A new twist on an old ligand: a [Mn\(_{16}\)] double square wheel and a [Mn\(_{10}\)] contorted wheel†

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Ligand design remains key to the synthesis of coordination compounds possessing specific topologies, nuclearities and symmetries that direct targeted physical properties. N,O-chelates based on ethanolamine have been particularly prolific in constructing a variety of paramagnetic 3d transition metal complexes with fascinating magnetic properties. Here, we show that combining three ethanolamine moieties within the same organic framework in the form of the pro-ligand 1,3,5-tri(2-hydroxyethyl)-1,3,5-triazacyclohexane (LH\(_3\)) leads to the formation of two highly unusual Mn wheels. Reaction of Mn(NO\(_3\))\(_2\)·6H\(_2\)O with LH\(_3\) in basic methanolic solutions leads to the formation of [Mn\(_{16}\)(µ-3-O)\(_6\)(µ-3-OMe)\(_2\)(µ-OMe)\(_2\)(L)\(_4\)(H\(_2\))\(_2\)NO\(_3\)\(_2\)(H\(_2\)O)\(_2\)] (1) and [Mn\(_{10}\)(µ-3-O)\(_2\)(µ-OMe)\(_2\)(µ-3-OMe)\(_2\)(µ-OMe)\(_2\)(L)\(_2\)(H\(_2\)O)\(_2\)]NO\(_3\)\(_2\) (2), the only difference in the synthesis being the ratio of metal:ligand employed. The structure of the former describes two offset [Mn\(_{16}\)(µ-3-O)] square wheels, linked through a common centre, and the latter a single [Mn\(_{10}\)] wheel twisted at its centre, such that the top half is orientated perpendicular to the bottom half. In both cases the L\(^3\)/LH\(^2\)– ligands dictate the orientation of the Jahn-Teller axes of the Mn\(_{10}\) ions which lie perpendicular to the triazacyclohexane plane. Direct current magnetic susceptibility and magnetisation data reveal the presence of competing exchange interactions in 1 and strong antiferromagnetic interactions in 2. Given the simplicity of the reactions employed and the paucity of previous work, the formation of these two compounds suggests that LH\(_3\) will prove to be a profitable ligand for the synthesis of a multitude of novel 3d transition metal complexes.

Introduction

The development of magneto-structural relationships in molecular coordination compounds can be traced back to measurements of copper(II) acetate and the basic metal(II) carboxylates, wherefrom their dinuclear and trinuclear structures, solved later, were predicted.\(^1,2\) Detailed, quantitative analyses of a variety of di-, tri- and tetracational 3d transition metal complexes followed, often revealing a complex relationship between exchange interactions/magnetic anisotropy, and the identity of the ligand, metal-metal distances, metal-ligand bond lengths, bond angles, torsions angles, and metal geometry – correlations now benefitting from detailed theoretical input.\(^3\)\(^–\)\(^5\) Interest in the magnetochemistry of Mn compounds in particular was boosted by the discovery of single-molecule magnets (SMMs), the first of which was a [Mn\(_{12}\)] complex\(^6\) whose structure was reported some years earlier.\(^7\) Magneto-structural studies were aided and abetted by magnetic measurements on the large library of low-nuclearity Mn compounds initially established as metalloenzyme model complexes, particularly those pertaining to the water oxidation centre in PSII.\(^8\) These proved vital in both the development of novel synthetic methodologies for the construction of new Mn compounds whose nuclearities now reach eighty four\(^9\) and in understanding the origin of the slow magnetisation relaxation dynamics.\(^10\)

Central to these studies has been the design of ligands capable of bridging between paramagnetic metal ions in a particular manner, be that within a rigid or flexible framework. One very successful class of ligands in the latter category are N,O-chelates including 2-(hydroxymethyl)pyridine (hmp\(_{\text{H}}\)),\(^1\)\(^1\) 2,6-pyridinemethanol (pdm\(_{\text{H}}\)),\(^1\)\(^2\) di- (R-dea\(_{\text{H}}\))\(^1\)\(^3\) and triethanolamine (tea\(_{\text{H}}\))\(^1\)\(^4\) which are all characterised by posses-
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Fig. 1 The N.O-chelates 2-(hydroxymethyl)pyridine (hmpH), 2,6-pyridinedimethanol (pdmH)2, di- (R-deaH) and triethanolamine (teaH)3, and 1,3,5-tri(2-hydroxyethyl)-1,3,5-triazacyclohexane (LH3) all of which contain one or more linked ethanolamine (eaH) moieties.

Experimental

General methods

All chemicals were obtained from commercial suppliers (Sigma-Aldrich) and were used without further purification/treatment.

Synthesis

LH3 was prepared as previously described.16

\[ \text{[Mn}^{III}{\text{Mn}}^{II}{\mu}_{3}-\text{O}]_{2}{[\mu-{\text{OH}}]}_{3}{[\mu-{\text{OMe}}]}_{3}{[\text{LH}]}_{4}{[\text{H}_{2}\text{O}]}_{10} \cdot \text{(NO}_{3}\text{)}_{2}{[\text{OH}]}_{2} \] (1). Mn(NO3)2·6H2O (143.5 mg, 0.5 mmol) and LH3 (103.58 mg, 0.5 mmol) were stirred in MeOH for 30 minutes in the presence of NEt3 (1 mmol). The solution was then filtered and allowed to stand. Crystals of 1 formed in ~5 days in a yield of 20%. Elemental analysis (%) calcd for C40H96Mn10N14O34 (M = 1866.70 g mol⁻¹), tetragonal, space group I4₁/a (no. 88), a = 18.2751(8) Å, c = 32.4453(3) Å, V = 10 836.1(13) Å³, Z = 4, T = 100.0 K, Bruker D8 Venture diffractometer, μ(CuKα) = 9.687 mm⁻¹, Dcalc = 1.144 g cm⁻³, 182 310 reflections measured (8.748° ≤ 2θ ≤ 149.676°), 5547 unique (Rint = 0.0634, Rsigma = 0.0144) which were used in all calculations. The final R1 was 0.0297 (I > 2σI) and wR2 was 0.0972 (all data). Neighbouring clusters in the structure of 2 pack so as to form large solvent/anion occupied spaces that are extremely disordered as evidenced by the presence of diffuse electron density. The presence of nitrate counterions in the large voids in the structure of 2 was confirmed by IR spectroscopy. Given the diffuse nature of the density in the difference map it is not possible (or sensible) to try and model this.

Magnetometry

Variable-temperature and variable-field magnetic measurements were carried out using a MPMS-XL Quantum Design magnetometer equipped with a 5 T magnet. Diamagnetic corrections were applied using Pascal's constants.

Powder-XRD measurements

Powder XRD measurements were collected on freshly prepared samples of the complexes on a PANanalytical X'Pert Pro MPD diffractometer.

Infra-red spectroscopy

FTIR-ATR (Fourier-transform infrared-attenuated total reflection) spectra were recorded on a PerkinElmer FTIR Spectrum BX spectrometer.

Results and discussion

The 1:1 reaction of LH3 with Mn(NO3)2·6H2O in a basic MeOH solution produces dark brown crystals of [Mn12Mn14][µ3-O]6[µ-OH]14[µ-OMe]12[LH]4[H2O]10[NO3]6[OH]2·2 (1; Fig. 2) after 5 days. 1 crystallises in a monoclinic cell and structure solution was performed in the space group P21/n. The asymmetric unit of 1 contains half the cluster complex and three nitrate/one hydroxide counter ions. The metallic skeleton of the asymmetric unit contains a non-planar, asymmetric...
xyethyl)-1,3,5-triazacyclohexane ligands are of two types, two around the outside of the wheel (Mn₃–Mn₇, Mn₆–Mn₈) and two around the inside of the square wheel (Mn₂, Mn₄, Mn₆, Mn₈). The deprotonated O-atoms further bridge to neighbouring MnIII ions, while the protonated arm remains terminally coordinated to MnI. The MnIII ions are all six-coordinate and in regular octahedral geometries, with MnI having one and MnV having three coordinated H₂O molecules.

There are several short intermolecular interactions. The H₂O molecule (O0) and terminally bonded O(L) atom (O2) on MnI are H-bonded to their symmetry equivalent atoms on neighbouring molecules (O⋯O, 2.63 Å) creating 1D chains of wheels down the c-axis of the crystal. (O0) is also H-bonded to a NO₃⁻ counter anion (O0⋯O20, 2.86 Å) which is further H-bonded to both H₂O solvent of crystallisation (O21⋯O31, 2.80 Å) and to the H₂O molecule (O22⋯O29, 2.63 Å) and terminal O(L) on MnI (O7⋯O21, 2.75 Å). The NO₃⁻ counter anions are also H-bonded to the H₂O molecules on Mn5 (O17⋯O25, 2.78 Å; O18⋯O23, 2.72 Å; O19⋯O27, 2.87 Å) and the μ-OH⁻ bridging between Mn6–Mn7. The result is a complicated network of interactions in all three dimensions.

Repeating the reaction that produces 1, but increasing the Mn:OH₃ ratio to 1:2 produces the complex [MnIII₄(H₂O)₄(μ-OMe)₄(L)₄(H₂O)₄][NO₃]₂ (2). 2 crystallises in the tetragonal space group P4/α (Fig. 3, top) with three MnIII ions, one O²⁻ (O5), one OMe⁻ (O4) and one OH⁻ (O7) ion in the asymmetric unit. The metallic skeleton of 2 describes a rather contorted [MnIII₄] square wheel of corner sharing [MnIII₃O] triangles, twisted at its centre such that the top half is orientated perpendicular to the bottom half (Fig. 3, bottom). There are two corner sharing [MnIII₃O] triangles in each [MnIII] half, each with a μ-O²⁻ at its centre and a μ-OMe⁻ along the Mn1–Mn3 edge (Mn1–O4–Mn01, 96.9°). The two halves of the molecule are connected via four μ-OH⁻ ions (Mn01–O7–Mn01, 138°), which are H-bonded to the μ-OMe⁻ ions (O7⋯O5, 2.895 Å). There are two L⁻ ligands in each [MnIII] half of the molecule bonding in an identical μ₄-fashion, N,O-chelating to the MnIII ions with just one of the three arms (O2) further bridging to a neighbouring metal centre. The MnIII ions are all in Jahn-Teller (JT) distorted octahedral geometries, again dictated by the Mn–N(L) bonds. The remaining coordination site on Mn1 is occupied by a H₂O molecule (O6) which, alongside O1(L), H-bond to the symmetry equivalent atoms on neighbouring molecules (O6⋯O1, 2.588 Å). The result is that the [MnIII₄O] clusters pack in an aesthetically pleasing brickwork-like fashion, forming large solvent filled channels (Fig. 4). O6 also forms an internal H-bond to one of the terminally bonded O(L) atoms (O6⋯O3, 2.579 Å).

There are several structural similarities between 1 and 2. Both are square wheels composed of corner-sharing [Mn₃O]⁻.
triangles as directed by the L\(^3\) and LH\(^2\) ligands. The N-atoms of the ligands also dictate the orientation of the JT axes (and hence the d\(_{z^2}\) orbital) of the Mn\(^{III}\) ions, which has important design consequences for tuning magnetic exchange and magnetic anisotropy.\(^{21}\) Both compounds possess terminally bonded H\(_2\)O molecules which mediate similar intermolecular interactions in the extended structure. Perhaps the biggest differences between the two compounds, despite the very similar synthetic procedures, is the high symmetry of 2 versus the asymmetry of 1, and the dimerization of wheels in 1 versus the single wheel in 2. The intricacies involved in driving these differences are unknown and will require a larger library of clusters to be synthesised and characterised. Given that 1 and 2 are just the second and third Mn complexes made with LH\(_3\), it would seem likely that many more species await discovery. It also suggests that other homo- and heterometallic 3d and 4f cluster compounds will be readily accessible. A search of the Cambridge structural database reveals that, bar [Mn\(_{16}\)Mn\(^{III}\)\(_{14}\)trz\(_{14}\)L\(_4\)(μ-3-O)\(_8\)(H\(_2\)O)\(_{10}\)]\(^{6+}\), there are no [Mn\(_{16}\)] or [Mn\(_{10}\)] molecules in the literature with similar topologies to 1 and 2.

### Magnetic properties

The direct current (dc) molar magnetic susceptibility, \(\chi\), of freshly prepared polycrystalline samples of 1 and 2 were measured in an applied field, \(B\), of 0.1 T, over the 2–300 K temperature, \(T\) range. The purity of the samples was verified by means of PXRD comparison with the simulated data from the single-crystal structure (Fig. S3†). The experimental results are showed in Fig. 5, in the form of the \(\chi T\) product, where \(\chi = M/B\), and \(M\) is the magnetisation of the sample. At room temperature the \(\chi T\) products of 1 (36.0 cm\(^3\) K mol\(^{-1}\)) and 2 (15.4 cm\(^3\) K mol\(^{-1}\)) are lower than the sum of the Curie constants expected for non-interacting [Mn\(^{III}\)\(_{12}\)Mn\(^{II}\)\(_4\)] (53.5 cm\(^3\) K mol\(^{-1}\)) and [Mn\(^{III}\)\(_{10}\)] (30 cm\(^3\) K mol\(^{-1}\)) units, respectively. As temperature decreases, the \(\chi T\) product for both complexes decreases rapidly and for 2 reaches a value close to 0 cm\(^3\) K mol\(^{-1}\) at \(T = 2\) K, clearly indicative of strong antiferromagnetic exchange and a diamagnetic ground state. For 1, there is a plateau in the value of \(\chi T\) ≈ 24 cm\(^3\) K mol\(^{-1}\) between \(T = 15–25\) K, before it decreases rapidly to a value of 10 cm\(^3\) K mol\(^{-1}\) at \(T = 2\) K. The plateau in \(\chi T\) is suggestive of the presence of competing ferro- and antiferromagnetic interactions which may, or may not, be related to the dimeric nature of the structure. Low-temperature variable-temperature-and-variable-field magnetisation data were measured in the temperature range 2–7 K, in magnetic fields up to 5.0 T (Fig. 6). At the lowest temperature and highest field measured, \(M\) reaches a
cages: a [Mn16] double square wheel and a [Mn10] contorted [Mn3O]n+ triangles dictated by the presence of N,O-chelating square wheel. Both are constructed from corner sharing nuclearity of the two compounds (and the structural chemistry of this ligand with any other paramagnetic 3d or 4f metal ions. There therefore remains much synthetic chemistry to be explored.

Conclusions

The first concerted effort at examining the coordination chemistry of LH3 with Mn has afforded two large and unusual cages: a [Mn16] double square wheel and a [Mn10] contorted square wheel. Both are constructed from corner sharing [Mn3O]n+ triangles dictated by the presence of N,O-chelating L3+ and LH2− ligands, which also direct the JT axes of the MnIII ions along the Mn–N(L) bonds. While 1 describes two linked, offset [MnIII2MnII] wheels, 2 is a single wheel but one in which the upper half is oriented perpendicular to the lower half. Magnetic measurements reveal the presence of strong antiferromagnetic interactions and a diamagnetic ground state in 2 and strong, competing exchange interactions in 1.

The simplicity of the synthetic procedures that produce 1 and 2 suggests that many more Mn coordination compounds constructed with LH3 await discovery. Variation in metal salt, oxidation state, base, solvent, co-ligands, temperature and pressure have proved enormously successful in the coordination chemistry of ethanalamine-based ligands with Mn to date.11–15 Building a library of such species is the first step to understanding what controls the self-assembly process, which, in turn, aids interpretation and exploitation of magneto-structural parameters. We also note that there is no coordination chemistry of this ligand with any other paramagnetic 3d or 4f metal ions.

Conflicts of interest

There are no conflicts to declare.

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