Introduction

The design and synthesis of coordination networks with interesting topologies and potential applications has attracted great attention during recent years. These new complexes thus synthesized are widely applied in catalysis, magnetism materials, gas storage, separation, ion exchange and optics. Entanglement is a very interesting phenomenon in coordination networks, and it is also a major factor contributing to the diversities of coordination networks.

The coordination networks of metal complexes containing flexible bidentate ligands are less predictable due to the possible occurrence of supramolecular isomerism involving the adoption of different ligand conformations. However, the flexible ligands with long spacer lengths are prone to form entangled structures due to their inclination to show large voids. Moreover, the dicarboxylate ligands that show diverse coordination ability can be used as auxiliary building blocks in the construction of coordination networks. Recently, we have reported two highly interpenetrated diamondoid nets of Zn(II) and Cd(II) coordination networks, \( \{[Zn(L^1)(1,4-BDC)] \cdot H_2O\}_n \) \( (L^1 = N,N'-di(4-pyridyl)adipamide; 1,4-H_2BDC = 1,4-benzenedicarboxylic acid) \) and \( \{[Cd(L^1)(1,4-BDC)] \cdot 2H_2O\}_n \) which show distorted cages with 9- and 9-fold interpenetrating modes. The combination of a long flexible \( L^1 \) ligand with short rigid 1,4-BDC\(^{2-} \) ligands reduced the number of interpenetration compared with the complex \( \{\text{CuSO}_4(L^1)(H_2O)\}_n \) featuring a 12-fold interpenetration. We have also shown that increasing the number of the backbone carbon atom of the neutral spacer ligand not only decreases the degree of interpenetration but also changes the structural type.

With this background information, we sought to investigate the influence of geometry and flexibility of the auxiliary dicarboxylate ligand on the structural diversity of coordination networks containing the flexible \( N,N'\text{-di(3-pyridyl)suberoamide} \) (L) ligands. The syntheses, structures and properties of \( \{[Cu(L)(1,2-pda)] \cdot H_2O\}_n \) \( [H_2(1,2-pda) = 1,2\text{-phenylenediacetic acid}] \),...
Crystals suitable for single-crystal X-ray diffraction. Yield: 0.026 g (43%). Anal. calcd for C_{28}H_{32}N_{4}CuO_{8} (MW = 616.12, for 2 days. Slow cooling of the reaction system afforded blue crystals suitable for single-crystal X-ray diffraction. Yield: 0.026 g (43%). Anal. calcld for C_{28}H_{32}N_{4}CuO_{8} (MW = 616.12, 1 + H_{2}O): C, 54.58; H, 5.24; N, 9.09%. Found: C, 55.28; H, 5.48; N, 8.78%. IR (cm^{-1}): 3442 (m), 2931 (s), 1611 (w), 1545 (w), 1481 (m), 1409 (m), 1272 (m), 1087 (s), 1040 (s), 742 (s). 

General procedures
IR spectra (KBr disk) were obtained using a JASCO FT/IR-460 plus spectrometer. Elemental analyses were performed using a PE 2400 series II CHNS/O analyzer or a HERAEUS VarioEL analyzer. Thermal gravimetric analyses (TGA) measurements were carried out with a TG/DTA 6200 analyzer from SII Nano Technology Inc. from 30 to 900 °C at a heating rate of 10 °C min^{-1} under nitrogen. Powder X-ray diffraction was carried out using a PANalytical PW3040/60 X’Pert Pro diffractometer with CuKα (λ = 1.54 Å) radiation.

Materials
The reagent Cu(OAc)$_2$·H$_2$O was purchased from SHOWA, 1,3-phenylenediacetic acid from Alfa Aesar, and 1,2-phenylenediacetic acid and 1,4-phenylenediacetic acid from ACROS. The ligand N$_2$N’-di(3-pyridyl)suberamide (L) prepared according to a published procedure.}

Preparation
[(Cu(L)(1,3-pda))·2H$_2$O]$_n$ 1. A mixture of Cu(CH$_3$COO)$_2$·H$_2$O (0.020 g, 0.10 mmol), L (0.033 g, 0.10 mmol), 1,2-H$_2$PDA (0.019 g, 0.10 mmol) and 15 mL of NaOH (0.04 M) solution was sealed in a 23 mL Teflon-lined stainless steel autoclave, which was then heated under autogenous pressure to 120 °C for 2 days. Slow cooling of the reaction system afforded blue crystals suitable for single-crystal X-ray diffraction. Yield: 0.026 g (43%). Anal. calcld for C$_{28}$H$_{32}$N$_{4}$CuO$_{8}$ (MW = 616.12, 1 + H$_2$O): C, 54.58; H, 5.24; N, 9.09%. Found: C, 55.28; H, 5.48; N, 8.78%. IR (cm$^{-1}$): 3442 (m), 2931 (s), 1611 (w), 1545 (w), 1481 (m), 1377 (w), 707 (s).

[(Cu(L)(1,3-pda))·2H$_2$O]$_n$ 2. Prepared as described for 1, except that Cu(BF$_4$)$_2$·2H$_2$O (0.023 g, 0.10 mmol) and 1,3-H$_2$PDA (0.019 g, 0.10 mmol) were used. Yield: 0.025 g (40%). Anal. calcld for C$_{28}$H$_{32}$CuN$_{4}$O$_7$ (MW = 600.12, 2 – H$_2$O): C, 56.04; H, 5.37; N, 9.34%. Found: C, 55.74; H, 5.20; N, 9.11%. IR (cm$^{-1}$): 3267 (s), 1705 (s), 1548 (w), 1483 (s), 1390 (m), 1287 (s), 1158 (s).

[(Cu(L)(1,4-pda))·2H$_2$O]$_n$ 3. Prepared as described for 1, except that 1,4-H$_2$PDA (0.021 g, 0.10 mmol) was used. Yield: 0.020 g (32 %). Anal. calcld for C$_{28}$H$_{32}$N$_{4}$CuO$_{8}$ (MW = 618.14): C, 54.40; H, 5.54; N, 9.06%. Found: C, 54.20; H, 5.07; N, 8.89%. IR (cm$^{-1}$): 3289 (m), 3069 (m), 2940 (m), 1712 (s), 1606 (m), 1553 (s), 1478 (m), 1380 (s), 1332 (m), 1294 (m), 1156 (m), 776 (m), 717 (m).

X-ray crystallography
The diffraction data for complexes 1–3 were collected with a Bruker AXS SMART APEX II CCD diffractometer at 22 °C, which was equipped with graphite-monochromated Mo Kα (λ = 0.71073 Å) radiation. Data reduction was carried out by standard methods with the use of a well-established computational procedure. The structure factors were obtained after Lorentz and polarization corrections. An empirical absorption correction based on “multi-scan” was applied to the data for all complexes. The positions of some of the heavier atoms were located by direct methods and the remaining atoms were found in a series of alternating difference Fourier maps and least-square refinements, except that the hydrogen atoms were added by using the HADD command in SHELXTL 5.10. In complexes 1 and 2, some of the carbon [C(6)–C(9)] and C(17)–C(18) for 1 and C(14) for 2] and oxygen [O(1) and O(17) for 1] atoms are disordered such that two orientations can be found for each disordered atom. Basic information pertaining to crystal parameters and structure refinement is summarized in Table 1. Selected bond distances and angles are listed in Table 2.

Results and discussion

Structure of 1
Crystals of 1 conform to the triclinic space group P1 with one Cu(II) cation, two halves of two different L ligands and one 1,2-PDA$^{2-}$ anion in the asymmetric unit. Fig. 1(a) shows the coordination environment of the Cu(II) center, which is five-coordinated by two pyridyl nitrogen atoms from two L ligands and three oxygen atoms from three 1,2-PDA$^{2-}$ ligands, resulting in a distorted square pyramidal geometry with a τ value of 0.32. The Cu(II) ions are linked together by the μ$_3$-κ$_1$-κ$_1$κ$^1$ coordination mode to afford 1D looped chains with dinuclear metal units (Fig. 1(b)) which are further connected by the L ligands through the pyridyl nitrogen atoms to form a 3D structure. In the structure, 1,2-PDA$^{2-}$ interact with the L ligand through N=H⋯O (H=O = 2.02 Å, N⋯O = 2.86 Å, <N–H–O = 162°) hydrogen bonds.

If the Cu(II) cations are defined as 5-coordinated nodes and the 1,2-PDA$^{2-}$ ligands as 3-coordinated nodes, topological analysis reveals that the structure of 1 can be regarded as a 3,5-connected net with the (4$^2$-6$^5$-8$^3$)(4$^4$-6)-3,5T1 topology (Fig. 1(c)), as determined using TOPOS. If the Cu$_2$(μ-COO)$_2$ dimer is considered as a node, the structure of 1 can be further simplified as a 6-connected net with a single (4$^2$-6$^3$)-pcu topology (Fig. 1(d)). The underlying 3,5T1 net is observed in 81 crystals (interpenetrated in 13) as can be found in TTD TOPOS database. Interestingly five compounds present M$_2$(μ-COO)$_2$ dimers that allow the same alternative cluster description of the net as pcu, with [Cd(adp)(bpfp)]$_n$ [adp = adipate; bpfp = bis(4-pyridylformyl)piperazine] the only one with two different ligands of very different lengths.

Structure of 2
Crystals of 2 conform to the monoclinic space group C2/c with the Cu(II) cation on the inversion center, with half of L...
Table 1  Crystal data for complexes 1-3

<table>
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<tr>
<th>Compound</th>
<th>Formula</th>
<th>Space group</th>
<th>α, Å</th>
<th>β, Å</th>
<th>γ, Å</th>
<th>αo</th>
<th>βo</th>
<th>γo</th>
<th>V, Å³</th>
<th>Z</th>
<th>Dcalc, g cm⁻³</th>
<th>μ, μV</th>
<th>ω</th>
<th>β, °</th>
<th>θ, °</th>
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<tr>
<td>1</td>
<td>C₂₄H₃₂N₄O₄Cu</td>
<td>P1</td>
<td>8.3326(4)</td>
<td>10.3780(5)</td>
<td>16.5929(9)</td>
<td>87.460(3)</td>
<td>77.874(3)</td>
<td>74.322(3)</td>
<td>1350.09(12)</td>
<td>2</td>
<td>1.476</td>
<td>1.445</td>
<td>1.438</td>
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<td></td>
</tr>
<tr>
<td>2</td>
<td>C₂₄H₃₂N₄O₄Cu</td>
<td>C2/c</td>
<td>29.5458(9)</td>
<td>6.3519(2)</td>
<td>18.5588(6)</td>
<td>90</td>
<td>125.320(2)</td>
<td>90</td>
<td>2841.88(15)</td>
<td>3</td>
<td>1.476</td>
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<tr>
<td>3</td>
<td>C₂₄H₃₂N₄O₄Cu</td>
<td>Pna</td>
<td>15.8723(8)</td>
<td>12.1103(6)</td>
<td>14.8539(7)</td>
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<td>90</td>
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<td>177.784(3)</td>
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Table 2  Selected bond lengths (Å) and angles (°) for complexes 1-3

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<td>Cu-O(6A)</td>
<td>1.959(4)</td>
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<td>Cu-N(1)</td>
<td>2.001(5)</td>
<td>2.001(5)</td>
<td>2.001(5)</td>
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<tr>
<td>Cu-O(5B)</td>
<td>2.173(4)</td>
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<tr>
<td>O(6A)-Cu-O(3)</td>
<td>150.3(2)</td>
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<tr>
<td>O(3)-Cu-N(1)</td>
<td>90.9(2)</td>
<td>90.9(2)</td>
<td>90.9(2)</td>
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<tr>
<td>O(3)-Cu-N(3)</td>
<td>90.9(2)</td>
<td>90.9(2)</td>
<td>90.9(2)</td>
</tr>
<tr>
<td>O(6A)-Cu-O(5B)</td>
<td>122.9(2)</td>
<td>122.9(2)</td>
<td>122.9(2)</td>
</tr>
<tr>
<td>N(1)-Cu-O(5B)</td>
<td>97.0(2)</td>
<td>97.0(2)</td>
<td>97.0(2)</td>
</tr>
</tbody>
</table>

and 1,3-PDA⁺⁻ ligands in the asymmetric unit. The coordination environment of the Cu(n) metal center is shown in Fig. 2(a), which is four-coordinated by two pyridyl nitrogen atoms from two L ligands and two oxygen atoms from two 1,3-PDA⁺⁻ ligands, resulting in a distorted square planar geometry. Two weak Cu⋯O interactions with a distance of 2.620(4) Å are also observed in the axial positions. The Cu(n) ions are linked together by the 1,3-PDA⁺⁻ ligands that adopt the μ₂-K⁺,K⁺,K⁺ coordination mode to afford 1D linear chains (Fig. 2(b)), which are further linked by the L ligands through the pyridyl nitrogen atoms to form a 3D structure.
Structure of 3

Crystals of 3 conform to the orthorhombic space group Pnma with the Cu(n) cation lying on a mirror plane, with half of L and 1,4-PDA \( \equiv \) ligand and two cocystallized water molecules in the asymmetric unit. Fig. 3(a) shows the coordination environment of the Cu(n) metal center, which is five-coordinated by two pyridyl nitrogen atoms from two L ligands and three oxygen atoms from three 1,4-PDA \( \equiv \) ligands, resulting in a distorted square pyramidal geometry with a \( \tau \) value of 0.06. The Cu(n) ions are bridged by the 1,4-PDA \( \equiv \) anions that adopt the \( \mu_3-k^1,k^1,k^2 \) coordination mode to form dinuclear units with a Cu...Cu distance of 3.309(1) Å, which are further linked by the L ligands and the 1,4-PDA \( \equiv \) ligands to form a 1D self-catenated coordination network (Fig. 3(b) and (c)). It can be seen that each 34-membered ring that involves two Cu(n) ions and two L ligands is penetrated by two 1,4-PDA \( \equiv \) ligands. Adjacent 1D chains are connected by water molecules through N–H⋯O (H⋯O = 2.14 Å, N⋯O = 3.00 Å, \( \angle \text{N–H}⋯\text{O} = 177^\circ \)) hydrogen bonds originating from the amine hydrogen atoms of the L ligands to the water molecules and O–H⋯O (H⋯O = 1.82–1.93 Å, O⋯O = 2.65–2.76 Å, \( \angle \text{O–H}⋯\text{O} = 164–168^\circ \)) hydrogen atoms to the carboxylate oxygen atoms of 1,3-PDA \( \equiv \) ligands (Fig. S5†).

Self-catenated nets are single nets that exhibit the peculiar feature of containing shortest rings through which pass other components of the same network.13 Catenation can be identified by the presence of edges that thread a ring; in other words, they share at least one point with a disc-like film bounded by the ring. Since infinite nets with three lattice vectors inevitably contain large cycles that are threaded, we restrict “rings” to smallest cycles in the net that cannot be decomposed into sums of still smaller cycle, such as “strong rings”13 if both the threaded edge and the ring belong to a single connected component of the structure, it is self-catenated. Most coordination networks showing the feature of self-catenation are 3D nets,14 and there are only 2D cases that are presently known.15 Complex 3 thus appears to be the first 1D coordination network showing self-catenation.

This network has an additional feature that is worth noting, namely, it is entangled. To see this, it is convenient to redraw complex 3 as a “ladder” (with single back- “\( / \)” and paired forward-slash “\( / / \)" stringers), as shown in Fig. 4(a). Here the structure has been redrawn to preserve its topology. Further, although its geometric details (such as vertex
locations and bonding geometries) are distorted, its tangled features, namely, the mutual catenation of edges and cycles, are also preserved. This means that the structure in Fig. 4(a) is an equivalent “tangled-isotope” that preserves both the topology and the edge crossings. Here we adopt the term “tangled-isotope” from the mathematical operation of “ambient isotopy”, corresponding to distortions of a structure that do not alter its inherent entanglement.16 The ring formed by a pair of forward-slashes – one above and one below – sharing two vertices is threaded by (two parallel) back-slashes, describing the self-catenation. A simpler, untangled-isotope is that whose pair of forward-slash stringers is either both above or both below the back-slash stringers, as illustrated in Fig. 4(b). The edges of this untangled-isotope reticulate a cylinder without any edge crossings (with forward- and back-slashes running on opposite faces of the cylinder). In contrast, the edges of complex 3 cannot reticulate a cylinder without edge crossings: a multi-handled surface made of a pair of parallel cylinders with connecting tubes is necessary to support a crossing-free reticulation. Since this latter surface is topologically more complex than the simple cylinder, complex 3 is (according to the definition advanced in ref. 15) entangled.

Closely related tangled-isotopes to complex 3 are knotted, with more common signatures of entangled networks in terms of constituent knots or links. For example, permutations of the crossings between forward- and back-slash stringers result in patterns containing threaded links, or pairs of catenated rings. Two of the simpler tangled-isotopes are those with Hopf links (Fig. 4(c)) and Solomon links (Fig. 4(d)). In stark contrast to complex 3, both of these are tangled-isotopes since they contain pairs of interwoven cycles that cannot be separated without breaking edges, as illustrated in the figures. They are distinguished by the crossings of one cycle (blue in the figures) with the interlocked (black) cycle. If we denote crossings of blue edges over/under black edges by the symbols “+/−”, an anticlockwise walk around a blue cycle made of four edges in Fig. 4(c) crosses black edges in the order “− + + +” (or cyclic permutations); the analogous walk in Fig. 4(d) is “− + − −”. These sequences of over and under crossings characterise Hopf and Solomon links, respectively (see, for example, ref. 17). However, the entanglement in complex 3 is a peculiar and hitherto unrecognised one. It has neither knots nor links. Further, it is also free of “ravels” – a subtle entanglement mode distinct from knots or links – that can be formed in tangled nets.19 To date, knots, links and ravels have all been identified in metal–organic complexes.2,19 Complex 3 is, to our knowledge, the first example of a tangled net that does not contain knots, links or ravels, thus exemplifying a new mode of entanglement.

Ligand conformations and structural comparisons

Twenty ligand conformations can be found for L if the A and G conformations are given when the C−C−C−C torsion angle (θ) is $0 \leq \theta \leq 90^\circ$ and $90 \leq \theta < 180^\circ$, respectively. In
addition, based on the relative orientation of the C=O groups, each conformation can adopt a cis or a trans arrangement. Moreover, due to the difference in the orientations of the pyridyl nitrogen atom positions, three more orientations, anti–anti, syn–anti and syn–syn, based on the relative positions of the pyridyl nitrogen and amide oxygen atom can also be derived (Fig. 5).

Table 3 lists all of the ligand conformations of the L ligand in complexes 1–3. The structural parameters of the reported complexes [Cu(L)(1,2-bdc)]n, (1,2-H2bdc = 1,2-benzenedicarboxylic acid) and [Cu(L)(1,3-bdc)(H2O)·3H2O]n (1,3-H2bdc = 1,3-benzenedicarboxylic acid) that contain L are also shown for comparison.20 A comparison of these structures shows that the flexibility of the dicarboxylate ligands, in which the pda2− ligands in 1–3 hold two more CH2 groups than the bdc2− ligands, significantly affects the structural diversity. In 1 and [Cu(1,2-bdc)(L)]n, the 1,2-pda2− and 1,2-bdc2− ligands adopt the μ3-x,y−,x,z−,y,z− mode and μ3-x,y−,x,z− mode, respectively, resulting in AAAAA trans anti–anti and GGAGG trans anti–anti conformations for the L ligands in 1 and AAAAA trans syn–syn conformation for the L ligands in [Cu(1,2-bdc)(L)]n, forming a 3D structure with 3,5T1 topology and a 2D wavelike structure, respectively. In contrast, the 1,3-pda2− and 1,3-bdc2− ligands of 2 and [Cu(1,3-bdc)(L)(H2O)·3H2O] adopt the same μ3-x,y−,x,z− coordination mode, but lead to the formation of AAAAA trans anti–anti conformation for the L ligands in 2 and GAAAG trans syn–syn and AAAAA trans syn–syn conformations for the L ligands in [Cu(1,3-bdc)(L)(H2O)·3H2O], resulting in 5- and 3-fold interpenetration, respectively. In each single cds structure, the distances between the Cu(ii) nodes are 12.08 Å (through 1,3-pda2−) and 21.69 Å (through L) for 2 significantly longer than those of 9.66 Å (through 1,3-bdc2−) and 19.46 Å (through L) for [Cu(1,3-bdc)(L)(H2O)·3H2O], indicating the importance of the length of the spacer ligand in determining the number of interpenetrations. The cis- and trans-conformations of the phenylenediacetate ligands have been investigated recently.21 For example, in the 1D ladder-like silver(i) complex [Ag3(cis-pda)(bipy)]6−·0.5n(trans-pda)·5H2O (bipy = 4,4′-bipyridine), both of the two different formations (cis- and trans) of the 1,2-pda2− ligands were observed.21a Inspection of the structures of the three isomeric pda2− ligands in 1–3 reveals that they all adopt the trans conformation.

Thermal properties

In order to estimate the stability of the frameworks, thermogravimetric analysis (TGA) of complexes 1–3 was carried out in nitrogen from 30 to 900 °C (Fig. S6 and Table S1†). The TGA curve of 1 shows the gradual weight loss of water molecules (calculated 5.02%; observed 5.35%) in 50–190 °C. The weight loss between 200 and 460 °C corresponds to the decomposition of the L ligand and 1,4-pda2− ligand (calculated 84.96%; observed 86.41%). The TGA curve of 2 shows the gradual weight loss of water molecules (calculated 3.00%; observed 2.16%) in 45–200 °C, and the weight loss of 86.19% between 200 and 510 °C corresponds to the decomposition of the L ligand and 1,4-pda2− ligand (calculated 86.41%). For complex 3, the TGA curve shows the gradual weight loss of water molecules (calculated 5.83%, observed 6.56%) in 50–200 °C. The weight loss of 83.26% between 200 and 460 °C can be ascribed to the decomposition of the L ligand and 1,4-pda2− ligand (calculated 83.89%). The TGA results show that the weight losses that are due to the removal of cocrystallized and the bonded water molecules

![Diagram of possible orientations for the pyridyl nitrogen atoms of the L ligand: (a) syn–syn (b) syn–anti (c) anti–anti.](image)

Table 3 Some important parameters of complexes 1–3, [Cu(L)(1,2-BDC)]n, and ([Cu(L)(1,3-BDC)(H2O)·3H2O]n

<table>
<thead>
<tr>
<th>Complex</th>
<th>Coordination mode of the dicarboxylate ligand</th>
<th>Conformation of L</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Cu(L)(1,2-pda)][H2O]n, 1</td>
<td>μ3-x,y−,x,z−,y,z−</td>
<td>AAAAA trans anti–anti</td>
<td>cds, 5-fold interpenetration</td>
</tr>
<tr>
<td>[Cu(L)(1,3-pda)][2H2O]n, 2</td>
<td>μ3-x,y−,x,z−</td>
<td>GGAGG trans anti–anti</td>
<td>1D self-catenation</td>
</tr>
<tr>
<td>[Cu(L)(1,4-pda)][2H2O]n, 3</td>
<td>μ3-x,y−,x,z−,y,z−</td>
<td>AAAAA trans anti–anti</td>
<td>sql</td>
</tr>
<tr>
<td>[Cu(L)(1,2-bdc)]n, 4</td>
<td>μ3-x,y−,x,z−,y,z−</td>
<td>AAAAA trans syn–syn</td>
<td>cds, 3-fold interpenetration</td>
</tr>
</tbody>
</table>

* Derived from the crystallographic data for [Cu(L)(1,2-BDC)]n and ([Cu(L)(1,3-BDC)(H2O)·3H2O]n.

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occurred in 45–200 °C, and the organic ligands decomposed at temperatures above 200 °C. The different structural types of 1–3 do not affect the decomposition of the organic ligands significantly. The PXRD patterns of complexes 1–3 heated at 150 °C for 3 hours shown in Fig. S1–S3 reveal that their structures are stable upon solvent removal. It is noted that the solvent-accessible volumes of complexes 1–3 calculated with PLATON22 after the removal of water molecules are as small as 67.5, 161.0 and 196.2 Å³, which are 5.0%, 5.7% and 6.9% of each of the unit cell volumes, respectively.

Conclusion

Three new coordination networks containing L and isomeric phenylenediacetate ligands have been successfully synthesized under hydrothermal conditions. The different structures in 1–3 can be attributed to the ligand isomerism of the PDA²⁻ ligands that result in different L conformations. Obviously, the L ligands are sufficiently flexible to adjust to the stereochemical requirements for the formation of complexes 1–3, which adopt the conformations that maximize their intra- and intermolecular forces. To our best knowledge, complex 1 is a rare example with two different ligands of very different lengths, with underlying 3,5T1 nets further simplified to pcu; complex 2 shows the maximum number of interpenetrations presently known for cdv coordination networks; and complex 3 is the first 1D self-catenated, entangled, unknotted and unravelled coordination network.

Acknowledgements

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References


