MIL-96, a Porous Aluminum Trimesate 3D Structure Constructed from a Hexagonal Network of 18-Membered Rings and $\mu_3$-Oxo-Centered Trinuclear Units

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Abstract: A new aluminum trimesate Al$_{12}$O(OH)$_{18}$([H$_2$O]$_3$)[Al$_2$O(OH)$_4$][btc]$_6$â•€•€24H$_2$O, denominated MIL-96, was synthesized under mild hydrothermal conditions (210 °C, 24 h) in the presence of a 1,3,5-benzenetricarboxylic acid (trimesic acid or H$_3$btc) in water. Hexagonal crystals, allowing a single-crystal XRD analysis, are grown from a mixture of trimethyl 1,3,5-benzenetricarboxylate (Me$_3$btc), HF, and TEOS. The MIL-96 structure exhibits a three-dimensional (3D) framework containing isolated trinuclear $\mu_3$-oxo-bridged aluminum clusters and infinite chains of AlO$_4$(OH)$_2$ and AlO$_2$(OH)$_4$ octahedra forming a honeycomb lattice based on 18-membered rings. The two types of aluminum groups are connected to each other through the trimesate species, which induce corrugated chains of aluminum octahedra, linked via $\mu_2$-hydroxo bonds with the specific cis–cis–trans sequence. The 3D framework of MIL-96 reveals three types of cages. Two of them, centered at the special positions 0 0 0 and $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{4}$, have estimated pore volumes of 417 and 635 Å$^3$, respectively, and encapsulate free water molecules. The third one has a smaller pore volume and contains disordered aluminum octahedral species (Al(OH)$_4$). The solid-state NMR characterization is consistent with crystal structure and elemental and thermal analyses. The four aluminum crystallographic sites are resolved by means of $^{27}$Al 3QMAS technique. This product is able to sorb both carbon dioxide and methane at room temperature (4.4 mmol g$^{-1}$ for CO$_2$ and 1.95 mmol g$^{-1}$ for CH$_4$ at 10 bar) and hydrogen at 77 K (1.91 wt % under 3 bar).

Introduction

In the past decade, there has been an increasing interest in the synthesis of porous metal–organic framework materials (MOF) or coordination polymers,1–3 based on the connection of metal ions (nodes) and organic ligands (linkers), such as amines or carboxylates, usually including one or several benzene rings. These solids exhibit novel fascinating three-dimensional (3D) topologies, which may potentially find applications as molecular sieves and gas adsorbents (for instance, hydrogen or methane storage4–6), ion exchange or heterogeneous catalysis.5,6 Some of them possess rigid open-frameworks, which are reminiscent to the functionalities observed in the microporous aluminosilicates networks (zeolites). The carboxylate family has attracted more attention since extra-large pore solids with relative high thermal stability have been synthesized. This series is well illustrated with the examples of the MOF-n solids (MOF-5,7 MOF-1778) by Yaghi, or MIL-1009 and MIL-10110 compounds by Férey. Usually, 3d divalent (Ni$^{2+}$, Cu$^{2+}$, Zn$^{2+}$) and trivalent (Sc$^{3+}$, V$^{3+}$, Cr$^{3+}$, Fe$^{3+}$) or rare earth metals are used as metallic centers, but the trivalent p elements such as Al$^{3+}$, Ga$^{3+}$ or In$^{3+}$ are much more rarely reported. Recently, a systematic investigation of the reactivity of indium with the benzene-based carboxylates was described by different groups.11–15 The use of aluminum or gallium for the production

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of MOFs materials is less common, despite a rich literature relating to the chemistry of molecular aluminum carboxylate complexes for their neurotoxic effects in biological systems or alumoxanes species resulting from the reaction of carboxylic acids with boehmite. On the other hand, the hydrolysis of aluminum in aqueous solution has been intensively studied for over 40 years. Besides the isolation of large purely inorganic aggregates such as Al₄³⁻⁵⁻ or Al₆²⁶⁻, polycations species, novel clusters such as the Al₁₃⁵⁻ or Al₂⁵⁻²⁹ species are formed in the presence of the carboxylic ligands heidi (H₄heidi = N(CH₂COOH)₃CH₂(CH₂OH)) and hpda (H₄hpda = HOCH₂-[CH₃N(CH₂COOH)₂]₂), respectively.

The group in Versailles focuses its attention on the reaction of aluminum or gallium with aromatic carboxylic acids under hydrothermal conditions. Up to now, two phases have been isolated with aluminum, MIL-53(Al) (with 1,4-benzenedicarboxylic acid or H₂btc) and MIL-69 (with 2,6-naphthalenedicarboxylic acid or H₂nadc), and one for gallium, MIL-61 (with 1,2,4,5-benzenehexacarboxylic acid or H₄btec). The MIL-53 compound was found to be a good candidate for H₂, CH₄, and CO₂ adsorption, and significant uptake values were observed for these different gases, partly due to the low density of these materials with aluminum being a relatively light element. We continue these series with the utilization of the trimesic acid ligand (1,3,5-benzenetricarboxylic acid or H₃btc).

Under some conditions, a ladder-like pure fluoride aluminum ([Al₄F₁₈])ₙ species intercalated by the tripotanion H₂btc molecules was previously obtained with the HF/pyridine solvent mixture. Here, this contribution addresses the synthesis and characterization of a new 3D framework, noted MIL-96 Al, Al₂O₃(OH)$_{3x}$[H₂O]ₙ[Al₂(OH)$_{2+}$]btc$^-$·24H₂O involving the aluminum octahedral units interacting with the btc ligand. We applied an alternative synthesis route for growing well-defined single crystals of such MOF solids by using the methyl ester form (trimethyl 1,3,5-benzenetricarboxylate or Me₃btc) of the trimesic acid together with HF and TEOS. The arrangement of the metallic cation Al$^{3+}$ in this particular solid is new and original for the chemistry of aluminum since isolated µ₁-oxo-centered trigonal units and a two-dimensional (2D) network of hexagonal 18-membered rings are observed. The MIL-96 (Al) compound is characterized by means of single-crystal X-ray analysis, ²⁷Al solid-state MAS NMR and gas adsorption was reported for CO₂, CH₄, and H₂.

**Experimental Section**

Synthesis. The aluminum trimesate Al₂O(OH)$_{3x}$[H₂O]ₙ[Al₂(OH)$_{2+}$]btc$^-$·24H₂O (MIL-96) was hydrothermally synthesized under autogenous pressure from a mixture of aluminum nitrate and 1,3,5-benzenetricarboxylic acid in water. The starting reactants were aluminum nitrate (Al(NO₃)$_₃$·9H₂O, Carlo Erba Regenti, 98%), 1,3,5-benzenetricarboxylic acid (C₆H₅(CO₂H)$_₃$, Aldrich, 95%, or H₃btc). In addition, different synthesis batches used trimethyl 1,3,5-benzenetricarboxylate (C₆H₅(CO₂CH₃)$_₃$, 98%, Aldrich, noted Me₃btc), dilute hydrofluoric acid (HF, Normapur, 4.8%) and tetraethylorthosilicate (Si(OCH₃)$_₄$, TEOS, Merck, >98%). Typically, the reaction mixture containing the molar ratio: 1 Al(NO₃)$_₃$·9H₂O (3.5 mmol, 1.314 g),0.14 H₂btc (0.5 mmol, 0.105 g)/80 H₂O (278 mmol, 5 mL) was placed in a 23-ML Teflon-lined steel Parr autoclave at 210 °C for 24 h (batch 1). The pH of synthesis was 1. After the hydrothermal treatment, a powdered product was obtained, which was filtered off, washed with deionized water, and dried in air at room temperature. Optical microscopy analysis indicated that the sample is composed of a mixture of a fine white powder (~1 µm size) of the title compound and large parallelepiped-shaped crystals of recrystallized H₂btc (C₆H₅O₃·0.83H₂O$^{37}$ form). Preliminary X-ray powder diffraction pattern showed that the fine powder is a novel phase. The MIL-96 (Al) phase could be obtained as a pure phase by using the trimethyl 1,3,5-benzenetricarboxylate as starting reactant (batch 2). The reaction mixture was 1 Al(NO₃)$_₃$·9H₂O (3.5 mmol, 1.314 g)/0.5 Me₃btc (1.75 mmol, 0.440 g)/80 H₂O (278 mmol, 5 mL), and a pure phase of MIL-96 (Al) is prepared after heating for 24 h at 210 °C under hydrothermal conditions. Suitable single-crystals for XRD analysis were obtained from a different mixture (batch 3). In this case, the reaction molar composition was 1 Al(NO₃)$_₃$·9H₂O (3.5 mmol, 1.314 g)/0.5 Me₃btc (1.75 mmol, 0.440 g)/0.4 HF (1.4 mmol, 0.6 mL)/0.2 TEOS (0.7 mmol, 0.16 mL)/320 H₂O (1111 mmol, 20 mL). HF$^{38}$ and TEOS$^{39}$ are used for their mineralizing effect under hydrothermal conditions, inducing the increased crystallinity. These reactants were placed in a 125-ML Teflon-lined steel Parr autoclave at 210 °C for 24 h, and flat hexagonal-shape crystals of 10–40 µm size were obtained in the absence of recrystallized H₂btc.

**Single-Crystal X-ray Structure Analysis.** A colorless hexagonal-shaped crystal (0.04 mm × 0.04 mm × 0.04 mm) was selected under polarizing optical microscope and glued on a glass fiber for a single-crystal X-ray diffraction experiment. X-ray intensity data were collected on a Bruker$^{38}$ KAPAX-2 CCD area-detector diffractometer using Mo Kα radiation (λ = 0.71073 Å). Four sets of narrow data frames (60 s per frame) were collected at different values of θ for 2 and 2 initial values of ϕ and Ω, respectively, using 0.3° increments of ϕ or Ω. Data reduction was accomplished using SAINT V7.03. The substantial redundancy in data allowed a semiempirical absorption correction (SADABS V2.10) to be applied, on the basis of multiple measurements of equivalent reflections. The structure was solved in the space group P6₅/mmc by direct methods, developed by successive difference Fourier

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Table 1. Crystal Data and Structure Refinement for MIL-96 (Al)

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<th>Identification code</th>
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<tr>
<td>Volume</td>
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<td>$F(000)$</td>
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<td>Largest diff. peak and hole</td>
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Solid-State NMR. All spectra were collected with a Bruker Avance 500 spectrometer using a 2.5-mm triple resonance MAS probe at room temperature. Powder samples were packed into 2.5-mm ZrO$_2$ rotors.

$^{31}$Al (130.3 MHz) spectra were referenced to external alum ($^{31}$NH$_2$Al(SO$_4$)$_2$:12H$_2$O) at −0.6 ppm relative to 1 M aqueous Al(NO$_3$)$_3$ as secondary reference. The radio frequency field strength was determined from nutation experiments on both cubic solid alum (liquidlike behavior) and solid nutation of berlinit AlPO$_4$. The single-pulse $^{27}$Al spectrum was collected at a spinning rate of 30 kHz. The experiment was recorded after excitation with π/12 pulses of 86 kHz rf power and repetition time of ~100 ms. The 3QMAS 2D NMR experiment$^{40,41}$ was achieved at a spinning rate of 25 kHz using a $z$-filter sequence$^{42}$ with two hard pulses of 4.1 μs (127°) and 1.6 μs (50°) and one selective pulse of 12.2 μs (90°). Additional 3QMAS experiment with rf pulses of higher power (~170 kHz) and fast amplitude modulation (FAM-II) method for conversion of observable coherence was run to optimize MQMAS efficiency for sites with larger quadrupolar interactions.$^{43}$ Simulations of spectra were done with the Bruker mNMR software package.

Adsorption Microcalorimetry. The apparatus used for microcalorimetry experiments is able to measure both isotherms and experimental enthalpies of adsorption in the temperature region from 25 to 150 °C.$^{44}$ The Tian-Calvet type microcalorimeter used to measure the enthalpies consists of two thermopiles mounted in electrical opposition. Each thermopile comprises ~500 chromel–alumel thermocouples. The adsorption isotherms are obtained using a manometric device built to withstand pressures up to 100 bar. The pressure gauge measures pressures up to 50 bar. A point-by-point introduction of gas is most adapted to this system. Each introduction of adsorbate to the sample is accompanied by an exothermic thermal effect, until equilibrium is attained. This peak in the curve of energy with time has to be integrated to provide an integral (or pseudodifferential) molar enthalpy of adsorption for each dose. The calorimetric cell (including the relevant amount of adsorbent and gas) is considered as an open system. In this procedure, it is important to consider that the gas is introduced reversibly. Under these conditions it is possible to determine the differential enthalpy of adsorption $\Delta_\text{ad}h$, via the following expression:

$$\Delta_\text{ad}h = \left(\frac{dQ_{\text{ads}}}{dn}\right)_T + V\left(\frac{dp}{dn}\right)_T$$

Here $dQ_{\text{ads}}$ is the heat reversibly exchanged with the surrounding environment at temperature $T$, as measured by the calorimeter, $dV$ is the amount adsorbed after introduction of the gas dose, $dp$ is the increase in pressure and $V$ is the dead space volume of the sample cell within the calorimeter itself (thermopile). The term $Vdp$ can be obtained via blank experiments. Prior to each adsorption experiment, the samples were outgassed using sample controlled thermal analysis, SCTA.$^{45,46}$ Around 0.3 g of each sample was thus heated under a constant residual vacuum pressure of 0.02 mbar up to a final temperature of 100 °C which was maintained until the residual pressure was less than 5 × 10$^{-3}$ mbar. The carbon dioxide and methane were obtained form Air Liquide (Alphagaz, France) and are of 99.998% purity. These greenhouse gases have been chosen with respect to their interest in several applications. From a more fundamental point of view, a comparison between a probe molecule with a significant quadrupole moment and house gases have been chosen with respect to their interest in several applications. From a more fundamental point of view, a comparison between a probe molecule with a significant quadrupole moment and house gases have been chosen with respect to their interest in several applications.

Figure 1. Projection of the structure of MIL-96 (Al) along the c axis, showing the hexagonal network of the aluminum octahedra (gray) containing 18-membered rings connected to the $\mu_3$-oxo-centered trinuclear units, via the trimesate ligand. Water molecules: black.

Hydrogen Storage Measurements. Hydrogen storage measurements were made using a volumetric device (Sieverts’ method) equipped with calibrated and thermalized volumes and pressure gauges. About 1 g of powder was transferred into a tight stainless steel sample holder. The sample was then outgassed at 150 °C during 18 h under primary vacuum. At the end of the experiment, the sample holder was transferred into a glovebox under purified argon and weighed again to measure the mass loss from the dehydrated sample. All weight capacities refer here to outgassed samples. For adsorption measurements at 77 K, the sample holder was immersed in liquid nitrogen and high-purity hydrogen (Alphagaz H2) was introduced step by step in the container up to 0.41 MPa. The pressure variations due to both gas cooling and hydrogen adsorption were measured as a function of time. Under these thermodynamic conditions (0 < P < 1 MPa, T = 77 K), the ideal gas law is no longer valid, and a different equation of state was used for this pressure and temperature range.47

Results and Discussion

Structure Description. The MIL-96 (Al) phase exhibits a 3D framework (Figure 1) consisting of the connection of octahedrally coordinated aluminum, which are linked through the trimesate ligand, \[ 3\text{-btc} \]. The presence of the \[ 3\text{-btc} \] ligand in the structure indicates that the trimethyl 1,3,5-benzenetricarboxylate ester is hydrolyzed into the carboxylate form, which further reacted with the aluminum cations under hydrothermal conditions. Here, the use of the methyl ester form of the carboxylic acid has a drastic effect on the synthesis conditions since no recrystallization of the trimesic acid is observed and a pure form of the MIL-96 (Al) is obtained. The method of the in situ carboxylate ligand synthesis was previously reported for the formation of 3D coordination polymers, and it was reported that the hydrothermal in situ slow hydrolysis reaction improved the crystallization process for the crystal engineering of such compounds.48,49 Using this strategy, analogous compounds of MIL-96 have been prepared with gallium50 and indium51 and will be discussed elsewhere.

Two distinct inorganic blocks have been found in the MIL-96 framework. The first one (Figure 2) is a discrete trinuclear moiety with the aluminum octahedra \[ \text{AlO}_6 \], corner-sharing a $\mu_3$-oxo centered group (corresponding to the crystallographic site Al1 (6h)). The four equatorial Al–O distances are 1.889- (2) Å while the Al–$\mu_3$O is shorter (1.834(2) Å). The remaining oxygen atom belongs to a coordinated water molecule, which is trans to the Al–$\mu_3$O bond. It corresponds to a longer Al–O distance (1.945(5) Å), and this is in good agreement with valence bond calculations52 (expected value for \[ \text{H}_2\text{O} \]: 0.4, calculated: 0.45). This type of trinuclear oxo-centered carboxylate-bridged cluster formulated as \([\text{Al}_3\text{O}(\text{O}_2\text{C})(\text{OH})_6\text{L}_3]^{1+}\) is quite well-known for many trivalent transition metals such as V, Cr, Mn, Fe, Co, Ru, Rh, Ir, and Pt.52 Most of these phases have been isolated using the acetate species, but more recently, such triangular metal-centered units have been encountered in phases crystallizing with aromatic polycarboxylates.49,50–55 However, to our knowledge, this is the first time that such a trinuclear configuration is observed in aluminum chemistry. Indeed, only one trinuclear complex was reported with aluminum.

The second inorganic part consists of a 2D network containing chains of aluminum \[ \text{Al}_2\text{O}_3\text{OH} \] and \[ \text{Al}_3\text{O}_3\text{OH} \] octahedra, interconnected with each other to generate a hexagonal 18-membered ring in the \((a,b)\) plane (Figure 1). There are two crystallographically inequivalent types of aluminum, Al2 and Al3, which are respectively coordinated to four and two oxygen atoms coming from the carboxylate groups of the trimesate molecule and four and two bridging $\mu_2$-hydroxo moieties: the

two oxo groups are in cis positions for Al2, whereas the hydroxo groups are in trans positions for Al3, with typical Al–O distances. The lengths of the Al2–OH bonds are in the range 1.837(1)–1.893(1) Å, and the Al2–O bond is longer at 1.906–2.0 Å (Al2–O3). The Al3–O6 distances correspond to the value of 1.862(1) Å, whereas the length of the Al3–OH bond is shorter at 1.836(1) Å (Al3–O2). The hydroxo groups are in a bridging position between two aluminum atoms. The valence bond calculations give values in the range 1.189–1.210 for oxygen atoms bridging aluminum, which are close to the expected value (1.2) for hydroxo species. This μ2-OH connection generates original corrugated infinite chains of aluminum octahedra, with a -cis-cis-trans- corner-sharing sequence (Figure 3). The sinusoidal shape of the octahedra file is induced by the ternary geometry of the trimesate molecule. Such a configuration was already reported in other metal carboxylates synthesized with the same trimesic acid. For instance in a cobalt-based solid,60 infinite chains of edge- and corner-sharing octahedra occur. In MIL-96, the particular corner-sharing octahedra connection mode differs from those previously observed in the tancote-like solids61,62 (straight chains with trans corner-sharing linkage [see for instance the MIL-5331/MIL-6932 topologies]) or KTP-like materials63,64 (zigzag chains with cis-trans- corner-sharing linkage). However, an identical arrangement of such corrugated corner-sharing chains was recently encountered in the titanium phosphate, Ti3P6O27âââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââââ â...
in this third cavity. Of the additional aluminum atom Al4, no empty space occurs would be surrounded by six hydroxo ligands. With the presence of the special position 12k. In accordance with the electroneutrality of the structure, the aluminum species Al4 one of the three possible sites can easily be done. It is based on a prior knowledge of the structural description. The signal, which has the smallest distance between the residues are quite short (2.7 Å), and two aluminum atoms on the Al4 site could not be face to face with each other because of the electrostatic repulsion. This involves the shift of the aluminum Al4 on one site over the three possible ones of the special position 12k. In accordance with the electroneutrality of the structure, the aluminum species Al4 would be surrounded by six hydroxo ligands. With the presence of the additional aluminum atom Al4, no empty space occurs in this third cavity.

NMR Investigations. The $^{27}$Al MAS NMR spectrum of Al$_2$O(OH)$_6$(H$_2$O)$_6$(Al$_2$(OH)$_6$)$_6$[btc]$_6$24H$_2$O (MIL-96) exhibited overlapped signals in the chemical shift range (from 20 to −40 ppm) for which aluminum is in an octahedral environment. This is in perfect agreement with the XRD structure determination, which indicated that all aluminum atoms were exclusively located on hexacoordinated sites. To gain more spectral resolution, MQMAS spectroscopy has been applied. This technique combines MAS and multiquantum spectroscopy that can remove completely the second-order quadrupolar interaction. Two experiments with different conditions and methods were performed. The first using the classical $\gamma$-filter three-pulse sequence with moderate rf power (~85 kHz) yielded three well-resolved signals with pure-absorption line shapes, shown in Figure 6. The 2D map is analyzed by modeling the quadrupolar line shape of the observed experimental signals. The second experiment used fast amplitude modulation (FAM) pulse sequence under strong rf power (~170 kHz) conditions. Significantly better sensitivity was afforded by the FAM technique enabling evidence of a fourth site with much larger quadrupolar coupling constant (see Supporting Information). The efficiency of multiple-quantum excitation and conversion is strongly dependent on the quadrupolar frequency ($\nu_Q$), resulting in spectra with intensities that may not quantitatively represent the relative site population. However, the resulting anisotropic line shapes that were severely distorted may be a result of the use of strong multiple-pulse rf irradiation. In this case, the lack of a full powder pattern prevented obtaining the quadrupolar parameters accurately via numerical fitting of the anisotropic line shape. It was thus possible to extract meaningful quadrupole parameters by the two combined MQMAS methods, which were used as a starting point for the simulation of the 1D MAS NMR spectrum to obtain site occupancy (Figure 7). Therefore, $^{27}$Al NMR resolved four distinct lines, which account for the four crystallographic sites Al1, Al2, Al3, and Al4.

The single-pulse 1D spectrum obtained with 30 kHz MAS (Figure 7) can be very well simulated with virtually identical dephasing $\delta_{CS},e^2\eta Qh,$ and $\eta$ values used to simulate the MQMAS spectra. Because the resolution is better, the MQMAS results provide the most meaningful values for the second-order quadrupolar powder parameters. Table 2 summarizes data of quadrupolar parameters used in the decomposition are shown.

According to these signal attributions, line integrals contribute to more disorder within their local environments. The most intense signal is attributed to the Al₂ site since this site adsorbs for each gas at 10 bar (4.4 mmol·L⁻¹) as well as for the adsorption of methane in the aluminosilicate zeolite NaY(17 kJ·mol⁻¹) as well as for the adsorption of methane in the aluminosilicate zeolite NaY(17 kJ·mol⁻¹). These values are all quite similar, emphasizing that the methane molecule, having no permanent moment, is not affected by the electric field gradient. One would expect differences due to strong confinement effects which do not seem to be the case.

Following the return to the baseline of the calorimetric signal gives an indication of the adsorption equilibrium. For the adsorption of carbon dioxide, complete adsorption equilibrium takes ~1 h for each point. It is noteworthy that the methane adsorption, > 5 h were required for the equilibrium of each point. For most porous solids, methane equilibrium is attained after 20 min per point. Such a long time to reach equilibrium is the first time, to our knowledge, that such behavior is observed and should be compared to the hydrogen experiments. Furthermore, the adsorption at room temperature can be compared with nitrogen adsorption at 77 K, and interestingly, almost no uptake.

The initial enthalpies of adsorption of carbon dioxide are very similar to those previously reported for MIL-53 (Al) and 32 kJ·mol⁻¹ for MIL-53 (Cr). These adsorption enthalpies observed are significantly lower than those obtained with aluminosilicate zeolites(72,73) (50–70 kJ·mol⁻¹) and in the same range as those observed for pure silica zeolites such as silicalite.72 This highlights the relative electronegativity of the MIL-96 framework as the quadrupole moment of the CO₂ molecule would engender stronger interactions, closer to those observed with the aluminosilicate zeolites.

Table 2. Data of Al Sites Present in Al₃O(OH)₁₈(H₂O)₉[Al₂(OH)₆] [btc]₂·24H₂O (MIL-96) from Simulation of the ²⁷Al MAS NMR Spectra Obtained in Both 3QMAS (Figure 6) and Single Pulse Experiments (Figure 7)

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* Accuracy: ±4%. 
* Quadrupole constant QCC = (e²Q)/h, with the z-component, V_z = eQ of ²⁷Al nuclei, and Planck’s constant, h. Accuracy: ±0.6 MHz. 
* Asymmetry parameter η = (V_x - V_y)/V_z as a function of the x-, y-, and z-components of the electric field gradient. Accuracy: ±0.03. 
* Isotropic chemical shift. Accuracy: ±0.8 ppm.

1.95 mmol·g⁻¹ for CH₄ is below many materials including zeolites,72–75 activated carbons,76–78 and other MOF materials.

The initial enthalpies of adsorption of carbon dioxide are very similar to those already observed for previous experiments on adsorption of carbon dioxide in the aluminosilicate zeolite NaY(17 kJ·mol⁻¹) as well as for the adsorption of methane in the aluminosilicate zeolite NaY(17–17.8 kJ·mol⁻¹). These values are all quite similar, emphasizing that the methane molecule, having no permanent moment, is not affected by the electric field gradient. One would expect differences due to strong confinement effects which do not seem to be the case.
was observed in the latter case, certainly due to the temperature differences of the experiments.

The above-mentioned behavior can be explained by the very small apertures of the pores. Carbon dioxide may be able to enter the apertures due to its smallest dimension of 2.5 Å and due to relatively favorable attraction to the free hydroxyls groups inside the pores. The adsorption of methane is probably hindered by its dimension of 3.8 Å which is very close to that of the pore apertures. It may also be limited by the residual water inside the cages. This presence of water may also explain the limited free volume for both of these probe molecules. Unfortunately, a higher heat treatment leads to a collapse of the structure and no significant adsorption. This is the first of a series of studies with MOF-type materials in which the presence of water either is beneficial or can significantly alter the adsorption of these gases. Nevertheless, it should be noted that saturation is not observed for carbon dioxide and methane adsorption, and a higher adsorption capacity can be expected at higher pressures.

Hydrogen Adsorption. Before the hydrogen adsorption experiments, the MIL-96 (Al) sample was thermally outgassed to remove the water molecules lying within the cages. After a thermal treatment at 150 °C overnight, a weight loss of 18.5% is observed in good agreement with the 19% measured by TGA at 200 °C. The hydrogen uptake was then measured at 77 K. Surprisingly, an unexpected behavior for physisorption was observed with a rather slow adsorption process, and the thermodynamic equilibrium was reached only after 6 to 7 h (Figure 9). This might be due to the small pore opening size of the cavity which would slow the hydrogen diffusion kinetic within the 3D network of the different cages in MIL-96. The adsorption isotherm shows a significant hydrogen uptake at moderate pressure and increases rapidly to reach 1.91 wt %, and a saturation plateau is observed above 0.3 MPa. The adsorption branch was tentatively measured down to low pressure, but the reversibility is rather poor, and the sample was not desorbed below 1.82 wt % (Figure 9). Again, the slow desorption kinetic might be attributed to poor hydrogen diffusion kinetics within the framework. The H2 uptake value is lower than that observed in the porous aluminum terephthalate MIL-53 (3.8% under 1.6 MPa) or some metal–organic framework compounds under higher pressures. Nevertheless, it is similar to the storage capacity of other microporous MOF solids such as Ni(dhtp)(H2O)2·8H2O, Prussian blue analogues, or the NaY zeolite.

Conclusions

Hydrothermal reaction of aluminum nitrate with 1,3,5-benzenetricarboxylic acid (or the methyl ester form, trimethyl 1,3,5-benzenetricarboxylate) in water led to a new aluminum trimesate compound: Al3+2(OH)3(H2O)3(Al3+2(OH)2)[btc]6·24H2O. The structure, solved by means of single-crystal X-ray diffraction, consists of a 3D framework containing isolated trimuclear μ3-oxo-bridged aluminum clusters and infinite chains of Al3+2(OH)2 octahedra forming a layerlike hexagonal network based on an 18-membered ring. As far as we are aware, this is the first time that such a μ3-oxo-centered trimuclear configuration is observed in aluminum chemistry. The organic molecules interact strongly with both the oxo-centered trimetric aluminum units and the 2D network of aluminum octahedra ensuring the cohesion of the structure. Solid-state NMR characterization agrees well with the structure description. The four crystallographic Al sites are resolved in 27Al spectra using 3QMAS. The compound exhibits a hydrogen storage capacity up to 1.91 wt % at 77 K under 3 bar and is able to sorb carbon dioxide (4.4 mmol·g−1 at 10 bar) and methane (1.95 mmol·g−1 at 10 bar).

The remarkable feature is the occurrence of the μ3-oxo-centered trimuclear aluminum motif in the structure of MIL-96. Such a trimeric building block was previously described in very promising MOF materials based on super-tetrahedral units for the construction of zeolite-type frameworks with very large pores in MIL-100 and MIL-101 or solids exhibiting very large swelling effects in the MIL-88 series. Future work will be focused on the isolation of such a triangular unit and its reactivity with the carboxylate ligands. The synthesis of the MIL-100 compound incorporating aluminum is under progress and will be reported soon.

Acknowledgment.

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Supporting Information Available:
Crystallographic data in CIF format and figures showing the FAM-II-MQMAS 27Al spectrum, IR spectrum, and TG curve. This material is available free of charge via the Internet at http://pubs.acs.org.

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