

Published on Web 10/02/2009

Synthesis and Hydrogen Storage Properties of Be₁₂(OH)₁₂(1,3,5-benzenetribenzoate)₄

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Owing to their convenient modular synthesis and high surface areas, metal-organic frameworks have come under intense recent scrutiny for potential applications in gas storage.¹ The dense storage of hydrogen for use in fuel cell vehicles presents a particularly difficult challenge as a consequence of its high fugacity, which has prompted development of insulated pressure tanks for cryogenic storage.² For an adsorbent within such a system, Zn₄O(1,4-benzenedicarboxylate)₃ (MOF-5) offers the best known hydrogen storage properties, exhibiting a total uptake of 10 wt % and 66 g/L at 77 K and 100 bar.3 One possible means of improving the gravimetric capacity for this type of material would be to replace the typically di- or trivalent transition metal centers with light main group ions such as Be²⁺, B³⁺, Mg²⁺, or Al^{3+, 4} For example, substitution of the lightest divalent metal, Be²⁺, for the Zn²⁺ ions in MOF-5 would be expected to result in a roughly 40% increase in both surface area and gravimetric hydrogen storage capacity.^{5,6} Although early experiments involving addition of dicarboxylate bridging ligands to beryllium-containing solutions were recognized as leading to threedimensional polymeric solids (i.e., metal-organic frameworks),⁷ the isolation of such materials as pure, crystalline phases has proven elusive. Herein, we report the synthesis and hydrogen storage properties of the first structurally characterized metal-organic framework based upon beryllium: $Be_{12}(OH)_{12}(1,3,5-benzenetribenzoate)_4$ (1).

The solvated form of compound 1 was obtained as colorless blockshaped crystals by heating a solution of beryllium nitrate and 1,3,5benzenetribenzoic acid (H₃BTB) in a mixture of DMSO, DMF, and water at 130 °C for an extended period of 10 days. X-ray analysis of an evacuated single crystal revealed a highly porous network structure consisting of [Be₁₂(OH)₁₂]¹²⁺ rings connected through the tritopic BTB³⁻ ligands (see Figure 1). The saddle-shaped ring units are composed of tetrahedrally coordinated Be²⁺ ions linked around the inside edge via bridging hydroxide anions and around the periphery via bridging benzoate groups. These rings are observed every 27 Å along the [001] direction in the structure and reside on a 4 symmetry site. Each benzoate group is disposed $\sim 90^{\circ}$ to its neighboring benzoate groups, alternating in an up and down fashion around the ring. To the best of our knowledge, this 12-metal ring structure is without precedent in beryllium chemistry, or indeed in any other metal-carboxylate chemistry.

The overall network structure of 1 consists of an unprecedented 3,12 net, in which 12-connected [Be₁₂(OH)₁₂]¹²⁺ ring nodes are linked through 3-connected BTB³⁻ ligand nodes. The resulting framework contains large pear-shaped cavities that are ~ 27 Å along their long axis and 21 Å in diameter at the widest portion of their base. These pores can be accessed through three different types of openings:

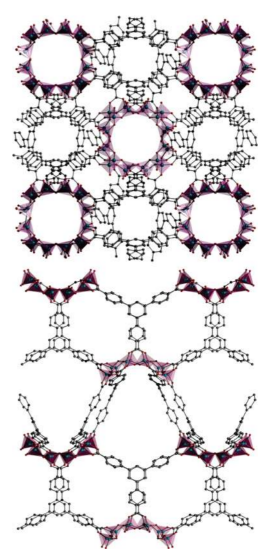


Figure 1. A portion of the crystal structure of $Be_{12}(OH)_{12}(BTB)_4$ (1), as viewed down the [001] (upper) and [010] (lower) directions. Turquoise, gray, and red spheres represent Be, C, and O atoms, respectively, while purple tetrahedra highlight the BeO₄ units; H atoms are omitted for clarity. Crystal data: $P\overline{4}c2$, a = 24.3013(9) Å, c = 54.570(3) Å. Selected interatomic distances (Å) and angles (deg): Be-O_{OH} 1.485(3)-1.662(2), Be-O_{BTB} 1.561(2)-1.721(2), Ве-О_{0Н}-Ве 114.7(1)-123.9(1), О_{ОН}-Ве-О_{ВТВ} 105.0(1)-117.1(1), Овтв-Ве-Овтв 101.6(1)-107.7(2), Ве-О-С 122.4(1)-129.8(1).

 $[Be_{12}(OH)_{12}]^{12+}$ rings with a diameter of ca. 6 Å (based on van der Waals radii), 7 Å wide hexagonal rings formed by two BTB3- ligands and two $[Be_2(OH)]^{3+}$ segments, and larger $14 \times 9 \text{ Å}^2$ arrowheadshaped rings involving three BTB³⁻ ligands, two Be²⁺ ions, and a

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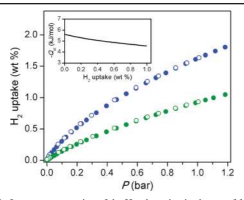


Figure 2. Low-pressure region of the H₂ adsorption isotherms of 1 recorded at 77 K (blue) and 87 K (green), with filled and open symbols representing adsorption and desorption data, respectively. Inset: Isosteric heat of H₂ adsorption for 1 calculated using the adsorption data.

[Be₂(OH)]³⁺ segment. The solvent-accessible volume calculated from the crystal structure using the PLATON routine⁸ is 77%, highlighting the extremely porous nature of the framework.

Low-pressure N2 adsorption measurements performed on 1 at 77 K afforded a type I isotherm characteristic of a microporous solid. Fits to the data gave a BET surface area of 4030 m²/g and a Langmuir surface area of 4400 m²/g. These values lie slightly above the accessible surface area of 3600 m²/g calculated from the crystal structure,⁹ indicating complete evacuation of the pores. Although the BET surface area of 1 exceeds the 3800 m²/g observed for MOF-5, it is still somewhat below that obtained for Zn₄O(BTB)₂ (MOF-177; 4750 m²/ g)^{1e} and at least two other frameworks with larger pore openings.^{1i,10} Nevertheless, it represents the highest value yet reported for a main group metal-organic framework or covalent organic framework.

The low-pressure H₂ adsorption isotherms for 1 at 77 and 87 K are shown in Figure 2. The total H₂ uptake of 1.6 wt % at 77 K and 1 bar is completely reversible, and the gradual rise of the isotherm indicates a small adsorption enthalpy. Indeed, a fit to the data yields an initial isosteric heat of adsorption of -5.5 kJ/mol, which gradually increases with loading. Hence, the $[Be_{12}(OH)_{12}]^{12+}$ rings do not serve as strong binding sites for H₂, resulting in a flat adsorption profile. Note that this behavior is actually favorable for a cryogenic storage material, since operation of the fuel cell at pressures above 1.5 bar means that H₂ retained at lower pressures would not ordinarily be utilized.

Consistent with its high surface area, compound 1 shows excellent performance as a hydrogen storage material at higher pressures (see Figure 3). At 77 K, the excess H₂ uptake reaches a maximum of 6.0 wt % at 20 bar, while the total uptake climbs to 9.2 wt % and 43 g/L at 100 bar. Although these capacities are among the highest recorded to date for a metal-organic framework,11 the volumetric storage density is still considerably below the 66 g/L achieved in MOF-5.3 This is due to the larger pores of 1, wherein a significant amount of H_2 gas can reside far from the influence of the framework walls. Interestingly, while the H₂ storage capacity is dramatically reduced at 298 K, reaching 2.3 wt % and 11 g/L at 95 bar, the combined values are arguably better than those of any other metal-organic framework. To our knowledge, only $Mn_3[(Mn_4Cl)_3(BTT)_8(MeOH)_{10}]_2$ (BTT³⁻ = 1,3,5benzenetristetrazolate), which possesses open metal coordination sites, has a higher volumetric capacity of 12 g/L at 90 bar and 298 K, although its gravimetric uptake is just 1.5 wt %.11 Here, the relatively good performance of 1 is likely a consequence of its structure, which contains many rings near the 7 Å diameter calculated as ideal for achieving a high storage density at 298 K.12

The foregoing results demonstrate reaction conditions under which the first pure, crystalline beryllium-based metal-organic framework could be isolated. It is likely that similar conditions may now supply

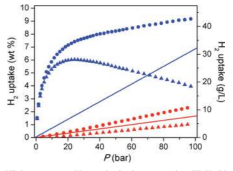


Figure 3. Higher-pressure H_2 uptake in 1 measured at 77 K (blue) and 298 K (red). Triangles and circles represent excess and total H2 uptake, respectively, while the solid lines show the density of pure H₂.

a range of new beryllium-based frameworks exhibiting exceptionally high surface areas. In particular, the use of 1,4-benzenedicarboxylate as a linker can be expected to afford frameworks with pore sizes near the 10 Å diameter sought for high-density hydrogen storage at cryogenic temperatures.¹² In addition, efforts are underway to functionalize the surfaces within 1 via substitution of the bridging hydroxide ligand protons.

Acknowledgment. This research was funded by DoE Grant DE-FG36-05GO15002. We thank Fulbright New Zealand for support of K.S.; the Australian Research Council Network for Advanced Materials and the CSIRO Chief Executive Science Leader Scheme for funding of M.R.H.; and Dr. A. DiPasquale, Dr. F. Hollander, Prof. S. Batten, and Prof. T. Düren for helpful advice.

Supporting Information Available: Full experimental details (PDF); an X-ray crystallographic file (CIF). This material is available free of charge via the Internet at http://pubs.acs.org.

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JA9072707