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In search of molecules displaying ferromagnetic exchange: multiple-decker $Ni₁₂$ and $Ni₁₆$ complexes from the use of pyridine-2-amidoxime†

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The use of pyridine-2-amidoxime (pyaoxH2) in Ni chemistry has provided access to a dodecanuclear complex and a hexadecanuclear Ni cluster, namely $[Ni_{12}(pyaox)_{6}(pyaoxH)_{6}(MeOH)_{2}Cl_{2}]Cl_{4}$ -5MeOH $(1.5MeOH)$ and $[Ni_{16}(pyaox)_{8}(pyaoxH)_{8}(MeOH)_{4}](SO₄)_{4}.10H₂O²6MeOH$ (2:10H₂O·26MeOH). Complex 1-5MeOH was isolated by the reaction of NiCl₂·6H₂O, pyaoxH₂ and NaOMe in a 1:1:2 molar ratio in MeOH in 60% yield. Treatment of NiSO₄·6H₂O with pyaoxH₂ and NEt₃ in a 1:1:2 molar ratio in MeOH afforded 2·10H₂O·26MeOH in good yield (65%). The two compounds display a multi-decker configuration based on stacked Ni₄ layers, {Ni₄(pyaox)₂(pyaoxH)₂)²⁺_x (x = 3, **1**·5MeOH; x = 4, **2**·10H₂O·26MeOH); each deck consists of two square planar and two octahedral Ni^{II} centres. The number of decks observed in 1.5MeOH and 2.10H₂O.26MeOH depends on the nature of the inorganic anion that is present in the reaction system, which provides elements of synthetic control towards new high nuclearity Ni^{II} species. 2.10H₂O·26MeOH is the first structurally characterized complex of any metal displaying a quadrupledecker configuration, being also the highest nuclearity metal cluster bearing pyaoxH₂ and the highest nuclearity Ni^{II} cluster with any type of 2-pyridyl oxime. Each cluster cation displays ferromagnetic exchange between the octahedral Ni^{II} ions resulting in a spin ground state of $S = 6$ for 1 and $S = 8$ for 2. Magnetothermal studies have been performed and discussed for both clusters. PAPER

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Introduction

It has been over 20 years since the discovery that the discrete $\left[\text{Mn}_{8}^{\text{III}}\text{Mn}_{4}^{\text{IV}}\text{O}_{12}(\text{O}_{2}\text{CMe})_{16}(\text{H}_{2}\text{O})_{4}\right]$ molecule could possess an energy barrier to its spin vector reversal that was large enough to observe magnetic hysteresis, the classical property of a magnet.¹ This discovery constituted the commencement of a

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new era in transition metal cluster chemistry, sparking off intense interest in the synthesis and characterization of new coordination compounds that could behave like magnets at very low temperatures in the absence of an applied magnetic field. Such species are now known as single-molecule magnets $(SMMs)^2$ with their properties originating from intrinsic structural and electronic characteristics of their metal cores. The potential applications of SMMs range from high-density information storage, molecular spintronics, to qubits for quantum computation.3–⁵ Furthermore, several other quantum mechanical phenomena have been identified in SMMs, such as spinphonon coupling,⁶ spin state entanglement,⁵ spin parity effects, 7 both thermally assisted and pure quantum tunneling of the magnetization (QTM) ,⁸ and quantum phase interference.^{7b,9}

Clusters containing Mn ^{III} ions have proven to be the most fruitful source of SMMs to date. This is not surprising considering that: (1) the first and most well-known SMM comes from Mn cluster chemistry, a fact that activated intense research in this area, and (2) the d^4 Mn^{III} ion displays significant anisotropy, an inherent requirement for the generation of new SMMs.2 However, SMMs are also known for 3d metal ions

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[†]Electronic supplementary information (ESI) available: Tables with bond distances and angles, hydrogen bonding details, magnetism and heat capacity plots for 2·10H₂O·26MeOH. CCDC 678537 and 1502969. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6dt03511f

other than Mn, e.g. V^{III} , Fe^{II}, Fe^{III}, Ni^{II}, lanthanide ions, actinide ions, and combinations of 3d with 4d, 5d or 4f paramagnetic metal ions.^{2,10} Among them, Ni^{II} has shown much promise in the synthesis of both $SMMs¹¹$ and spin phonon traps, 12 due to its significant single-ion anisotropy and its paramagnetic nature when confined within a highly symmetric cage.

The isolation of new SMMs remains a hot research field, not only for a better understanding of the phenomenon but also for bringing these species closer to technological applications. In fact, the chances of identifying novel types of coordination clusters with improved or new magnetic properties will be increased by the development of new reaction systems with suitable organic ligands or combinations of organic and inorganic ligands. The organic ligand often dictates the cluster symmetry, metal topology and the number of metal ions that are present, which in turn affects the nature of the magnetic exchange interactions and the intermolecular interactions between the polynuclear molecules in the crystal. One fertile approach towards the synthesis of new polynuclear metal complexes with interesting magnetic properties is the use of oxime ligands, since oximates are good bridging groups fostering formation of coordination clusters. 13 We and others have been investigating a number of oxime-based ligands, and one broad family of the latter has been the 2-pyridyl oximes (Scheme 1). 14 The anions of 2-pyridyl oximes are versatile ligands for a variety of research objectives; the activation of 2-pyridyl oximes by 3d-metal centres towards further reaction is also becoming an exciting area of research. 15 Such ligands have been key "players" in several areas of single-molecule and single-chain magnetism.¹⁶ Poper

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Restricting further discussion to nickel, our groups have been engaged in the use of 2-pyridyl oximate ligands for Ni^H cluster synthesis for many years. The first step of this approach included the synthesis and characterization of complexes of pao⁻,¹⁷ mpko⁻¹⁸ and ppko⁻,^{17e,18c,d,19} *i.e.*, complexes bearing anionic ligands in which R contains no additional donor site. In a second step, we have been studying Ni^H complexes with anionic ligands in which R contains a donor site, i.e., clusters of (py)pko^{-17a,20} and pyaoxH⁻/pyaox²⁻²¹ In particular, the ligand pyaoxH₂ (pyridine-2-amidoxime; IUPAC name: N-hydroxy-pyridine-2-carboxamidine) belongs to the class of amidoximes. Although the versatile chemistry of conventional oximes has been extensively studied over the years in both the organic chemistry and coordination chemistry literature,

Scheme 1 General structural formula and abbreviations of the 2-pyridyl oximes discussed in the text.

amidoximes, $RC(NH₂)$ = NOH, have so far received substantially less attention.²² The presence of the amino functionality is expected – due to its coordination capability, potential for deprotonation and hydrogen bonding effects – to alter the coordination behaviour of this ligand (and hence the identity of the resultant Ni^{II} complexes) in comparison with that of the pao−, mpko−, ppko−, (py)pko−, etc., ligands.

In 2008, our groups employed pyaox H_2 in Ni cluster chemistry and proved that it can be a fruitful method for the synthesis of high nuclearity species. 21 Nevertheless, the use of pyaoxH2 for the synthesis of polynuclear Ni compounds remains limited. $21,23-25$ In particular, so far there have been reports of only salts of the cations $\left[Ni_4(pyaox)_2(pyaoxH)_2L_4\right]^{2+}$ (L = imidazole, 1-methylimidazole, pyridine, 3-methylpyridine, 4-methylpyridine),^{23,24} [Ni₈(pyaox)₄(pyaoxH)₄L'₄] (L' = H₂O, imidazole),^{23,24} [Ni₁₂(pyaox)₆(pyaoxH)₆(H₂O)₂(N₃)₂¹⁴ and $\left[Ni_{12}(pyaox)_{6}(pyaoxH)_{6}(H_{2}O)_{2}(N_{3})_{2}\right]^{4+}$ and $[Ni_{12}(pyaox)_{6}(pyaoxH)_{6}(H_{2}O)_{2}]^{6+}.$ ^{21,23,24} The cations have single-(Ni₄), double-(Ni₈) and triple-decker (Ni₁₂) structures, where each deck is composed of the same tetranuclear cationic unit $\{Ni_4(pyaox)_2(pyaoxH)_2\}^{2+}$ (proved to be present in solution by ESI-MS spectrometry^{23,24}) and features two square planar and two octahedral Ni^{II} centres.

In this work, we expand the above mentioned family of multiple-decker Ni clusters by reporting the syntheses, crystal structures and magnetic and heat capacity properties of two such species, namely $[Ni_{12}(pyaox)_6(pyaoxH)_6(MeOH)_2Cl_2]$ Cl_4 ·5MeOH (1·5MeOH) and $[Ni_{16}(pvaox)_8(pyaoxH)_8(MeOH)_4]$ $(SO_4)_4$ ·10H₂O·26MeOH (2·10H₂O·26MeOH). The two clusters are high-spin featuring both the singly (pyaoxH−) and doubly deprotonated (pyaox^{2−}) forms of pyaoxH₂. 2·10H₂O·26MeOH is the first example of a metal cluster with a quadruple-decker topology, being also the highest nuclearity metal cluster, in general, containing any form of the pyaox $H₂$ ligand and the highest nuclearity Ni^H cluster with any type of 2-pyridyl oxime. In addition, its nuclearity is the highest that has been observed so far in metal clusters bearing 2-pyridyl oximes.^{17c,26} The number of the decks in 1.5MeOH and 2.10H₂O·26MeOH, and hence the nuclearity of the products, depend on the nature of the inorganic anion that is present in the reaction system, providing elements of synthetic control towards new ferromagnetic species. Portions of this work have been previously communicated.²¹

Experimental section

Materials and physical measurements

All manipulations were performed under aerobic conditions using materials (reagent grade) and solvents as received. The organic ligand pyaox H_2 was synthesized following the reported method.²⁷ Elemental analyses (C, H, N) were performed by the University of Patras microanalysis service. IR spectra (4000–400 cm−¹) were recorded using a Perkin Elmer 16PC FT-IR spectrometer with samples prepared as KBr pellets. The magnetic susceptibility measurements were carried out on a Quantum Design SQUID magnetometer MPMS-XL 7 operating

between 1.8 and 300 K for dc-applied fields ranging from 0 to 5 T. Microcrystalline samples were dispersed in vaseline in order to avoid torquing of the crystallites. The sample mulls were contained in a calibrated gelatin capsule held at the center of a drinking straw that was fixed at the end of the sample rod. Alternating current (ac) susceptibility measurements were carried out under an oscillating ac field of 3.5 Oe and frequencies ranging from 0.1 to 1500 Hz. The heat capacity measurements were carried out for temperatures down to 0.3 K by using a Quantum Design 14T-PPMS, equipped with a ³He cryostat. The experiments were performed on thin pressed pellets (ca. 1 mg) of a polycrystalline sample, thermalized by ca. 0.2 mg of Apiezon N grease, whose contribution was subtracted by using a phenomenological expression.

Synthetic details

Preparation of the complex $[Ni_{12}(pyaox)_6(pyaoxH)_6(MeOH)_2Cl_2]$ - $Cl₄·5MeOH$ (1·5MeOH). Solid NaOMe (0.054 g, 1.00 mmol) was added to a colourless solution of $pyaoxH₂$ (0.068 g, 0.50 mmol) in MeOH (15 mL); the solid soon dissolved. Solid $\text{NiCl}_{2} \cdot 6\text{H}_{2}\text{O}$ (0.119 g, 0.50 mmol) was then added and the resulting slurry was stirred for 15 min at room temperature. The dark brown slurry was filtered and the solution was left to stand undisturbed in a closed flask for a period of 3–4 days. Dark brown crystals appeared which were collected by filtration, washed with $H₂O$ (1 mL), cold EtOH (1 mL) and Et₂O (3 mL), and dried in air. Yield: 60%. The dried solid was analyzed satisfactorily as 1. Calcd (found): C 34.07 (33.84), H 2.86 (2.90), N 19.33 (19.21) %. IR (KBr, cm−¹): 3328b, 1654m, 1608s, 1577m, 1542s, 1483s, 1448w, 1416m, 1303w, 1263w, 1158m, 1103w, 1076m, 1023m, 992w, 938w, 782m, 729w, 702w, 645m, 517w, 480w. Datton Transactions

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Preparation of the complex $\left[\text{Ni}_{16}\right(pyaox)_8(pyaoxH)_8(MeOH)_4\right]$ $(SO_4)_4$ ·10H₂O·26MeOH (2·10H₂O·26MeOH). NEt₃ (0.139 mL, 1.00 mmol) was added to a colourless solution of $pyaoxH₂$ (0.068 g, 0.50 mmol) in MeOH (20 mL). Solid NiSO₄.6H₂O (0.130 g, 0.50 mmol) was then added and the resulting solution was stirred under heating $(45 \degree C)$ for 15 min. The dark brown solution was left to stand undisturbed in a closed flask for a period of 3–4 days. Dark brown crystals appeared which were collected by filtration, washed with cold EtOH (1 mL) and $Et₂O$ (3 mL), and dried in air. Yield: 65%. The dried solid was analyzed satisfactorily as $2.8H_2O.22MeOH$. Calcd (found): C 32.78 (32.52), H 4.69 (4.25), N 15.04 (15.17) %. IR (KBr, cm−¹): 3332b, 1654m, 1606m, 1580w, 1542m, 1482m, 1415m, 1300w, 1263w, 1100s, 1022w, 981w, 933w, 777m, 747w, 700w, 646m, 617s, 515w, 468m.

Single-crystal X-ray crystallography

Suitable single-crystals of 1.5MeOH and 2.10H₂O·26MeOH were mounted on a Hampton Research CryoLoop using FOMBLIN Y perfluoropolyether vacuum oil (LVAC 25/6) purchased from Aldrich, 28 with the help of a Stemi 2000 stereomicroscope equipped with Carl Zeiss lenses. Data were collected at 150(2) K on a Bruker X8 Kappa APEX II charge-coupled

device (CCD) area-detector diffractometer (Mo K_{α} graphitemonochromated radiation, $\lambda = 0.71073$ Å) controlled by the APEX2 software package, 29 and equipped with an Oxford Cryosystems Series 700 cryostream monitored remotely using the software interface, Cryopad. 30 Images were processed using the software package SAINT+ $,31$ and data were corrected for absorption by the multi-scan semi-empirical method implemented in SADABS. 32 The structures were solved by the direct methods of SHELXS-97, 33 and refined by full-matrix least squares on F^2 using SHELXL-97.³⁴

All non-hydrogen atoms were directly located from difference Fourier maps and successfully refined with anisotropic displacement parameters. For both compounds, hydrogen atoms attached to carbon and to the hydroxido groups were located at their idealised positions and included in the structural model in subsequent refinement cycles in riding-motion approximation with $U_{\text{iso}} = 1.2$ (aromatic carbon) or 1.5 (-CH₃ and -OH groups) of U_{eq} of the parent atom. Hydrogen atoms associated with the nitrogen atoms were markedly visible in difference Fourier maps and were included in the structures with the N–H and H⋯H distances restrained to 0.90(1) and 1.50(1) Å, respectively, and with $U_{\text{iso}} = 1.5 \times U_{\text{eq}}(N)$.

For $2.10H₂O₂6MeOH$ the hydrogen atoms belonging to the uncoordinated water molecules could not be directly located from difference Fourier maps and no attempt was made to place these in approximate calculated positions; they have been included in the empirical formula of the compound.

Beside the solvent molecules located in both the structures $(5 \text{ MeOH} \text{ in compound 1, as well as 10 H₂O and 26 MeOH in }$ 2), considerable smeared-out electron densities were found in the empty spaces available in the two structures, which prevented a sensible location and refinement of other solvent molecules. A search for the total potential solvent area using the software package $PLATOR^{35}$ revealed the presence of large voids housing the disordered molecules. The two data sets were then analysed using the Squeeze,³⁶ and the reflection lists arising from this procedure were used for the final structural refinements.

Full details can be found in Table 1 and the cif files.

Results and discussion

Synthetic parameters and IR spectra

Our groups have strong interest in the employment of 2-pyridyl oxime ligands as a route to high nuclearity 3d transition metal clusters and SMMs.^{14,17-21} As a part of this program, we recently decided to explore pyaox H_2 in Ni^{II} cluster chemistry. Thus, a variety of experiments were performed to study how the different synthetic parameters affect the identity and/or crystallinity of the isolated product. In particular, treatment of NiCl₂·6H₂O with one equivalent of pyaoxH₂ and two equivalents of NaOMe in MeOH gave a dark brown solution from which the dodecanuclear cluster $\left[\text{Ni}_{12}\right]$ pyaox H_6 pyaox H_6

Table 1 Crystallographic data for compounds 1·5MeOH and 2·10H₂O·26MeOH

	1.5MeOH	2.10H ₂ O.26MeOH	$(pyaoxH)_{8}(MeOH)_{4}[(SO_{4})_{4} (2)$ was obtained in 65% yield. Its formation can be summarized by eqn (2):
Formula ^a $M_{\rm w}$ Crystal system Space group $a/\text{\AA}$ $b/\text{\AA}$ $c/\text{\AA}$ a ^o β /° γ / \circ V/A ³ Ζ $T\!/\mathrm{K}$ λ^b/\mathring{A} $D_{\rm c}/g~{\rm cm}^{-3}$ μ (Mo Ka)/mm ⁻¹ Reflections collected Independent reflections $R_{\rm int}$ R_1 $\overline{wR_2}^d$ Goodness of fit on \mathbb{F}^2 $\Delta \rho$ max/min/e Å ⁻³ ^{<i>a</i>} Including solvent molecules. ^{<i>b</i>} Graphite monochromator. ^{<i>c</i>} $R_1 = \sum (F_o - F_c)/\sum F_o $. ^{<i>d</i>} w $R_2 = [\sum [w(F_o^2 - F_c^2)^2]/\sum [wF_o^2)^2]]^{1/2}$.	$C_{79}H_{94}Cl_6N_{36}Ni_{12}O_{19}$ 2769.12 Triclinic ΡĪ 14.8718(8) 15.1738(7) 15.3809(8) 60.525(2) 73.084(3) 69.854(2) 2805.3(3) $\mathbf{1}$ 150(2) 0.71073 1.639 2.182 147 901 14885 0.0422 0.0596 0.2006 1.047 $4.818/-0.908$	$C_{126}H_{228}N_{48}Ni_{16}O_{72}S_4$ 4635.16 Triclinic РĪ 16.6742(9) 17.3665(17) 19.7073(10) 106.175(3) 113.293(2) 99.202(4) 4792.3(6) $\mathbf{1}$ 150(2) 0.71073 1.606 1.672 321514 21715 0.0375 0.0461 0.1148 1.138 $1.194/-0.700$	$16N$ iSO ₄ ·6H ₂ O + 16 pyaoxH ₂ + $24NEt_3$ + $4MeOH$ $\left[\text{Ni}_{16}\text{(pyaox)}_{8}\text{(pyaoxH)}_{8}\text{(MeOH)}_{4}\right]\text{(SO$_4$)}_{4}$ (2) $+ 12(NHEt3)2SO4 + 96H2O$ The reaction mixture which leads to 2 should be heated for a few minutes to facilitate the dissolution of $Niso_4.6H_2C$ (presumably by reaction with the organic ligand) in MeOH We observed that reactions with small variations in the $NiII$: pyaox $H2$ ratio gave the same Ni ₁₆ compound. The nature of the base and the crystallization method are not crucial for the identity of the product and affect only its crystallinity; we were able to isolate 2 (IR evidence) by using a plethora of bases such as NaOMe, NaOH, NH ₃ etc. The IR spectra of dried 1.5MeOH and $2.10H_2O$ 26MeOH exhibit a strong broad band at \sim 3330 cm ⁻¹ assignable to the $\nu(OH)$ vibration of coordinated MeOH molecules, as well as solvent MeOH and H ₂ O molecules, and to the $\nu(NH_2)/\nu(NH_1)$ vibrations of the pyaoxH/pyaox ²⁻ ligands. The broadness and low frequency of these bands are both indicative of strong hydrogen bonding. There are medium intensity bands at 1566
(MeOH) ₂ Cl ₂]Cl ₄ ·5MeOH (1·5MeOH) was obtained in 60% yield. Its formation can be summarized by eqn (1) :			and 1116 cm^{-1} in the spectrum of the free pyaoxH ₂ ligand which are assigned to ν (C=N) _{oxime} and ν (N-O) _{oxime} , respec tively. ²⁷ The 1116 cm^{-1} band is shifted to a lower wavenumber $(1103 \text{ cm}^{-1} \text{ in } 1.5 \text{MeOH}; 1100 \text{ cm}^{-1} \text{ in } 2.10 \text{H}_2\text{O} \cdot 26 \text{MeOH})$
$12NiCl_2 \cdot 6H_2O + 12pyaoxH_2 + 18NaOMe$ MeOH) ₂ Cl ₂ Cl ₄ (1) 1			This shift is attributed to the coordination of the oximate nitrogen. ³⁷ The band is strong in the spectrum of the $Ni10$ cluster, because it also has a $\nu_3(F_2)$ [$\nu_d(SO)$] character due to
$+ 18$ NaCl + 72H ₂ O + 16MeOH The methoxide ions act as proton acceptors to facilitate			the ionic sulfates $(T_d$ symmetry). Somewhat to our surprise the 1566 cm^{-1} band is shifted to a higher wavenumber in the complexes (1577 cm ⁻¹ in 1.5MeOH, 1580 cm ⁻¹ in 2.10H ₂ O

$$
12\text{NiCl}_{2} \cdot 6\text{H}_{2}\text{O} + 12\text{pyaoxH}_{2} + 18\text{NaOMe}
$$
\n
$$
\xrightarrow{\text{MeOH}} [\text{Ni}_{12}(\text{pyaox})_{6}(\text{pyaoxH})_{6}(\text{MeOH})_{2}\text{Cl}_{2}] \text{Cl}_{4}
$$
\n
$$
1 + 18\text{NaCl} + 72\text{H}_{2}\text{O} + 16\text{MeOH}
$$
\n
$$
(1)
$$

The methoxide ions act as proton acceptors to facilitate formation of pyaox^{2−} and pyaoxH[−] ligands. The reaction procedure involved a slight excess of NaOMe over that required by eqn (1), and this might have proven beneficial in shifting a possible equilibrium to the right. The reaction mixtures which lead to 1 should be filtered before crystallization to remove NaCl which is insoluble in MeOH. The product also needs to be washed with a small amount of H_2O to ensure complete removal of NaCl.

The determination of the crystal structure of 1 with single crystal X-ray crystallography revealed that the twelve metal ions have an unprecedented triple-decker topology with each "deck" containing two square planar and two octahedral Ni^{II} atoms. The intriguing structural and magnetic features of 1 (vide infra) prompted us to further study this reaction system exploring the ability of $pyaoxH₂$ to deliver other novel polynuclear Ni^{II} compounds. In the context of this investigation, we decided to employ different counterions possessing higher charge than Cl[−] hoping that they could stabilize higher nuclearity clusters. Indeed, this turned out to be the case and the treatment of $Niso_4.6H_2O$ with one equivalent of pyaox H_2 and two equivalents of $NEt₃$ in MeOH gave a dark brown solution

$$
16NiSO4 \cdot 6H2O + 16pyaoxH2 + 24NEt3 + 4MeOH
$$

\n
$$
\longrightarrow \frac{[Ni16(pyaox)8(pyaoxH)8(MeOH)4](SO4)4}{2}
$$

\n+ 12(NHEt₃)₂SO₄ + 96H₂O (2)

The IR spectra of dried 1.5 MeOH and $2.10H₂O₂₆$ MeOH exhibit a strong broad band at ∼3330 cm−¹ assignable to the $\nu(OH)$ vibration of coordinated MeOH molecules, as well as solvent MeOH and H₂O molecules, and to the $\nu(NH_2)/\nu(NH)$ vibrations of the pyaoxH/pyaox^{2−} ligands. The broadness and low frequency of these bands are both indicative of strong hydrogen bonding. There are medium intensity bands at 1566 and 1116 cm⁻¹ in the spectrum of the free pyaoxH₂ ligand, which are assigned to ν (C=N)_{oxime} and ν (N-O)_{oxime}, respectively.²⁷ The 1116 cm⁻¹ band is shifted to a lower wavenumber $(1103 \text{ cm}^{-1} \text{ in } 1.5 \text{MeOH}; 1100 \text{ cm}^{-1} \text{ in } 2.10 \text{H}_2\text{O} \cdot 26 \text{MeOH}).$ This shift is attributed to the coordination of the oximate nitrogen.³⁷ The band is strong in the spectrum of the Ni₁₆ cluster, because it also has a $\nu_3(F_2)$ [$\nu_d(SO)$] character due to the ionic sulfates $(T_d$ symmetry). Somewhat to our surprise, the 1566 cm^{-1} band is shifted to a higher wavenumber in the complexes (1577 cm⁻¹ in 1·5MeOH, 1580 cm⁻¹ in 2·10H₂O· 26MeOH), overlapping with an aromatic stretch. This shift may be indicative of the oximate nitrogen coordination.³⁷ Extensive studies on Schiff-base complexes (which also contain a C=N bond) have shown³⁸ that a change in the s character of the nitrogen lone pair occurs upon coordination such that the s character of the nitrogen orbital, which is involved in the C $=N$ bond, increases; this change in hybridization produces a greater $C=N$ stretching force constant relative to the free neutral ligand. The in-plane deformation band of the 2-pyridyl ring of free pyaoxH₂ at 632 cm⁻¹ shifts upwards in 1⋅5MeOH (646 cm^{-1}) and 2⋅10H₂O⋅26MeOH (645 cm−¹), confirming the involvement of the ring-N atom in coordination.39

Description of structures

Representations of the molecular structure and the structural core of compound 1·5MeOH are shown in Fig. 1. Selected interatomic distances and angles are listed in Table 2. Compounds 1.5MeOH and 2.10H₂O.26MeOH display related structures based on parallel $Ni₄$ layers, $\{Ni₄(pvaox)\}$ $(pyaoxH)_2^{2+}$ $(x = 3, 1.5MeOH; x = 4, 2.10H_2O.26MeOH)$. Thus,

Fig. 1 Representations of the molecular structure of the cation of 1 (top) and its triple-decker structural core (bottom).

Ni1–O4	2.0279(18)	Ni3-N11	2.250(2)
Ni1-O2	2.0293(19)	$Ni4-O4$	1.8306(18)
Ni1-N10'	2.048(2)	$Ni4-N5$	1.853(2)
Ni1-N11'	2.070(2)	$Ni4-N12$	1.861(2)
Ni1-O1	2.1705(19)	$Ni4-N4$	1.903(2)
Ni1–N17	2.278(2)	$Ni5-O5$	1.8263(19)
Ni2-O3	1.998(2)	$Ni5-N15$	1.846(2)
Ni2–O5	2.0033(19)	$Ni5-N8$	1.852(2)
Ni2-N16	2.031(2)	$Ni5-N7$	1.887(2)
Ni2–N17	2.089(2)	$Ni6-O6$	1.8227(18)
Ni2-O2	2.1729(19)	Ni6-N18	1.846(2)
Ni2–Cl1	2.5713(10)	$Ni6-N2'$	1.846(2)
Ni3-N14	2.013(2)	$Ni6-N1'$	1.888(2)
Ni3-O6	2.0320(18)	$Ni2 \cdots Ni3$	3.891(2)
Ni3-O1'	2.0589(19)	$Ni2 \cdots Ni5$	3.323(2)
Ni3–N13	2.076(2)	$Ni2 \cdots Ni6$	4.670(2)
Ni3-O1 M	2.113(2)	$Ni1\cdots Ni3$	3.141(2)
O4-Ni1-N10'	178.74(8)	$N5-Ni4-N12$	174.14(10)
O2-Ni1-N11'	172.77(8)	$O4-Ni4-N4$	172.22(9)
O1-Ni1-N17	175.06(7)	$N12-Ni4-N4$	102.89(10)
O5-Ni2-N16	175.01(9)	N15-Ni5-N8	173.30(11)
O2-Ni2-Cl1	172.98(5)	$O5-Ni5-N7$	170.12(10)
O3-Ni2-N17	169.24(8)	$N15-Ni5-N7$	102.09(11)
O6-Ni3-N13	173.75(9)	N18-Ni6-N2'	174.26(10)
O1M-Ni3-N11	172.46(9)	O6-Ni6-N1'	172.83(10)
N14-Ni3-N11	89.73(9)	$N18-Ni6-N1'$	102.40(10)

^a Symmetry transformations used to generate equivalent atoms: \therefore -x, $-y, -z + 1.$

only the structure of compound 1·5MeOH will be discussed in detail and compared to that of $2.10H₂O₂₆MeOH$.

Complex 1·5MeOH crystallizes in the triclinic space group $P\overline{1}$. Its structure consists of $[Ni_{12}(pyaox)]_6(pyaoxH)_6$ Cl₂(MeOH)₂]⁴⁺ cations, Cl[−] counterions and MeOH lattice molecules. The centrosymmetric cation of 1·5MeOH is composed of six distorted octahedral (Ni1, Ni2, Ni3 and symmetry equivalents) and six square planar (Ni4, Ni5, Ni6 and symmetry equivalents) Ni^{II} atoms, which are held together through two $η¹:η¹:η¹:μ$ and four $η¹:η¹:η²:μ₃$ pyaoxH⁻ ligands, as well as two $\eta^1:\eta^1:\eta^2:\mu_3$ and four $\eta^1:\eta^2:\eta^1:\eta^2:\mu_4$ pyaox²⁻ groups (Scheme 2). The coordination sphere of the Ni^H atoms is completed by two terminal chlorido ligands on Ni2 and Ni2′, and two terminal MeOH molecules on Ni3 and Ni3′. The twelve metal ions in 1·5MeOH are arranged into three layers, which are stacked one above the other forming an unusual, for 3dmetal cluster chemistry, triple-decker topology. Each "deck" contains two square planar and two octahedral Ni^H atoms, whereas the linkage of the central deck with each of the outer decks is achieved through two bridging oxygen atoms from two η¹:η¹:η²:μ₃ pyaoxH⁻ ligands and two bridging oximate nitrogen atoms from two $η^1:η^2:η^1:η^2:μ_4$ pyaox²⁻ groups. The four Ni^{II} atoms of the two outer layers are arranged in an almost planar topology with the largest deviation from the plane being 0.14 Å for Ni3 and Ni6, whereas the metal ions in the central layer are perfectly coplanar. The distance between two adjacent decks is ∼2.70 Å. The chromophores in 1.5MeOH are $Ni(1,1')(N_{py})(N_{ox})_2(O_{ox})_3$, $Ni(2,2')(N_{py})(N_{ox})$ $(O_{ox})_3$ (Cl), Ni(3,3')(N_{py})(N_{ox})₂(O_{ox})₂(O_{MeOH}) and Ni(4,4'/5,5'/6,6') $(N_{\text{pv}})(N_{\text{ox}})(N_{\text{am}})(O_{\text{ox}})$, where the abbreviations "py", "ox" and "am" are for the 2-pyridyl, oximate and deprotonated amino donor atoms, respectively. The strong ligand field, deprotonated amino donor atoms are coordinated only to the square planar Ni^{II} atoms and may be considered responsible for their geometry. The average $Ni(4,5,6)$ -N_{py}, N_{ox}, O_{ox} bond distances are 1.893, 1.850 and 1.827 Å, respectively, while the average Ni(1,2,3)–N_{py}, N_{ox}, O_{ox} bond lengths are 2.052, 2.140 and 2.062 Å, respectively. The longer bonds involving $Ni(1,2,3)$ compared to those involving $Ni(4,5,6)$ are a consequence of the higher coordination number of Ni1, Ni2 and Ni3. Dation Tennesctions

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> The $\eta^1:\eta^2:\eta^1:\eta^2:\mu_4$ coordination mode of pyaox²⁻ found in 1·5MeOH is extremely rare in the coordination chemistry of pyridine-2-amidoxime. 21 In fact, this can be rationalised by considering the C– N_{ox} bond lengths; the C– N_{ox} and C– N_{am} distances for the $\eta^1:\eta^2:\eta^1:\eta^2:\mu_4$ pyaox²⁻ ligands (average values: 1.345 and 1.322 Å) are longer and shorter, respectively, than those for the $\eta^1:\eta^1:\eta^1:\eta^2:\mu_3$ pyaox²⁻ ligands (1.318 and 1.336 Å),

Scheme 2 The crystallographically established coordination modes of pyaoxH[−] and pyaox2[−] in complexes 1 and 2.

indicating a decreased double bond character for the $C-N_{ox}$ bond and an increased double bond character for the $C-N_{am}$ bond in the former.

The crystal structure of 1·5MeOH is stabilized by strong intramolecular hydrogen bonds between the O atoms of the terminally ligated MeOH molecules (O1M), which act as donors, and the terminal chlorido ligands, which act as acceptors $[O1 \ M \cdots Cl1 = 3.199(2) \ \AA$, H1 $M \cdots Cl1 = 2.256(2) \ \AA$, O1 M–H1 M–Cl1 = $167.19(2)°$. In addition, there is a network of intermolecular H bonds that involve: (1) the Cl[−] counterions (acceptors) and the $NH₂$ groups (donors) of the organic ligands $[N3\cdots C]$ = 3.245(2), H3B \cdots Cl3 = 2.360(2) Å, N3-H3B-Cl3 = 166.93°; N9…Cl3 = 3.282(2) Å, H9B…Cl3 = 2.421(2) Å, N9–H9B–Cl3 = 159.83°] resulting in the formation of 1D chains, and 2) the coordinated chlorido ligands (acceptor) and the MeOH lattice molecules (donors) [O3 M…Cl1 = 3.237(2) Å, H9 M⋯Cl1 = 2.422(2) Å, O3 M–H9 M–Cl1 = 163.49°]. The cations of 1 are in close proximity with the shortest metal…metal separation between the adjacent $Ni₁₂$ units being $4.797(2)$ Å (Ni5…Ni5).

Representations of the molecular structure and the structural core of compound 2·10H₂O·26MeOH are shown in Fig. 2. Selected interatomic distances and angles are listed in Table S1 in the ESI.†

Complex $2.10H₂O₂₆$ MeOH crystallizes in the triclinic space group \overline{PI} . Its structure consists of $\overline{[Ni_{16}(pyaox)]_8}(pyaoxH)_8$ $(MeOH)_4]^8$ ⁺ cations, SO_4^2 ⁻ counterions, and MeOH and H₂O lattice molecules. Similarly to 1.5MeOH , the Ni^{II} atoms in 2·10H2O·26MeOH are stacked in four layers, which form an unprecedented quadruple-decker topology. In particular, the inner decks in 2·10H₂O·26MeOH are identical with the central deck of 1·5MeOH; thus, figuratively speaking, an internal deck has been added to the $Ni₁₂$ cation of the latter to form a $Ni₁₆$ cluster ($2.10H₂O₂₆MeOH$). In addition, the external decks in $2.10H₂O·26MeOH$ are similar to the corresponding decks in 1·5MeOH, with their only difference being the replacement of the chlorido ligands in the latter by terminal MeOH molecules in the former. The octahedral metal centres are Ni1, Ni2, Ni3 and Ni4 (and their symmetry equivalents) and the square planar ones are Ni5, Ni6, Ni7 and Ni8 (and their symmetry equivalents). The presence of many lattice molecules and counterions in the crystal structure of $2.10H₂O₂₆MeOH$ results in the formation of a network of strong intermolecular hydrogen bonding interactions. Details of these interactions are listed in Table S2 in the ESI.† As it was the case for 1·5MeOH, the cations of 2·10H2O·26MeOH, are in close proximity with the shortest metal⋯metal separation between the adjacent Ni₁₆ units being 4.671(2) Å (Ni1…Ni7).

1·5MeOH and 2·10H2O·26MeOH are new additions in the small family of high nuclearity, high spin Ni compounds with multiple-decker metal topology.^{23,24} In particular, $2.10H_2O$ 26MeOH is the first structurally characterized complex of any metal displaying a quadruple-decker configuration. Compound 1·5MeOH can be described as a "trimer-of-tetramers", while 2·10H₂O·26MeOH as a "tetramer-of tetramers", where the tetramer in both cases is the structural unit

Fig. 2 Representations of the molecular structure of the cation of 2 (top) and its quadruple-decker structural core (bottom).

 ${Ni_4(pyaox)_2(pyaoxH)_2}^{2+}$, which has been previously observed in a discrete tetranuclear cluster. $23,24$ The number of the decks, and hence the nuclearity and spin, of 1·5MeOH and $2.10H₂O₂₆$ MeOH is affected by the nature (charge, shape) of the inorganic anion that is present in the reaction system providing the elements of synthetic control towards new high nuclearity Ni^{II} clusters. Both compounds contain the singly and doubly deprotonated pyridine-2-amidoxime ligand, with 2·10H2O·26MeOH being (1) the highest nuclearity metal cluster bearing this ligand, (2) one of the few highest nuclearity hexadecanuclear metal clusters with any 2-pyridyl oxime, in general,^{17c,26} and (3) the highest nuclearity Ni^H cluster with 2-pyridyl oximes. Furthermore, complex $2.10H_2O.26MeOH$ joins only a very small family of $Ni₁₆$ clusters with O- or/and N-ligation.⁴⁰

Magnetic susceptibility studies

In order to probe the magnetic properties of the clusters, direct current (dc) magnetic susceptibility measurements were performed on polycrystalline samples of compounds 1 and 2 in the temperature range 2–300 K, with an applied magnetic field of 0.1 T.

The collected data are plotted as $\chi_M T$ versus T in Fig. 3, which reveal room-temperature $\chi_M T$ values of 7.16 and 9.03 cm³ K mol⁻¹ for 1 and 2, respectively. These are in good agreement with the $\chi_M T$ value expected (g = 2.10) for six noninteracting octahedral Ni^{II} ions (6.615 cm³ mol⁻¹ K) in 1, and eight non-interacting octahedral Ni^{II} ions (8.82 cm³ mol⁻¹ K) in 2. For both compounds, the value of χ_MT steadily increases to a maximum value (14.5 cm³ mol⁻¹ K at 14 K, 1; 21.7 cm³ mol⁻¹ K at 10 K, 2), and then sharply decreases to 10.8 and 14.0 cm³ mol⁻¹ K at 2.0 K for 1 and 2, respectively. The profile of the $\gamma_M T$ versus T plots is indicative of dominant ferromagnetic interactions between the paramagnetic ions within the cations of 1 and 2. The low temperature decrease of the $\gamma_M T$ value is attributed to intermolecular antiferromagnetic interactions and/or zero-field splitting. The former is consistent with the crystal structure of the two compounds (vide supra). **Dation Transaction**
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 If the care of proportion access proportion access Components are are $\frac{1}{2}$ **and** $\frac{1}{2}$ **and** $\frac{1}{2}$ **and** $\frac{1}{2}$ **and** $\frac{1}{2}$ **and \frac{**

In order to fit the χ_{M} T data for 1.5MeOH, we modeled the $Ni₆$ magnetic core with the 2*J*-model described in Fig. 4, left, (the octahedral Ni^{II} atoms define two triangles related by an inversion centre) employing the isotropic spin-Hamiltonian $(1):$

$$
\hat{H} = -2J_{1}(\hat{S}_{1} \cdot \hat{S}_{3} + \hat{S}_{1} \cdot \hat{S}_{1'} + \hat{S}_{2} \cdot \hat{S}_{3} + \hat{S}_{1'} \cdot \hat{S}_{3'} + \hat{S}_{2'} \cdot \hat{S}_{3'}) \n-2J_{2}(\hat{S}_{1} \cdot \hat{S}_{2} + \hat{S}_{1} \cdot \hat{S}_{3'} + \hat{S}_{1'} \cdot \hat{S}_{3} + \hat{S}_{1'} \cdot \hat{S}_{2'}) + \Sigma_{i} \mu_{B} \mathbf{g}_{N i} \vec{H} \hat{S}_{i}
$$
\n(1)

where J is the isotropic exchange interaction parameter, i is the index for each Ni^H ion in the Ni₆ core, μ_B is the Bohr magneton, g_{Ni} = 2.10, the g-factor of the Ni^{II} ions, B is the applied magnetic field and \hat{S} is a spin operator. The data was numeri-

 22 20 18 χ_{M} T (cm³ K mol⁻¹) 16 14 12 10 **PARADO** 8 Ò 1 6 $\mathbf 0$ 50 100 150 200 250 300 $T(K)$

Fig. 3 Plot of $\chi_M T$ as a function of T for complexes 1 (\square) and 2 (\square); the solid lines represent the fit of the data to the theoretical models given in the text.

Fig. 4 Schematic presentation of the coupling scheme used to model susceptibility data for the cations of 1 (left) and 2 (right).

cally fitted using the simplex algorithm, 41 with intermolecular interactions being taken into account in the frame of meanfield theory, by the use of the Curie–Weiss temperature, θ . The fit affords the parameters $J_1 = +5.14$ cm⁻¹, $J_2 = +3.01$ cm⁻¹ and θ = −5.73 K. This results in an S = 6 ground state with the nearest excited states being $S = 5$, lying 8.38 cm⁻¹ higher in energy, and $S = 4$, lying at 17.11 cm⁻¹. A plot of energy versus the total spin state is given in Fig. S1, left, in the ESI.† Similarly, the 2J-model described in Fig. 4, right, was used in order to fit the data for the $Ni₈$ magnetic core of 2·10H2O·26MeOH using the isotropic spin-Hamiltonian (2):

$$
\hat{H} = -2J_{1}(\hat{S}_{1} \cdot \hat{S}_{4'} + \hat{S}_{2} \cdot \hat{S}_{3'} + \hat{S}_{2} \cdot \hat{S}_{4'} + \hat{S}_{3} \cdot \hat{S}_{3'} + \hat{S}_{1'} \cdot \hat{S}_{4} + \hat{S}_{2'} \cdot \hat{S}_{3} + \hat{S}_{2'} \cdot \hat{S}_{4}) \n-2J_{2}(\hat{S}_{1} \cdot \hat{S}_{2} + \hat{S}_{2} \cdot \hat{S}_{3} + \hat{S}_{3} \cdot \hat{S}_{4} + \hat{S}_{1'} \cdot \hat{S}_{2'} + \hat{S}_{2'} \cdot \hat{S}_{3'} + \hat{S}_{3'} \cdot \hat{S}_{4'}) \n+ \Sigma_{i} \mu_{B} g_{Ni} \vec{H} \hat{S}_{i}
$$
\n(2)

The fit afforded the parameters J_1 = +5.45 cm⁻¹, J_2 = +2.00 cm⁻¹ and θ = −3.43 K. This results in an S = 8 ground state with the nearest excited states being $S = 7 (3.95 cm^{-1})$ and $S = 6 (8.07 cm^{-1})$ lying relatively close in energy, Fig. S1,† right. We have also collected the alternating current (ac) magnetic susceptibility in zero applied dc field for both complexes, which revealed no frequency dependence (Fig. S2 in the ESI†).

Fig. S3 in the ESI† shows the dc-field magnetization data $M(H)$ collected in the range 0–7 T at three temperatures between 2.3 and 20 K for 1 and 2. In both cases, $M(H)$ initially rises quickly at the lowest temperatures, as expected for ferromagnetic exchange, and then keeps increasing slowly even at the highest applied fields, without reaching saturation. In accordance with the susceptibility data, this behaviour is attributed to intermolecular antiferromagnetic interactions, likely combined with some degree of magnetic anisotropy. For comparison, the calculated Brillouin curves were also plotted assuming uncorrelated molecules with $S = 6(1)$ and $S = 8(2)$ for the representative temperatures. For both compounds, it is observed that the experimental data tend to lie below the calculated curve, suggesting the influence either from antiferromagnetic intermolecular correlations or from low-lying excited spin states. The latter is more evident in 2 with respect to 1, as it is revealed from the larger deviation of the calculated curve and experimental data, which indicates that 2 does not display a well-defined ground state spin value. Keeping the values of S and g stable, detectable improvements are obtained only by adding an abnormally high uniaxial magnetic anisotropy term D in the case of 1. Since this applies also at the lowest

investigated temperatures, in which the influence of excited states should be negligible, we conclude, in agreement with the susceptibility data, that antiferromagnetic interactions between cations are important for the two complexes.

It has recently been reported that the dominant factor which affects the magnetic properties of the oximato Ni^{II} compounds is the length of the bridging N–O bond.^{23,42} In particular, DFT calculations on an oximato bridged Ni^{II} compound revealed that N–O distances larger than 1.394 Å result in ferromagnetic interactions, while shorter ones propagate antiferromagnetic interactions. This observation has been explained by considering that with the decrease of the N–O bond, the spin population on the N and O atoms increases, thus enhancing the antiferromagnetic coupling between Ni^{II} ions. In compounds 1·5MeOH and 2·10H2O·26MeOH, the N–O distances are in the range of 1.401–1.425 Å, and thus the experimentally obtained ferromagnetic interactions are in excellent agreement with the previously reported theoretical calculations.

Specific heat measurements

Fig. 5 shows the collected specific heat $C(T,H)$ data of 1 as a function of temperature for several applied fields. No particular feature is detected except for a very broad anomaly around 2 K at zero applied field, whose magnetic nature is proven by the fact that it quickly moves towards higher temperatures as the field H is increased. As typical for moleculebased materials, 43 the contribution to the total C arising from lattice vibrations dominates as T increases above that of liquid helium.

The lattice contribution can be described by the Debye model (dashed line in Fig. 5), which simplifies to a $C/R = aT³$ dependence $(R$ is the gas constant) at the lowest temperatures, where $a = 3.9 \times 10^{-3} \text{ K}^{-3}$ for 1. The magnetic contribution C_m (see inset in Fig. 5) is then obtained by subtracting the lattice contribution to the total C. This analysis reveals a bump centered at ≈9 K, that we associate with a Schottky-like anomaly, *i.e.*, crystal-field splitting of the $S = 6$ multiplet. By decreasing the temperature, magnetic fluctuations within the 2D layers show up in the anomaly centered at \approx 2 K. To prop-

Fig. 5 Specific heat C for 1 normalised to the gas constant R vs. T in the indicated temperature and field ranges. Inset: Magnetic contribution C_m as obtained by subtracting the lattice contribution (dashed line) to C.

erly associate these features with the magnetic contribution, we estimate the zero-field magnetic entropy $S_{\rm m} = \int C_{\rm m}/T dT$, shown in Fig. S4 in the ESI.† This tends at high-T to the value of 2.8 R, which is in fair agreement with the magnetic entropy expected, $R \ln(2S + 1) = 2.56 R$ (provided the spin value is $S = 6$) and consistent with the magnetization data. Since the lattice contribution is sensibly larger than C_m , the latter is affected by an important uncertainty. This inhibits us from providing anything other than a qualitative analysis of the experimental data.

Specific heat data have been also collected for 2 and these are shown in Fig. S5 in the ESI† as a function of temperature for several applied magnetic fields. As it is the case for 1, the specific heat at high temperatures is dominated by a non-magnetic contribution, which is due to the thermal vibrations of the lattice. Furthermore, there is a clear magnetic contribution, which extends over a broad temperature range, i.e., from the lowest accessed T to well inside the range mainly determined by the lattice contribution to the specific heat. The fact that 2 does not display a well-defined ground state spin value ($S \approx 8$, *vide supra*) prevents to determine reliably the magnetic contribution by subtracting the lattice specific heat from the total measured signal, as in the case of the $Ni₁₂$ complex. As a result, a valuable analysis of the specific heat data is not feasible for 2. Poper

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Conclusions

The employment of pyaox H_2 in Ni^{II} chemistry has provided access to the new multiple-decker Ni complexes $\left[Ni_{12}\right(pyaox)\right]_6$ (pyaoxH)₆(MeOH)₂Cl₂]Cl₄·5MeOH (1·5MeOH) and [Ni₁₆(pyaox)₈ $(pyaoxH)_8(MeOH)_4[(SO_4)_4 \cdot 10H_2O \cdot 26MeOH$ (2·10H₂O·26MeOH) comprising stacked Ni₄ layers, $\{Ni_4(pyaox)_2(pyaoxH)_2\}^{2+}(x=3,$ 1.5MeOH; $x = 4$, 2.10H₂O·26MeOH), with each layer containing two square planar and two octahedral Ni^{II} centres.

Compound 1·5MeOH is a new addition to the small family of Ni_{12} clusters with a triple-decker topology.^{23,24} Compound 2·10H2O·26MeOH displays a quadruple-decker configuration, which has never been observed before in metal cluster chemistry. In addition, $2.10H₂O₂₆MeOH$ is the highest nuclearity metal cluster bearing $pyaoxH₂$ and joins only a very small family of $Ni₁₆$ clusters with O- or/and N-ligation. It also belongs to the very small family of the hexadecanuclear homometallic or/and heterometallic clusters, which display the highest nuclearity that has been observed so far for compounds containing 2-pyridyl oximes, being the only Ni^H cluster in this category.17c,25

Dc and ac magnetic susceptibility and heat capacity measurements were performed on 1 and 2, which revealed that both display ferromagnetic exchange interactions resulting in a spin ground state of $S = 6$ for 1 and $S = 8$ for 2. Intercationic interactions, as revealed by single crystal X-ray crystallography, affect the magnetic properties preventing an in-depth analysis of the heat capacity measurements.

The number of decks in 1 and 2 is affected by the use of different inorganic anions with the bulkier and of a higher charge anion $(SO_4^2$ ⁻ vs. Cl⁻) being able to stabilize the higher nuclearity cluster. This could lead to the development of a rational method towards the increase of the nuclearity and the spin of ferromagnetic species. Based on this, we believe that we have seen just the tip of the iceberg in the $Ni^H/pyaoxH₂$ compounds with a multiple decker topology. We have every reason to believe that higher nuclearity such species (e.g., $Ni₂₀$, $Ni₂₄, etc.$) are waiting to be discovered. Further studies towards this direction are currently in progress including the employment of higher charge counterions $(e.g., PO₄³⁻)$. These studies will be reported in due course.

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Notes and references

- 1 (a) R. Sessoli, H. L. Tsai, A. R. Schake, S. Wang, J. B. Vincent, K. Folting, D. Gatteschi, G. Christou and D. N. Hendrickson, J. Am. Chem. Soc., 1993, 115, 1804; (b) R. Sessoli, D. Gatteschi, A. Caneschi and M. A. Novak, Nature, 1993, 365, 141.
- 2 For reviews and books, see: (a) R. Bagai and G. Christou, Chem. Soc. Rev., 2009, 38, 1011; (b) C. Papatriantafyllopoulou, E. Moushi, G. Christou and A. J. Tasiopoulos, Chem. Soc. Rev., 2016, 45, 1597; (c) M. Murrie and D. J. Price, Annu. Rep. Prog. Chem., 2007, 103, 20; (d) G. Aromi and E. K. Brechin, Struct. Bonding, 2006, 122, 1; (e) D. Gatteschi, R. Sessoli and J. Villain, Molecular Nanomagnets, Oxford University Press, Oxford, 2006 ; (f) C. J. Milios and R. E. P. Winpenny, Struct. Bonding, 2015, 164, 1.
- 3 (a) M. Affronte, J. Mater. Chem., 2009, 19, 1731; (b) L. Bogani and W. Wernsdorfer, Nat. Mater., 2008, 7, 179.
- 4 (a) G. A. Timco, T. B. Faust, F. Tuna and R. E. P. Winpenny, Chem. Soc. Rev., 2011, 40, 3067; (b) P. C. E. Stamp and A. Gaita-Ariño, J. Mater. Chem., 2009, 19, 1718.
- 5 (a) S. Hill, R. S. Edwards, N. Aliaga-Alcalde and G. Christou, Science, 2003, 3**02**, 1015; (b) R. Tiron, W. Wernsdorfer, D. Foguet-Albiol, N. Aliaga-Alcalde and G. Christou, Phys. Rev. Lett., 2003, 91, 227203 (1–4).
- 6 (a) A. Kovalev, L. Hayden, G. Bauer and Y. Tscherkovniak, Phys. Rev. Lett., 2011, 106, 147203; (b) D. A. Garanin and E. Chudnovsky, *Phys. Rev.*, 2011, 1, 011005;
- (c) M. Ganzhorn, S. Klyatskaya, M. Ruben and W. Wernsdorfer, Nat. Nanotechnol., 2013, 8, 166.
- 7 (a) W. Wernsdorfer, S. Bhaduri, C. Boskovic, G. Christou and D. N. Hendrickson, Phys. Rev. B: Condens. Matter, 2002, 65, 180403; (b) W. Wernsdorfer, N. E. Chakov and G. Christou, Phys. Rev. Lett., 2005, 95, 037203.
- 8 (a) G. Gatteschi and R. Sessoli, Angew. Chem., Int. Ed., 2003, 42, 268; (b) E. Del Barco, A. D. Kent, S. Hill, J. M. North, N. S. Dalal, E. M. Rumberger, D. N. Hendrickson, N. E. Chakov and G. Christou, J. Low Temp. Phys., 2005, 140, 119; (c) P. C. E. Stamp, Nature, 1996, 383, 125.
- 9 (a) W. Wernsdorfer and R. Sessoli, Science, 1999, 284, 133; (b) W. Wernsdorfer, M. Soler, G. Christou and D. N. Hendrickson, J. Appl. Phys., 2002, 91, 7164.
- 10 (a) J. D. Rinehart and J. R. Long, Chem. Sci., 2011, 2, 2078, and references therein; (b) D. N. Woodruff, R. E. P. Winpenny and R. A. Layfield, Chem. Rev., 2013, 113, 5110; (c) K. R. Meihaus and J. R. Long, Dalton Trans., 2015, 44, 2517 (Perspective).
- 11 (a) A. Ferguson, J. Lawrence, A. Parkin, J. Sanchez-Benitez, K. V. Kamenev, E. K. Brechin, W. Wernsdorfer, S. Hill and M. Murrie, Dalton Trans., 2008, 6409; (b) E.-C. Yang, W. Wernsdorfer, L. N. Zakharov, Y. Karaki, A. Yamaguchi, R. M. Isidoro, G.-D. Lu, S. A. Wilson, A. L. Rheingold, H. Isimoto and D. N. Hendrickson, Inorg. Chem., 2006, 45, 529; (c) A. Bell, G. Aromi, S. J. Teat, W. Wernsdorfer and R. E. P. Winpenny, Chem. Commun., 2005, 2808; (d) M. Moragues-Cánovas, M. Helliwell, L. Ricard, E. Riviére, W. Wernsdorfer, E. Brechin and T. Mallah, Eur. J. Inorg. Chem., 2004, 2219; (e) E.-C. Yang, W. Wernsdorfer, S. Hill, R. S. Edwards, M. Nakano, S. Maccagano, L. N. Zakharov, A. L. Rheingold, G. Christou and D. N. Hendrickson, Polyhedron, 2003, 22, 1727; (f) S. T. Ochseinbein, M. Murrie, E. Rusanov, H. Stoeckli-Evans, C. Sekine and H.-U. Güdel, Inorg. Chem., 2002, 41, 5133; (g) H. Andres, R. Basler, A. J. Blake, C. Cadiou, G. Chabboussant, C. M. Grant, H.-U. Güdel, M. Murrie, S. Parsons, C. Paulsen, F. Semadini, V. Villar, W. Wernsdorfer and R. E. P. Winpenny, *Chem. - Eur. J.*, 2002, 8, 4867. Datton Transactions

The authority of the bublished on 2 is lighter to the published on 2 is lighter to the published on 2 is lighter to the authority of the common Access Commons Article. Commons Articles. The access Art
	- 12 S. Carretta, P. Santini, G. Amorretti, M. Affronte, A. Candini, A. Chirri, I. S. Tidmarsh, R. H. Laye, R. Shaw and E. J. L. McInnes, Phys. Rev. Lett., 2006, 97, 207201.
	- 13 (a) P. Chaudhuri, Coord. Chem. Rev., 2003, 243, 143; (b) A. J. L. Pombeiro and V. Yu Kukushkin, in Comprehensive Coordination Chemistry II, ed. J. A. McCleverty and T. J. Meyer, Elsevier, Amsterdam, 2004, vol. 7, pp. 631–637; (c) C. J. Milios, S. Piligkos and E. K. Brechin, Dalton Trans., 2008, 1809 (Perspective); (d) T. Pathmalingam, S. I. Gorelsky, T. J. Burchell, A.-C. Bédard, A. M. Beauchemin, R. Clérac and M. Murugesu, Chem. Commun., 2008, 2782; (e) C. J. Milios, A. Vinslava, W. Wernsdorfer, S. Moggach, S. Parsons, S. P. Perlepes, G. Christou and E. K. Brechin, J. Am. Chem. Soc., 2007, 129, 2754; (f) L. F. Jones, A. Prescimone, M. Evangelisti and E. K. Brechin, Chem. Commun., 2009, 2023.
- Paper **Dalton Transactions** (Paper Section 2018) **Paper** 2014 **Contractions (Paper Section 2018)**
- 14 For a comprehensive review, see: C. J. Milios, T. C. Stamatatos and S. P. Perlepes, Polyhedron, 2006, 25, 134 (Polyhedron Report).
- 15 (a) H. Kumagai, M. Endo, M. Kondo, S. Kawata and S. Kitagawa, Coord. Chem. Rev., 2003, 237, 197; (b) C. J. Milios, P. Kyritsis, C. P. Raptopoulou, A. Terzis, R. Vicente, A. Escuer and S. P. Perlepes, Dalton Trans., 2005, 501; (c) K. F. Konidaris, C. D. Polyzou, G. E. Kostakis, A. J. Tasiopoulos, O. Roubeau, S. J. Teat, E. Manessi-Zoupa, A. K. Powell and S. P. Perlepes, Dalton Trans., 2012, 41, 2862.
- 16 (a) T. C. Stamatatos, D. Foguet-Albiol, C. C. Stoumpos, C. P. Raptopoulou, A. Terzis, W. Wernsdorfer, S. P. Perlepes and G. Christou, Polyhedron, 2007, 26, 2165; (b) T. C. Stamatatos, D. Foguet-Albiol, C. C. Stoumpos, C. P. Raptopoulou, A. Terzis, W. Wernsdorfer, S. P. Perlepes and G. Christou, J. Am. Chem. Soc., 2005, 127, 15380; (c) A. Escuer, G. Vlahopoulou and F. A. Mautner, Inorg. Chem., 2011, 50, 2717; (d) A. M. Mowson, T. N. Nguyen, K. A. Abboud and G. Christou, Inorg. Chem., 2013, 52, 12320; (e) T. Nguyen, W. Wernsdorfer, K. A. Abboud and G. Christou, J. Am. Chem. Soc., 2011, 133, 20688; (f) T. N. Nguyen, M. Shiddiq, T. Ghosh, K. A. Abboud, S. Hill and G. Christou, J. Am. Chem. Soc., 2015, 137, 7160; (g) T. N. Nguyen, W. Wernsdorfer, M. Shiddiq, K. A. Abboud, S. Hill and G. Christou, Chem. Sci., 2016, 7, 1156. Paper Commutes are c. J. Million, 19 (d. G. Efferics Articles articles Commutes Commutes Articles. Articles and C. Paperbard Commutes Articles. Articles. Articles. Articles. Articles. Articles. Articles. Articles. Article
	- 17 (a) T. C. Stamatatos, A. Escuer, K. A. Abboud, C. P. Raptopoulou, S. P. Perlepes and G. Christou, Inorg. Chem., 2008, 47, 11825; (b) T. C. Stamatatos, K. A. Abboud, S. P. Perlepes and G. Christou, Dalton Trans., 2007, 3861; (c) C. Papatriantafyllopoulou, T. C. Stamatatos, C. G. Efthymiou, L. Cunha-Silva, F. A. A. Paz, S. P. Perlepes and G. Christou, Inorg. Chem., 2010, 49, 9743; (d) S. Zhang, L. Zhen, B. Xu, R. Inglis, K. Li, W. Chen, Y. Zhang, K. F. Konidaris, S. P. Perlepes, E. K. Brechin and Y. Li, Dalton Trans., 2010, 3563; (e) T. C. Stamatatos, C. Papatriantafyllopoulou, E. Katsoulakou, C. P. Raptopoulou and S. P. Perlepes, Polyhedron, 2007, 26, 1830.
	- 18 (a) C. D. Polyzou, H. Nikolaou, C. Papatriantafyllopoulou, V. Psycharis, A. Terzis, C. P. Raptopoulou, A. Escuer and S. P. Perlepes, Dalton Trans., 2012, 13755; (b) C. Papatriantafyllopoulou, G. Aromi, A. J. Tasiopoulos, V. Nastopoulos, C. P. Raptopoulou, S. J. Teat, A. Escuer and S. P. Perlepes, Eur. J. Inorg. Chem., 2007, 2761; (c) C. G. Efthymiou, C. P. Raptopoulou, A. Terzis, S. P. Perlepes, A. Escuer and C. Papatriantafyllopoulou, Polyhedron, 2010, 29, 627; (d) C. Papatriantafyllopoulou, M. Estrader, C. G. Efthymiou, D. Dermitzaki, K. Gkotsis, A. Terzis, C. Diaz and S. P. Perlepes, Polyhedron, 2009, 28, 1652; (e) D. Dermitzaki, C. P. Raptopoulou, V. Psycharis, A. Escuer, S. P. Perlepes and T. C. Stamatatos, Dalton Trans., 2014, 14520; (f) C. Papatriantafyllopoulou, C. P. Raptopoulou, A. Terzis, J. F. Janssens, S. P. Perlepes and E. Manessi-Zoupa, Z. Naturforsch., B: Chem. Sci., 2007, 62, 1123.
- 19 (a) C. G. Efthymiou, A. A. Kitos, C. P. Raptopoulou, A. Escuer, S. P. Perlepes and C. Papatriantafyllopoulou, Polyhedron, 2009, 28, 3177; (b) C. Papatriantafyllopoulou, C. G. Efthymiou, C. Raptopoulou, A. Terzis, E. Manessi-Zoupa and S. P. Perlepes, Spectrochim. Acta, Part A, 2008, 70, 718; (c) C. Papatriantafyllopoulou, T. C. Stamatatos, W. Wernsdorfer, S. J. Teat, A. J. Tasiopoulos, A. Escuer and S. P. Perlepes, Inorg. Chem., 2010, 49, 10486.
- 20 (a) E. Moushi, C. G. Efthymiou, S. P. Perlepes and C. Papatriantafyllopoulou, Int. J. Inorg. Chem., 2011, 606271; (b) T. C. Stamatatos, E. Diamantopoulou, C. P. Raptopoulou, V. Psycharis, A. Escuer and S. P. Perlepes, Inorg. Chem., 2007, 46, 2350.
- 21 C. Papatriantafyllopoulou, L. F. Jones, T. D. Nguyen, N. Matamoros-Salvador, L. Cunha-Silva, F. A. Almeida Paz, J. Rocha, M. Evangelisti, E. K. Brechin and S. P. Perlepes, Dalton Trans., 2008, 3153.
- 22 D. S. Boltin, N. A. Bokach and V. Yu. Kukushkin, Coord. Chem. Rev., 2016, 313, 62.
- 23 H.-Z. Kou, G.-Y. An, C.-M. Ji, B.-W. Wang and A.-L. Cui, Dalton Trans., 2010, 9604.
- 24 C.-M. Ji, H.-J. Yang, C.-C. Zhao, V. Tangoulis, A.-L. Cui and H.-Z. Kou, Cryst. Growth Des., 2009, 9, 4607.
- 25 G.-Y. An, H.-B. Wang, A.-L. Cui and H.-Z. Kou, New J. Chem., 2014, 38, 5037.
- 26 (a) L. Alcazar, M. Font-Bardia and A. Escuer, Eur. J. Inorg. Chem., 2015, 1326; (b) J. Jankolovits, C. M. Andolina, J. W. Kampf, K. N. Raymond and V. L. Pecoraro, Angew. Chem., Int. Ed., 2011, 50, 9660.
- 27 E. Bernasek, J. Org. Chem., 1957, 22, 1263.
- 28 T. Kottke and D. Stalke, J. Appl. Crystallogr., 1993, 26, 615.
- 29 APEX2, Data Collection Software Version 2.1-RC13, Bruker AXS, Delft, The Netherlands, 2006.
- 30 Cryopad, Remote monitoring and control, Version 1.451, Oxford Cryosystems, Oxford, United Kingdom, 2006.
- 31 SAINT+, Data Integration Engine v. 7.23a ©, Bruker AXS, Madison, Wisconsin, USA, 1997–2005.
- 32 G. M. Sheldrick, SADABS v.2.01, Bruker/Siemens Area Detector Absorption Correction Program, Bruker AXS, Madison, Wisconsin, USA, 1998.
- 33 G. M. Sheldrick, SHELXS-97, Program for Crystal Structure Solution, University of Göttingen, 1997.
- 34 G. M. Sheldrick, SHELXL-97, Program for Crystal Structure Refinement, University of Göttingen, 1997.
- 35 A. L. Spek, Acta Crystallogr., Sect. A: Fundam. Crystallogr., 1990, 46, C34; A. L. Spek, J. Appl. Crystallogr., 2003, 36, 7.
- 36 P. Van der Sluis and A. L. Spek, Acta Crystallogr., Sect. A: Fundam. Crystallogr., 1990, 46, 194.
- 37 C. Papatriantafyllopoulou, C. P. Raptopoulou, A. Terzis, E. Manessi-Zoupa and S. P. Perlepes, Z. Naturforsch., B: Chem. Sci., 2006, 61, 37.
- 38 J. Lopez-Carriga, G. T. Badcock and J. F. Harrison, J. Am. Chem. Soc., 1986, 108, 7241.
- 39 A. B. P. Lever and A. Mantovani, Inorg. Chem., 1971, 10, 817.
- 40 (a) F.-R. Dai, D. C. Becht and Z. Wang, Chem. Commun., 2014, 5385; (b) B. Biswas, S. Khanra, T. Weyhermuller and P. Chaudhuri, Chem. Commun., 2007, 1059; (c) K. V. Shuvaev, S. S. Tandon, L. N. Dawe and L. K. Thompson, Chem. Commun., 2010, 4755; (d) H. Jiang, T. Sheng, S. Bai, S. Hu, X. Wang, R. Fu, P. Yu and X. Wu, Inorg. Chem., 2013, 52, 12305; (e) K. Su, F. Jiang, J. Qian, Y. Gai, M. Wu, S. M. Bawaked, M. Mokhtar, S. A. AL-Thabaiti and M. Hong, Cryst. Growth Des., 2014, 14, 3116. Oattoo Transactions

2016 P. E. Download, C. Horsel, S. Kunstn, T. Westmanshed on 2014, S. 2024 3:33, (b) B. Bossel, S. S. 2024 3:30, (b) B. E. Channel Commons, Channel Commons, 2014, Unported University Commons, 2014, Car
	- 41 W. H. Press, S. A. Teukolsky, W. T. Vetterling and B. P. Flannery, Numerical Recipes in C: The Art of Scientific

Computing, Cambridge University Press, Cambridge, 2nd edn, 1992.

- 42 G.-Y. An, C.-M. Ji, A.-L. Cui and H.-Z. Kou, Inorg. Chem., 2011, 50, 1079.
- 43 (a) M. Evangelisti, F. Luis, L. J. De Jongh and M. Affronte, J. Mater. Chem., 2006, 16, 2534; (b) M. Evangelisti, Molecular Magnets, NanoScience and Technology, in Molecule-based Magnetic Coolers: Measurement, Design and Application, ed. J. Bartolomé, F. Luis and J. F. Fernandez, Springer-Verlag, Berlin, Heidelberg, 2014, pp. 365–387.