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A supramolecular  $Co(II)_{14}$ -metal-organic cube in a hydrogen-bonded network and a Co(II)-organic framework with a flexible methoxy substituent

In situ imidazolate-4,5-diamide-2-olate (L3) linker generation leads to the formation of a  $[{\rm Co}_{14}({\rm L3})_{12}({\rm O})({\rm OH})_2({\rm DMF})_4]$  metal organic cube (MOC), wherein a  ${\rm Co}_6$  octahedron inscribed in a  ${\rm Co}_8$  cube. The MOCs connect by amide–amide H-bonds in a 8-c body-centered cubic (bcu) net, forming a 3D supramolecular network. Moreover, a new imidazolate-4-amide-5-imidate (L2) based honeycomb-like MOF, IFP-8, is also generated, having flexible methoxy groups, which act as molecular gates for guest molecules. This allows selective  ${\rm CO}_2$  sorption over  ${\rm N}_2$  and  ${\rm CH}_4$  gases.





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## A supramolecular Co(||)<sub>14</sub>-metal-organic cube in a hydrogen-bonded network and a Co(||)-organic framework with a flexible methoxy substituent†

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The reaction of 4,5-dicyano-2-methoxyimidazole (L1) with  $Co(NO_3)_2$ ·  $6H_2O$  under solvothermal conditions in DMF, a MOF, IFP-8 and a hydrogen-bonded network consisting of tetradecanuclear  $Co(II)_{14}$ -metal organic cube (1) are achieved. 1 shows the bcu net with 14 cobalt atoms.

In the design of new solid-state materials, supramolecular chemistry is of great interest because it takes advantage of self-assembly to synthesize new materials by virtue of cooperative interactions such as ion–ion interactions, hydrogen bonding, dipole–dipole interactions and aromatic  $\pi$ – $\pi$  interactions.  $^{1-6}$  Moreover, the flexible or soft porous networks, also known as the third generation of porous coordination polymers, receive much attention because of their interesting properties.  $^{7,8}$  Flexible metal–organic frameworks (MOFs) are extremely interesting for applications in selective gas adsorption/ separation or chemical sensing.  $^8$  Such frameworks exhibit guest dependent structural transformation and breathing effects.  $^9$ 

However, our group succeeded in using a particular approach to obtain the zinc based isoreticular IFP series (IFP = imidazolate framework Potsdam) by replacing the substituent of the R-group of a 2-substituted imidazolate 4-amide-5-imidate linker (–R = CH<sub>3</sub>,  $C_2H_5$ , Cl, Br). To extend the number of frameworks at the isoreticular series of IFP, recently, we reported a Zn-based material, known as IFP-7 that showed the gate-effects due to the flexible methoxy substituent of the linker. In addition to IFP-7, an *in situ* imidazolate-4,5-diamide-2-olate (L3) linker based Zn<sub>14</sub>-molecular building block (MBB) is also formed. These MBBs or specifically metal–organic cubes (MOCs) contain the peripheral H-bonding substituents that construct an effective way to obtain H-bonded supramolecular networks with channels and pores. Hence, to

Fig. 1 (a) Tetradecanuclear cobalt MOC of 1; (b) hydrogen-bonded supramolecular assembly of 1, view along a axis; (c) space filling model of 1; (d) hexagonal channels in IFP-8, the methoxy substituent at the linker L2 is presented in a space filling mode. The structure is based on density functional ab initio calculations (pink Co, blue N, red O, dark gray C, light gray H).

achieve such a kind of rigid and directional single-metal-ion based MBB based supramolecular assemblies, the solvothermal reactions with such linker precursor and other metal salts can be explored. Herein, we report a cube-like  $\text{Co}(\pi)_{14}\text{-MOC}$  are engaged into amideamide H-bonds (denoted as 1), forming a supramolecular network (Fig. 1b), which is similar to  $\text{Zn}_{14}\text{-MBB}$  and an imidazolate-4-amide5-imidate based MOF (named as IFP-8, Fig. 1d) shows flexibility due to the methoxy substituent.

A H-bonded MOCs based supramolecular network (1) is formed *via in situ* hydrolysis of the ligand precursor 4,5-dicyano-2-methoxyimidazole (L1). Under solvothermal conditions in N,N'-dimethylformamide (DMF), partial hydrolysis of the cyano

a) b) c) d)

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Scheme 1 Syntheses of  $\bf 1$  and IFP-8. In situ imidazolate-4,5-diamide-2-olate (L3) linker, synthesis with indication of its cobalt coordination (Fig. S3, ESI $\dagger$ ) and H-bonds in  $\bf 1$ . See ESI $\dagger$  for experimental details.

groups to amide groups and of the methoxy to the hydroxy group followed by twofold deprotonation generates the L3 linker (Scheme 1 and Scheme S1, in the ESI†). The linker L3 is only stable in the deprotonated and metal-coordinated state because the free ligand H<sub>2</sub>L3 irreversibly transforms into a stable tautomeric keto-form. Under these reaction conditions, in addition to 1, due to another in situ functionalization 2-methoxyimidazolate-4-amide-5-imidate (L2) linker based material, IFP-8, is formed as the main product (Scheme 1 and ESI†). IFP-8 was separated using the sieving technique, 14 wherein 1 was trapped by a mesh while IFP-8 filtered through it. After several attempts, we could not find a suitable crystal of IFP-8 for single X-ray diffraction. Hence, the structure was determined by a combination of Powder X-ray diffraction (PXRD), structure modelling and IR spectroscopy (Fig. 1d). The structural model of IFP-8 was constructed by using the single-crystal X-ray structure determination for IFP-1,10a and was further optimized by using a density functional theory ab initio method (see ESI†). The PXRD pattern of the optimised IFP-8 structure shows very good agreement with the experimental data (Fig. S10, ESI†).

Compound 1 was characterized by single-crystal X-ray diffraction as  $[Co_{14}(L3)_{12}(O)(OH)_2(DMF)_4] \cdot (DMF)_{16}$  for the asymmetric unit. <sup>15</sup> The degree of in situ hydrolysis of the cyano groups of L1 into the corresponding L3 linker was studied using infrared (IR) spectroscopy (see ESI†). Compound 1 crystallizes in the high-symmetry space group I4/m of the tetragonal crystal system. Twelve L3 ligands, one oxide ion, two hydroxide ions and four DMF molecules assemble with fourteen cobalt ions to form a tetradecanuclear Co(II)<sub>14</sub>-MOC with peripheral amide groups (Fig. 1a). However, the H-bonded Zn<sub>14</sub>-MBB network possesses the space group Ia3d (No. 230), having highest crystallographic symmetry and contains four water molecules, instead of four DMF. 13 The oxide ion (O1) is located in the centre of the MOC, surrounded by four Co1 and two Co2 atoms in an exactly octahedral coordination environment (Fig. 1a and Fig. 4Sb, ESI†). Each of these four Co1 atoms is further coordinated by four olate oxygen atoms of four imidazolate ligands (two O2 and two O3), one O2- (O1) and one DMF (O4) forming a distorted octahedral coordination geometry (Fig. S5, ESI†). The two Co2 centres are surrounded by four olate oxygen atoms (O2), one O2- (O1) and one DMF (O5) to form a distorted octahedral coordination geometry (Fig. S5, ESI†). Eight Co3 atoms are each coordinated by three nitrogen atoms (N1, N2 and N5) from two imidazolate ligands, two amide oxygen atoms (O6 and O8) and an olate ion (O2) as the linker in a twofold facecapped tetrahedron (Fig. S5, see ESI†). The metal ion at the IFP-8 structure is penta-coordinated by the L2 linkers to form a distorted

trigonal-bipyramidal geometry (Fig. S7, ESI $\dagger$ ). The structure possesses 1D hexagonal channels running along 0, 0, z; 1/3, 2/3, z and 2/3, 1/3, z. The methoxy groups protrude into the open channels and determine their accessible diameter (Fig. 1d). By considering the van der Waals radii, a probable channel diameter of the channels was estimated to be 1.9 Å (see ESI $\dagger$  for details).

The MOC of 1 contains coordinated and free amide groups at its vertices and edges. These are involved in intramolecular hydrogen bonds of L3 and intermolecular hydrogen bonds between the MOCs or protrude into the channels of 1, the intramolecular hydrogen bonds (N3-H3B···O7 = 2.68 Å) additionally stabilizing the MOC. Each MOC is connected through its vertices with eight MOCs by intermolecular N-H···O hydrogen bonds (N(4)-H(4)A···O(7) =2.90 Å) between the peripheral amide groups, generating the 3D-supramolecular assembly of 1 (Fig. 1b). Hence, L3 acts as a bridging linker because of its unique potential to offer the amide groups, N-donor sites of imidazole and olate ion when chelated to a metal ion and additionally, some amides of the MOC were employed in hydrogen-bonding with other MOCs, which are necessary for supramolecular assemblies. However, the framework exhibits two types of infinite channels. The first type of channel running along the crystallographic c axis has small openings with an approximate diameter of 1.7 Å (Fig. S6, ESI†), while the second type of accessible channel running along the a axis can accommodate a sphere with a maximum diameter of 3.2 Å given the van der Waals radii of the nearest atoms (Fig. 1c). Hence, guest molecules such as DMF can be hydrogen-bonded by potential donors and acceptors with the amide functionality.

The topology of 1 can be described as a hydrogen-bonded 8-c body-centered cubic (**bcu**) net with nodes  $Co(\pi)_{14}$ -MOCs (Fig. S9, ESI†). Moreover, as the  $Co(\pi)_{14}$ -MOC could be inscribed in a cube, one can also describe the net as the augmented version of **bcu** (= **bcu-a**), which is called polycubane (**pcb**). In contrast,  $Co^{2+}$  ions at **IFP-8** and bridging ligands L2 act as 3-connected topological species forming a net with a rare uninodal topology, named *etb*. The topology of IFP-8 is classified by the vertex symbol 3.6.10.15. In

The PXRD pattern of activated **IFP-8** exhibited the sharp diffraction peaks similar to that of the as-synthesized sample. This indicates that the porous framework maintains the crystalline integrity even without solvent molecules (Fig. S10, ESI†).

The activated IFP-8 is expected to show the gas-sorption selectivity towards small polar molecules due to its polar and flexible methoxy side chains. The gas sorption isotherms are recorded for N2, H2, CH4, and CO2 gases at various temperatures at 1 bar (Fig. 2). Moreover, we are unable to examine the gas uptake capacities as well as other physical properties for compound 1 in the present study due to low yield. We are in the process of optimizing the synthetic conditions for obtaining a better yield. As indicated in Fig. 2a, IFP-8 barely adsorbed N2, which can be attributed to a gating of the pores by the pendant methoxy groups, while the CO2, CH4 and H<sub>2</sub> sorption isotherms show very different sorption behaviors. 9,11 The CO<sub>2</sub> sorption measurements at 195, 273 and 298 K show typical type I isotherms and a small hysteresis is visible in all the desorption branches, indicating that the framework structure contains the flexible substituent. Such hysteretic behavior of IFP-8 is similar to zinc based imidazolate-4-amide-5-imidate (L2) frameworks ChemComm

Fig. 2 Gas sorption isotherms for activated IFP-8. Adsorption and desorption branches are indicated by closed and open symbols, respectively.

IFP-7 and other MOFs, having flexible alkoxy substituents. 9,11 The uptake of CO<sub>2</sub> by IFP-8 at 298 K and 1 bar is 27 cm<sup>3</sup> g<sup>-1</sup> and the uptake of CO<sub>2</sub> by IFP-7 is 40 cm<sup>3</sup> g<sup>-1</sup>. However, 2-methylimidazolate-4-amide-5-imidate linker based isostructural frameworks Zn-based IFP-1<sup>10a</sup> and Co-based IFP-5 were reported. 12 The channel diameter of IFP-5 (3.8 Å) is slightly lower than that of IFP-1 (4.2 Å); hence, the gas uptake capacities and BET surface area are slightly lower than IFP-1. Similarly, the channel diameter of Co-based IFP-8 is slightly narrower than that of the Zn-based IFP-7. Hence, it can be inferred that the gas uptake capacities for IFP-8 are much less than IFP-7. IFP structures contain the 1D hexagonal channel and the gas uptake capacities depend on the size of the channel. In contrast, the CO2 uptake capacities of Co-based ZIFs (Co-ZIF-68, -69, -81) are higher than their isostructural Zn based analogues (Zn-ZIF-68, -69 and -81).<sup>16</sup> In such frameworks, the interaction with CO<sub>2</sub> presumably increases inside the Co based ZIFs compared to their Zn based analogues due to the decreased ionic radius [Zn+2 (0.68 Å) and Co+2 (0.67 Å)] and the low density of Co-ZIFs. 16 The isosteric heats of adsorption were calculated from the CO2 adsorption isotherms at 273 K and 298 K (Fig. 2a). At zero loading the  $Q_{\rm st}$  value ( $-\Delta H$ ) for **IFP-8** is 37 kJ mol<sup>-1</sup> (Fig. S12, ESI†), comparable to other MOFs.<sup>8b</sup> Upon increasing the loading the  $Q_{\rm st}$  value decreases to 24 kJ mol<sup>-1</sup>. The high  $Q_{\rm st}$  value can be attributed to the highly polar framework and the effect of the small pore size effect.

Remarkably, the desorption branches for CH<sub>4</sub> isotherms show a wide desorption hysteresis which is very similar to IFP-7. IFP-8 adsorbs 11 cm $^3$  g $^{-1}$  CH $_4$  at 195 K and 1 bar (Fig. 2b). The CH $_4$ desorption isotherm at 195 K confirms that 85% of the adsorbed CH<sub>4</sub> is trapped in the framework when the pressure is reduced from 760 mmHg to 75 mmHg, and 71% of the adsorbed CH<sub>4</sub> remains when the pressure is further reduced to 25 mmHg. Such a broad desorption behaviour for the CH<sub>4</sub> isotherm at atmospheric pressure is rarely observed in microporous MOFs. 11,17 After CH<sub>4</sub> uptake the PXRD pattern of IFP-8 maintained its structural integrity. The structural transformation did not occur (Fig. S10, ESI†). Another agreeable example that has proven the flexibility of the framework is by H<sub>2</sub> sorption. **IFP-8** adsorbs 9.3 cm<sup>3</sup> g<sup>-1</sup> of H<sub>2</sub> at 77 K and 1 bar (Fig. 2a), showing a wide desorption hysteresis. Moreover, we determined the initial slopes in the Henry region of the adsorption isotherms of IFP-8 (Fig. S13, ESI†). The adsorption selectivities of IFP-8 are 34:1 and 9:1 for CO<sub>2</sub>/N<sub>2</sub> and CO<sub>2</sub>/CH<sub>4</sub>, respectively,

wherein the adsorption selectivities of IFP-7 are 37:1 and 7:1 for CO<sub>2</sub>/N<sub>2</sub> and CO<sub>2</sub>/CH<sub>4</sub>, respectively at 273 K and 1 bar. 11

In conclusion, not only could zinc form the H-bonded supramolecular assembly but cobalt also yielded a large Co(II)14-MOC via in situ functionalization of unusual linker generation under solvothermal conditions. The cobalt atoms in the Co(II)14-MOC formed the Co<sub>6</sub> octahedron inscribed distorted Co<sub>8</sub> cube (Co<sub>6</sub>@Co<sub>8</sub>). However, 1 shows the bcu net with 14 cobalt atoms and due to the flexible methoxy substituent, the hysteretic sorption behavior for IFP-8 indicates a flexible MOF. IFP-8 exhibits gas sorption selectivity.

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