the materials permanent state in the large pore form.

To further analyse the MIL-53(Al) materials’ structural properties, $^{27}$Al MAS NMR was performed on activated samples and compared to samples in the presence of DMF. The $^{27}$Al spectra for activated sDMF$_{120}$ exhibits a quadrupolar pattern with $\delta_{27A}$ = 3.3 ppm, $C_Q$ = 8.2 MHz, $\eta_0$ = 0.2, sDMF$_{220}$ features a well-defined quadrupolar pattern with $\delta_{27A}$ = 3.6 ppm, $C_Q$ = 8.4 MHz, $\eta_0$ = 0.13, while the spectrum for sH$_2$O exhibits a quadrupolar pattern with $\delta_{27A}$ = 2.9 ppm, $C_Q$ = 8.1 MHz, $\eta_0$ = 0.08 and no trace of any other aluminium-containing phase, confirming there is no remaining amorphous Al(OH)$_3$. In order to analyse the breathing transition of sDMF$_{220}$ and sH$_2$O, the activated materials were saturated with DMF to place them in their lp form. The $^{27}$Al MAS spectrum for sDMF$_{120}$ features a single quadrupolar pattern with $\delta_{27A}$ = 11 ppm, $C_Q$ = 10.1 MHz, $\eta_0$ = 0.16, sDMF$_{220}$ features a quadrupolar pattern with $\delta_{27A}$ = 5 ppm, $C_Q$ = 10.1 MHz, $\eta_0$ = 0.098, while the spectrum for sH$_2$O exhibits a quadrupolar pattern with $\delta_{27A}$ = 3.6 ppm, $C_Q$ = 10.1 MHz, $\eta_0$ = 0.1. The smaller increase in the quadrupolar coupling constant upon saturation with DMF for sDMF$_{220}$, 8.4–10.1, and sDMF$_{120}$, 8.2–10.1 compared to the increase in $C_Q$ for sH$_2$O, 8.1–10.1, can be attributed to a decrease in the strain on the AlO$_6$(OH)$_2$ octahedra in the materials synthesized in DMF. As the octahedra experience less strain in the materials synthesized in DMF, they are able to gradually transition from closed to open pore forms in the case of sDMF$_{220}$ or remain in the open pore form permanently in the case of sDMF$_{120}$ as is confirmed with CO$_2$ adsorption experiments.

After analysing the PXRD and solid-state NMR results and concluding the MIL-53(Al)$_{DMF,120,220}$ samples are unique MIL-53(Al) materials, further investigation of their structure and behaviour was made with high pressure, pure-component CO$_2$ and CH$_4$ adsorption experiments. The CO$_2$ and CH$_4$ adsorption isotherms for sDMF$_{120}$ shown in Fig. 5 exhibit Type I isotherms with no hysteresis. CO$_2$ is more strongly adsorbed than CH$_4$ as it has a higher quadrupole moment than the nonpolar methane. The CH$_4$ adsorption isotherms for sH$_2$O and sDMF$_{220}$ shown in Fig. 5 also exhibit Type I behaviour. The CO$_2$ adsorption isotherm for sH$_2$O exhibits strong Type IV behaviour with large hysteresis upon desorption, which is characteristic of the traditional breathing behaviour seen in literature. The CO$_2$ isotherm for sDMF$_{220}$ exhibits a weak Type IV behaviour with small hysteresis upon desorption characteristic of a gradual structural transition, lacking the quick step-like change observed for CO$_2$ adsorption in sH$_2$O, resulting in the observation of only a slight breathing behaviour. The observation of this gradual breathing behaviour during CO$_2$ adsorption combined with the results from $^{27}$Al NMR showing reduced strain on the octahedral metal centers while in the lp form supports the hypothesis presented previously that the sDMF$_{220}$ material is indeed able to undergo a complete structural transition, but with a more gradual movement between open and closed forms. In addition, the presence of the material in the np form after activation suggests the number of coordinated DMF molecules is not enough to induce a transition to the lp form without guest adsorption.

![Graph](image)

Fig. 5 CO$_2$ and CH$_4$ adsorption isotherms at 25°C for sDMF$_{220}$ (red stars and pentagons), sDMF$_{120}$ (blue triangles and circles) and sH$_2$O (green squares and left triangles) as a function of pressure ranging from 0-20 bar. Desorption isotherms are omitted except for sH$_2$O (open green squares), which shows large hysteresis.

The CO$_2$ and CH$_4$ adsorption observed for sH$_2$O, 8.2 mmol/g and 4.3 mmol/g, match that found in the literature, while the overall CO$_2$ loading for sDMF$_{120}$ is more than 1 mmol/g higher than sH$_2$O, 9.3 mmol/g. The CH$_4$ loading is only slightly increased by 0.5 mmol/g to 4.8 mmol/g. The CO$_2$ loading for sDMF$_{220}$ is more than 1.5 mmol/g higher than sH$_2$O, 9.8 mmol/g, while the CH$_4$ loading is the same as sDMF$_{220}$, 4.8 mmol/g. However, a much greater increase over sH$_2$O is observed for both materials synthesized in DMF in the 1-5 bar region. It is in this region that sH$_2$O is still present in the np form, and is unable to have additional significant adsorption, while sDMF$_{120}$ is stabilized in the lp form and has continued adsorption throughout this region. It is evident from the adsorption data that the sDMF$_{120}$ material is indeed present in the lp form throughout adsorption of CO$_2$ as it closely follows the desorption curve for sH$_2$O in the 2-5 bar region, where the sH$_2$O material is still present in the lp form before relaxing to the np form as the pressure drops below 2 bar. Contrary to the observations for the material synthesized at low temperatures in DMF in this intermediate pressure region, sDMF$_{220}$ undergoes a gradual structural transition to the lp form, as seen in Fig. 5 rather than the step breathing transition.
observed in the sH₂O material, allowing for greatly increased adsorption throughout the region. The coordinated DMF molecules allow retention of the breathing ability of the framework and greatly increased CO₂ adsorption capacity in the transition region. Due to the large increase in CO₂ adsorption with a negligible increase in CH₄ adsorption in this region, the sDMF₂₂₀ and sDMF₁₂₀ materials offer a three-fold increase in working capacity for processes run in the 1-5 bar region, 3.6 mmol/g vs 1.2 mmol/g, in addition to increased CO₂ adsorption at 1 bar for sDMF₂₂₀, 2.7 mmol/g vs 2.1 mmol/g.

Fig. 6 H₂O adsorption isotherms at 25°C and 1 bar for sDMF₁₂₀ (blue circles), sDMF₂₂₀ (red squares) and sH₂O (green stars) as a function of relative humidity.

Due to the discovery of the non-breathing behaviour of sDMF₁₂₀, and the gradual breathing behaviour of sDMF₂₂₀ a further investigation of the materials’ performance in the presence of water was performed in comparison to that of sH₂O. Water adsorption isotherms were obtained at 1 bar with relative humidity ranging from 0-90%. The water adsorption isotherm for the MIL-53(Al) material synthesized by traditional water synthesis shown in Fig. 6 (sH₂O) exhibits Type I behaviour typical of a small pore material with very little increase in loading after the initial adsorption of 4 mmol/g at 10 %RH. The material synthesized at low temperatures in DMF, sDMF₁₂₀, however, exhibits a Type V isotherm characteristic of a large-pore material, with very low initial loading until >50 %RH. sDMF₁₂₀ adsorbs less than 1 mmol/g of water vapour until the humidity rises above 30 %RH, and maintains a lower loading than sH₂O until 50 %RH. This low adsorption along with the absence of any change in PXRD pattern (Fig. S4) or BET surface area after routine exposure to laboratory atmosphere of similar humidity leads us to believe that the material is stable in humid conditions less than 50 %RH. The material synthesized at high temperatures in DMF, sDMF₂₂₀, however, exhibits an approximately linear isotherm, with very low loading across the entire range of humidity. sDMF₂₂₀ adsorbs less than 2 mmol/g of water vapour at 90 %RH, less than half the loading of sH₂O at the same humidity level.

Coincident with the observation of DMF molecules remaining after activation in the FTIR spectra and the gradual breathing behaviour of sDMF₂₂₀ during CO₂ adsorption it is likely that the few coordinated DMF molecules prevent or reduce water adsorption around their locations in the framework resulting in low water adsorption and increased hydrophobicity, as well as continued low adsorption in the high humidity range similar to sH₂O with the np form preventing the clustering and adsorption of large amounts of water seen in sDMF₁₂₀. Similar exposure to laboratory air did not yield any change in the PXRD pattern or BET surface area of sDMF₂₂₀ as was expected from the ¹H NMR results demonstrating its greater hydrophobicity in room conditions than sH₂O.


In addition, we are able to gain insight into the water stability of the lp and np conformations of MIL-53(Al). It is evident that the continued water stability at higher relative humidity levels of the traditionally synthesized material stems from the immediate and complete pore filling with low water adsorption at initial humidity levels due to the presence of inaccessible areas of the pore volume while in the np form. Conversely, while sDMF₁₂₀ is in the lp form, water experiences weaker interactions with the pore surface at low humidity levels, as illustrated by the low initial adsorption until 50 %RH. However, as seen in experimental and computational studies of the effects of water on other MOFs, beyond this point water clusters form and fill the pore, interacting more with the metal-ligand bond and leading to a breakdown of the structure as the material begins to adsorb large amounts of water, with an ultimate loading of 13.3 mmol/g. In addition, the presence of defects, indicated by FTIR analysis of this material, allows easy access to the metal-ligand bond and subsequent collapse of the structure. The large increase in adsorption in sDMF₁₂₀ corresponding to a breakdown of the MOF structure is
confirmed through comparison of PXRD patterns before and after water adsorption experiments shown in Fig. 7, and BET surface areas before and after water adsorption experiments. It is evident that a large change has occurred in the structure of sDMF$_{220}$ as the structure no longer resembles the activated pattern or the simulated lp or np form patterns and the BET surface area was reduced from 1472 m$^2$/g to 370 m$^2$/g.

sDMF$_{220}$, while appearing to behave similarly to sH$_2$O, is found to not be water stable, with the pattern after water adsorption experiments closely resembling its as-synthesized pattern shown in Fig. 3 (8), suggesting there are free BDC and Al(OH)$_3$ molecules present after being removed by attacking water molecules. The reduction of the BET surface area from 1570 m$^2$/g to 402 m$^2$/g confirms the breakdown of the framework in the presence of high humidity. The instability of this material, although present in a narrow-pore form, suggests that the coordinated DMF molecules may induce the observed larger np form which allows for enough water to be adsorbed to degrade the material. On the other hand, there is little change in the PXRD pattern for MIL-53(Al)$_{120}$ after water adsorption, with the pattern closely resembling the activated pattern as well as the simulated np form. In addition, BET analysis showed a negligible change in surface area before and after water adsorption for MIL-53(Al)$_{120}$, 1410 m$^2$/g to 1408 m$^2$/g, confirming the water stability of the material while in the np form. These results lead to the hypothesis that traditionally synthesized MIL-53(Al) materials as well as the large-pore stabilized materials are susceptible to water effects at high humidity levels while in pore forms any larger than the traditional np form, which would have a negative impact on CO$_2$ adsorption applications at higher pressures, where all types of these materials would be present in the large-pore form.

**Conclusions**

In conclusion, we have reported solvothermal syntheses of large-pore stabilized MIL-53(Al) MOFs that do not possess the standard breathing behaviour found in literature. Synthesis at lower temperature produces a material with no breathing behaviour, while synthesis at higher temperature results in a material with a gradual, slight breathing behaviour. The different breathing behaviour or lack thereof and increased stability of the large-pore form allows for a large increase in CO$_2$ adsorption in the 1-5 bar region, and overall loading for the large-pore only material, and a large increase throughout the entire pressure range for the gradually breathing MOF. Water adsorption studies of the sDMF$_{120}$ and sDMF$_{220}$ materials suggest that at humidity levels less than 50 %RH, the materials are stable and have much lower water loading than the sH$_2$O framework. It is evident that understanding and modulating the breathing behaviour of metal-organic frameworks offers unique opportunities for enhancing applications in gas adsorption, chemical sensing, and other areas of interest.

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**Notes and references**

School of Chemical and Biomolecular Engineering, Georgia Institute of Technology, 311 Ferst Drive NW, Atlanta, Georgia 30332, United States walton.chbe.gatech.edu; Email: krista.walton@chbe.gatech.edu

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