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The remarkable influence of N,O-ligands in the assembly of a bis-calix[4]arene-supported [MnIV2MnIII10MnII8] cluster

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Calix[4]arenes are versatile ligands, capable of supporting the formation of a wide variety of polymetallic clusters comprising 3d, 4f or 3d-4f metal ions. Calixarene-based metal ion fragments act as both bridging and structure capping moieties in these systems, and this behaviour is systematically extended upon moving to bis-calix[4]arene, a relatively new ligand in which two calix[4]arenes are tethered at the methylene bridge position. N,O-ligands greatly influence cluster formation with bis-calix[4]arene, affording a remarkable mixed-valence [MnIV2MnIII10MnII8] cluster that displays coordination chemistry typical of each ligand type, but also new structure capping behaviour for the latter.

The ability to control both the structure and composition of large, multi-component polymetallic clusters of paramagnetic metal ions is a challenging synthetic goal; this has concomitant effects on the resulting magnetic properties of the system. Ligand design has played a crucial role in this regard as it allows one to exert influence through the incorporation of well-known coordination modes.1 Amongst the plethora of ligands available for the synthesis of such high-spin clusters, the utilisation of tripodal supports bearing disparate functionalities represents a logical choice. N,O-Chelates have been widely employed in the synthesis of polymetallic clusters, taking advantage of their common coordination modes. This includes the use of unsubstituted and N-substituted diethanolamines2 to obtain, for example, a heptanuclear, six-membered mixed-valence MnIII10 wheel in which MnII and MnIII cations alternate around a central MnII cation (1, Fig. 1A).2a Close inspection of these cluster topologies suggests that many can be visualised as being constructed from distorted-fused cubanes or, alternatively, fused butterflies.

The latter represents a topology often encountered in our research in which we have used methylene-bridged calix[4]arenes as ligands / cluster supports. These molecules (e.g. p-Bu-calix[4]arene and p-H-calix[4]arene, hereafter collectively termed H2C[4]s) have emerged as versatile platforms for the synthesis of polymetallic clusters, capable of affording a wide variety of different structural topologies depending on the nature of the metal ions present; markedly different structures result from the incorporation of 3d,4f,5 or 3d and 4f metal ions.5 From our work we have established a vast library of C[4]-supported metal clusters in which a recurring structural theme is that [C[4]TM]n, [C[4]TM]2 or [C[4]Ln]n moieties act as bridges to metal ions within the cluster through their phenolate groups, but also as polyhedral capping fragments.6 With respect to the present contribution, particularly noteworthy systems resulting from these studies include a family of [MnIII10MnIV8(C[4])2] single molecule magnets2b,c (SMMs, 2, Fig. 1B) that feature the aforementioned butterfly-like cluster; in this case the oxidation state distribution is the reverse of those reported in the literature for other ligand types.7 A family of [MnIII10LnIII8(C[4])2] clusters was obtained in the presence of both TM and Ln ions, and these can be best described as four [MnIII(C[4])]: moieties capping the edges of a LnIII square (Ln = Gd, Tb Dy).8 Finally, [LnIII8(C[4])2] octahedra were isolated when the latter reaction was carried out in the absence of TM ions.4

It should be noted that the vast majority of the examples of C[4]-supported clusters outlined above were obtained under mild conditions,3b-d,4,5 but similar species can also be obtained using microwave3e,f or solvothermal3h,8 synthesis. Markedly different structural topologies can be obtained when using Schlenk-line conditions, as this generally involves the use of bases that promote the incorporation of additional metal ions (e.g. alkali metals), dramatically affecting the resulting cluster topology.9 Thia-, sulfinyl-, and sulfonylcalix[4]arenes have also been used widely in cluster synthesis, and again, the prevailing coordination chemistry is drastically different to that found when using C[4]s as cluster supports; this is due to the

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d Electronic Supplementary Information (ESI) available: Additional figures relevant to discussion, table of bond lengths and bond valence sum (BVS) calculations. CCDC 1573336. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/S0000000x
incorporation of additional donor atoms in the ligand framework, the result of which is formation of common structural fragments comprising the relevant calixarene and four 3d, 4f or 3d-4f metal ions. A detailed discussion of this vast coordination chemistry is far beyond the scope of this introduction, and for the sake of brevity the reader is directed to recent reviews on the topic, as well as examples of sulfonyl-C[4]-supported manganese clusters.

With the aforementioned coordination chemistry of C[4] in mind, we recently turned our attention to bis-p-Bu-calix[4]arene (BisTBC[C4], H2L1, Fig. 1C), a relatively new ligand in which two H2C[4]s are directly linked via a methylene bridge position. Being conformationally mobile, we envisaged that it would act in a similar manner to H2C[4], but that it would exhibit double structure capping behaviour and thus considerably enhance the reactivity of any resulting polymetallic cluster. We recently reported the formation of a [MnII4MnIII4(L1)2] cluster (3, Fig. 1D) in which the polymetallic core can be thought as two fused and highly distorted [MnII2MnIII2] butterflies. Perhaps the most important feature of this structure is the positions occupied by MnII ions, located in binding pockets between the constituent C[4] lower-rims that are generated by ligand inversion. A remarkably similar topology is encountered in a related [MnII4MnIII4Lnx(L1)2] cluster, obtained via interchange of two MnII with LnIII ions. Control over reactant stoichiometry afforded new clusters of formula [MnII4MnIII4(L1)2] and [MnII5MnIII4Lnx(L1)2] (4, Fig. 1E), and these were obtained via ‘insertion’ of two additional methoxy-bridged MnII ions to the [MnII4MnIII4(L1)2] or [MnII4MnIII3LnIII2(L1)2] cores. Our exploratory chemistry with H2L1 also recently afforded a new family of clusters of general formula [TMnIIxLnIIIy(L1)2]x where the TM was Cu, Fe or Mn, the Ln was Gd, Tb or Dy, and the x:y ratio was 4:0, 5:4, 4:5 or 0:4 (e.g. 5, Fig. 1F). Although these new clusters have rather different topologies, they all share common features that follow established binding rules and structural capping behaviours. In this contribution we report our initial findings from multi-component cluster-forming reactions incorporating an N,O-co-ligand as well as H2L1 in a one-pot synthesis. The remarkable Mn20 cluster formed displays coordination modes that are typical of both ligand types, but also features new structural behaviour for L1 which we attribute to the competitive metal ion binding nature of the co-ligand employed.

Reaction of 1:10:4 equivalents of H2L1 with manganese(II) chloride tetrahydrate and diethanolamine (H2DEA) in a DMF/MeOH mixture (to aid solubility) and in the presence of Et3N (XS) as a base, afforded dark purple single crystals of [MnII8MnII4MnIII4(L1)(DEA)(µ-O)]6(µ- O)(dmf)(Cl)(H2O)](dmf)(MeOH)(H2O)]2 (6, Fig. 2) upon vapour diffusion of Et2O into the mother liquor. The crystals were found to be in a monoclinic cell and structure solution was carried out in the space group C2/c. The asymmetric unit (ASU) comprises half of the reported formula, with ten Mn cations, two of which reside in the tetra-phenolato cavities in perfect accordance with the established binding rules for both C[4] and L1. The remaining eight cations propagate horizontally from L1, forming a polymetallic core supported by DEA molecules and bridging hydroxides/chlorides (Fig. 3A).

![Fig. 1](Image) Single crystal X-ray structures of 1 (A), 2 (B), H2L1 (C), 3 (D), 4 (E), and 5 (F). Colour code C – grey, O – red, N – royal blue, MnII – pale blue, MnIII – purple, CuII – light blue, LnIII – green. Hydrogen atoms (except those involved in lower-rim H-bonding in C), Bu groups of C[4] and H2L1 / L1 (except those shown in C), ligated solvent molecules in D, E and F, and co-crystallised solvent/anions omitted for clarity. Figures not to scale.

![Fig. 2](Image) Asymmetric unit in the single crystal X-ray structure of 6 with selected atoms labelled according to discussion. Colour code C – grey, O – red, N – royal blue, MnII – pale blue, MnIII – purple, MnIV – orange, Cl – dark green. H atoms, ‘Bu groups of L1, ligated solvent and co-crystallised solvent/anions omitted.}

Analysis of the structure shows a variety of coordination modes and oxidation states of the metal centres, the latter being confirmed by BVS calculations (Supporting Information).
Given the complexity of the structure it is necessary to include a detailed description of the coordination environment of the constituent metal ions. As expected, Mn1 is in the 3+ oxidation state and is bound in a C[4] lower-rim pocket described by O1–O4 (Mn–O distances in the range 1.892(4) – 1.967(3) Å). The coordination sphere of Mn1 is completed by a ligated dmf molecule residing in the C[4] cavity (Mn1–O10, 2.240(4) Å) and a µ-oxide (Mn1–O8, 2.141(3) Å) which bridges to Mn3, Mn4 and Mn6 (Mn3–O8, 2.417(4) Å; Mn4–O8, 1.896(3) Å; Mn6–O8, 1.858(4) Å). The Jahn-Teller axis deviates from linearity and is defined by the O8–Mn1–O10 vector (170.79(15)°). Mn2, also in the 3+ oxidation state, has an analogous coordination environment, being bound to the lower-rim phenolic oxygens of one of the C[4] constituents of the L1 octa-anion (Mn2–O range 1.893(4) – 1.974(4) Å), a ligated dmf molecule (Mn2–O11, 2.250(3) Å) and a µ-oxide (Mn2–O2, 2.117(3) Å), the latter of which bridges to Mn3, Mn4 and Mn5 (Mn3–O9, 2.504(4) Å; Mn4–O9, 1.888(3) Å; Mn5–O9, 1.848(4) Å). The Jahn-Teller axis shows similar deviation from linearity and is defined by the O11–Mn2–O9 vector (173.88(15)°). Mn3 is in the 2+ oxidation state, is hepta-coordinate, has distorted octahedral geometry and, in addition to the aforementioned µ-oxide ligands, is also bound to two phenolic oxygens belonging to the C[4] moieties of L1 (Mn3–O1, 2.209(3) Å and Mn3–O7, 2.209(3) Å), a ligated dmf molecule (Mn3–O14, 2.200(3) Å) and two oxygens from different DEA ligands (Mn3–O15, 2.180(3) Å and Mn3–O17, 2.168(3) Å). Mn4 is in the 3+ oxidation state, is penta-coordinate and has a distorted square pyramidal geometry. The base of the pyramid is defined by the O8 and O9 µ-oxides, as well as two µ-oxides (Mn4–O12, 1.866(3) Å and Mn4–O13, 1.866(3) Å), whilst an oxo bridge is located at the apex (Mn4–O16, 2.284(3) Å). Mn5 is in the 3+ oxidation state, has distorted octahedral geometry and is bound to a µ-phenolate moiety (Mn5–O6, 2.232(3) Å), the O9 µ-oxide, a chloride anion (Mn5–Cl1, 2.781(15) Å) that bridges to Mn6 (Mn6–Cl1, 2.827(15) Å) and a tridentate DEA ligand (Mn5–O17, 1.864(3) Å; Mn5–N4, 2.019(4) Å; Mn5–O18, 1.891(4) Å). The Jahn-Teller axis is located along the O6–Mn5–Cl1 vector and shows significant deviation from linearity (165.27(10)°). The coordination spheres of Mn2 (3+) and Mn6 (3+) are nearly identical to those of Mn1 (3+) and Mn5 (3+) respectively, with only negligible differences observed (Table S1); the Jahn-Teller axis for Mn6 is defined by the O2–Mn6–Cl1 vector (164.79(11)°), again showing deviation from linearity. Mn7 is in the 2+ oxidation state, is penta-coordinate and has distorted square pyramidal geometry. It is bound to a phenolic oxygen atom (Mn7–O5, 2.136(4) Å), a µ-oxide (Mn7–O13, 2.028(3) Å), an oxygen of a ligated DEA (Mn7–O18, 2.135(4) Å), a ligated dmf molecule (Mn7–O19, 2.136(4) Å) and a µ-Cl (Mn7–Cl2, 2.521(14) Å) that bridges to Mn9. Mn8, Mn9 and their symmetry equivalent (s.e.) ions form a DEA-supported butterfly located at the centre of the large cluster, acting as a link between the two Mn8 moieties (Fig. 3A). Mn8 is hexa-coordinate, is in a distorted octahedral geometry, and is in the 4+ oxidation state. Although not directly interacting with any calixarene moiety, this represents the first example in which a Mn4+ is incorporated in an L1-supported cluster, along with frequently observed MnIII/O ions. Mn9 is bound to three µ-oxides (Mn9–O12, 1.820(3) Å; Mn9–O13, 1.828(3) Å; Mn9–O20, 1.917(3) Å) and a ligated DEA molecule (Mn9–O21, 1.925(4) Å; Mn9–N7, 2.041(4) Å; Mn9–O22, 1.923(3) Å). Mn9 is in the 2+ oxidation state, has distorted octahedral geometry, and is bound to two µ-oxides (Mn9–O20, 2.239(3) Å and Mn9–O22, 2.254(3) Å), two oxygens from DEA ligands (Mn9–O21, 2.110(4) Å and Mn9–O22, 2.109(4) Å) and two µ-chlorides (Mn9–Cl2, 2.497(14) Å and Mn9–Cl3, 2.492(14) Å) that bridge to Mn7 and Mn10 (Mn7–Cl2 distance given above, Mn10–Cl3, 2.531(15) Å). Finally, Mn10 is in the 2+ oxidation state, has distorted trigonal bipyramidal geometry, and is bound to the µ-chloride, a µ-oxide (Mn10–O12’, 2.029(3) Å), an oxygen atom of a DEA ligand (Mn10–O25, 2.152(4) Å), a phenolic oxygen (Mn10–O23, 2.122(4) Å) and a ligated dmf molecule (Mn10–O24, 2.126(4) Å).

Symmetry expansion of the ASU affords the entire cluster as shown in Figure 3A, which can be visualised as two MnIII/O3 moieties linked via the central DEA-supported MnIII/O2 fragment. This fragment displays coordination modes that are typical for DEA and compound 6 can be considered an excellent example in which the nuclearity of clusters supported by one ligand type can be augmented (or greatly enhanced) via the incorporation of complementary and / or competing co-ligands; the complex topology of 6 (Fig. 3B) displays several recurring coordination modes typical for both L1-10 and DEA-supported clusters (Fig. S1). Capping behaviour is observed with four [MnIII(C[4])2] moieties (Mn1,
Mn2 and s.e.) ‘encapsulating’ one MnIV cation (Mn3) located in one of the binding pockets generated upon inversion of L1, whereas the other binding site remains unoccupied. This is unprecedented, and we propose that the strongly competing nature of DEA gives rise to this new structural feature; the presence of two additional DEA-supported MnIV cations (Mn5 and Mn6) appears to create steric hindrance in that region of the cluster, preventing it from being occupied by a metal ion. This also reflects on the tetrahedral geometry of Mn4, which is unusual and unprecedented in L1-supported cluster chemistry thus far. A CSD search for Mn20 clusters returned 18 hits with disparate topologies, the metallic cores of which are supported by a variety of ligands including phosphonate,18 triethanolamine,19 pivalate20 and other N-containing polydentates.21 These clusters possess Mn ions in oxidation states between 1+ and 3+ in either mixed or non-mixed valence arrangements, but none were found to contain MnIV ions. The topology of 6 is thus found to be unique, as is the oxidation state distribution and mixed-valence arrangement. Inspection of the expanded structure of 6 shows that neighbouring clusters are well-isolated, with no significant intermolecular interactions being observed between symmetry equivalents; the shortest metal-metal inter-cluster distance is found between Mn1 and Mn7 ~12.1 Å along the crystallographic b axis (Fig. S2).

The dc (direct current) molar magnetic susceptibility, \( \chi_M \), of a freshly filtered polycrystalline sample (Euan???) of 6 was measured in an applied magnetic field, \( B \), of 0.1 T, over the \( T = 5-300 \) K temperature range. The experimental results are shown in Figure S3 in the form of the \( \chi_M T \) product versus \( T \), where \( \chi_M = M / B \), and \( M \) is the magnetisation of the sample. At room temperature, the \( \chi_M T \) product is approximately 68 cm³ K mol⁻¹, in good agreement with the sum of Curie constants for a [MnIV]₅MnII₉MnIII₄] unit (68.75 cm³ K mol⁻¹, \( g = 2.0 \)). Upon cooling, the \( \chi_M T \) product decreases to a minimum value of 53.4 cm³ K mol⁻¹ at \( T = 34 \) K, then rises to a value of 58.3 cm³ K mol⁻¹ at \( T = 8 \) K, before decreasing again to a value of 57.5 cm³ K mol⁻¹ at \( T = 5 \) K. This behaviour is consistent with competing (weak) ferromagnetic and antiferromagnetic exchange interactions between the Mn centres. The decrease in \( \chi_M T \) below 8 K can be attributed to zero-field splitting effects and / or the presence of antiferromagnetic inter-molecular interactions. The large nuclearity and complex topology of the [Mn20] core prevents any quantitative analysis of the exchange constants, but the overall behaviour is entirely consistent with that observed for other C[n]-constructed, O-atom bridged, heterovalent Mn cages, where typical coupling constants are of the order \(| J | \leq 10 \) cm⁻¹.22 Note also that \(| J | \) is likely of the same order of magnitude as \(| D_{\text{Dipole}} \). The low temperature variable-temperature-and-variable-field (VTVB) magnetisation data collected in fields of up to \( B = 7 \) T (Fig. S3, inset) are also consistent with this picture, showing \( M \) increasing in a near linear fashion with \( B \), without saturating (\( M = 44.7 \mu_B \) at \( T = 2 \) K and \( B = 7 \) T). Complex 6 does not display any frequency-dependent signals in out-of-phase (\( \chi'' \)) susceptibility measurements, and thus is not an SMM.

In conclusion, we have shown that the use of BisTBC[4] with DEA as a co-ligand leads to the formation of a very high nuclearity mixed-valence Mn cluster that displays typical coordination modes relative to each ligand type. The competitive metal ion binding of DEA appears to influence that of BisTBC[4], and this delicate balance in ligand composition may be a route towards unearthing a large library of such cluster species in C[n] chemistry, whilst simultaneously increasing / enhancing metal ion composition. This strategy may also allow one to control the incorporation of metal ions with coordination numbers and oxidation states that are otherwise unobtainable using just BisTBC[4] as a ligand; this may also apply to all C[n] ligands. This will have concomitant effects on the magnetic properties for these multi-component systems, the results of which will be reported in due course.

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Notes and references

HaL1 was synthesised according to literature procedure.12 1°Synthesis of [MnIV₅MnII₉MnIII₄]L1)[(DEA)₂(μ-Cl)(μ-Cl)(μ-Cl)(μ-Cl)I][Cl] (M = 5659): C, 50.3%; H, 6.23%; N, 4.46%. Found: C, 49.97%; H, 5.89%; N, 4.16%. Yield 404 mg (18.3 %).

Crystal Data for 6 (CCDC 1573336): C₂₃H₂₅NO₂S₆Cl₆, \( M = 5655.83 \) g/mol, monoclinic, space group C2/c (no. 15), a = 18.395(7) \( \AA \), b = 45.6912(18) \( \AA \), c = 40.7908(16) \( \AA \), \( \beta = 99.212(2) \), \( V = 3843(3) \) \( \AA^3 \), \( Z = 4 \), \( T = 100(2) \) K, 16355 reflections measured (4.198° ≤ 2Θ ≤ 58.238°), 34860 unique \( (R_{int} = 0.0610, R_{free} = 0.0498) \) which were used in all calculations. The final \( R_1 \) was 0.0828 \( (I > 2\sigma(I)) \) and \( wR_2 \) was 0.2223 (all data).


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17 Crystals grew most readily from a 1:10:4 ratio of the reactants as described in the text, this being achieved through a combinatorial screen.


