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Double-walled pyr topology networks from a novel fluoride-bridged heptanuclear metal cluster†

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Two isostructural metal–organic materials, **Tripp-1-M** (**Tripp** = 2,4,6-tris(4-pyridyl)pyridine; M = Co, Ni), that exhibit binodal 3,6-connected **pyr** network topology have been prepared and characterized. **Tripp-1-M** are based upon a novel $M_7F_{12}^{2+}$ cluster that possesses 12 connection points but, because of double cross-linking by 3-connected **Tripp** ligands, it functions as a 6-connected supermolecular building block (SBB).

Introduction

Metal-organic materials (MOMs)1 have attracted rapidly increasing attention from the scientific community in the last two decades thanks to their inherent modularity. This modularity can promote diversity of composition, amenability to systematic design² and control over certain bulk properties.³ However, not all MOMs are well suited to serve as prototypal platforms for the generation of families of materials with the same topology. Such platforms are important because they enable systematic fine-tuning of both pore size (e.g. organic linkers with different lengths) and pore chemistry (e.g. functional group substitution at the linker or metal substitution at the node). Most platforms are built from single metal ion or small cluster (molecular building block, MBB) nodes and are exemplified by platforms sustained by carboxylate clusters such as 4-connected (4-c) M₂(RCOO)₄ (e.g. HKUST-1, PCN-6, MOF-2,6 DMOF-1 7 and NU-100 8), 6-c M₃O(RCOO)₆ (e.g. MIL-88,9 MIL-101 10 and PCN-600 11) and 6-c M₄O(RCOO)₆ (e.g. MOF-5 12 and MOF-177 ¹³). The exploitation of larger, high symmetry clusters offers the possibility of much higher levels of connectivity and even greater control over topology. Such "supermolecular building blocks", SBBs, are exemplified by 12-connected (12-c) $Zr_6O_4(OH)_4(RCOO)_{12}$ and 24-c "nanoball" $Cu_{24}(1,3-bdc)_{24}$ clusters.15 High connectivity mixed carboxylate/N-donor clusters have also been utilised in this context.¹⁶ In addition to affording greater control over topology because of fewer possible topological outcomes, higher connectivity nodes can result in greater robustness.17

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In this contribution, we introduce a new inorganic SBB of formula $M_7F_{12}^{2+}$ (M = Co, Ni, Fig. 1) and demonstrate that it can serve as a 6-c SBB through double cross-linking of its 12 connection points by a facile to prepare 3-c ligand, 2,4,6-tris-(4-pyridyl)pyridine, **Tripp** (Scheme 1). A new type of double-walled **pyr** topology (Fig. 2) material which exhibits permanent porosity is thereby generated.

In previous work, double^{18,19} or quadruple²⁰ cross-linking of carboxylate²¹ or oxide²² based SBBs has been shown to represent a suitable approach to build MOMs with well-known²³ or hitherto novel²⁴ topology. However, the use of fluoride-bridged MBBs and SBBs as nodes for the construction of three-dimensional MOMs remains rare.²⁵ This is despite the fact that discrete fluoride-bridged metal clusters are known^{26,27} and that such structures can exhibit interesting magnetic properties.²⁸

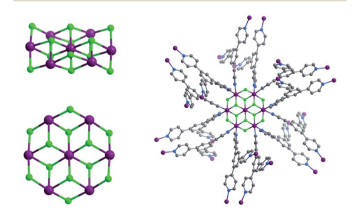


Fig. 1 (Left) Perspective and above views of the novel $M_7F_{12}^{-2+}$ cluster that sustains **Tripp-1-M**. (right) Illustration of the 12 connection points of the Co_7F_{12} SBB in **Tripp-1-Co** (Co, F, N and C atoms in purple, green, blue and grey). Solvent molecules, hydrogen atoms and counter-ions are omitted for the sake of clarity.

Scheme 1 Synthesis of 2,4,6-tris(4-pyridyl)pyridine (Tripp)

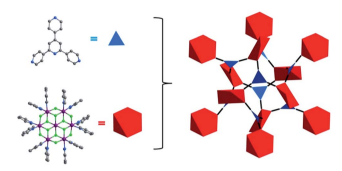


Fig. 2 Illustration of the topology of Tripp-1-M. The Tripp ligand is represented by a 3-c triangle (blue) whereas the ${\rm M_7F_{12}}^{2+}$ SBB is reduced to a 6-c octahedron (red).

Results and discussion

Tripp was prepared by the cyclization reaction of 4-acetylpyridine and 4-pyridinecarbaldehyde (Scheme 1).29 Single crystals of Tripp-1-Co were initially obtained by solvothermal reaction between $Co(NO_3)_2 \cdot 6H_2O$, Tripp and $(NH_4)_2SiF_6$ in DMF (for full details of synthetic procedures see ESI†). Tripp-1-Co crystallizes in the cubic space group $Pa\bar{3}$. A crystallographic 3-fold axis runs through the centre of the **Tripp** node and the disordered atoms of the central pyridine ring were therefore refined as 2/3 carbon and 1/3 nitrogen. All atoms of central pyridine ring are presented as carbon atoms for clarity in Fig. 1 and 3. In Tripp-1-Co, every Tripp ligand links three M₇F₁₂²⁺ SBBs and every SBB is connected by 12 Tripp ligands. However, the arrangement of the 12 connection points enables double cross-linking by pairs of **Tripp** ligands (Fig. 1) meaning that the connectivity is effectively reduced to 6. Therefore, each pair of Tripp ligands can be simplified to a single node, the M₇F₁₂ SBB can be treated as a 6-c node and the **Tripp** ligand as a 3-connected node. The outcome of this connectivity is structure that exhibits binodal 3,6-connected **pyr** topology (Fig. 2), of which there are relatively few examples^{22b,30} when compared to other types of 3,6 nets such as those with rtl, ant, sit or qom topology.

The structure of **Tripp-1-M** is comprised of two types of double-walled cages; tetrahedral cages (TCs) and octahedral cages (OCs) with face-sharing configurations (Fig. 3). The $M_7F_{12}^{2+}$ SBB is located at the vertex of every TC and OC, while the double-walls are constructed from **Tripp** ligands. The pore diameter of the TC and OC are *ca.* 7.7 and 7.6 Å, respectively. Every OC is surrounded by eight adjacent TCs and every TC is surrounded by four OCs. However, there are two faces and one

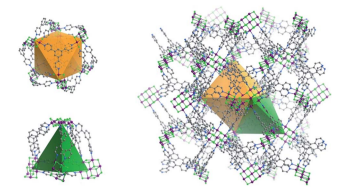


Fig. 3 The two types of double-walled cages (OC, left above; TC left, below) found in Tripp-1-M (right).

face capped by double walls of Tripp ligands in the OCs and TCs, respectively. Therefore, each OC connects with only six TCs and each TC crosslinks with three OCs, which corresponds to the required 3,6-connectivity of a pvr net. The guest-accessible porosity (considering the presence of counter ions) of Tripp-1-Co is 53.5%, based on Platon software.31 Two other structures with pyr topology using similar tripyridyl-based 3-c ligands, 2,4,6-tri(4-pyridyl)-1,3,5-triazine (TPT) and 1,3,5-benzene tricarboxylic acid tris[N-(4-pyridyl)amide (TPBTC), with 6-c metal ions Hg2+ and Cd2+, were reported by Robson and Kitagawa, respectively.32 These tripyridyl-based ligands can also be utilized as 3-c organic nodes to construct nets with high porosity.33 Further, some discrete cages consisting of 4-c Pd2+ ions linked by TPT and cage-based three-dimensional nets were reported by Fujita and co-workers.34 In contrast, the facile to prepare Tripp ligand has not been as widely studied as TPT and TPBTC.

The double-walled nature of **Tripp-1-M** is unusual and, to our knowledge, such as structure has not been prepared using a single ligand and a single SBB. However, Bu *et al.* recently reported two isostructural double-walled cage-based MOMs that were designed using a strategy based upon size-matching between two tritopic ligands (TPT and 2,4,6-tris[1-(3-carboxylphenoxy)-yl-methyl]mesitylene) connected by with same paddle-wheel unit.¹⁸

Analysis of crystal structure of **Tripp-1-Co** revealed that the Co-F distances in heptanuclear cluster lie the range from 2.036 (5) to 2.121 (7) Å, which are consistent with the values found in other fluoride-bridged Co(II) structures, e.g. [Co₅F₂- $(\text{tetrazolate})_4(H_2O)_4]^{35}$ and $[\text{Co}_{12}(\text{RCOO})_6(\text{PO}_4)_4F_4(H_2O)_6](\text{NO}_3)_2.^{36}$ The crystal structure of Tripp-1-Co also revealed that the charge of each heptanuclear cobalt cluster is balanced by one SiF₆² anion that exhibits three-fold F···F (distance: 2.68 (2) Å) interactions with three bridging F⁻ anions (Fig. S1†). Energy-dispersive X-ray spectroscopy verified the presence of Si and F in crystals of Tripp-1-Co (Fig. 4). There are previous reports concerning the generation of F by decomposition of PF6 and BF₄⁻, ^{26,37} so we speculated that the source of the bridging fluoride anions in the Co₇F₁₂²⁺ SBBs was in situ decomposition of SiF₆²⁻. Further, there are two drawbacks to the use of this synthetic method: the relatively low solubility of (NH₄)₂SiF₆ in

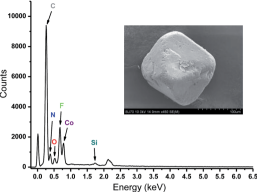
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DMF; the requirement to decompose SiF₆²⁻ anions before the

SBB can form. These drawbacks mean that unreacted (NH₄)₂SiF₆ is isolated in a physical mixture with Tripp-1-Co crystals, mitigating against phase purity and also resulting in low product yield (ca. 10%). Therefore, we tested a different synthetic approach involving reflux of starting materials in DMF/MeOH with NH₄F instead of (NH₄)₂SiF₆ as the F⁻ source. This method facilitated an increase in yield to 80%. The composition of Tripp-1-Co was changed since SiF₆²⁻ counterions are no longer present. Rather, two NO₃⁻ anions from the Co(NO₃)₂ starting material balance charge as indicated by the presence of two diagnostic peaks measured using FT-IR at around 1320 and 1400 cm⁻¹ (Fig. S4†).³⁸ The relatively high solubility of NH₄F in MeOH enabled subsequent isolation of pure reaction product. The purity of bulk product was established by powder X-ray diffraction (PXRD) patterns of as-synthesized samples, which are good matches to those calculated from the crystal structure of Tripp-1-Co (Fig. S5 \dagger). To further verify the composition of the $M_7F_{12}^{2+}$ cluster, X-ray photoelectron spectroscopy (XPS) analysis of a sample prepared from NH₄F was conducted, and a molar ratio for F: Co of 1.44 (expected 1.71) was observed (Fig. 4). A series of control experiments conducted without using NH₄F as a source of F- revealed that different concentrations of metal and ligand and different reaction temperatures failed to afford the desired Tripp-1-Co product. Nevertheless, these experiments confirm the essential role that F⁻ plays in construction of the M₇F₁₂²⁺ SBB and subsequently the overall MOM framework. The isostructural nickel analogue of Tripp-1-Co, Tripp-1-Ni, was obtained via the same modified synthetic protocol with a yield of ca. 70% as verified by PXRD (Fig. S6†).

To address the thermal stability of Tripp-1-M, thermo-gravimetric analysis (TGA) was conducted for both as-synthesized and MeOH exchanged samples. The resulting TGA plots reveal that solvent guest molecules in the as-synthesized samples can be fully exchanged with MeOH, which in turn can be removed below 110 °C. No further weight loss until after 300 °C was observed, which we presume is a consequence of framework decomposition (Fig. S8†). Framework integrity was also verified by PXRD experiments conducted after desolvation of MeOHexchanged samples at 120 °C. Furthermore, samples exposed to the air under ambient conditions for two months, were observed to exhibit PXRD patterns conforming to those calculated from single-crystal data (Fig. S5 and S6†). These results demonstrate that both Tripp-1-Co and Tripp-1-Ni possess good thermal and air/moisture stability, which we attribute to some extent to the high connectivity of the $M_7F_{12}^{2+}$ SBB.

The permanent porosities of **Tripp-1-Co** and **Tripp-1-Ni** were established by measuring CO_2 sorption isotherms at 195 K (Fig. 5). The apparent BET surface area was calculated to be 822 and 1149 m² g⁻¹ for **Tripp-1-Co** and **Tripp-1-Ni**, respectively. Pore volumes of 0.358 and 0.516 cm³ g⁻¹ for **Tripp-1-Co** and **Tripp-1-Ni** were calculated by assuming liquid filling of CO_2 at saturated state, which are close to the value of 0.587 cm³ g⁻¹ estimated from the crystal data for **Tripp-1-Co**. The relatively lower uptake of **Tripp-1-Co** might be attributed to partial collapse of the framework during activation, which does not appear to occur for **Tripp-1-Ni**. CO_2 and N_2 sorption isotherms



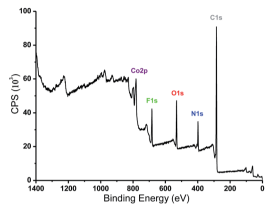


Fig. 4 Energy-dispersive X-ray spectroscopy (top) and X-ray photoelectron spectroscopy (bottom) of Tripp-1-Co.

of Tripp-1-M at 273, 283 and 293 K were also measured. As shown in Fig. 6, CO_2 uptakes of 76.7, 61.4 and 49.2 cm³ g⁻¹ at 273, 283 and 293 K, respectively, are much higher than the values of 3.7, 3.2 and 2.5 cm 3 g $^{-1}$ observed for N $_2$ in **Tripp-1-Co**. Tripp-1-Ni exhibits higher CO₂ uptakes of 99.3, 79.8 and 64.3 $\text{cm}^3 \text{ g}^{-1}$ at 273, 283 and 293 K, respectively, than those observed for Tripp-1-Co. Meanwhile, N2 uptakes at 273, 283 and 293 K are only 5.2, 4.3 and 3.5 cm³ g⁻¹ for Tripp-1-Ni. The high CO₂ uptakes of Tripp-1-M could be attributed to high pore volume and polarized pore surface originating from the heptanuclear cluster and central pyridine ring of the Tripp ligand. These results indicate that both Tripp-1-M variants exhibit good selectivity for CO2 over N2. Preliminary CO2/N2 selectivity for **Tripp-1-Co** and **Tripp-1-Ni**, calculated from the uptakes of CO₂ at 0.15 bar and N2 at 0.85 bar, are 41.4 and 40.8, 36.8 and 36.7, and 36.2 and 32.9 at 273, 283 and 293 K. These uptakes and selectivities are comparable to many well-known MOMs containing polar functional groups and/or open metal sites, both of which are absent in Tripp-1-M materials.39

To assess the strength of interaction between CO_2 and framework, the CO_2 isotherms measured at 273, 283 and 293 K were fitted using the virial equation (Fig. S9†), and the isosteric heat of adsorption ($Q_{\rm st}$) was calculated using the Clausius–Clapeyron equation. The enthalpies at zero loading for **Tripp-1-Co** and **Tripp-1-Ni** are 25.6 and 26.3 kJ mol⁻¹, respectively (Fig. S10†). These values are also consistent with those observed

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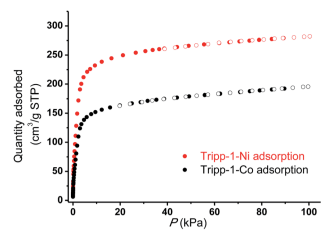


Fig. 5 CO_2 adsorption isotherms (filled symbols) and desorption (empty symbols) for Tripp-1-Co (black) and Tripp-1-Ni (red) conducted at 195 K.

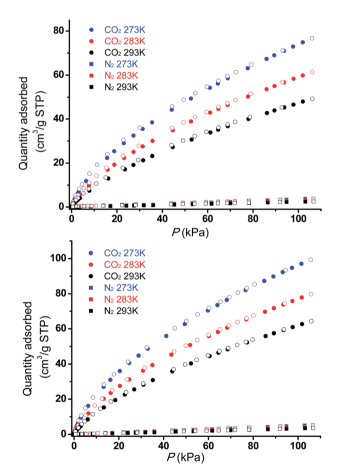


Fig. 6 CO_2 and N_2 adsorption (filled symbols) and desorption (empty symbols) isotherms for Tripp-1-Co (top) and Tripp-1-Ni (bottom) at three temperatures, 273 K, 283 K and 293 K.

in other classes of MOMs such as MOF-5 (34 kJ mol^{-1}),⁴⁰ HKUST-1 (35 kJ mol^{-1}),⁴¹ MAF-25 (26 kJ mol^{-1}),⁴² InOF-1 (29 kJ mol^{-1}),⁴³ NOTT-140 (25 kJ mol^{-1}).⁴⁴

Conclusions

In summary, we report a novel fluoride-bridged heptanuclear metal cluster-based SBB which has not been previously observed as a discrete structure. This cluster has 12-connection points, but 3-c **Tripp** ligands doubly cross-link to adjacent SBBs in order to form **Tripp-1-M**, two isostructural MOMs with binodal 3,6-connected **pyr** network topology. Good thermal and air/moisture stabilities were observed and gas sorption experiments demonstrate that both **Tripp-1-Co** and **Tripp-1-Ni** exhibit permanent porosity. The novel $M_7F_{12}^{2+}$ SBB reported herein has the potential to serve as an SBB for a wider range of MOMs with tailored pore sizes and surface chemistries. Follow-on studies on this platform will address properties related to gas sorption, catalysis and magnetism and are currently underway.

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