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Structure-directing factors when introducing hydrogen bond functionality to metal-organic frameworks†

Ross S. Forgan,^{*a} Ross J. Marshall,^a Mona Struckmann,^a Aurore B. Bleine,^a De-Liang Long,^a María C. Bernini^b and David Fairen-Jimenez^{*c}

The introduction of H-bond donor/acceptor functionality into metal-organic frameworks (MOFs) can have a beneficial effect on their molecular recognition, uptake selectivity and catalytic properties. The changes in ligand geometry induced by incorporation of functional groups may also affect the topology and composition of the resultant MOFs. Herein, we present a comprehensive study of functional group incorporation into MOFs, linked by either Zn^{2+} paddlewheel units or monomeric Zn^{2+} corners, which exhibit **pcu** and **dia** topology, respectively. Crystallographic analysis shows that amide groups can be easily incorporated into isoreticular **pcu** pillared-MOFs, whilst integration of urea units results in materials with **dia** topology. Molecular simulations allow the examination of hypothetical structures with differing constitutions and topologies, and highlight the influence of the urea units in generating the experimentally observed topologies. Noncovalent interactions between independent nets may be significant structure-directing influences, a finding which has great implications for the design of MOFs containing more complex functional groups.

1. Introduction

Metal-organic frameworks (MOFs),¹ crystalline network structures comprised of multtopic organic ligands, or struts, linked by metal ions or metal clusters, have found application in many areas as a result of their tunable functionality and porosity, including, but not limited to, gas storage,² molecular separations,³ catalysis,⁴ drug delivery⁵ and sensing.⁶ Enhancement of desired properties is often achieved through functionalization of the MOF, either by post-synthetic modification⁷ or by installing functional groups into the constituent ligands prior to MOF assembly. Pendant nucleophilic groups, such as $-NH_2$ and OH moieties, are known to enhance⁸ CO_2 capture through strong interactions with the quadrupolar moment of the gaseous substrate, and, in some cases, through formation⁹ of transient covalent bonds between the MOF and the guest. Amide moieties incorporated

into MOF struts give a convenient synthetic route to larger linkers, and are known¹⁰ to enhance CO_2 capture through favourable H-bonding and dipole-quadrupole interactions. A diverse range of amide-containing MOFs has been prepared¹¹ in the last few years to take advantage of this improved affinity for CO_2 . Incorporation¹² of urea units into MOFs generates heterogeneous organocatalytic species, which act as H-bond donor centres to the substrate(s) in question.

Incorporation of hydrogen bond functionality directly into MOF struts is clearly an attractive method for tuning and enhancing desirable properties, and an isoreticular series of frameworks containing differing functional groups would offer insight into the individual effect of each moiety on properties such as gas uptake and molecular recognition. Amide and urea units, however, bring the added complication of altering ligand geometry when compared to unsubstituted control ligands, with deviations of the disposition of the donor units from linearity that may influence the structure of the resultant MOFs.

In order to examine the effect of introducing hydrogen bonding units into MOFs on their topologies in a sequential manner, we have initially chosen the family of so-called **pillared**¹³ MOFs – structures comprised of two dimensional sheets of diacids linked by dimeric $Zn^{(II)}$ -paddlewheel secondary building units (SBUs) that are connected in the third dimension by nitrogen donor pillars in the **pcu** topology – as a system that allows the sequential incorporation of increasing numbers of functional groups. A series of

^a School of Chemistry, University of Glasgow, Joseph Black Building, University Avenue, Glasgow, G12 8QQ, UK.

E-mail: ross.forgan@glasgow.ac.uk; Web: <http://www.forganlab.com>

^b Facultad de Química, Bioquímica y Farmacia, Universidad Nacional de San Luis, Instituto de Tecnología Química-CONICET, San Luis, Argentina

^c Department of Chemical Engineering and Biotechnology, University of Cambridge, Pembroke St., Cambridge CB2 3RA, UK.

E-mail: df334@cam.ac.uk; Web: <http://people.ds.cam.ac.uk/df334>

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functionalised diacid ligands (**L1**·2H-**L3**·2H) and pillars (**P1**-**P3**) has been collated (Fig. 1a) and, using the known¹⁴ compound $[\text{Zn}_2(\text{L1})_2(\text{P1})]_n$ as a basic starting point (Fig. 1b), the structures of four new resultant MOFs are reported. Of these structures, two of them show a **pcu** topology, whereas the two other structures, obtained with a pillar (**P3**) that includes the urea unit, exhibit a **dia** topology. In addition to these four materials obtained experimentally, we have built a series of new hypothetical structures of varying compositions and topologies in order to analyse the geometries and interactions between ligands and pillars present in both **pcu** and **dia** topologies. Density functional theory calculations have been used previously to examine the energy landscapes of MOFs¹⁵ as well as their physical properties.¹⁶ However, since these calculations are computationally expensive, we followed¹⁷ a classical simulation approach that allowed us to easily obtain the new hypothetical constructs and examine their structures.

2. Results and discussion

In addition to the known structure $[\text{Zn}_2(\text{L1})_2(\text{P1})]_n$, four further MOFs were isolated¹⁸ after solvothermal treatment (see experimental section) of mixtures of $[\text{Zn}(\text{NO}_3)_2] \cdot 6\text{H}_2\text{O}$, a diacid ligand and a pillar in *N,N'*-dimethylformamide (DMF), and their structures elucidated by single crystal X-ray diffraction (Table 1). While the first two structures using the unfunctionalised and monoamide functionalised ligands and pillars showed the same Zn-paddlewheel units and **pcu** topology as the parent $[\text{Zn}_2(\text{L1})_2(\text{P1})]_n$ precursor, the two other structures, using the urea-functionalised **P3** pillar, showed a monomeric Zn^{2+} metal corner and a change into a **dia** topology.

The first compound, 1, $[\text{Zn}_2(\text{L1})_2(\text{P2})]_n$, forms through the combination of the unsubstituted ligand **L1** and the

monoamide functionalised pillar **P2** with Zn^{2+} , and exhibits the same **pcu** topology as the base structure, with paddlewheel SBUs connecting the diacids in two-dimensional sheets along the *ab* crystallographic plane and dipyridyl pillars linking the sheets along the *c* axis. In common with the parent, unfunctionalised species, the MOF shows two-fold interpenetration, but the introduction of the amide moiety into the pillar results in a change in orientation of the SBUs with respect to one another. The near rectangular arrangement of SBUs seen across the Zn^{2+} -**L1** sheets in the unfunctionalised analogue is distorted to a parallelogram arrangement in $[\text{Zn}_2(\text{L1})_2(\text{P2})]_n$, with inter-SBU angles of 75.8° and 104.2° . The offset introduced (Fig. 2a) by the amide unit in **P2** also results in distortions down the *c* axis, along which the pillars link the sheets, with inter-SBU angles of 74.6° , 79.7° , 100.3° and 105.4° deviating significantly from right angles. Each amide unit forms (Fig. 2b) a hydrogen bond to an ordered DMF solvent molecule, from the amido NH to the formamide CO, at a $\text{N}\cdots\text{O}$ distance of 2.79 \AA , with flanking $\text{CH}\cdots\text{O}$ interactions from the β -aromatic CH units of each ring to the formamide oxygen atom ($\text{C}\cdots\text{O} = 3.12\text{ \AA}$, $\text{C}\cdots\text{O} = 3.38\text{ \AA}$) supporting the molecular recognition of the substrate. The carbonyl oxygen of the amide moiety in **P2** is also involved in a $\text{CH}\cdots\text{O}$ interaction, with **P2** linkers from individual nets forming reciprocal dimers ($\text{C}\cdots\text{O} = 3.35\text{ \AA}$) that ensure the close proximity of the two interpenetrated nets of **pcu** topology (Fig. 2c).

Compound 2, $[\text{Zn}_2(\text{L2})_2(\text{P1})]_n$, has an amide functionality incorporated in its acid linker, **L2**, rather than in the pillar, which is the unfunctionalised **P1**. A structure analogous to the previous examples again forms; a 2-fold interpenetrated **pcu** material (Fig. 2c), with Zn^{2+} paddlewheel SBUs linking the ligands. In a manner similar to 1, the arrangement of SBUs in the Zn^{2+} -**L2** plane is distorted from the parent structure, with inter-SBU angles of 70.1° , 70.3° , 109.1° and 110.4° resulting from the offset nature of amido-substituted **L2**. The unsubstituted pillar **P1** links the sheet at angles close to 90° , however, in a manner similar to the parent structure (Fig. 2d). The sheets of acid ligands adopt an alternating arrangement (Fig. 2e), likely induced by a complementary pair of short $\text{CH}\cdots\text{O}$ contacts ($\text{C}\cdots\text{O} = 3.34\text{ \AA}$) between the amide oxygen of one molecule of **L2** and an aromatic CH of another from the second, interpenetrating net. Half of the **L2** units are involved in this interaction, while the other half form a bifurcated hydrogen bond to an ordered DMF guest, with the amido NH and one aromatic β -CH forming interactions with the formamide CO ($\text{N}\cdots\text{O} = 2.96\text{ \AA}$, $\text{C}\cdots\text{O} = 3.34\text{ \AA}$).

In contrast to the previous structures, compound 3, $[\text{Zn}(\text{L1})(\text{P3})]_n$, obtained through the introduction of the urea-functionalised pillar **P3** to the synthesis, results in an altogether different structure and topology. Instead of the dimeric paddlewheel SBU, a monomeric Zn^{2+} cation coordinates (Fig. 3a) two carboxylic groups of two unsubstituted **L1** molecules and two N donors of two molecules of **P3**, forming a distorted octahedral coordination sphere that yields a distorted tetrahedral 4-connected node, with inter-SBU angles of 95.1° ,

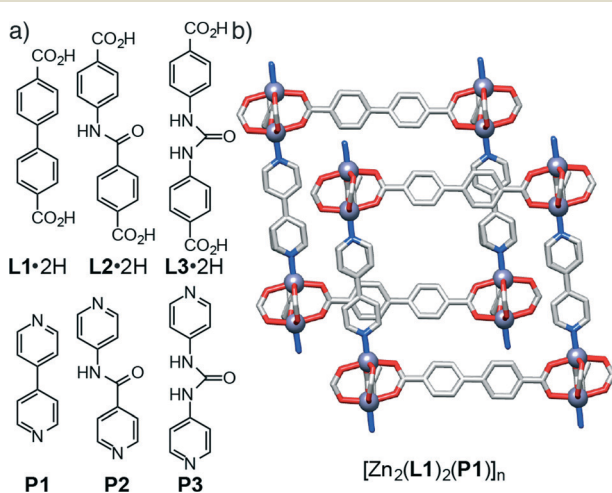


Fig. 1 a) Structures of the diacid ligands (**L1**·2H-**L3**·2H) and di-pyridyl pillars (**P1**-**P3**) examined in this study. b) Portion of the solid-state structure of $[\text{Zn}_2(\text{L1})_2(\text{P1})]_n$, the basic framework upon which the work is based. Redrawn from CCDC deposition FEFCUQ with hydrogen atoms and disorder omitted for clarity.



Table 1 Crystallographic data for compounds 1–4

	Compound 1	Compound 2	Compound 3	Compound 4
Formula	$C_{57}H_{67}N_9O_{15}Zn_2$	$C_{64}H_{82}N_{12}O_{18}Zn_2$	$C_{37}H_{46}N_8O_9Zn$	$C_{38}H_{47}N_9O_{10}Zn$
Mass (FM)	1248.93	1438.15	812.19	855.21
Crystal size (mm)	$0.408 \times 0.356 \times 0.142$	$0.536 \times 0.262 \times 0.145$	$0.100 \times 0.050 \times 0.040$	$0.110 \times 0.080 \times 0.050$
T (K)	150(2)	150(2)	150(2)	150(2)
Crystal system	Triclinic	Monoclinic	Monoclinic	Monoclinic
Space group	$P\bar{1}$	$P2_1/c$	$P2_1/n$	$P2_1/c$
a (Å)	15.1431(6)	19.7220(5)	16.381(3)	14.529(2)
b (Å)	15.2322(6)	28.0588(6)	14.170(3)	16.543(2)
c (Å)	16.0597(4)	28.0835(4)	19.415(4)	21.125(3)
α (°)	98.484(2)	90	90	90
β (°)	102.682(2)	94.423(2)	104.25(3)	104.305(9)
γ (°)	104.206(3)	90	90	90
V (Å ³)	3423.1(2)	15 494.4(6)	4367.9(16)	4919.8(12)
Z	2	8	4	4
ρ (g cm ⁻³)	1.212	1.233	1.235	1.155
$F(000)$	1304	6032	1704	1792
θ range (°)	2.818–25.772	2.880–26.000	1.926–25.764	1.971–25.132
Reflections	54 284	125 275	70 868	26 904
Independent reflections	13 023	30 417	8546	12 472
R_{int}	0.0360	0.0624	0.0960 ^a	0.0876 ^a
GOOF (F^2)	1.062	1.054	1.086	1.098
Final R indices [$I > 2\sigma(I)$]	$R_1 = 0.0662$ $wR_2 = 0.2062$	$R_1 = 0.0812$ $wR_2 = 0.2480$	$R_1 = 0.0905$ $wR_2 = 0.2450$	$R_1 = 0.1026$ $wR_2 = 0.2844$
Final R indices (all data)	$R_1 = 0.0759$ $wR_2 = 0.2157$	$R_1 = 0.1393$ $wR_2 = 0.2952$	$R_1 = 0.1336$ $wR_2 = 0.2956$	$R_1 = 0.1688$ $wR_2 = 0.3282$

^a Twin abs correction applied.

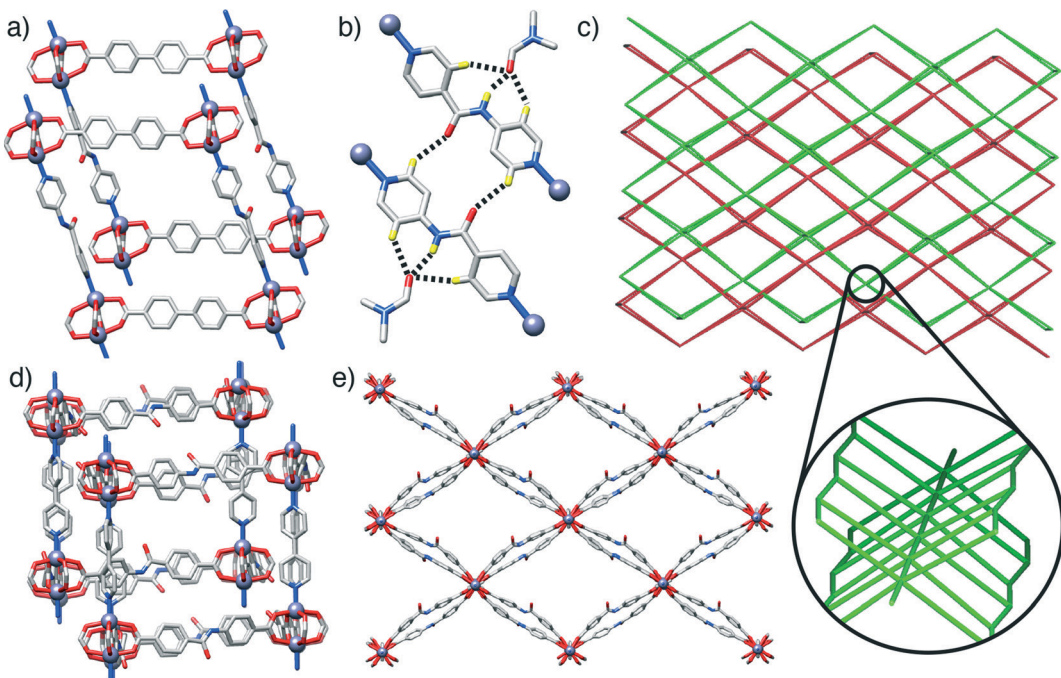


Fig. 2 a) Portion of the solid-state structure of $[Zn_2(L1)_2(P2)]_n$, showing the disposition of the SBUs induced by the amide pillar. b) The molecular recognition of a DMF solvate molecule by the amide unit of P2, which also forms a complementary H-bonded pair with a P2 unit of a separate net. c) The pcu topology present in both 1 and 2 with an inset showing the connection in the third dimension. d) Portion of the solid-state structure of 2, which has a more regular arrangement of SBUs compared to 1. e) Packing diagram of one of the two interpenetrated nets of $[Zn_2(L2)_2(P1)]_n$, viewed down the crystallographic c axis, showing the alternating pattern of L2 linkers.

95.1°, 101.6°, 110.0°, 110.0° and 140.0° deviating significantly from the ideal 109.5°. Four identical diamondoid (dia) nets

are interpenetrated (Fig. 3b), with DMF solvent molecules occupying significant rectangular channels (Fig. 3c) running



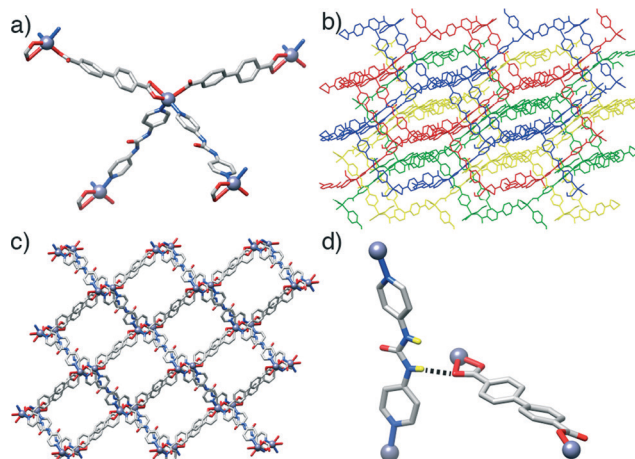


Fig. 3 Representations of the crystal structure of $[Zn(L1)(P3)]_n$. a) The four-connected Zn^{2+} node which connects the ligands. b) Packing structure showing the four-fold interpenetration of the material, with each net coloured differently. c) An alternative view of the packing structure to show the one dimensional channels that run down the crystallographic *b* axis. d) The hydrogen bond interaction between a urea proton of $P3$ and a carboxylic oxygen atom of $L1$ in an adjacent net.

down the crystallographic *b* axis, which may impart permanent porosity on the framework if successfully removed without collapse. Interestingly, the structure shows an important interaction between the interpenetrated nets (Fig. 3d), through a H-bond between a urea NH of each $P3$ and a carboxylic O of each $L1$ ($N \cdots O = 2.97 \text{ \AA}$). This is also manifested in the distorted Zn -O bond lengths, with distances of 2.564 and 1.951 \AA for the H-bonded and the non-interacting oxygens of the carboxylate, respectively.

In common with 3, compound 4, $[Zn(L2)(P3)]_n$, forms an interpenetrated **dia** topology, with a distorted octahedral Zn^{2+} coordination sphere consisting of two molecules each of $L2$ and $P3$. The amido-functionalised ligand $L2$ is disordered across two positions, with the offset of the amide moiety resulting in large thermal ellipsoids for the adjacent aromatic carbon atoms of the ligand. Compared to 3, the structure is also four-fold interpenetrated, with slightly smaller but still distinct one-dimensional channels running down the crystallographic *a* axis. Disorder in both the solvent and the amide unit make intermolecular interactions difficult to ascertain but, as in compound 4, a bifurcated interaction between the NH units of the urea pillar $P3$ and a carboxylic O atom from $L2$ in an adjacent net is observed ($N \cdots O = 2.76$, $N \cdots O = 2.85 \text{ \AA}$). Again, the H-bonding is reflected in the Zn -O bond distances, with values of 1.979 \AA and 2.566 \AA for the non-interacting and H-bonded oxygen atoms, respectively. Different nets also interact via π - π stacking between pyridine rings of adjacent molecules of $P3$, at a distance of 3.27 \AA .

Some trends are noticeable from the structural data of these four compounds. The introduction of the urea-functionalised pillar $P3$ gives MOFs of a dramatically different **dia** topology compared to the **pcu** topology seen for both the unfunctionalised parent structure and the two amide-containing frameworks. Presumably, the bent nature of the

urea moiety, which results in the non-linear (*ca.* 160°) disposition of the N-donors of the pyridyl units of $P3$, makes the **dia** topology more favourable. While the four compounds have structures comprising interpenetrating nets, those based on the longest pillar $P3$ lead to the formation of a more open framework and a higher degree of interpenetration, *i.e.* four-fold interpenetrated **dia** nets, classified as **IIIA** type. On the contrary, those composed of shorter pillars $P1$ and $P2$, give only two-fold interpenetrated **pcu**-nets, corresponding to class **IIa**.¹⁹ This is in agreement with the origin of interpenetration, since it can be ascribed to the presence of large free voids in a single network²⁰ and it is well known that one effective way to increase the number of interpenetrating folds is to extend the length of the ligand sustaining the framework (Fig. 4).²¹

The introduction of functional groups into the frameworks results in H-bonds between the urea NH of each $P3$ and a carboxylic O of each $L2$ or $L1$, for compounds 3 and 4 respectively, of adjacent nets in the solid-state interpenetrated structures, possibly acting as structure-directing effects.²² Unmasking these functional groups to act as substrate binding sites would necessitate breaking of these interactions during activation. It should also be noted that choice of solvent can affect the structures formed by these functionalised ligands and pillars. A hydrothermal synthesis involving $L1$, $P1$, Zn^{2+} and base has been shown, by Rietveld refinement of powder X-ray diffraction data, to yield²³ a non-porous coordination polymer, with five-fold interpenetrated **dia** topology, while a solvothermal synthesis using $L2$, $P1$ and Zn^{2+} in aqueous DMF gave²⁴ a triply interpenetrated structure linked by an alternative paddlewheel SBU into an elongated **pcu**

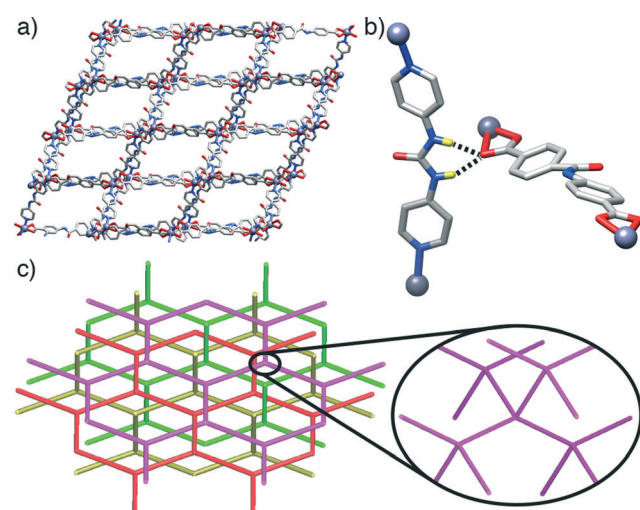


Fig. 4 a) Crystal packing in the solid-state structure of $[Zn(L2)(P3)]_n$ viewed down the crystallographic *a* axis to show one-dimensional pores. b) A bifurcated hydrogen bonding interaction in the crystal structure, between the urea unit of $P3$ and a carboxylic oxygen atom of a molecule of $L2$ from an adjacent net. c) The four-fold interpenetrated **dia** topology, with inset, adopted by both $[Zn(L1)(P3)]_n$ and $[Zn(L2)(P3)]_n$.



topology with overall formula $[\text{Zn}(\text{L2})(\text{P1})]_n$. In this regard, the **pcu** topology has a higher degree of interconnectivity compared to the **dia** topology, and therefore higher metal-ligand interactions that in the absence of additional external factors may provide robustness to the framework as well as minimise the energy. Higher metal-ligand interactions and, consequently, higher node-connectivity are frequently associated with increasing synthesis temperature,²⁵ which, in turn, favours the formation of thermodynamically more stable phases.²⁶

In an attempt to rationalise the formation of the different topologies and to understand the effect of functional group incorporation on materials properties, we turned to molecular simulations in order to build hypothetical new structures corresponding to the synthesised materials with swapped linkers, *i.e.* allowing us to examine the arrangement of ligands and pillars in topologies in which they are not observed experimentally. That is, compound **1** was used as a parent structure of three new hypothetical ones using the ligand and pillar combinations of compounds **2–4**, and this was repeated for each experimental structure. Therefore, from the initial 4 structures we built 12 new ones, named as $[\text{X-(L)(P)}]^*$, where **X** is the mother structure, **L** and **P** correspond to the ligands and pillars used according to Fig. 1, and $*$ denotes a computational structure. The strategy is similar to the methodology we have used previously for **rht** topology¹⁷ and alternative pillared MOFs.^{13e,f} We start with the asymmetric unit of every mother structure, maintaining the metal cluster (*i.e.* Zn paddle wheel or Zn), the **pcu** or **dia** topology as well as the corresponding group symmetry, but we swap the organic linkers. After the asymmetric units are modified, the symmetry operations of the space group allow the construction of the new structures. The unit cells are then subject to energy minimisation and geometry optimisation based on molecular mechanics, which include van der Waals and electrostatic interactions, modifying the size of the unit cells and the atomic coordinates of the new structures.

Interestingly, **pcu** topology hypothetical structures built with the **L1** and **L2** ligands and **P3** pillar (*i.e.* those with the constituents of the **dia** topology structures **3** and **4**) do not show any H-bonding between the urea moieties of each **P3** and carboxylic units of either **L1** or **L2**. This observation contrasts with the experimental **dia** compounds **3** and **4**, which show these close interactions. Fig. 5 illustrates how, in the case of the hypothetical $[\text{1-(L1)(P3)}]^*$ structure, the urea NH groups from **P3** point to the centre of the main cavities, precluding any possible interaction with **L1**. This observation reinforces our hypothesis about the importance of the H-bonds between adjacent nets present in compounds **3** and **4** in the formation and stabilisation of MOFs with **dia** topology instead of **pcu**. Indeed, it suggests that **dia** topology is effectively allowed and preferred when the urea moiety is incorporated into the pillar.

The fact that compounds **3** and **4** show a higher degree of interpenetration is not unexpected, since the **dia** topology is

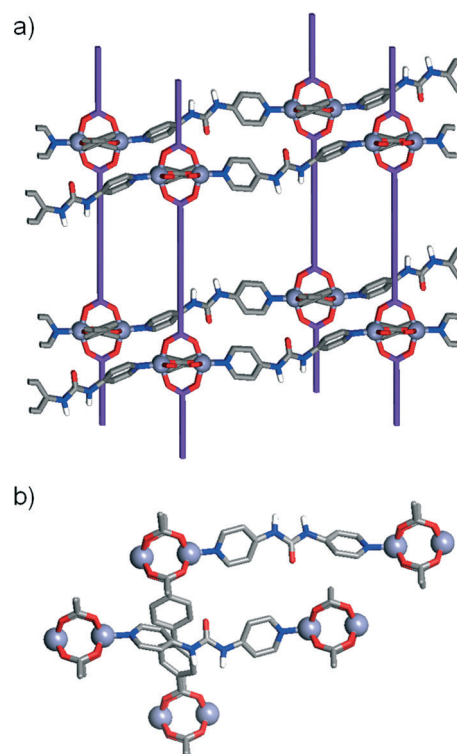


Fig. 5 a) Crystal packing of and b) detail of the solid-state structure of $[\text{1-(L1)(P3)}]^*$. Ligands **L1** have been simplified for clarity and are represented in purple in (a). Note that the urea protons of **P3** are pointing to the centre of the cavity and not to the carboxylic group of **L1**.

the most preferred one for interpenetrating frameworks. This is because these arrays can be more easily assembled from tetrahedral coordination centres than from octahedral (in particular, most of the arrays with high interpenetration degree $Z > 6$ are diamondoid). The most abundant topologies are the most symmetrical ones because they provide the most isotropic medium during crystallization. For instance, the diamond net has the automorphism group isomorphic to the $Fd\bar{3}m$ space group and may have an infinite number of distorted geometrical realizations (embeddings) of the space group symmetries corresponding to subgroups of $Fd\bar{3}m$. As it was demonstrated, any pair of nets can interpenetrate in some embedding, but in crystals there are physical reasons (like the condition of close packing), which essentially restrict possible combinations of interpenetrating nets.²⁷ These observations support once more the importance of these interactions between interpenetrated structures and the possibility of the urea acting as a structure-directing agent.

To investigate the propensity for urea moieties to form noncovalent interactions with carboxylate units in the solid state, we examined existing structures deposited with the Cambridge Structural Database.²⁸ Using a fragment-based search approach, 16 crystal structures, ranging from 1D coordination polymers to 3D MOFs, were identified, all of which containing both a carboxylate group coordinated to a metal and a *N,N*-disubstituted urea function in an extended network. Of these 16 structures, 10 contained H-bonding



interactions between urea NH units and carboxylate oxygen donors; eight involved interactions between individual ribbons of one dimensional coordination polymers²⁹ and two between distinct two dimensional grids.^{29c} Significant hydrogen bonding interactions were also present in the remaining structures, between different coordination polymer strands (three by urea-urea hydrogen bonded tapes,³⁰ and one through hydrogen bonding with a terminal water ligand^{29c}) as well as H-bonding to solvent (two examples with sterically constrained ligands^{12a,31}) and one example of hydrogen bonding to nitrate counterions as well as to carboxylate units.^{29c} In all but two of these cases, the urea groups of individual ribbons, sheets and nets form significant H-bonding interactions with adjacent ribbons, sheets and nets. The abundance of these noncovalent contacts in the solid-state structures of coordination polymers indicates they represent a significant structure directing influence, and may explain the preference for 3 and 4 to adopt the **dia** topology.

In order to compare the textural properties of the synthesised and hypothetical compounds, we first measured their geometrical surface area³² by rolling a 3.681 Å-diameter sphere, which corresponds to a nitrogen molecule, across the surface of the material. The pore size distributions (PSD) were calculated by recording the largest sphere that can fit in a random point within a structure without overlapping the van der Waals surface of the framework.³³ In addition, we calculated the N₂ adsorption isotherms at 77 K of the synthesised materials using grand canonical Monte Carlo (GCMC) simulations³⁴ to predict their potential adsorption performance. Table S1† shows the geometrical surface area of all the materials. Fig. 6 shows the simulated N₂ adsorption isotherms and the geometric PSD. Geometrical surface areas of all the synthesised compounds are very similar, with values ranging from 2000 and 2400 m² g⁻¹, and all four compounds present microporous cavities with diameters ranging from 6 to 8 Å. The simulated N₂ isotherms each present a type I shape and show total pore volumes between 0.65 and 0.77 cm³ g⁻¹. The textural properties (e.g. PSD, geometrical surface area and total pore volume) are directly related, with compound 2 showing the largest PSD, geometrical surface area and total pore volume, followed by 1 and 4. In addition, the N₂ adsorption isotherm of compound 2 is displaced to higher relative pressures when compared with the other isotherms. The shape of the adsorption isotherms is related to the balance between the strength of the fluid–fluid and fluid–solid interactions, which in turn is a strong function of the length of the linker and therefore the pore size.³⁵

3. Conclusions

We have utilised a combination of crystallographic and computational analyses to investigate the systematic incorporation of amide and urea functional groups directly into metal-organic frameworks with **pcu** and **dia** topology. Introduction of amide units is tolerated within the **pcu** topology (compounds

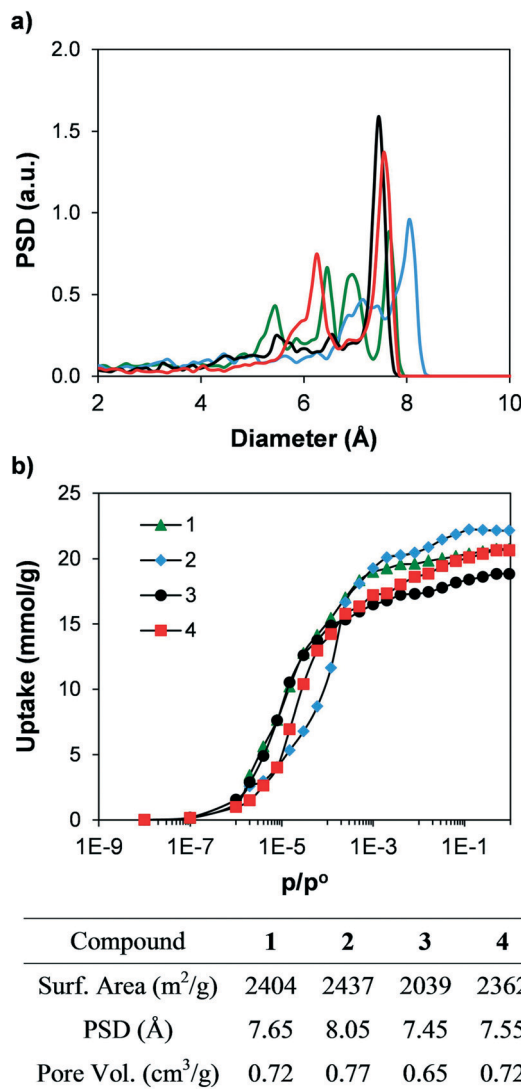


Fig. 6 a) GCMC simulated adsorption isotherms of N₂ at 77 K, b) geometric pore size distribution of the synthesised compounds. The table shows the geometrical surface area, pore size distribution (PSD) and total pore volume of the synthesised compounds.

1 and 2). However, integration of urea functional groups was not found to be possible under the synthetic conditions examined and led to the isolation of four-fold interpenetrated networks of **dia** topology with different compositions and SBUs (compounds 3 and 4). The comparison of hypothetical computationally created and experimental structures has shown that the presence of significant hydrogen bonding interactions between urea moieties of the pillars and carboxylate groups of the acidic linkers may contribute to the formation of the **dia** topology found in structures 3 and 4. Prediction of MOF topology and structure from a certain combination of ligands and metals remains difficult but possible,³⁶ with control over variables such as solvent, temperature and concentration required. Our results demonstrate that noncovalent interactions between linkers may be significant structure-directing factors, which has implications



for the integration of functional groups into MOFs to alter molecular recognition and catalytic properties.

4. Experimental

All the frameworks described were prepared by dissolution of the ligand, pillar and $[\text{Zn}(\text{NO}_3)_2] \cdot 6\text{H}_2\text{O}$ in DMF, with crystallisation induced by heating the colourless solution in a sealed vial for a period of approximately 2–5 days. Detailed descriptions, alongside ligand syntheses and computational details, are given in the ESL.† CCDC 1011797–1011800 contain the supplementary crystallographic data for this paper.

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