Structural and magnetic variations in tetranuclear Ni\textsuperscript{II} clusters: the effect of the reaction solvent and ligand substitution on product identity\textsuperscript{†}

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Three structurally and magnetically different tetranuclear Ni\textsuperscript{II} complexes have been isolated and magnetically characterized, emphasizing the effect of the reaction solvent and organic ligand substitution on the chemical identity of cluster compounds.

The synthesis of new polynuclear 3d-metal complexes (coordination clusters or simply clusters) is definitely one of the most fascinating areas of inorganic and coordination chemistry.\textsuperscript{1} The interest mainly stems from the relevance of molecular cluster compounds in bioinorganic chemistry, catalysis, optics and magnetism.\textsuperscript{2} Numerous synthetic strategies and approaches have been developed over the last three decades for the construction of structurally and physically novel 3d-metal clusters.\textsuperscript{3} To this end, the self-assembly route has been proved to be the most fruitful, yielding high-nuclearity compounds with beautiful mosaics and impressive magnetic properties such as high-spin molecules,\textsuperscript{4} single-molecule magnets (SMMs)\textsuperscript{5} and magnetic refrigerants\textsuperscript{6} for low-temperature cooling applications.

In general lines, self-assembly refers to the unpredictable, one-pot reaction of a metal ion precursor with a single bridging/chelating organic ligand or a combination of more than one such ligands. The unpredictable nature of this approach arises from the coordination affinity and versatility of the bridging organic ligand(s) and the flexibility of the 3d-metal ion toward various coordination geometries and topologies.\textsuperscript{3} Hence, the choice of the polydentate organic bridging/chelating ligand is an important task, albeit not the only significant

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Since then, we have been seeking new routes to NiII cluster compounds and we thus turned our attention to ring-substituted derivatives of saphH₂ with both donor and non-donor groups. We had initially decided to replace the H-atom at position 4 of the ω-aminophenol moiety with a methyl group, aiming at the positive effect of the –CH₃ functionality on the crystallization of the resulting products. We then attempted to replace the –OH donor group of the ω-aminophenol moiety with a carboxylate (–COOH) functionality which could potentially coordinate to more metal centers than saphH₂ through the two O donor atoms and therefore foster formation of very high nuclearity metal species. In addition, we included a non-donor chloro group in place of a phenyl H atom at position 5 which could, in principle, differentiate the electronic and steric properties, and hydrogen bonding effects. The resulting ligands N-salicylidene-4-methyl-ω-aminophenol (samphH₂, Scheme 1) and N-salicylidene-2-amino-5-chlorobenzoic acid (sacbH₂, Scheme 1) have a limited previous use in metal cluster chemistry, restricted only to a recently reported Ni₁₁ compound featuring both Bu₃CH₂CO₂⁻ and sacb²⁻ groups. In the present work, we show how the thorough investigation of a reaction system can lead to three Ni₄ compounds with different structures and magnetic properties, emphasizing the effect of two synthetic variables on the products’ identities.

The reaction of the [Ni₂(H₂O)(O₂CBu')₄(HO₂CBu')₂]₄⁻ precursor and samphH₂ in a 1:2 molar ratio in EtOH gave a brown solution that upon slow evaporation at room temperature gave olive-green crystals of [Ni₂(samph)₂(EOH)₂]₂·0.7EOH (1·0·7EOH) in 35% yield.† Complex 1·0·7EOH crystallizes in the triclinic space group P1 and has a virtual S₃ symmetry. Its structure consists of four distorted octahedral NiII ions bridged by four deprotonated μ₁-O atoms from the methyl-aminophenol part of four samph²⁻ ligands (Fig. 1); the latter all bind in an μ₁:μ₂:μ₃:μ₄ mode (Scheme 1). The tetranuclear cluster molecule has a distorted cubane [Ni₄(μ₁-OR)₄]²⁺ core (Fig. S1†) with the μ₁-O atoms occupying alternate vertices of the cube. Thus, the molecule consists of two interpenetrating concentric tetrahedra, one having four metal ions and the other, four triply bridging oxygen atoms. Peripheral ligation about the core is provided for by four terminal EtOH molecules each lying on a NiII ion. The six faces of the cubane are not equivalent. Using the notation introduced by Williams, Decurtins and coworkers for cubanes,¹⁰ we code 1 as a whole by Ni₄/2s₄/m₄. The symbol “s” indicates that the bridging atoms of samph²⁻ are supported, the number “2” designating that one arm of samph²⁻ links the bridging O atom to two terminal donor atoms (the imino N and phenolate O atoms); the abbreviation “m₄” indicates that four monodentate, terminal ligands (EtOH groups) complete the coordination spheres of the four NiII ions.

Within the concept of chemical reactivity on cluster compounds, an important synthetic factor that is worth investigating is the effect of the solvent on the structural identity of the complexes. The volatility, polarity, rigidity and coordination affinity of reaction solvents are some of the features which directly affect the identity of a product. To that end, the reaction that led to complex 1 has been repeated in DMF (instead of EtOH) under exactly the same conditions. The resulting dark orange solution afforded orange crystals of a new [Ni₄(samph)₂(DMF)₂]·2DMF (2·2DMF) compound in the yield of 40%.† Complex 2·2DMF crystallizes in the triclinic space group P1. Its structure consists of centrosymmetric [Ni₄(samph)₂(DMF)₂]₂ (Fig. 2) and solvate DMF molecules; the latter are not further discussed. The four NiII ions are located at the four vertices of a defective dicubane, i.e. two cubanes sharing a face [Ni(2O3Ni(2′O3′)] and each missing one metal vertex. The NiII centers are bridged by two μ₁-O (O₃, O₃′) and four μ-O (O₁, O₁′, O₂, O₂′) atoms from two μ₁:μ₂:μ₃:μ₄ and two μ₁:μ₁:μ₂:μ₃ samph²⁻ ligands, respectively (Scheme 1). Thus, the core is {Ni₄(μ₁-OR)₂(μ₂-OR)₂}²⁺ (Fig. S1†), where RO⁻ = samph²⁻. Peripheral ligation is provided by two terminally bound DMF molecules on Ni2 and Ni2′ and the imino N

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**Fig. 1** Labelled plot of complex 1 emphasizing the (Ni₄(μ₁-OR)₄)²⁺ cubane core. Color scheme: NiII green, O red, N blue, C gray. H-atoms are omitted for clarity.

**Fig. 2** Labelled plot of complex 2 emphasizing the (Ni₄(μ₁-OR)₂(μ₂-OR)₂)²⁺ defective dicubane core. Color scheme as in Fig. 1. H-atoms are omitted for clarity. Symmetry operation for the primed atoms: 1 – x, 2 – y, –z.
atoms of four samph\textsuperscript{2−} ligands, each bound to a Ni\textsuperscript{II} ion. As a result, the latter atoms are six-coordinate and near-octahedral, whereas Ni1 and Ni1′ are five-coordinate with highly distorted coordination geometries (τ = 0.43, where τ = 0 and 1 for perfect square-pyramidal and trigonal-bipyramidal geometries\textsuperscript{11} respectively).

Complex 2 can alternatively be described as containing a butterfly of Ni\textsuperscript{II} sites with the ‘body’ of the butterfly (Ni2 and Ni2′) bridged by two deprotonated μ\textsubscript{3}-O atoms from two η\textsuperscript{1}:η\textsuperscript{1}:μ\textsubscript{3} samph\textsuperscript{2−} ligands; these O atoms also bridge to Ni\textsuperscript{II} ‘wing-tip’ sites (Ni1 and Ni1′). Each of the four edges of the closed-type butterfly is bridged by one μ−O atom from two in total η\textsuperscript{2}:η\textsuperscript{2}:μ\textsubscript{3} samph\textsuperscript{2−} ligands. In the crystal structures of both 1 and 2 there are no significant intermolecular H-bonding or π−π stacking interactions. There are four intra-molecular H-bonds in 1 (Fig. S2, Table S5\textsuperscript{†}). 1 and 2 are the first cluster compounds bearing any form (neutral, singly- or doubly-deprotonated) of samphH\textsubscript{2}.

The interesting behavior of the Ni\textsuperscript{II}/samphH\textsubscript{2} system prompted us to replace the samphH\textsubscript{2} Schiff base ligand with the sacbh\textsubscript{2} ligand that is closely related, but possessing enhanced coordination capabilities. Thus, the 1:2 reaction of [Ni\textsubscript{2}(H\textsubscript{2}O)(O\textsubscript{2}CBu\textsubscript{3})\textsubscript{4}(HO\textsubscript{2}CBu\textsubscript{3})\textsubscript{4}] and sacbh\textsubscript{2} in EtOH led to a dark green solution from which were isolated green crystals of the Ni\textsubscript{4} compound in yields as high as 60%. Complex 3-6EtOH crystallizes in the triclinic space group PI with the Ni\textsubscript{4} molecule lying on an inversion center located at the middle-point of the Ni2−Ni2′ vector (Fig. 3). The Ni\textsubscript{4} compound comprises a non-linear, zigzag array of four Ni\textsuperscript{II} atoms (Ni1−Ni2−Ni2′ = 108.56°) with each Ni\textsubscript{3} pair bridged by two μ-O atoms from the deprotonated alkoxido (O1, O1′, O11, O11′) and/or carboxylate (O12, O12′) functionalities of four sacb\textsuperscript{2−} ligands. The latter are of two types (Scheme 1): two are binding in an η\textsuperscript{2}:η\textsuperscript{1}:μ−μ mode, acting as Ni\textsubscript{2}O,O-tridentate chelates to a Ni\textsuperscript{II} atom (Ni1, Ni1′) and simultaneously bridging an additional Ni\textsuperscript{II} atom (Ni2 or Ni2′) through the phenolate O atom, and the remaining two adopt the more complex η\textsuperscript{2}:η\textsuperscript{1}:η\textsuperscript{1}:μ\textsubscript{4} binding mode, chelating the central Ni\textsuperscript{II} atoms and bridging all four metal ions through the phenolate and both carboxylate O atoms. Thus, the core of 3 is \{Ni\textsubscript{3}(μ−OR)\textsubscript{6}\}\textsuperscript{2+} (Fig. S1†), with peripheral ligation provided by four terminal EtOH molecules. The central Ni2−O12−Ni2′−O1′ rhombus is strictly planar as a result of the inversion center, but the other two rhombus are quite distorted, with the Ni1−O1−Ni2−O11 torsion angle being 17.1°. The Ni\textsuperscript{II} atoms are all six-coordinate with distorted octahedral geometries. Finally, in the crystal structure of 3-6EtOH there are H-bonding interactions between the carboxylate and phenolate moieties of sacb\textsuperscript{2−}, and the bound EtOH and lattice EtOH solvate molecules (Fig. S3, Table S6†). In addition, weak π−π stacking interactions between the aromatic rings of sacb\textsuperscript{2−} ligands serve to link the Ni\textsubscript{4} clusters into a 2-D network (Fig. S4†). Although there are hundreds of tetranuclear Ni\textsuperscript{II} complexes structurally and magnetically characterized, compound 3 is only the fourth example of a Ni\textsubscript{4} cluster with a zigzag topology\textsuperscript{12}.

Variable-temperature dc magnetic susceptibility measurements were performed on freshly-prepared microcrystalline solids of 1, 2-DMF, and 3-2EtOH in the temperature range 2–300 K; a dc field of 0.3 T was applied from 30 to 300 K and a weak dc field of 0.03 T was applied from 2 to 30 K to avoid saturation effects. The data are shown as \(\chi_M T\) versus \(T\) plots in Fig. 4. The values of the \(\chi_M T\) product at 300 K are 5.53 \(\pm 1\) (1), 5.35 (2-DMF) and 5.80 (3-2EtOH) cm\textsuperscript{3} mol\textsuperscript{−1} K, slightly higher than the value of 4.84 cm\textsuperscript{3} mol\textsuperscript{−1} K (calculated with \(γ = 2.2\)) expected for four non-interacting Ni\textsuperscript{II} (\(S = 1\)) atoms. For the cubane and zigzag complexes, 1 and 3-2EtOH, the value of \(\chi_M T\) steadily increases in the 300−7 K and 300−17 K ranges, respectively, and then slightly (for 1) or more rapidly (for 3-2EtOH) decreases with decreasing \(T\) to reach the values of 6.68 (1) and 4.40 (3-2EtOH) cm\textsuperscript{3} mol\textsuperscript{−1} K at 2 K. For the defective dicubane complex 2-DMF the magnetic response is distinctly different; the \(\chi_M T\) product remains essentially constant in the 300−50 K region and then decreases sharply, reaching a value of

![Fig. 3](image-url) Partially-labelled representation of the structure of complex 3 emphasizing the \{Ni\textsubscript{3}(μ−OR)\textsubscript{6}\}\textsuperscript{2+} zigzag core. Color scheme as in Fig. 1; Cl cyan. H-atoms are omitted for clarity. Symmetry operation for the primed atoms: 1 − x, 1 − y, −z.

![Fig. 4](image-url) \(\chi_M T\) versus \(T\) plots of 1, 2-DMF, and 3-2EtOH. The solid lines are the fits of the data; see the text for the fit parameters. (inset) J-coupling scheme employed for the elucidation of the magnetic exchange interactions in 1 (a) and 2/3 (b); see the text for the corresponding spin-Hamiltonians.
0.53 cm$^3$ mol$^{-1}$ K at 2 K. The data and shape of plots indicate predominant ferromagnetic (for 1 and 3-EtOH) and antiferromagnetic (2-DMF) exchange interactions, thus suggesting $S = 4$ and $S = 0$ ground state spin values, respectively.

In order to quantify the nature of the magnetic exchange within 1–3, theoretical expressions of $\chi_M$ as a function of $T$ were derived and fitted to the experimental data. Structural data indicate that the core of complex 1 consists of a distorted cubane with different metric parameters in the six faces. However, opposite faces show similar Ni–O–Ni bond angles and thus, in order to reduce the number of coupling constants in the fitting process, only three coupling constants were employed according to the simplified coupling scheme shown in Fig. 4a (inset). The centrosymmetric complex 2 shows three sets of Ni–O–Ni bond angles and hence three coupling constants were applied according to Fig. 4b (inset). Complex 3 exhibits a different zigzag structure, but magnetically it can be reduced to the same coupling scheme as 2 (vide infra). On the basis of Fig. 4 (inset), the applied Heisenberg spin-Hamiltonians for complexes 1 and 2/3 are shown in eqn (1) and (2), respectively.

$$\mathcal{H} = -J_1(\hat{S}_1 \cdot \hat{S}_2 + \hat{S}_3 \cdot \hat{S}_4) - J_2(\hat{S}_1 + \hat{S}_2 \cdot \hat{S}_4) - J_3(\hat{S}_1 \cdot \hat{S}_2 + \hat{S}_3 \cdot \hat{S}_4)$$

(1)

$$\mathcal{H} = -J_1(\hat{S}_1 \cdot \hat{S}_2) - J_2(\hat{S}_1 + \hat{S}_2 \cdot \hat{S}_3) - J_3(\hat{S}_1 \cdot \hat{S}_3 + \hat{S}_2 \cdot \hat{S}_4)$$

(2)

The PHI program was used to fit the susceptibility and magnetization data. The quality of the fits is quantified by the agreement factor $R$, defined as $(\chi_M \text{calc} - \chi_M \text{exp})^2/(\chi_M \text{exp})^2$. Preliminary fits for 1, applying the spin-Hamiltonian of eqn (1), were not satisfactory and therefore the zero-field splitting parameter, $D_{on}$, was included in the model. An excellent fit of the experimental data gave the following as best-fit parameters: $J_1 = -0.1$ cm$^{-1}$, $J_2 = +8.4$ cm$^{-1}$, $J_3 = +1.3$ cm$^{-1}$, $D = +3.8$ cm$^{-1}$ and $g = 2.28$ (R = 7.2 x 10$^{-5}$) for 1; $J_1 = -10.0$ cm$^{-1}$, $J_2 = +14.9$ cm$^{-1}$, $J_3 = +10.4$ cm$^{-1}$ and $g = 2.30$ (R = 7.3 x 10$^{-5}$) for 2; and $J_1 = +12.5$ cm$^{-1}$, $J_2 = -5.2$ cm$^{-1}$, $J_3 = +1.8$ cm$^{-1}$ and $g = 2.35$ (R = 6.7 x 10$^{-5}$) for 3.

Magnetization versus field measurements were performed at 2 K and the corresponding plots (Fig. S5†) showed a continuous increase up to a non-saturated value equivalent to 7.0 and 7.2 electrons for 1 and 3, respectively, and 3.6 electrons for 2. These values are in agreement with the predominant ferro- and antiferromagnetic interactions observed in susceptibility measurements. To evaluate the accuracy of the susceptibility data, the magnetization data were also fitted using the spin-Hamiltonians expressed in eqn (1) and (2), and including the $D_{on}$ term. Best-fit parameters are: $J_1 = 0.0$ cm$^{-1}$, $J_2 = +9.4$ cm$^{-1}$, $J_3 = +2.4$ cm$^{-1}$, $D = +4.7$ cm$^{-1}$ and $g = 2.24$ (R = 1.5 x 10$^{-5}$) for 1; $J_1 = -9.1$ cm$^{-1}$, $J_2 = +15.0$ cm$^{-1}$, $J_3 = +10.8$ cm$^{-1}$, $D = -0.98$ cm$^{-1}$ and $g = 2.27$ (R = 8.1 x 10$^{-6}$) for 2; and $J_1 = +10.4$ cm$^{-1}$, $J_2 = -3.2$ cm$^{-1}$, $J_3 = +1.6$ cm$^{-1}$, $D = +3.9$ cm$^{-1}$ and $g = 2.34$ (R = 1.1 x 10$^{-5}$) for 3, in good agreement with the susceptibility data. Both measurements confirmed an $S = 4$ ground state for 1, an $S = 0$ ground state (population 80.3%) for 2, with very close in energy $S = 1$ (gap 2.0 cm$^{-1}$, population 19.0%) and $S = 2$ (gap 6.6 cm$^{-1}$, population 0.7%) excited states, and a quasi-frustrated situation for 3 with a mixture of all possible spin states ($S = 4$ to 0) being very close in energy and almost equally populated at 2 K (see the ESI†). The $S = 4$ spin state for 3 is still not fully saturated under a 5 T field, but it tends to saturate to 8 electrons at higher fields.

Rationalization of the magnetic interactions in zigzag Ni$_4$ compounds is unprecedented and thus urged us to further investigate the quasi-frustrated complex 3. The $J_3$ coupling constant, which is associated with the next-nearest neighbor interactions of Ni1⋯Ni2‘ and Ni1’⋯Ni2 mediated by the carboxylate functionality of sabc$^2$, is undoubtedly necessary and cannot be ignored in order to avoid overparametrization effects. If $J_3$ was discarded from the fitting model, the ground state would be zero as a result of two external ferromagnetic Ni2 pairs and a central, antiferromagnetically coupled Ni2 pair; this is totally against the experimental susceptibility and magnetization data. Finally, none of the complexes show out-of-phase ac magnetic susceptibility signals down to 1.8 K, suggesting that these are not SMMs.

The most important parameter in the magnetostructural correlations of tetranuclear clusters possessing the [Ni$_4$(µ$_3$-OR)$_4$]$^{++}$ cubane or [Ni$_4$(µ$_3$-OR)$_3$(µ-OR)$_4$]$^{++}$ defective dicubane cores has been reported to be the average Ni–O–Ni angle of the cubane faces.$^{10,11}$ A ferromagnetic exchange is expected for Ni–O–Ni angles lower than 99°, and the positive coupling constant value increases as the angle decreases. On the other hand, Ni–O–Ni angles in the vicinity of, and larger than, 99° lead to antiferromagnetic interactions, and the absolute value increases as the angle increases. The mean Ni–O–Ni angles for the three Ni$_4$ pairs described by $J_1$, $J_2$ and $J_3$ in 1 are 101.6°, 94.4° and 95.5°, respectively; thus, the $J_2$ coupling is predicted to be the most ferromagnetic, and indeed, it is. Clusters 2 and 3 possess diamagnetic and quasi-frustrated ground states, respectively, as a result of the highly competing ferromagnetic and antiferromagnetic interactions. The obtained signs of $J$ constants for 2 agree with the corresponding Ni–O–Ni angles ($J_1$ for the Ni2⋯Ni2’ pair of a 99.4° mean angle, $J_2$ for the Ni1⋯Ni2 and Ni1’⋯Ni2’ pairs of a 97.5° mean angle, and $J_3$ for the Ni1⋯Ni2’ and Ni1’⋯Ni2 pairs of a 97.8° mean angle). The fact that the absolute values of $J$ constants are very close to each other is reasonably attributed to the Ni–O–Ni angles which are close to the ferro-/antiferromagnetic border. In the case of the zigzag complex 3, $J_1$ and $J_2$ constants were employed for the two outer (Ni1–Ni2 and Ni1’–Ni2’) and one central (Ni2–Ni2’) interaction with average Ni–O–Ni angles of 93.8° and 100.2°, respectively. Both $J_1$ and $J_2$ were reasonably found to be ferromagnetic and antiferromagnetic, respectively, according to the calculated mean angles.

**Conclusions**

In conclusion, we have reported three tetranuclear NiII complexes with cubane (1), defective dicubane (2) and rare zigzag
(3) topologies, all resulting from similar one-pot reactions of a Ni^{II} precursor with structurally related Schiff base ligands. Besides the structural differences, complexes 1–3 also show variations in their magnetic behaviour; 1 is ferromagnetically coupled with an $S = 4$ ground state opposite to cluster 2 which possesses a diamagnetic ground state, and different from 3 which is a quasi-frustrated magnetic system with a mixture of spin states populated at very low temperatures. The key point of this work is the need to insist on the examination and trial of as many synthetic variables as possible in a given reaction system when seeking ways to isolate new polynuclear metal complexes. The search for completely new ligand types is not the only route to obtain structurally unique compounds; re-consideration and thorough investigation of various synthetic factors such as the reaction solvent and ligand substituent(s) can also offer alternative pathways for the emergence of beautiful and magnetically interesting 3d-metal clusters.

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Notes and references


