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A Metal–Organic Framework with Entatic Metal Centers Exhibiting High Gas Adsorption Affinity

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The lack of an effective, economic, and safe on-board vehicular gas (hydrogen or methane) storage method is one of the major technical barriers preventing fuel-cell driven automobiles to compete with traditional ones.¹ Recently, the gas-storage properties of metal—organic frameworks (MOFs) have been actively explored.² In the area of hydrogen storage, suggested strategies to increase hydrogen uptake include using pore sizes comparable to gas molecules³ and introducing coordinatively unsaturated metal centers.⁴ Both strategies boil down to the enhancement of gas affinity of the material. Recent inelastic neutron scattering studies suggest that high-affinity H₂ binding sites are metal-based.⁵ These recent findings prompted us to search for new ways to increase the gas affinity of metal centers in MOFs.

In metalloproteins, such as hemoglobin, reversible O_2 binding is achieved by utilizing an iron active center (Figure 1a), where a porphyrin ligand and a histidine residue force the iron center into a square pyramidal geometry, generating an open coordination site for gas binding. Similarly, in Vitamin B₁₂ (VB₁₂), the central cobalt is surrounded by a corrin and a proximal ligand, and the distal position is open for substrate binding.⁶ Herein we report a MOF with a novel secondary building unit (SBU, Figure 1b), containing four hemoglobin/VB₁₂-like cobalt centers bundled through sharing a μ_4 -oxo bridge, producing four metal centers at an entatic state (vide infra) for gas binding. This MOF exhibits exceptional gas adsorption affinity for a number of gases.

In bioinorganic chemistry, an entatic state is a state in which an unusual geometry is *imposed* by the protein polypeptides on a metal center whose reactivity in electron transfer, substrate binding, or catalysis is enhanced.⁷ Similarly, due to the specific geometric requirements of the ligands and SBUs in a MOF, the metal centers can be forced into an entatic state to enhance their affinity toward gases.

Entatic metal centers (EMCs) represent a biomimetic approach to coordinatively unsaturated metal centers (UMCs).⁸ The latter are frequently obtained by removal of one or more ligands from a metal center to achieve coordinative unsaturation; the ligand removal required to achieve UMCs may collapse the entire network.⁹ Conversely, most of the EMCs are ready for substrate binding without the need for ligand removal.

Herein we report a MOF with EMCs, $H_2[Co_4O(TATB)_{8/3}]$, designated PCN-9 (Porous Coordination Network) for convenience. TATB (4,4',4"-*s*-triazine-2,4,6-triyltribenzoate) is a ligand developed in this laboratory for its planar conformation, potential ability to bind additional metal atoms, and high thermal stability of resulting MOFs.^{4b,10} PCN-9 crystals were grown under solvothermal conditions by a reaction between H₃TATB and cobalt nitrate in DMSO at 135 °C.

X-ray single-crystal analysis¹¹ revealed that PCN-9 crystallizes in the $Im\bar{3}m$ space group. It adopts a square-planar Co₄(μ_4 -O) SBU, with a μ_4 -oxo residing at the center of a square of four Co atoms. All four Co atoms in the SBU are five-coordinate with square-



Figure 1. (a) Schematic drawing of the active center of hemoglobin. The gold sphere represents an Fe atom. (b) The $Co_4(\mu_4-O)(carboxylate)_4$ SBU found in PCN-9. Color scheme: C, gray; Co, aqua; and O, red.



Figure 2. (a) Crystal structure of PCN-9 showing an octahedral cage; the red sphere represents void inside the cage. (b) View of the crystal structure of PCN-9 along the *a*-axis; the two interpenetrated networks are shown in gold and pink, respectively. All atoms are shown in arbitrary scales.

pyramidal geometry. The Co $-\mu_4$ -O distance is 2.351(2) Å. If one of the five-coordinate Co atoms is compared to the active center in hemoglobin, the μ_4 -O is analogous to the proximal ligand, and on the opposite side of the square-pyramidal base is the distal position of the Co, which is below the plane of the four O atoms in an entatic state, and ready to bind a substrate to achieve octahedral coordination. Although a tetrahedral μ_4 -oxo bridge is common,¹² a square-planar μ_4 -oxo bridge is rare. Only a few examples have been found in iron,13a copper,13b vanadium,13c and niobium^{13d} compounds. The square-planar μ_4 -oxo bridge found in PCN-9 is unique in a MOF. Every $Co_4(\mu_4-O)$ SBU connects eight trigonal-planar TATB ligands, and every TATB ligand connects three $Co_4(\mu_4-O)$ SBUs to form a (8,3)-net. Alternatively, this (8,3)net can also be described as a three-dimensional network formed by corner sharing of octahedral cages (Figure 2a); the O_h -cage is defined by six $Co_4(\mu_4-O)$ SBUs at the corners and eight TATB ligands on the faces. Overall, the structure of PCN-9 can be obtained by the interpenetration of two (8,3)-nets, the second being generated through translation along [1 1 0].

The volume of the octahedral cage is 2740 Å³. Each open window of the cage is 6.7×6.7 Å (atom to atom distance) and possesses two EMCs with distal sites pointing toward the void. The solvent accessible volume of PCN-9 calculated using PLA-TON¹⁴ is 58.8%. The permanent porosity of PCN-9 is confirmed by its N₂ adsorption isotherm. After desolvation, PCN-9 exhibits a Langmuir surface area of 1355 m²/g and a pore volume of 0.51 cm³/g.

To determine the gas affinity of PCN-9, adsorption isotherms were collected at various temperatures, and the data were fit using the Langmuir–Freundlich equation.¹⁵ Isosteric adsorption enthalpies as a function of the quantity of gas adsorbed were calculated using a variant of the Clausius–Clapeyron equation.^{16,18a,b}

At low coverage, PCN-9 exhibits an O_2 adsorption enthalpy of 17.8 kJ/mol, comparable to that of Co exchanged zeolites (15.1–18.5 kJ/mol).¹⁷ Similarly, at low coverage, the CO adsorption enthalpy of PCN-9 reaches 21.0 kJ/mol. These results represent the first measurement of O_2 or CO adsorption enthalpy in MOFs.

 H_2 adsorption isotherm at 77 K indicates that PCN-9 has an uptake of 1.53 wt % at 760 Torr. Hydrogen adsorption isotherms at two other temperatures were collected and used to calculate the heat of adsorption. At low coverage, the heat of adsorption of PCN-9 for H_2 is 10.1 kJ/mol, which is higher than that of MOF-5 (5.2 kJ/mol), Prussian blue (7.4 kJ/mol), MOF-74 (8.3 kJ/mol), or HKUST-1 (6.6 kJ/mol) and is comparable to that of IRMOF-11 (9.1 kJ/mol) or a magnesium MOF (9.5 kJ/mol).¹⁸

In VB₁₂, the active center can bind a methyl or an alkyl group.⁶ It is expected that PCN-9, with its SBUs structurally similar to the active center of VB₁₂, should have a high methane affinity. The CH₄ isotherms of PCN-9 at three temperatures were collected, corresponding to an adsorption enthalpy of 23.3 kJ/mol at low coverage. To the best of our knowledge, this is by far the highest CH₄ adsorption enthalpy found among MOFs (IRMOF-6 = 12.1 kJ/mol, MIL-53 = 17 kJ/mol).¹⁹

To further confirm that this remarkable enhancement of gas affinity is due to the EMCs, freshly prepared PCN-9 crystals (or a guest-free sample) were soaked in a dilute tetrabutylammonium cyanide solution in THF (1 mg/mL) for 2 days until the color of the crystals changed from violet to dark red. An IR spectra using this dark-red polycrystalline sample showed a sharp peak at 2105 cm⁻¹, a +50 cm⁻¹ shift of the C=N stretch relative to that of free cyanide, consistent with the literature value for terminal cyanides bound to Co atoms.²⁰

Aware that the bulkiness of the counterion may prevent cyanide ions from permeating into the channels, we decided to use CO as an IR probe. A guest-free PCN-9 sample was kept under a CO atmosphere (3 bar) overnight. IR spectra of the sample showed a peak at 2044 cm⁻¹ (compared to 2143 cm⁻¹ for free CO),²¹ indicating terminal CO binding to the EMCs. The same sample was then kept under dynamic vacuum for 0.5 h to remove CO guests. The 2044 cm⁻¹ feature disappeared, demonstrating the reversibility of EMC binding, consistent with CO adsorption measurements.

In summary, EMCs have been created based on the geometric requirements of a MOF. This approach is analogous to the formation of protein EMCs, which are enforced by surrounding polypeptides. The EMC-containing MOF possesses exceptionally high affinity to a number of commercially relevant gases. In particular, the H_2 and CH_4 adsorption enthalpies of PCN-9 are among the highest reported thus far. The biomimetic strategy presented here will be generally applicable in the search of new adsorptive materials to meet the requirements of on-board vehicular gas storage.

Additionally, the EMCs inside the open channels of PCN-9 make them ideal candidates for size- and/or shape-selective catalysis. This line of research is currently underway in our laboratory. Acknowledgment. This work was supported by the National Science Foundation (CHE-0449634) and Miami University. H.C.Z. acknowledges the Research Corporation for a Research Innovation Award and a Cottrell Scholar Award.

Supporting Information Available: Detailed experimental procedures, X-ray structural data, thermogravimetric analysis, and gas adsorption data. This material is available free of charge via the Internet at http://pubs.acs.org.

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