

Hydrogen Storage in a Microporous Metal–Organic Framework with Exposed Mn²⁺ Coordination Sites

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Abstract: Use of the tritopic bridging ligand 1,3,5-benzenetristetrazolate (BTT³⁻) enables formation of [Mn-(DMF)₆]₃[(Mn₄Cl)₃(BTT)₈(H₂O)₁₂]₂·42DMF·11H₂O·20CH₃OH, featuring a porous metal-organic framework with a previously unknown cubic topology. Crystals of the compound remain intact upon desolvation and show a total H₂ uptake of 6.9 wt % at 77 K and 90 bar, which at 60 g H₂/L provides a storage density 85% of that of liquid hydrogen. The material exhibits a maximum isosteric heat of adsorption of 10.1 kJ/mol, the highest yet observed for a metal-organic framework. Neutron powder diffraction data demonstrate that this is directly related to H_2 binding at coordinatively unsaturated Mn^{2+} centers within the framework.

Introduction

The utilization of hydrogen in automobiles as a clean-burning substitute for fossil fuels relies in part upon the development of a viable on-board storage system.^{1,2} Recently, a new class of ordered, three-dimensional extended solids composed of metal ions and organic linkers, known as metal-organic frameworks, has emerged as a promising storage alternative to high-pressure and liquefied hydrogen tanks, metal hydrides, and carbon-based adsorbents.3-25 Due to their porous nature and unusually high

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surface areas, these new materials show exceptional H₂ uptake by mass but are also characterized by very weak H₂ adsorption energies (typically 4-7 kJ/mol), such that cryogenic temperatures are required to observe significant H₂ uptake. Clearly, frameworks exhibiting stronger binding interactions are needed to facilitate H₂ adsorption at higher temperatures; indeed, a binding energy of 15 kJ/mol has been predicted to maximize the amount of adsorbed H₂ accessible at 298 K within the pressure range 1.5-20 bar.²⁶

Two main strategies have been pursued for enhancing H₂ binding within metal-organic frameworks. First, by forming frameworks with very narrow pores, wherein overlapping potentials from two or more pore walls interact with a single H₂ molecule, H₂ binding energies of up to 9.5 kJ/mol have been observed.15 More promisingly, it has been proposed that the

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presence of coordinatively unsaturated metal centers could increase the affinity for H₂ through strong metal—H₂ interactions.^{14,27,28} Our strategy has involved the use of polytetrazolate bridging ligands with an N donor atom arrangement deemed favorable for creating rigid frameworks with exposed metal coordination sites.¹⁹ The present work demonstrates the success of this approach, with use of 1,3,5-benzenetristetrazolate (BTT^{3–}) in producing a stable, microporous material exhibiting a high surface area and exceptional H₂ uptake. Moreover, neutron powder diffraction is employed to confirm that the high H₂ adsorption enthalpy observed for this material is due to H₂ binding directly to unsaturated metal centers within the pores of the framework.

Experimental Section

All reagents were obtained from commercial vendors and, unless otherwise noted, were used without further purification. Methanol was distilled over Mg/I_2 prior to use.

1,3,5-Tris(2H-tetrazol-5-yl)benzene Hydrochloride (H3BTT· 2HCl). A mixture of 1,3,5-tricyanobenzene²⁹ (3.06 g, 20.0 mmol), NaN₃ (11.7 g, 180 mmol), and triethylamine hydrochloride (24.7 g, 180 mmol) in 150 mL of toluene and 30 mL of methanol was heated at reflux in a 500-mL round-bottom flask for 3 days. Upon cooling to room temperature, 100 mL of an aqueous solution of NaOH (1 M) was added, and the mixture was stirred for 30 min. The aqueous layer was treated with ca. 100 mL of diluted HCl (1 M) until no further white precipitate formed. The precipitate was then collected by filtration, dried in the air, and dissolved in aqueous NaOH (1 M). The resulting clear, colorless solution was titrated with ca. 75 mL of diluted HCl (1 M) until the pH of the solution was 4-5. The ensuing white precipitate was washed with successive aliquots of distilled water (3 \times 50 mL), methanol (2 \times 50 mL), and acetone (50 mL) to afford 7.1 g (87%) of product. Anal. Calcd for C₉H₆N₁₂•2HCl•H₂O•CH₃OH: C, 29.64; H, 3.48; N, 41.48. Found: C, 29.84; H, 3.49; N, 41.84. IR (neat): 3347 (s, br), 3066 (m), 2329 (m, br), 1889 (m, br), 1636 (m, br), 1430 (s), 1409 (s), 1377 (s), 1321 (w), 1204 (m), 1176 (m), 1116 (m), 1053 (s), 896 (s), 746 (s) cm⁻¹. ¹H NMR (dmso- d_6): δ 8.79 ppm (s, 3H). This compound is soluble in DMF and dimethylsulfoxide but is insoluble in nonpolar organic solvents; it can be recrystallized from hot methanol or hot water.

Caution! Metal azides are water sensitive and potentially explosive and should be handled with care. Although we did not encounter any incidents while handling H₃BTT or compounds **1**, **1'**, **1m**, and **1m'** under the conditions of our experiments, previous reports suggest that handling H₃BTT at temperatures above 160 °C is potentially unsafe and that melting with decomposition (explosion) occurs at 280 °C.²⁹

 $[Mn(DMF)_6]_3[(Mn_4Cl)_3(BTT)_8(H_2O)_{12}]_2 \cdot 42DMF \cdot 11H_2O \cdot 20CH_3OH$ (1). A solution of MnCl₂ · 4H₂O (200 mg, 1.0 mmol) in 5 mL of methanol was added to a solution of H₃BTT · 2HCl · H₂O · CH₃-OH (45 mg, 0.11 mmol) in 5 mL of DMF in a Teflon-capped 20-mL scintillation vial. The pH of the solution was then adjusted to ca. 3 using an aqueous solution of HCl (1 M), and the resulting mixture was heated to 70 °C. Colorless cube-shaped crystals suitable for X-ray analysis were collected from the walls of the vial after 24 h. The crystals were collected by filtration, washed with successive 20-mL aliquots of DMF and methanol, and then quickly dried on a piece of filter paper to afford 65 mg (40%) of product. Anal. Calcd for C₃₄₄H₆₁₈Cl₆-Mn₂₇N₂₅₂O₁₁₅: C, 34.95; H, 5.27; N, 29.86. Found: C, 34.81; H, 4.97; N, 30.15. IR (neat): 1651 (s), 1414 (m), 1387 (w), 1101 (w), 790 (m), 749 (w), 675 (w) cm⁻¹. The solvent content of this compound was

confirmed by thermogravimetric analysis (TGA), and phase purity was confirmed by powder X-ray diffraction. Crystals of **1** become opaque after prolonged exposure to the atmosphere and were therefore stored under dinitrogen. The partially desolvated form of this compound, [Mn-(DMF)₆]₃[(Mn₄Cl)₃(BTT)₈(DMF)₁₂]₂ (**1**'), was generated by heating **1** at 150 °C under dynamic vacuum for a minimum of 2 h.

 $[Mn(CH_3OH)_6]_3[(Mn_4Cl)_3(BTT)_8(CH_3OH)_{12}]_2 \cdot 42CH_3OH$ (1m). Crystals of 1 were immersed in freshly distilled methanol and soaked for 1 day inside an N2-filled glovebag. The solvent was exchanged three times, such that the total soaking time was 4 days. The methanolexchanged crystals were then collected by filtration inside the glovebag. Anal. Calcd for C₂₂₈H₃₈₄Cl₆Mn₂₇N₁₉₂O₈₄: C, 30.93; H, 4.37; N, 30.37. Found: C, 30.91; H, 4.14; N, 30.67. IR (neat): 1618 (w, br), 1417 (s), $1230 (w), 1187 (w), 1016 (s), 901 (w), 789 (s), 751 (m), 698 (w) cm^{-1}$. Crystals of 1m become opaque and turn light yellow-brown in color upon prolonged exposure to the atmosphere and were therefore stored in sealed glass ampules. The desolvated form of the compound, $Mn_3[(Mn_4Cl)_3(BTT)_8(CH_3OH)_{10}]_2$ (1m'), was generated by heating 1m at 150 °C under dynamic vacuum for a minimum of 2 h. Anal. Calcd for C₁₆₄H₁₂₈Cl₆Mn₂₇N₁₉₂O₂₀ (1m'): C, 28.95; H, 1.90; N, 39.52. Found: C, 28.46; H, 1.66; N, 39.13. IR (neat): 1587 (w, br), 1417 (s), 1236 (w), 1189 (w), 1015 (w), 790 (s), 751 (m), 692 (w) cm⁻¹.

Low-Pressure Gas Adsorption Measurements. Gas adsorption isotherms for pressures in the range 0-1.2 bar were measured using a Micromeritics ASAP2020 instrument. Samples of 1 and 1m were transferred under a dinitrogen atmosphere to preweighed analysis tubes, which were then capped with a Transeal to prevent intrusion of oxygen and atmospheric moisture during transfers and weighing. The samples were evacuated by heating at 150 °C under dynamic vacuum until the outgas rate was less than 2 mTorr/min (0.27 Pa/min). The evacuated analysis tubes containing degassed samples were then carefully transferred to an electronic balance and weighed to determine the mass of sample (typically 50-200 mg). The tube was transferred back to the analysis port of the gas adsorption instrument. The outgas rate was again confirmed to be less than 2 mTorr/min (0.27 Pa/min). For all isotherms, warm and cold free space correction measurements were performed using ultra-high-purity He gas (UHP grade 5.0, 99.999% purity); H₂ and N₂ isotherms at 77 K were measured in liquid nitrogen baths using UHP-grade gas sources. H₂ isotherms at 87 K were measured in liquid argon baths. Oil-free vacuum pumps and oil-free pressure regulators were used for all measurements to prevent contamination of the samples during the evacuation process or of the feed gases during the isotherm measurement.

Derivation of the Isosteric Heats of Adsorption. A virial-type expression of the following form was used to fit the combined isotherm data for a given material at 77 and 87 K:

$$\ln P = \ln N + 1/T \sum_{i=0}^{m} a_i N^i + \sum_{i=0}^{n} b_i N^i$$
(1)

Here, *P* is the pressure expressed in Torr, *N* is the amount adsorbed in mmol/g, *T* is the temperature in K, a_i and b_i are virial coefficients, and *m* and *n* represent the number of coefficients required to adequately describe the isotherms. The equation was fit using the **R** statistical software package;³⁰ *m* and *n* were gradually increased until the contribution of extra added *a* and *b* coefficients was deemed to be statistically insignificant toward the overall fit, as determined using the *t*-test. The values of the virial coefficients a_0 through a_m were then used to calculate the isosteric heat of adsorption using the following expression:

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⁽³⁰⁾ Download, instructions, and further details on the use and capabilities of this software package area available online at http://www.r-project.org.

$$Q_{\rm st} = -R \sum_{i=0}^{m} a_i N^i \tag{2}$$

Here, Q_{st} is the coverage-dependent isosteric heat of adsorption and *R* is the universal gas constant.

High-Pressure Gas Adsorption Measurements and Analysis. Crystals of 1 and 1m were loaded into sample holders under an argon atmosphere. The samples were evacuated under a pressure of less than 10^{-4} Torr (0.013 Pa) in two stages: first heating at 40-50 °C for 5 h, and then at 150 °C for at least 10 h. Hydrogen excess adsorption analyses were performed using an automated controlled Sieverts' apparatus (PCT-Pro 2000 from Hy-Energy LLC) over a pressure range of 0-50 bar for 1' and 0-90 bar for 1m'. At least 300 mg of adsorbent was used in each experiment, and UHP-grade hydrogen and helium (99.999% purity) were used for all measurements. Volumetric measurements at 77 K were carried out by submerging the sample holder in a liquid nitrogen bath, for which the fill level was maintained constant throughout the experiment.

The volume of the sample holder and the connecting gas manifold was previously determined using H_2 and He at 298 K and at 77 K. Hydrogen was used to determine the dead-space volume correction for a nonporous sample of known volume; this correction accounted for the change in effective sample volume observed when cooling the sample holder from room temperature to 77 K. The adsorbent was then introduced into the sample holder, and He gas was used to determine the volume of the sample at room temperature. Since He gas penetrates the pores of the sample without being adsorbed onto the surface, the volume measured with He corresponds to the volume of the framework walls, also referred to as the framework skeleton. Consequently, the skeletal density of the material, d_{sk} , can be obtained from the following expression:

$$d_{\rm sk} = m/V_{\rm sk} \tag{3}$$

Here, *m* is the sample mass expressed in g, and $V_{\rm sk}$ is the sample volume in cm³, as determined using He expansion at room temperature. In order to obtain an accurate assessment of the value of the skeletal density, the volume of the sample, $V_{\rm sk}$, was measured 20 times, and the average of these was used as the final value.

Excess adsorption is defined as the amount of gas taken up by the surface of a porous adsorbent above and beyond the quantity of gas that would have occupied the adsorbent pore volume under the same temperature and pressure conditions in the absence of an adsorbent. The total storage capacity, however, is the sum of the capacity due to adsorption on the surface and the capacity due to compression within the void space of the pores of the adsorbent. The total adsorption capacity can therefore be expressed as follows:

$$C_{\rm tot} = C_{\rm exc} + \frac{100 \times d_{\rm g} V_{\rm pore}}{1 + d_{\rm g} V_{\rm pore}} \tag{4}$$

Here, C_{tot} is the total adsorption capacity expressed in weight percent (wt %), C_{exc} is the excess adsorption in wt % and is the quantity being measured, d_{g} is the density of the compressed gas (here H₂) at a given temperature and pressure in g/cm³, and V_{pore} is the pore volume in cm³/g.

In eq 4, the second term of the sum on the right-hand side represents the contribution of the compressed H₂ inside the pores to the total adsorption. Note that the denominator must contain the term d_gV_{pore} for the capacity to be expressed in units of wt %. Omitting this term would correspond to units of grams of H₂ adsorbed per 100 g of adsorbent, and not to wt % of H₂ adsorbed. The compressed gas density, d_g , is the density of H₂ gas as a function of pressure at a given temperature. In the current experiments, the software package GasPak (v. 3.41) was used to plot the density of H₂ at 77 K as a function of pressure, as shown in Figure S8 (Supporting Information). If the bulk density of the sample, d_{bulk} , is known (typically, d_{bulk} is the crystallographic density of the sample), the pore volume can then be calculated as follows:

$$V_{\text{pore}} = \frac{d_{\text{sk}} - d_{\text{bulk}}}{d_{\text{sk}} d_{\text{bulk}}}$$
(5)

The volumetric density of H_2 adsorbed inside the sample can be obtained simply by multiplying the adsorbed quantity with the bulk density of the sample, d_{bulk} :

$$C_{\rm vol} = Q_{\rm ads} d_{\rm bulk} \tag{6}$$

Here, $C_{\rm vol}$ is the volumetric H₂ capacity expressed in g/L, and $Q_{\rm ads}$ is the total quantity of H₂ taken up in mmol/g. The excess quantity of H₂ adsorbed may instead be used in this equation if "excess" volumetric capacity is the desired quantity.

X-ray Structure Determinations. Crystals of 1 and 1m' were coated in Paratone-N oil, attached to Kapton loops, transferred to a Siemens SMART APEX diffractometer, and cooled in a dinitrogen stream. Lattice parameters were initially determined from a least-squares analysis of more than 100 centered reflections; these parameters were later refined against all data. None of the crystals showed significant decay during data collection. The raw intensity data were converted (including corrections for background, Lorentz, and polarization effects) to structure factor amplitudes and their esd's using the SAINT 4.15 program. An empirical absorption correction was applied to each data set using SADABS. Space group assignment was based on systematic absences, E-statistics, and successful refinement of the structures. Structures were solved by direct methods with the aid of difference Fourier maps and were refined against all data using the SHELXTL 5.0 software package. Hydrogen atoms were inserted at idealized positions and refined using a riding model with an isotropic thermal parameter 1.2 times that of the attached carbon atom. For 1, thermal parameters for all non-hydrogen atoms in the framework skeleton and for the N and O atoms from bound DMF were refined anisotropically. The C atoms in each DMF molecule were disordered over four equivalent sites around the crystallographic 4-fold rotation axis and were consequently refined with 0.25 of their natural occupancies. All the interatomic distances in each DMF molecule were constrained to predefined values [O-Ccarbonyl, 1.22(1) Å; Ccarbonyl-N, 1.31(2) Å; N-C_{methyl}, 1.46(2) Å; C_{methyl}-C_{methyl}, 2.47(4) Å; C_{carbonyl}-C_{methyl}, 2.41-(3) Å] and refined as such. The van der Waals contacts between the Cmethyl atoms of the DMF molecules and the Cl and N atoms in the framework lie between 3.8 and 4.0 Å. The coordination environment around the half-occupied Mn²⁺ was assigned to six DMF molecules using TGA and elemental microanalysis. Extensive disorder both locally, around the Mn²⁺ ion, and over neighboring unit cells prevents the refinement of any other atoms other than six O atoms disordered over 14 sites. Extra electron density in the pores was modeled as partially occupied oxygen atoms, with occupancies varying between 0.25 and 0.5 of the natural occupancy for a given set of crystal coordinates. For 1m', thermal parameters for all non-hydrogen atoms except for C and O in methanol molecules were refined anisotropically. Residual methanol molecules in the structure of 1m' were disordered, and the total occupancy, including all disordered atoms, was refined best at 83.3% of the site occupancy. Hence, there is 0.83 of a methanol molecule for each Mn²⁺ site in the framework skeleton.

Neutron Diffraction Experiments. Neutron powder diffraction data were collected on the High Resolution Neutron Powder Diffractometer BT-1 at the NIST Center for Neutron Research (NCNR) with a Ge-(311) monochromator and using in-pile collimation of 15 min of arc, corresponding to a wavelength of 2.0787 Å. Measurements were taken as a function of deuterium loading at a temperature of 3.5 K with measurement times of ca. 9 h.



Figure 1. Portions of the crystal structure of 1: (a) molecular structure of the tritopic ligand H_3BTT , (b) a square-planar Mn_4Cl cluster surrounded by eight tetrazolate rings, (c) a sodalite cage-like unit encasing a $[Mn(DMF)_6]^{2+}$ complex, and (d) a cube of eight such units sharing square Mn_4Cl faces. Hydrogen atoms and solvent molecules are omitted for clarity. Selected interatomic distances (Å) and angles (°): Mn-Cl 2.736(1), Mn-N 2.227(3), Mn···Mn 3.869-(2), Mn-Cl-Mn 90.0, N-Mn-N 87.6(1), 91.4(1), Mn-N-N 125.3(2).

All sample transfers were performed in a helium-filled glovebox equipped with water and oxygen monitors. Initial sample activation was performed in a glass tube with a packless bellows valve attached. The sample was evacuated using a turbo pump (10^{-5} Torr) and heated to 150 °C with a ramp speed of ca. 1 °C per minute. After degassing at 150 °C for 24 h, the sample was cooled, pulverized using an agate mortar and pestle, transferred to a cylindrical vanadium can (i.d. = 0.95 cm) equipped with a capillary gas line and a packless valve, and sealed with a lead O-ring. The sample was mounted onto a sample stick equipped with a stainless-steel gas line with an additional valve for a top-loading closed-cycle helium refrigerator. The sample was further degassed in situ for ca. 20 min under high vacuum (8.5 $\times 10^{-7}$ to 7.5×10^{-6} Torr) at 150 °C. During the experiments, a known amount of hydrogen (deuterium) gas was loaded into the sample (907 mg), which was maintained at a temperature of 77 K until no pressure drop was observed for at least 1 min. The sample was then cooled down to the base temperature of 3.5 K over a period of 1 h in order to perform measurements. In all cases, the outgas pressure reading was zero well before reaching 25 K.

A diffraction pattern was collected for 1m' prior to dosing with D₂. Subsequent Rietveld analysis indicated the existence of extra neutron density close to the Mn²⁺ ions in the framework skeleton, as expected due to the presence of residual methanol molecules. Accordingly, the extra neutron density was modeled using disordered O and C atoms, and this corrected "bare" model was used as a baseline for all subsequent measurements involving D₂. This extra scattering source proved to be absolutely necessary in order to achieve a low goodnessof-fit value for the bare material. The presence of methanol was confirmed by single-crystal X-ray diffraction³³ (see Figure S1), infrared spectroscopy (Figure S4), and elemental analysis. For X-ray diffraction, a single crystal was used, and, as such, modeling the disordered methanol molecules could be accomplished more easily. In contrast, the neutron powder diffraction experiments involved large amounts of powdered crystalline samples, and the methanol disorder in one crystallite may have differed from that in other crystallites, making it difficult to model the disorder over the entire sample. Consequently, partially occupied C and O atoms were introduced to account for the extra scattering densities in the bare material. In addition, owing to the extended disorder of the extraframework Mn²⁺ ions and to the small scattering factor of manganese, these ions could not be located using neutron diffraction; as such, their contribution to the fitted models was omitted.

Neutron scattering diffraction patterns were analyzed using the Rietveld refinement method.³¹ The program EXPGUI was used to perform all refinements.³² The model of the bare material was refined first, and it was used as the starting point for subsequent refinements of the D₂-loaded samples. Deuterium molecules are treated as point scatters with double occupancy since they are expected to be quantum mechanically spherically averaged in the ground state. As mentioned, during the refinement of the model for the bare material, extra C and O atoms were introduced to account for the scattering intensities from the residual methanol molecules bound to the intraframework Mn²⁺ ions. The information on the extra atoms was obtained from the

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diffraction pattern of the bare material and was fixed at these values when analyzing the cases with deuterium molecules loaded. The coordinates of all other atoms and the Debye–Waller factors were allowed to vary during the refinement of each deuterium loading case. Based on the structure obtained from the diffraction pattern of the bare material, the diffraction pattern of the case with 12 D₂/f.u. was analyzed by neglecting the D₂ molecules. The Fourier difference map was calculated and clearly indicated the positions of D₂ adsorption sites. Accurate values for the D₂ locations and occupancy numbers were then obtained by Rietveld refinement. For each successive D₂ loading, the Fourier difference map was calculated on the basis of the results of the previous D₂ loading and used to identify new D₂ adsorption sites.

Other Physical Measurements. Infrared spectra were collected on a Nicolet Avatar 360 FTIR spectrometer with an attenuated total reflectance accessory. ¹H NMR spectra were obtained using a Bruker AVQ-400 instrument. Carbon, hydrogen, and nitrogen analyses were obtained from the Microanalytical Laboratory of the University of California, Berkeley. Thermogravimetric analyses were carried out at a ramp rate of 1 °C/min in a nitrogen flow with a TA Instruments TGA 2950. X-ray powder diffraction data were collected using Cu Kα ($\lambda = 1.5406$ Å) radiation on a Siemens D5000 diffractometer. Prior to collecting X-ray powder diffraction data, the as-synthesized sample was ground using an agate mortar and pestle.

Results and Discussion

Reaction of H₃BTT with MnCl₂·4H₂O in an acidic solution of N,N-dimethylformamide (DMF) and methanol at 70 °C afforded cube-shaped crystals of [Mn(DMF)₆]₃[(Mn₄Cl)₃(BTT)₈-(H₂O)₁₂]₂·42DMF·11H₂O·20CH₃OH (1) in 40% yield. X-ray analysis³³ of a crystal of **1** revealed a cubic structure in which chloride-centered square-planar [Mn₄Cl]⁷⁺ units are linked via BTT³⁻ ligands to form the anionic, three-dimensional framework shown in Figure 1. The N2 and N3 atoms of two tetrazolate rings on individual BTT³⁻ ligands bridge pairs of Mn²⁺ ions along the edges of a square, such that each $[Mn_4Cl]^{7+}$ moiety is surrounded by eight BTT³⁻ ligands. In turn, each BTT³⁻ ligand connects three [Mn₄Cl]⁷⁺ units so that, overall, six squares and eight planar BTT³⁻ ligands define the faces of a truncated octahedron, reminiscent of a sodalite cage. Each cage shares its square faces with six neighboring cages, defining a network structure closely related to that of sodalite.³⁴ Alternatively, the square-planar [Mn₄Cl]⁷⁺ units surrounded by eight tetrazolates can be considered as tetragonally distorted 8-connected nodes that, together with the trigonal planar nodes presented by the BTT³⁻ ligands, define a three-dimensional 3,8connected net (see Figure 2). Notably, this is the first example of a compound displaying such a "Moravia" net, the existence of which was only recently predicted by theory.35 Within the anionic framework skeleton, formulated as $[(Mn_4Cl)_3(BTT)_8(H_2O)_{12}]^{3-}$, water ligands occupy the sixth coordination site on each Mn²⁺ ion, while charge balance is provided by [Mn(DMF)6]2+ complexes situated inside both the sodalite cage-like units and the larger three-dimensional cavities formed by eight such units. Note that a single $[Mn(DMF)_6]^{2+}$ complex fits snugly within a cage, suggesting that it may in fact act as a template for the in situ formation of the framework.



Figure 2. 3,8-Connected three-dimensional net derived from the structure of **1**. Gray 3-connected nodes represent BTT^{3-} ligands, while green 8-connected nodes represent square-planar Mn₄Cl clusters surrounded by eight tetrazolate rings, as depicted in Figure 1a and b, respectively.

Thermogravimetric analysis of **1** showed a weight loss of 22.0% up to 160 °C, suggesting that methanol, water, and some DMF molecules are evacuated at this temperature (21.9% expected for 20 methanol, 35 water, and 18 DMF molecules). Framework decomposition, observed above 200 °C, precluded evacuation of the remaining 42 DMF molecules, which were expected to occupy all 18 coordination sites on the extraframework cations and all 24 sites on the intraframework cations, thus prohibiting access to unsaturated Mn²⁺ centers and reducing the accessible surface area. To attain a more complete evacuation, DMF was exchanged with methanol by soaking crystals of 1 in methanol for 4 days. While other organic solvents can be exchanged inside the framework, methanol was chosen for its relatively low boiling point and ease of potential removal via evacuation. As indicated by infrared spectroscopy and elemental analysis, the resulting material is water- and DMFfree, with formula [Mn(CH₃OH)₆]₃[(Mn₄Cl)₃BTT₈(CH₃OH)₁₂]₂. 42CH₃OH (1m). Thermogravimetric analysis of 1m showed a weight loss step below 100 °C, corresponding to methanol loss, followed by a plateau between 100 and 200 °C, beyond which framework decomposition occurred.

Remarkably, crystals of **1m** remained single upon heating to 150 °C, enabling a crystal structure determination for the desolvated compound.³³ Here, X-ray analysis revealed an intact framework skeleton, with some residual methanol (20 molecules per formula unit). The charge-balancing, extraframework Mn^{2+} ions, initially complexed by DMF in **1**, are now ligated by the N1 and N4 atoms of adjacent tetrazole rings and are disordered over 24 symmetry-equivalent positions (see Figure S1). The residual methanol molecules coordinate 83% of the framework Mn^{2+} ions, leaving a small fraction of these metal centers with open coordination sites. Despite numerous attempts, evacuation conditions for removing all of the remaining methanol without degrading the framework have not yet been discovered.

The partially desolvated forms of **1** and **1m**, denoted **1'** and **1m'**, adsorb significant amounts of N₂: 310 and 550 cm³/g, respectively, with both displaying type I adsorption isotherms

⁽³³⁾ Crystal and structure refinement parameters for 1: T = 166 K, space group Pm3m, Z = 2, a = 19.116(1) Å, V = 6985.2(7) Å³, R₁ = 0.0771, wR₂ = 0.2508. For 1m': T = 161 K, Pm3m, Z = 2, a = 19.009(1) Å, V = 6868.3-(7) Å³, R₁ = 0.0486, wR₂ = 0.1417.

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Figure 3. Isotherms for the adsorption of N_2 within 1' (red) and 1m' (blue) at 77 K.

typical of a microporous material (see Figure 3).³⁶ Fitting the BET equation to the N₂ isotherms gave estimated surface areas of 1100 and 2100 m²/g for 1' and 1m', respectively. For comparison, crystalline zeolites and activated carbons can have surface areas of up to 904 and 2800 m²/g, respectively.^{37,38} The highest surface area yet reported for a microporous metalorganic framework is 5900 m²/g,³⁹ although most values fall below 1000 m²/g.⁴⁰ Note that replacement of the DMF molecules with methanol, followed by evacuation at 150 °C, results in almost a doubling of the available surface area, suggesting that H₂ adsorption should also be enhanced in **1m'** relative to 1'.

The hydrogen storage properties of 1' and 1m' were evaluated using volumetric gas adsorption measurements. The resulting isotherms, depicted in Figure 4a, show fully reversible uptakes of 1.7 wt % in 1' and 2.2 wt % in 1m' at 77 K and 1.2 bar. The excess H₂ adsorption, defined as the amount of H₂ adsorbed in excess of the bulk gas that would occupy the pores of the adsorbent,⁴¹ reaches 2.8 wt % in 1' and 5.1 wt % in 1m' at saturation, as shown in Figure 4b. A more informative quantity for practical applications, however, is the total H₂ uptake, which is defined simply as the total amount stored in the bulk volume of the adsorbent, as determined using the crystallographic density of the material (see Table S2). Thus, within a fuel tank maintained at 77 K, the materials are expected to show total uptakes of 3.9 wt % for 1' at 50 bar and 6.9 wt % for 1m' at 90 bar. Note that a higher gravimetric capacity has recently been reported for $Zn_4O(1,3,5$ -benzenetribenzoate)₂, which exhibits an excess adsorption of 7.0 wt % and is expected to exhibit an even higher total adsorption at 60 bar and 77 K.18

Together with the weight requirements, H₂ storage in mobile applications imposes limitations with respect to the tank volume. Volumetric capacity thus becomes a critical parameter when comparing different storage materials. Significantly, the excess volumetric adsorption in 1m' reaches 43 g/L at 90 bar and 77 K, 9 g/L higher than that observed within $Zn_4O(1,3,5-benzen$ etribenzoate)₂,¹⁸ while the total volumetric uptake reaches 60

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Figure 4. H_2 adsorption isotherms (a) below 1.2 bar and (b) up to 90 bar within 1' (red) and 1m' (blue) at 77 K, and within 1m' at 298 K (green). Triangles and circles represent excess and total H₂ adsorption, respectively, while filled and open symbols represent adsorption and desorption data, respectively.



Figure 5. Isosteric heat of adsorption curves for H_2 uptake in 1' (red) and 1m' (blue).

g/L. The latter value is only 11 g/L lower than the density of liquid hydrogen (71 g/L at 1 bar and 20 K), suggesting the possibility of converting existing liquid hydrogen-based storage systems for use with a liquid nitrogen coolant. We attribute this high volumetric storage density to the relatively compact nature of the metal-organic framework and, as discussed below, a reduction in the volume taken up by the strongly adsorbed H_2 molecules.

To probe how the presence of open Mn²⁺ coordination sites influences the binding enthalpy, H₂ adsorption isotherms in the range 0-1.2 bar were also measured at 87 K. The 77 and 87 K



Figure 6. Rietveld refinement of neutron powder diffraction data for a loading of 12 D₂ molecules per formula unit of **1m'**. Green lines, crosses, and red lines represent the background, calculated, and experimental diffraction patterns, respectively. The blue line shows the difference between experimental and calculated patterns. The final Rietveld fit parameter was $\chi^2 = 0.985$.



Figure 7. Initial D_2 adsorption sites within 1m'. Light blue spheres represent D_2 centroids, while the transparent orange sphere shows the position of a partially occupied, extraframework Mn^{2+} ion site. Hydrogen atoms and methanol molecules are omitted for clarity.

isotherms were then fit using a virial equation⁴² of the type previously used to model adsorption in microporous materials with heterogeneous pore surfaces.^{21,43,44} Virial coefficients obtained from the fits were then employed in calculating the isosteric heats of adsorption for the two evacuated materials. The ensuing enthalpies lie in the range 5.7–7.6 kJ/mol for 1', depending on degree of H₂ loading, while for **1m'** the upper limit increases to 10.1 kJ/mol (see Figure 5). This represents the highest value reported for a metal–organic framework.^{15,21} Notably, the heat of adsorption curves for **1'** and **1m'** show a large gap at low H₂ coverages and then overlay at coverages above ca. 0.8 wt %. Thus, strong H₂ adsorption sites become available in **1m'** that are not present in DMF-passivated **1'**.

Neutron powder diffraction experiments⁴⁵ were performed as a direct check of whether the high binding enthalpy in **1m'** is associated with coordination of H₂ at exposed Mn²⁺ sites. Figure 6 shows the diffraction pattern observed at 3.5 K upon adsorption of 12 D₂ molecules per formula unit. Using a Rietveld profile analysis, the two strongest binding sites at this low loading could be associated with sites I and II in Figure 7. Significantly, site I is located just 2.27 Å from the Mn²⁺ ions in the framework skeleton, providing the first neutron diffraction evidence for a metal-H₂ interaction within a metal-organic framework.⁴⁶ To our knowledge, this is the first example of H₂ binding to a Mn^{2+} ion, although many examples of H_2 binding to Mn⁺ complexes and other transition metal species exist.⁴⁷⁻⁵⁰ The D₂ molecules at site II are situated inside the sodalite-like cages and exhibit van der Waals contacts of 3.47 and 3.66 Å with a chloride anion and four equidistant tetrazolate rings, respectively. The sum of the five corresponding attractive interactions is likely responsible for a significant adsorption enthalpy at this site as well. Indeed, when the total D2 loading is increased to 48 molecules per formula unit, site II becomes fully occupied.

Increased D₂ loadings also permitted identification of the next strongest binding site, labeled III, and a possible binding site labeled IV in Figure 7. The D2 molecules at site III are situated inside the larger framework cavities and have van der Waals contacts of 3.26 Å with two equidistant tetrazole rings. At site IV, only 5 D_2 molecules could be identified; these are located inside the smaller cavities, and are in closest contact (4.72 Å) with four carbon atoms from neighboring tetrazolate rings. Out of 48 D₂ molecules refined from a model derived from the difference scattering length map, 7.0 reside at site I, 10.7 at site II, 23.9 at site III, and 5.0 at site IV. Importantly, the H_2 uptake corresponding to complete filling of sites I and II is 0.6 wt %, coinciding with the region where heat of adsorption curves for 1' and 1m' separate, as shown in Figure 5. This demonstrates that the two strongest H₂ binding sites are indeed likely responsible for the increase in binding energy upon going from

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1' to 1m'. Experiments intended to locate weaker adsorption sites and to determine the rotational potential dynamics at the H_2 adsorption sites (using inelastic neutron scattering) are underway.

Outlook

A key and challenging aspect of hydrogen storage research is the ability to control the H₂ binding energy, which governs the adsorption in materials ranging from microporous solids to metal hydrides. In this regard, a very attractive property of metal-organic frameworks is that their pores can be engineered at the atomic scale, thereby enabling some control over the H₂ binding interaction. The foregoing results show that, on a materials characteristic basis, metal-organic frameworks with unsaturated metal centers can exceed the 2010 DOE H₂ storage targets of 6.0 wt % and 45 g/L.² Meeting these targets for a complete storage system, however, will require further efforts toward raising the H₂ binding energy, such that storage at closer to ambient conditions can be accomplished. Ultimately, this may be possible in materials related to those reported here through adjustment of the electronic structure of the metal ions and exposure of a higher concentration of coordination sites.

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Supporting Information Available: Full table of X-ray refinement statistics, tables of refinement statistics and atomic parameters from neutron diffraction, tables of gas adsorption data, further views of the crystal structure of **1m**', thermogravimetric analysis plots for **1** and **1m**', additional plots of isotherm data and plots of the virial fits including the values of virial coefficients a_i and b_i , powder X-ray diffraction patterns for **1**, a Fourier difference neutron density map for the case of $12 \text{ D}_2/\text{ f.u., and the structure of$ **1m**' as determined using neutron powder diffraction (PDF); X-ray crystallographic files (CIF). This material is available free of charge via the Internet at http://pubs.acs.org.

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