The Influence of Isomerism on Crystallization in Aluminum Pyridinedicarboxylate Coordination Compounds

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Dedicated to Professor Wolfgang Bensch on the Occasion of his 65th Birthday

Abstract. The reactions of three isomeric pyridinedicarboxylic acid linkers – 2,4-, 2,6-, and 3,5-pyridinedicarboxylic acid – with various aluminum salts were investigated by high-throughput solvothermal synthesis. Two coordination compounds were obtained from the symmetrical linkers (2,6- and 3,5-pyridinedicarboxylic acid). The former is based on molecular Al–O dimers, similar to those reported in the coordination polymer CAU-16. The latter forms a porous extended framework and is a member of the CAU-10 family of compounds. Full characterization of both compounds is reported. Crystal structures were determined from high-resolution synchrotron powder diffraction data. The number of guest water molecules determined crystallographically is in excellent agreement with the number found by thermogravimetric and elemental analyses. It was found that the CAU-10 analogue undergoes a reversible, displacive phase transition from a non-centrosymmetric to a centrosymmetric space group on loss of guest water molecules, due to guest-framework interactions.

Introduction

The application of metal-organic frameworks (MOFs) as water sorbents for de-/humidification and water production applications has grown in popularity in recent years.[1–3] A precondition for such use is long term stability under operating conditions, which is usually investigated by repeated cycles of adsorption and desorption under hydrothermal conditions.[4] MOFs constructed from high-valence metals, such as Zr4+ or Cr3+, demonstrate good stability under these conditions, compared to the significantly larger number of MOFs reported with divalent metal cations.[5] Among these higher valent compounds, a number of Al3+-based MOFs have demonstrated notably better stability and water sorption performances. Aluminum fumarate [Al(OH)(O2C–C6H4–CO2)], also known as CAU-10 (CAU = Christian-Albrechts-Universität),[10] has shown even greater stability and water sorption performances. Aluminum fumarate undergoes a reversible, displacive phase transition from a non-centrosymmetric to a centrosymmetric space group on loss of guest water molecules, due to guest-framework interactions.

It should be noted that all three of these compounds have a structural motif of unidirectional channels, deriving from the μ2-hydroxo corner-sharing chains, which are recognized as a chemically robust secondary building unit (SBU).[5]

A particular feature of the synthetic chemistry of MOFs is the ease with which compounds with the same network topology, but incorporating different SBUs may be prepared. The approach, referred to as either isoreticular synthesis[14] or scale chemistry,[15] has been widely applied to the replacement of ligands, for example in the case of DUT-4 or DUT-5, which both have MIL-53 structures, but where terephthalate is replaced by 2,6-naphthalenedicarboxylate or 4,4′-biphenyldicarboxylate, respectively;[16] and it has been applied to the partial or complete replacement of metal cations/clusters, as for example with Zr4+ and Ce4+ in the compounds MIL-66.[17]

During the initial characterization of CAU-10, the compound demonstrated a strong influence of the functional group on the sorption properties,[10] and moreover was also found to show good stability against hydrolysis. The inclusion of an additional coordinating functional group might be expected to change the host-guest interactions or lead to a different synthetic pathway and hence a new product, thanks to different coordinative bonding possibilities.[18] The isomeric organic molecules 2,4-pyridinedicarboxylic acid (H22,4pydc), 2,6-pyridinedicarboxylic acid (H22,6pydc) and 3,5-pyridinedicarboxylic acid (H23,5pydc) all have the same meta-substitution pattern of the carboxylic acid groups relative to one another and are hence structural analogues of isophthalic acid, which is used in the synthesis of CAU-10 (Scheme 1). We also note that H22,6pydc has an arrangement related to that of 2,4,6-pyridine–dicarboxylic acid (H3PTC). On reaction with Al3+, H3PTC forms first a molecular compound, [Al(μ–OH)(H2O)(PTC)]2, and subsequently a wine-rack chan-
nel structure, labeled CAU-16 ([Al(μ-OH)₆]Al(μ-OH)(H₂O)·(PTC)₁₂),[¹⁹] CAU-16 forms through coordination of Al³⁺ cations by the free carbonylate group of the molecular dimer, which is thought to be a structural intermediate.

Scheme 1. The three isomeric pyridinedicarboxylic acid linkers investigated.

Herein we report a synthetic study of the complex-forming chemistry of three isomeric pyridinedicarboxylic acid linkers, determining the conditions necessary to obtain a pyridine functionalized form of CAU-10 (CAU-10-pydc). Whilst the water sorption properties of CAU-10-pydc have been reported previously,[¹²] herein we present a full structural characterization of the compound and report a new dimeric Al–O cluster compound, closely related to CAU-16.

Results and Discussion

Synthesis

The reaction of Al³⁺ salts with three isomeric pyridine dicarboxylic acid linkers (Scheme 1) was investigated using an in-house developed 24 reactor multiclave.[²⁰] Reactions of H₂ 3,5pydc with six Al³⁺ salts [AlCl₃, Al(NO₃)₃, Al(OH)₃, Al(OH)(AcO)₂, Al₂O₃ and Al₂(SO₄)₃]; full hydration states of all salts given in Experimental Section] in a DMF:H₂O mixture (1:1) were first investigated. From Al(NO₃)₃, AlCl₃, and Al₂(SO₄)₃, a highly crystalline bright yellow powder was obtained, labelled 1-AP and later found to be isostructural with the known compound CAU-10 (CAU-10-pydc; see Structural Analysis). The most crystalline product was obtained from Al(NO₃)₃. A subsequent high-throughput optimization array was used to determine the optimum stoichiometry and also the DMF:H₂O solvent ratio. This was obtained with an Al(NO₃)₃: H₂O:H₂ 3,5pydc ratio of 1.5:1 and a DMF:H₂O ratio of 9:1.

(Figure 2), the synthesis of which was successfully scaled up (see Experimental Section: Optimized scale-up synthesis of 2). With H₂ 2,4pydc only poorly crystalline products were obtained and further synthetic optimization was therefore not pursued.

Structural Analysis

Two crystalline compounds were obtained from the synthetic study (see Section Synthesis); both were characterized by powder X-ray diffraction and for compound 2 a single-crystal X-ray diffraction study could also be performed. The first, CAU-10-pydc, could only be obtained as a microcrystalline powder and for the as-prepared form (1-AP) it was not possible to reliably determine the nature and composition of
Table 1. Crystallographic parameters from the final Rietveld refinements. See Supporting Information for complete list of refinement parameters.

| Conditions     | 1            | 1-H$_2$O     | 2            |
|----------------|--------------|--------------|--------------|
| a /Å           | 21.5100(3)   | 21.4002(3)   | 7.19193(4)   |
| b /Å           | 10.71549(6)  | 10.67754(17) | 10.87754(6)  |
| c /Å           | 21.5100(3)   | 21.4002(3)   | 7.19193(4)   |
| β /°           | 97.5836(4)   | 97.5836(4)   | 97.5836(4)   |
| V /Å$^3$       | 4865.11(16)  | 4889.98(14)  | 830.946(8)   |
| Space Group    | I$_4$_1/amd  | I$_4$_1/md   | I$_2$/m      |
| $R_{wp}$ /%    | 2.99         | 2.44         | 9.62         |
| $R_{Bragg}$ /% | 0.91         | 1.80         | 4.37         |
| $\chi^2$      | 22.735 a)    | 14.948       | 9.672        |
| $\lambda$ /Å  | 0.826952     | 0.826952     | 0.8260285    |

a) Higher $\chi^2$ for 1 is due to textural effects in certain reflections, which could not be fitted with strain broadening models. The origin of this is under investigation.

...the guest species in the pores. Thus, the structures of the activated compound, under vacuum at 150 °C (1) and the structure of the hydrated compound (after removal of the guest molecules by calcinations, 1-H$_2$O) are here reported and compared. For 2 only the crystal structure of the as-prepared compound is reported, since no guest molecules are present and dehydration results in a non-crystalline product.

[Al(OH)(3,5pydc)] (1)

The structure of dehydrated (activated) [Al(OH)(3,5pydc)] (1) was obtained by heating a calcined sample of 1 to 150 °C under vacuum for 30 mins, to remove physisorbed water. Synchrotron powder X-ray diffraction (SR-PXRD) data were indexed in the tetragonal space group I$_4$_1/amd, which is also adopted by other desolvated members of the CAU-10 family of compounds (e.g. CAU-10-CH$_3$[10] and MIL-160;[13] see Supporting Information for details of systematic absences). An initial model was developed from the desolvated form of CAU-10-CH$_3$ and Rietveld refined against the SR-PXRD data (see Experimental Section for details). Details of the crystallographic parameters of the final refinement are given in Table 1 and a Rietveld plot for the final cycles of refinement is shown in Figure 3a.

Each central Al$^{3+}$ ion is octahedrally coordinated by six oxygen atoms, two of which – O1 – are shared in a bridging fashion with neighboring Al$^{3+}$ sites along the c-direction. The shared O1 and the Al site are slightly displaced in the ab plane with respect to the 41 screw axis, thus helical chains are developed parallel to the c-direction (Figure 4a); it should also be noted that the helices of neighboring chains turn in opposite senses. Chains are linked together through two symmetry independent 3,5pydc$^{2-}$ ligands to form a pore network of corrugated unidirectional channels parallel to the c-direction, with the N-heteroatom of the ligand ring pointed into the cavity (cf. MIL-160; Figure 4b). The channels consist of larger cavities (~5.79 Å free diameter) linked by narrower windows (3.76 Å free diameter; pore size distribution plot and details of PoreBlazer[21] calculation given in the Supporting Information). For charge balance, the O1 sites must be protonated; evidence for this can be seen in the IR spectra (see Infrared Spectroscopic Studies).

[Al(OH)(3,5pydc)]·3.7H$_2$O (1-H$_2$O)

The structure of [Al(OH)(3,5pydc)]·3.7H$_2$O (1-H$_2$O) (Figure 5), was confirmed by Rietveld refinement of SR-PXRD data collected on a sample of CAU-10-pydc, which had been calcined and adsorbed water on cooling (1-H$_2$O). Data were collected under ambient conditions and indexed in the noncentrosymmetric space group I$_4$_1/md, a sub-group of I$_4$_1/amd adopted by the desolvated compound, which gave the best fit to the observed systematic absences (see Supporting Infor-
Figure 4. The crystal structure of CAU-10-pydc, 1, showing (a) the structure of the helical chains of corner-sharing AlO$_6$ polyhedra – the pink dashes indicate the $4_1$ axis; and (b) a view along the $c$-direction, showing the unidirectional channels. C atoms in black, N atoms pale blue, O atoms red, Al atoms/polyhedral darker blue. Dashed blue line indicates unit cell edges. Hydrogen atoms are omitted for clarity and as they were not part of the final refinement.

Figure 5. Structure of 1-H$_2$O, showing the distribution of H$_2$O molecules within the channels. Hydrogen-bonding interactions are indicated between guests (red dashes) and guests and framework (blue dashes). Partially filled spheres indicate partially occupied O atoms (H$_2$O molecules). Hydrogen atoms are omitted for clarity and as they were not part of the final refinement.
In the present compound 2, the asymmetric unit consists of one Al site, half a 2,6pydc$^{2-}$ unit, one μ-OH, and one H$_2$O molecule. The center of mass of all four of these components is located on the mirror plane. Thus, each Al site is octahedrally coordinated by five O atoms – two from monodentate carboxylic acid groups, two from μ-OH groups and one from a terminal H$_2$O molecule – and one N atom from the pyridinyl ring. Typically the coordination chemistry of Al$^{3+}$ is dominated by oxophilic interactions.[24,25] In the structure of 2, coordination by the pyridinyl N atom occurs due to the chelating nature of the O/N/O pocket. Two of these monomeric [Al(μ-OH)(H$_2$O)(2,6pydc)]$^2-$ units link in an edge-sharing fashion through the two μ-OH groups to form dimers. Dimers interact through weaker and stronger hydrogen-bonding interactions between the carboxylate O2 and terminal water O3 sites above and below the plane of the 2,6pydc linkers [2.554(4) Å and 3.112(4) Å, respectively]. Dimers are densely packed within the cell and no porosity is developed.

**Thermoanalytical Studies**

Both compounds 1 and 2 were analyzed by thermogravimetric analysis (TGA) and elemental analysis to understand the composition and importantly the solvent molecules occluded in their pore space.

\[\text{[Al(OH)(3,5pydc)]·X (1-AP)}\]

The TGA trace for 1-AP shows two principal weight losses, the first, beginning at 25 °C and complete by 270 °C, is associated with the loss of guests from the pore space, and the second, starting at 415 °C and finished by 550 °C, is related to destruction of the framework (Figure 7, top). The trace of 1-H$_2$O shows a similar pattern of weight losses, except that the first step, the desolvation process, is complete by 80 °C (see below).

In 1-AP, the desolvation process occurs in two steps: 25 °C to 110 °C is attributed to the loss of 3.4 molecules of water from the pores (20.2 wt%; water content calculated assuming a final product of Al$_2$O$_3$); and 140 °C to 270 °C, assigned to the loss of DMF or breakdown products thereof (8.0 wt%, equating to 0.3 molecules of DMF per Al). As ethanol is also used in the washing of the product, it is likely that some is present as guest molecules. Due to the additional complexity of modeling this however, it has not been considered in the analysis of the TGA data. No further weight losses are observed up to 415 °C, showing the high stability of the framework. Above 415 °C, collapse of the framework occurs, also in a two-step process: between 415 °C and 440 °C, a weight loss of 8 wt% occurs, which is assigned to dehydroxylation of the helical chains (cf. for example, MIL-91[26]). This is rapidly followed by a second weight loss from 465 °C to 550 °C (42.4 wt%) corresponding to the loss of linker molecules (observed: 0.78 molecules per Al; expected: 1.0; discrepancy due to retention of some O atoms from linker in final product).

\[\text{[Al(OH)(3,5pydc)]·3H$_2$O (1-H$_2$O)}\]

To better understand the thermal behavior of compound 1 it was deemed necessary to remove the occluded DMF/break-
down products. To that end a sample of 1-AP was heated to 250 °C in a tube furnace in a flow of nitrogen gas for 3 h. The final product was found to have a pale cream color and a PXRD pattern showed no significant changes (see Structural Analysis), confirming the thermal stability of the framework.

1-H2O shows a similar pattern of weight losses during TGA measurements as 1-AP (Figure 7, bottom): an initial weight loss attributed to desolvation begins at 25 °C and is complete by 80 °C; whilst a second group of weight losses occur from 380 °C to 570 °C. In 1-H2O, desolvation proceeds through a single step process, attributed to the loss of 3.7 molecules of water per Al atom (observed: 23.7 wt %; expected: 24.8 wt % – calculated for [Al(OH)(C7H3NO4)]·3.7H2O). This is in excellent agreement with the crystallographically determined number of water molecules. No weight losses are observed between 80 and 380 °C, further emphasizing the thermal stability of the framework. However the onset of framework decomposition is approximately 35 °C degrees lower than for 1-AP. This difference is thought to be due to the introduction of defects in the framework during the activation procedure: that is the thermal cleavage of some framework bonds, but without the liberation of framework components. 1-H2O shows a similar two-step decomposition process to 1-AP, with the first step (380 to 425 °C; 7.1 wt %, expected: 6.7 wt %) again attributed to the dehydroxylation of the framework and the second step (450 to 570 °C: 44.1 wt %, expected 43.6 wt % – assuming one O atom retained in oxide product).

Elemental analysis data are in good agreement with TGA and gave C 29.60 , H 4.49 , N 5.04 % (calculated, assuming composition Al(OH)(C7H3NO4)·3.7H2O: C 30.49 , H 4.17 , N 5.07 %).

Z. Anorg. Allg. Chem. 2018, 1816–1825

Figure 7. TGA plots for as-prepared 1 (1-AP) (top) and 1 rehydrated after calcination at 250 °C (1-H2O) (bottom) under flowing air from room temperature to 800 °C. Principal mass losses are indicated.

Infrared Spectroscopic Studies

Infrared spectra were measured for both the as-prepared compound 1 and for the compound after calcination at 300 °C (when it contains only H2O guests). Both spectra show the same principle absorption bands (Figure 9; see Supporting Information for complete assignment). Differences between the two spectra are due to the different guest species present: in 1-AP guests are unknown breakdown products from DMF, water and possibly ethanol (the latter two from washing of the product after synthesis); in 1-H2O, H2O molecules are the only guest species. In 1-AP, there are two distinct peaks at 3625
and 3591 cm$^{-1}$ which are attributed to an N–H stretch and the chain OH group respectively. In the same region in 1-H$_2$O, the peak at 3603 cm$^{-1}$ is attributed to the chain OH group, the slight shift being due to changes in the interactions of this group with other guest species. The observation of an OH stretch for the framework of 1-H$_2$O confirms that corner-shared O atoms of the chains must be protonated. The broad adsorption in both spectra, centered around 3350 cm$^{-1}$, is narrower and reduced in intensity in 1-H$_2$O, reflecting the reduced diversity of hydrogen-bonding interactions when only H$_2$O is present compared to the mixture of H$_2$O and unknown guests in 1-AP. In the lower wavenumber region of the spectra, further differences are observed with peaks at 1663 and 1101 cm$^{-1}$ absent from the spectrum of 1-H$_2$O. These peaks can be assigned to stretching vibrations expected for amides (e.g. C=O or C–N). Given the observed differences in the spectra and the reaction conditions, it is hypothesized that the unknown guest species are likely either DMF, dimethylamine, formic acid, or other related DMF breakdown products.

\[\text{[Al(µ-OH)(H}_2\text{O)(2,6pydc)]}_2 \ (2)\]

The infrared spectrum of 2 is similar to the reported compound [Al(µ-OH)(H$_2$O)(HPTC)]$_2$.\textsuperscript{19} A broad absorption band centered around 3540 cm$^{-1}$ is assigned to an hydrogen bonding OH group, as anticipated from the crystal structure (Figure 10). This compares to 3432 cm$^{-1}$ in the reported compound, indicating the stronger nature of the bond in the present compound. Absorption bands for the ligand are also similar to those of [Al(µ-OH)(H$_2$O)(HPTC)]$_2$, e.g. 1651 cm$^{-1}$ – the coordinated C=O stretch; and 1585 cm$^{-1}$, the O–C–O anti-symmetric stretch (see Supporting Information for complete assignment).

**N$_2$ Sorption Studies**

CAU-10-pydc, compound 1, has a unidirectional channel structure, similar to that of the other members of the CAU-10 family of compounds. To assess the porosity of this compound, a N$_2$ sorption isotherm at 77 K was measured on a sample of 1-H$_2$O. The sample was activated by heating to 150 °C for 3 h under vacuum of 1×10$^{-4}$ mbar to ensure all guests were removed. The adsorption isotherm shows a Type I shape up to \(p/p_0 \approx 0.06\). Above this point, a second adsorption step occurs, the origin of which is under investigation (Figure 11). No hysteresis is observed during desorption, including over the step region.

Using the first adsorption step, a BET surface area of 884 m$^2$·g$^{-1}$ was determined, which is in good agreement with other members of the CAU-10 family of compounds.\textsuperscript{10} Dubinin-Radushkevitch analysis for N$_2$ at 77 K gives a micropore volume of 0.34 cm$^3$·g$^{-1}$ (\(N_0 = 9.69 \text{ mmol·g}^{-1}\)).\textsuperscript{27} This compares to a theoretical micropore volume, calculated using PoreBlazer\textsuperscript{21}, of 0.40 cm$^3$·g$^{-1}$. Differences between the calculated and observed micropore volume are attributed to pore blocking within the real framework and that the simulation uses the Universal Force Field\textsuperscript{28} as a model for interactions, which may not be applicable in all cases.
Conclusions

From three isomeric pyridinedicarboxylic acid linkers, in which the carboxylic acid groups are meta-substituted relative to one another, two new aluminum carboxylate coordination compounds were prepared. The substitution position of the pyridinyl N atom relative to the carboxylate groups strongly influences the composition and structures formed. In the lowest symmetry case of 2,4-pyridinedicarboxylic acid, only poorly crystalline products could be obtained. For the high-symmetry, 3,5-pyridinedicarboxylic acid, an extended coordination polymer, 1, isostructural to CAU-10 is obtained. For the equally high symmetry 2,6-pyridinedicarboxylic acid, the O/N chelating pocket forces N to engage in Al–N bonding and leads to the formation of a non-porous, molecular compound, 2, consisting of dimeric Al–O units. The structure is similar to the dimeric linkers observed in CAU-16 and the structure of [Al(μ-OH)(H₂O)(H₂PTC)]₂ and is also a structural analogue of two transition metal complexes, formed with Cr₃⁺ and Mn₃⁺ ions and 2,6-pyridinedicarboxylic acid.[19,22,23]

The CAU-10 analogue, CAU-10-pydc, crystallizes as a bright yellow compound (1-AP), which is thermally stable to the removal, at high temperature, of unknown guest molecules (derived from the synthetic procedure). On cooling the compound adsorbs aerial water to yield 1·H₂O. Hydrated CAU-10-pydc (1·H₂O) exhibits symmetry consistent with the non-centrosymmetric space group I₄₁,dmd. Such behavior is known for other members of the CAU-10 family of compounds. The driving force for the phase transition in CAU-10-pydc is the interaction of guest water molecules with the framework pyridinyl N atoms, which cause a distortion of the framework. The de-solvated form of 1 is porous to N₂ with a BET surface area of 884 m²·g⁻¹ and a pore-volume of 0.34 cm³·g⁻¹, in good agreement with simulation.

Experimental Section

High Throughput Discovery Synthesis: For the discovery array with H₂3,5pydc, the linker and six aluminum salts [AlCl₃·6H₂O, Al(NO₃)₃·9H₂O, Al(OH)₃, Al(OH)(AcO)₂, Al₂O₃, and Al₂(SO₄)₃·18H₂O] were loaded into Teflon® liners of a 24 reactor multiclave in a 1:1 stoichiometric ratio of 0.4 mol·L⁻¹ and a pore-volume of 0.34 cm³·g⁻¹, in good agreement with simulation.

Reaction Optimization

Al(NO₃)₃ with H₂3,5pydc: Al(NO₃)₃·9H₂O and H₂3,5pydc in stoichiometric ratio 1:1 were loaded into Teflon® reactors of a 24 reactor multiclave. To this were added DMF and H₂O solvents in ratios 10:0, 8:3:1.67, 6.67:3.33, 5:5, 3:3:6.67, 1:6.7:8.33 and 0:10 (1500 μL total volume). In addition to these, three reactions with Al(NO₃)₃·9H₂O and H₂3,5pydc ratios 1:1.5, 1:5, and 2:1 with DMF and H₂O in a constant ratio of 9:1 (1500 μL total volume) were also undertaken, yielding a total of 10 reactions studied in parallel. The reaction concentrations were kept constant (0.4 mol·dm⁻³). The reactor was heated to 150 °C for 12 h before cooling back to room temperature. The most crystalline product was obtained for the stoichiometric ratio 1:1.5 and solvent ratio 9:1 (Figure 1). Reactions with more DMF than H₂O gave crystalline 1-AP as the product; reactions with equal amounts of DMF and water or more water gave X-ray amorphous products or recrystallized ligand (Figure 1).

The reaction temperature was optimized by taking the optimized conditions and performing three further identical reactions at 80, 120, and 160 °C in culture tubes. A temperature of 120 °C was found to produce the most crystalline product.

Al(NO₃)₃ with H₂2,4pydc or H₂2,6pydc: Two high-throughput optimization arrays were used to investigate reactions of Al(NO₃)₃·9H₂O with H₂2,4pydc and H₂2,6pydc. Al(NO₃)₃·9H₂O and the respective ligand were loaded into Teflon® reactors of a 24 reactor multiclave in stoichiometric ratios varying between 2:1 and 1:2. DMF:H₂O solvent mixtures in ratios of 9:1, 1:1, or 1:9 were added; with H₂2,6pydc pure ethanol and 2-propanol were also investigated as solvents. The reactions had volumes of 1500 μL (half-filling the reactors) and concentrations of 0.4 mol·dm⁻³, and were heated to 120 °C for 18 h before cooling back to room temperature.

With H₂2,6pydc large, white blocky crystals of a phase labeled 2 were obtained from all reactions. The most crystalline products were obtained with an Al(NO₃)₃·H₂2,6pydc stoichiometry of 1:1.5 and DMF:H₂O ratio of 1:9; therefore these conditions were used for the scale-up synthesis. By contrast, with H₂2,4pydc only poorly crystalline products were obtained.

Optimized Scale-Up Syntheses

Synthesis of 1: A mixture of solid H₂3,5pydc (0.80 g, 4.80 mmol) and Al(NO₃)₃·9H₂O (1.20 g, 3.20 mmol) was loaded into a 100 mL Duran glass screw-top jar. To this was added DMF (18 mL) and distilled water (2 mL) to yield a total reaction volume of 20 mL (concentration: 0.4 mol·dm⁻³). The reaction was stirred until clear before being sealed. The reaction was then heated at 120 °C for 18 h in a convection oven, after which time a fine, bright yellow precipitate was observed at the bottom of the reactor. The reactions were allowed to cool naturally to room temperature. The solid was separated by centrifugation in the mother liquor (3000 rpm, 30 min). The supernatant was carefully decanted off. Afterwards, the solid was washed with ethanol (20 mL) and centrifuged again (3000 rpm, 10 min) to remove remaining DMF. Washing was repeated once more before the solid was dried in vacuo at 100 °C to remove remaining ethanol. The dry yellow solid was then transferred to an alumina boat and calcined at 250 °C in flowing N₂ for 3 h. The final cream-colored product was allowed to cool back to room temperature overnight in flowing N₂. The final solid had a mass of 0.270 g (90.90 mmol (ex. solvent); yield: 28 %).

Synthesis of 2: A mixture of Al(NO₃)₃·9H₂O (0.375 g, 1.00 mmol) and H₂2,6pydc (0.167 g, 1.00 mmol) was loaded into a 20 mL microwave vial. To this was added DMF (0.5 mL) and H₂O (4.5 mL) to give a final ratio of 1:9 and a final concentration of 0.4 mol·dm⁻³. The vial was sealed and heated in a convection oven to 120 °C for 15 h before being cooled back to room temperature. The final product of white
blocky crystals was separated by vacuum filtration and washed with distilled water (2 × 10 mL). The final solid had a mass of 0.218 g (0.48 mmol; yield: 96 %, based on Al).

Details of Crystallographic Studies

[Al(OH)3,5pydc]·3.7H2O (1-H2O) and [Al(OH)3,5pydc] (1): Synchrotron powder X-ray diffraction data for both the hydrated (1-H2O) and desolvated forms of 1 were collected at beamline I11 at Diamond Light Source (Oxon., UK). A sample of hydrated CAU-10-pydc (1-H2O) was ground and loaded into a 0.5 mm quartz glass capillary tube. The sample was then mounted on an H11 gas cell, mounted on the diffractometer and connected to the H11 gas-handling system.[29] Data were collected at a wavelength of λ = 0.826952 Å using the Position Sensitive Detector (PSD) over the range 0–90° 2θ with a step size of 0.004° in two steps (δ = 2.00° and δ = 2.25°) to allow for the gaps between detector modules; these scans were merged as part of the data acquisition. Data for the dehydrated form of 1 were collected using the same procedure on the same sample after heating to 150 °C under vacuum for 1 h.

Data were cut off above 40° 2θ as above this diffraction angle peak intensities were too low to distinguish from the background. Data for the desolvated form of CAU-10-pydc, 1, were analyzed first. Data were indexed, using the routines available in TOPAS-Academic,[10] in the tetragonal space group I4_1/amdd and Le Bail fitted to give a cell a = 21.5064(3) Å, c = 10.51189(16) Å. As this is the same symmetry as CAU-10-CH3,[10] a model for 1 was built by replacing 5-methylisophthalate linkers with 3,5-pyridinedimethylcarboxylate units and geometry optimizing the resulting structure in the refined cell, using the Forcite routine of Materials Studio.[31] The resulting structure was refined by the Rietveld method using TOPAS-Academic. Final refinement parameters are given in Table 1 and a Rietveld plot is shown in Figure 3a (see Supporting Information for complete list of refinement parameters).

For the hydrated form of CAU-10, 1-H2O, indexing gave a cell in the space group I4_1/mcd with Le Bail fitted lattice parameters a = 21.3963(2) Å, c = 10.6740(2) Å. The symmetry of desolvated 1 was lowered using the program POWDER CELL,[32] and the resulting structure geometry optimized using the Forcite routine of Materials Studio. H2O molecules were located within the pores of the resulting model by Fourier difference maps as a final map in series of Rietveld refinement cycles, performed using TOPAS-Academic V6. The occupancies and positions of each molecule were allowed to refine freely. Finally eight symmetry independent water molecules were identified, three of which are fully occupied. Details of the final refinement parameters are given in Table 1 and a Rietveld plot is shown in Figure 3b (see Supporting Information for complete list of refinement parameters).

[Al2.6pydc(µ-OH)(H2O)3] (2): Single crystal X-ray diffraction data were measured at beamline I19 at Diamond Light Source (Oxon., UK). A crystal was selected and mounted on an imide loop and placed on the diffractometer before being cooled to 80 K. Data were collected using a Pilatus 300K detector at a wavelength of λ = 0.6889 Å. Data were reduced using the CrystAlisPRO software.[33] Systematic absences were consistent with a monoclinic unit cell (a = 7.1605(2) Å, b = 10.7407(5) Å, c = 10.8433(4) Å, β = 96.975(3)°; space group: I2/m), however it was noticed that a number of satellite peaks were present in the data, possibly indicating a modulated structure. The frames were integrated although this gave a high Rint (Rint = 6.06 %) – attributed in part to the satellite peaks. The structure was solved and refined using the routines available in the SHELX suite, within OLEX2,[34,35] but with relatively large final R factors (R1 = 7.07 % and wR2 = 18.93 %). Moreover in the final structure (see Supporting Information), the thermal ellipsoids of the Al1, O1 and O3 sites have a strongly prolate shape, indicating a distribution of atomic positions and hinting at the nature of the modulation present in the structure.

To more accurately determine the structure, room temperature synchrotron powder X-ray diffraction data were collected at beamline I11 (Diamond Light Source, Oxon, UK) using the multi-analyzer crystal detector.[36] Data were collected over the range 0–150° 2θ with wavelength λ = 0.8260285 Å. Data were cut off above 65° 2θ as the intensity of the diffraction peaks became difficult to distinguish from the background. Data were successfully indexed in the triclinic space group P1 (a = 8.12758(9) Å, b = 7.19259(4) Å, c = 8.12851(7) Å, α = 69.2738(6)°, β = 82.4838(4)°, γ = 69.2461(6)°) using TOPAS-Academic.[10] This triclinic cell is related to the I2/m cell from single crystal diffraction by a simple transformation, which removes the centering and reduces the cell volume by half. The structure obtained from single crystal diffraction was thus transformed using PLATON[37] and successfully refined by the Rietveld method using TOPAS-Academic. Errors on the isotropic displacement parameters were found to be noticeably large in the final refinement and when the final structure was tested for missing symmetry, using the ADDSYM routine of PLATON, a new monoclinic cell in the space group I2/m was suggested [a = 7.19139(4) Å, b = 10.71549(6) Å, c = 10.87754(6) Å, β = 97.5836(4)°], which is very similar to the cell found from the single crystal study. The structure was transformed into this new cell and re-refined by the Rietveld method. The final refinement gave a fit of Rwp = 9.62 % (R2 = 9.672). In the final refinement all peaks are fitted (i.e. no satellite peaks were observed); the slightly high values of the R factor is typical for the refinement of MAC data on a coordination polymer. Final refinement parameters are given in Table 1 and a Rietveld plot is shown in Figure 3c (see Supporting Information for complete list of refinement parameters).

Crystallographic data (excluding structure factors) for the structures in this paper have been deposited with the Cambridge Crystallographic Data Centre, CCDC, 12 Union Road, Cambridge CB2 1EZ, UK. Copies of the data can be obtained free of charge on quoting the deposition numbers CCDC-1861790 (1), CCDC-1861789 (1-H2O), and CCDC-1861788 (2) (Fax: +44-1223-336-033; E-Mail: deposit@ccdc.cam.ac.uk, http://www.ccdc.cam.ac.uk).

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Supporting Information (see footnote on the first page of this article): Thermogravimetric and elemental analyses, IR spectroscopy, summary of crystallographic results (including systematic absence comparisons, Rietveld plots, single crystal diffraction study) and Pore Size Distribution plots.

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