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# Changes in magnetic order through two consecutive dehydration steps of metal-phosphonate diamond chains†

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Hydrothermal reactions of the multitopic ligand 1-hydroxy-1-(piperidin-4-yl)methylidenebisphosphonic acid (hpdpH<sub>4</sub>) with cobalt or nickel sulfates afforded two new isostructural metal phosphonates,  $M_3^{II}(hpdpH)_2(H_2O)_6\cdot 4H_2O)$  [M = Co (Co-10H<sub>2</sub>O), Ni (Ni-10H<sub>2</sub>O)]. Their structures consist of parallel diamond chains of three MO<sub>6</sub> octahedra bridged by the PO<sub>3</sub>C tetrahedra. Six of the seven oxygen atoms of the ligand are involved in coordination; for two ligands that amounts to 12 bonds for 3 MO<sub>6</sub> and the remaining six are occupied by terminal water molecules. In addition, four water molecules sit in between the chains providing H-bonds to the formation of a 3D-net. Thermal analyses show identical two-step dehydration processes involving first the departure of six water molecules followed by the remaining four. A detailed study of the ac- and dc-magnetization as a function of temperature, field and frequency reveals associated drastic changes. The virgin form Co-10H<sub>2</sub>O is a paramagnet while its partial dehydrated form Co-4H<sub>2</sub>O is an antiferromagnet displaying canting below  $T_N = 4.7$  K and the fully dehydrated form Co is a ferrimagnet ( $T_C = 12$  K). Ni-10H<sub>2</sub>O and Ni-4H<sub>2</sub>O exhibit long-range ordered antiferromagnetism ( $T_N = 2.7$  and 4.0 K, respectively) and also become ferrimagnets ( $T_C = 9.4$  K) when fully dehydrated to Ni. The dehydrated samples can be fully rehydrated with the complete recovery of both the structures and magnetic properties.

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# Introduction

The chemistry of metal phosphonates has been receiving an increasing amount of attention because of their potential application in catalysis, ion-exchange, and gas adsorption, as well as having optical, magnetic and proton conducting properties. The derivatives of methylenediphosphonates  $(H_2O_3-PC(R')(R'')PO_3H_2)$  are capable of binding metal ions in different coordination modes, thus resulting in metal phosphonates with versatile architectures and interesting associated physical properties. For example, a series of metal diphosphonate compounds based on 1-hydroxylidene-diphosphonate [hedpH4,  $CH_3C(OH)(PO_3H_2)_2$ ] have been prepared which show structures ranging from 0D mononuclear species or clusters, 1D chains, 2D layers to 3D frameworks. The modification of the organic

groups may easily alter the structures and, consequently, their

magnetic changes to external stimuli are of great interest as candidates for switchable molecular magnets and as potential sensors. Such changes can be induced by temperature, 10 light, 11 redox,12 desorption/adsorption of guest molecules or coordinated water molecules.13 Although dehydration induced magnetic changes render ground-state switching in a number of coordination systems,14 it has rarely been described for cobalt15,16 or nickel17 phosphonate systems. Examples include a layer compound  $Co(2\text{-pmp})(H_2O)_2$  (2-pmpH<sub>2</sub> = 2-pyridylmethylphosphonic acid) with tuneable metamagnetic behaviour, and  $[CoLa(notp)(H_2O)_4] \cdot nH_2O$   $[notpH_6 = 1,4,7-1]$ triazacyclononane-1,4,7-triyl-tris(methylenephosphonic acid)] exhibiting oxidation state change of cobalt ion during the dehydration process.16 Furthermore, only a few examples are known to change the magnetic ground state upon desolvation and in general, rarely in two consecutive desolvation processes.14

In this paper, we report two new metal phosphonates based on 1-hydroxy-1-(piperidin-4-yl)methylidenebisphosphonic acid

physical properties. Thus when the methyl group in hedpH<sub>4</sub> is replaced by pyridyl-, imidazole- or other groups, new types of chain or layer architectures can be obtained showing interesting magnetic or optical properties.<sup>7-9</sup>

On the other hand, cobalt compounds showing drastic

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 $<sup>\</sup>dagger$  Electronic supplementary information (ESI) available: TGA, IR spectra, and additional magnetic property. CCDC 1901442 (Co-10H<sub>2</sub>O) and 1920728 (Ni-10H<sub>2</sub>O). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9ra05722f

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 $[(3-C_5H_{10}N)C(OH)(PO_3H_2)_2, hpdpH_4]$  (Fig. 1a) using two divalent first row transition metals (Co and Ni) for their different magnetic anisotropies, namely, Co<sub>3</sub>(hpdpH)<sub>2</sub>(H<sub>2</sub>O)<sub>6</sub>·4H<sub>2</sub>O (Co- $10H_2O$ ) and Ni<sub>3</sub>(hpdpH)<sub>2</sub>(H<sub>2</sub>O)<sub>6</sub>·4H<sub>2</sub>O (Ni-10H<sub>2</sub>O) as a comparison. The hpdpH4 was selected because the amino group in piperidine can not only serve as a potential coordination donor but also be protonated to adjust the overall charge of the phosphonate ligand, thus providing possibilities to construct new architectures of metal diphosphonates. Both compounds show a one-dimensional chain structure. Interestingly, the two compounds can be dehydrated in two-steps, accompanied with distinct magnetic changes going from paramagnet to canted antiferromagnet/antiferromagnet to ferrimagnet. Moreover, the fully dehydrated samples can be rehydrated with the recovery of both the structures and magnetic properties.

# Experiment section

#### **Materials**

1-Hydroxy-1-(piperidin-4-yl)methylidenebisphosphonic acid (hpdpH<sub>4</sub>) was prepared according to the literature method. <sup>18</sup> All the other starting materials were obtained commercially as reagent grade chemicals and used without further purification.

Synthesis of  $Co_3(hpdpH)_2(H_2O)_6 \cdot 4H_2O$  (Co-10H<sub>2</sub>O). A mixture of CoSO<sub>4</sub>·7H<sub>2</sub>O (0.1 mmol, 0.029 g), hpdpH<sub>4</sub> (0.1 mmol, 0.028 g) and  $H_2O$  (8 cm<sup>3</sup>), adjusted to pH = 3.5 with 1 M NaOH, was kept in a Teflon-lined autoclave (25 cm<sup>3</sup>) at 140 °C for 72 h. After cooling the autoclave to room temperature, red block crystals of Co-10W were collected as a monophasic material, judged by powder X-ray diffraction pattern (Fig. S1†). Yield: 8.7 mg (30% based on Co). Elemental analyses calcd for

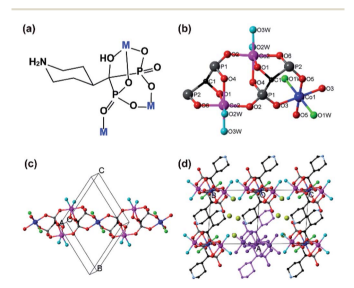


Fig. 1 (a) The coordination mode of hpdpH<sup>3-</sup> ligand to three metal centres in compounds Co-10H2O and Ni-10H2O. Structure represented by Co-10H2O: (b) a building unit with atomic labelling scheme, (c) fragment of a chain showing the octahedra bridged by PO<sub>3</sub>C tetrahedra, and (d) packing of the chains where one is highlighted in purple. The coordinated water molecules are identified by different colours, cyan in cis-positions of Co2, green in trans-positions of Co1 and yellow for non-coordinated ones of the lattice.

C<sub>12</sub>H<sub>44</sub>Co<sub>3</sub>N<sub>2</sub>O<sub>24</sub>P<sub>4</sub> (%): C, 15.99; H, 4.92; N, 3.11. Found: C, 15.43; H, 4.81; N, 2.98. IR (KBr, cm<sup>-1</sup>): 3403br, 3067m, 1619s, 1486w, 1458w, 1384m, 1331w, 1274w, 1123vs, 1094s, 1076s, 1047s, 1012s, 953s, 925m, 811m, 642m, 574s, 478w, 444m.

Synthesis of  $Ni_3(hpdpH)_2(H_2O)_6 \cdot 4H_2O$  (Ni-10H<sub>2</sub>O). This compound was prepared following a similar procedure to that for Co-10H<sub>2</sub>O except that NiSO<sub>4</sub>·7H<sub>2</sub>O (0.1 mmol, 0.026 g) was used instead of CoSO<sub>4</sub>·7H<sub>2</sub>O as the starting material and at 160 °C. Green needle crystals of Ni-10H2O were obtained as a monophasic material, judged by powder X-ray diffraction pattern (Fig. S2†). Yield: 8.9 mg (30% based on Ni). Elemental analyses calcd for C<sub>12</sub>H<sub>44</sub>Ni<sub>3</sub>N<sub>2</sub>O<sub>24</sub>P<sub>4</sub> (%): C, 16.01; H, 4.93; N, 3.11. Found: C, 16.01; H, 5.09; N, 3.04. IR (KBr, cm<sup>-1</sup>): 3415br, 3070m, 1619s, 1488w, 1461w, 1386m, 1326w, 1126s, 1105s, 1081s, 1051s, 1010s, 956m, 925w, 908w, 860m, 815w, 647m, 574s, 518m, 476m.

#### Physical measurements

The elemental analyses for C, H and N were performed in a PE240C elemental analyzer. The infrared spectra were recorded on a Bruker Tensor 27 spectrometer with ATR mode. Thermal analyses were performed in nitrogen with a heating rate of 5 °C min<sup>-1</sup> on a TGA-DTA V1.1b Inst 2100 instrument. The powder XRD patterns were recorded on a Shimadzu XD-3A X-ray diffractometer. Magnetization were measured using a Quantum Design SQUID VSM magnetometer. The magnetization data were corrected for the diamagnetic contributions of both the sample holder and the compound obtained from Pascal's constants.19

#### X-ray crystallographic studies

Single crystals of dimensions  $0.20 \times 0.15 \times 0.10 \text{ mm}^3$  for Co- $10H_2O$  and  $0.12 \times 0.10 \times 0.08 \text{ mm}^3$  for Ni- $10H_2O$  were selected for indexing and intensity data collection on a Bruker SMART APEX CCD diffractometer using graphite monochromatized Mo Kα radiation ( $\lambda = 0.71073 \text{ Å}$ ) at room temperature. A hemisphere of data was collected in each case in the  $\theta$  range 2.275– 25.242° for Co-10H<sub>2</sub>O and 2.277-25.242° for Ni-10H<sub>2</sub>O using a narrow-frame method with scan widths of  $0.30^{\circ}$  in  $\omega$  and exposure time of 10 s per frame. Numbers of observed and unique reflections are 11 163 and 2594 ( $R_{\rm int}=0.0139$ ) for Co- $10H_2O$ , 7534 and 2605 ( $R_{\text{int}} = 0.0255$ ) for Ni- $10H_2O$ . The data were integrated using the Siemens SAINT program,20 with the intensities corrected for Lorentz factor, polarization, air absorption, and absorption due to variation in the path length through the detector faceplate. Absorption corrections were applied. The structures were solved by direct methods and refined on F2 by full matrix least squares using SHELXTL.21 All non-hydrogen atoms were located from the Fourier maps and refined with anisotropic thermal parameters. All the hydrogen atoms not located from the Fourier maps were set at calculated positions and refined with isotropic parameters related to the non-hydrogen atom to which they are bonded. Crystallographic and refinement details are listed in Table 1. The selected bond lengths and angles for compounds Co-10H2O and Ni-10H2O are given in Tables S1 and S2,† respectively.

Table 1 Crystallographic data for Co-10H<sub>2</sub>O and Ni-10H<sub>2</sub>O

	Co-10H <sub>2</sub> O	Ni-10H <sub>2</sub> O
Empirical formula	$C_{12}H_{44}Co_3N_2O_{24}P_4$	$C_{12}H_{44}Ni_3N_2O_{24}P_4$
$F_{ m w}$	901.16	900.50
Crystal system	Triclinic	Triclinic
Space group	$P\bar{1}$	$Par{1}$
a (Å)	9.064(4)	9.056(3)
b (Å)	9.536(4)	9.430(3)
$c(\mathring{A})$	9.686(4)	9.605(3)
α (°)	114.883(6)	114.954(5)
β (°)	97.803(7)	97.790(5)
$\gamma$ (°)	90.634(6)	90.883(6)
$V(\mathring{A}^3), Z$	750.3(5), 1	734.4(4), 1
$D_{\rm c}$ (g cm <sup>-3</sup> )	1.994	2.036
$\mu  (\mathrm{mm}^{-1})$	1.953	2.224
F(000)	463	466
$R_1^a$ , w $R_2^b$ $[I > 2\sigma(I)]$	0.0374, 0.1187	0.0561, 0.1141
$R_1^a$ , w $R_2^b$ (all data)	0.0398, 0.1223	0.0767, 0.1226
Goodness-of-fit	1.005	1.001
$(\Delta \rho)$ max, min (e Å <sup>-3</sup> )	0.58, -0.83	0.86, -0.69
CCDC number	1901442	1920728

 $<sup>^{</sup>a}R_{1} = \sum ||F_{0}| - |F_{c}||/\sum |F_{0}|.$   $^{b}WR_{2} = [\sum W(F_{0}^{2} - F_{c}^{2})^{2}/\sum W(F_{0}^{2})_{2}]^{1/2}.$ 

# Results and discussion

#### **Synthesis**

Crystals of Co-10H2O and Ni-10H2O were prepared by the reaction of the metal sulfates and hpdpH4 under hydrothermal conditions. The M: ligand ratio, pH and reaction temperature could play critical roles in yielding pure phases. Co-10H2O with good crystal quality was prepared as a pure phase at 140 or 160 °C, pH of 3.0-3.5 and the Co: hpdpH<sub>4</sub> molar ratio of 1 : 1-2. At a low pH (<3.0), only clear solution was obtained. At moderate pH (5-7), powder materials were obtained which showed the same PXRD patterns with that of Co-10H<sub>2</sub>O (Fig. S3†). For Ni-10H<sub>2</sub>O, a pure phase with good crystal quality was obtained at 160 °C when the pH ranged between 2.5 and 4.0 and the Ni: hpdpH<sub>4</sub> molar ratio is 1:1. Reactions in the pH range 5.0-7.0 resulted in crystals of Ni-10H<sub>2</sub>O together with a large amount of powdered materials which had the same PXRD patterns with that of Ni-10H<sub>2</sub>O (Fig. S3†). When the Ni: hpdpH<sub>4</sub> molar ratio is lowered to 1:0.5, bigger block-shaped crystals of Ni-10H<sub>2</sub>O can be obtained but in a rather low yield. Notably, although the powder samples obtained at higher pH show identical PXRD patterns in both cases, the presence of impurities in the powder products cannot be fully excluded. Therefore, only crystals of M-10H2O were selected to be used for further physical measurements.

#### **Description of the structures**

The key feature of the two isostructural compounds differing only in the metals, Co and Ni, is the diamond chain of the metals along the [011] direction of the triclinic cell (Fig. 1 and Table 1). The piperidine alternately points above and below along the chains. The chains are packed parallel to one another forming layers separated by the piperidine and four water of

crystallization per formula unit, M3<sup>II</sup>(hpdpH)<sub>2</sub>(H<sub>2</sub>O)<sub>6</sub>·4H<sub>2</sub>O. The asymmetric unit contains 1.5 metal, 1 ligand and 5H<sub>2</sub>O. Thus, the ligand has lost three of its five protons to hpdpH<sup>3-</sup>. As we anticipated with our design of the ligand, it uses a high number of six of the eight potential coordinating atoms to connect the metal centers. It makes three different modes of coordination: a six-membered chelate M1-O-P-C-P-O-M1, a capping tripod to M2 through two five-membered chelates involving the alcohol oxygen atom, and a single terminal bond to M2. This leaves one phosphonate oxygen and the nitrogen atoms free from coordination.

The two independent metal ions adopt slightly distorted octahedral geometries where four of their oxygen atoms come from the phosphonates and the other two from coordinated water molecules. The latter are in trans-position for M1 but they are in cis-position for M2. The metal-oxygen bond lengths and angles are all normal for M(II) octahedra (Tables S1 and S2†). All the metals within a trimeric unit are connected through O-P-O bridges. The asymmetric triangle is disposed of one M1-M2 (4.883 Å for Co-10H<sub>2</sub>O and 4.836 Å for Ni-10H<sub>2</sub>O) edge consisting of two anti-anti O-P-O, one M1-M2 (6.365 Å for Co-**10H<sub>2</sub>O** and 6.297 Å for **Ni-10H<sub>2</sub>O**) with a single anti-anti O-P-O and one M2-M2 (4.658 Å for Co-10H2O and 4.617 Å for Ni-10H<sub>2</sub>O) with two equivalent syn-anti O-P-O. The piperidine nitrogen (N1) is protonated. One interchain H-bond is found between the piperidine nitrogen and phosphonate oxygen (O5) atoms (N1···O5: 2.769 Å for Co-10H2O and 2.774 Å for Ni-10H<sub>2</sub>O), leading to a layer in the (0 1-1) plane (Tables S3 and S4†). The interlayer space is occupied by lattice water molecules, extensively H-bonded through the coordinated and lattice water molecules as well as the phosphonate oxygen atoms, leading to three-dimensional supramolecular network structure (Fig. S4†).

It should be mentioned that the structures of Co-10H2O and Ni-10H<sub>2</sub>O are distinguished from that of a related Cu<sub>3</sub>-<sup>II</sup>(hpdpH)<sub>2</sub>(H<sub>2</sub>O)<sub>4</sub>·4H<sub>2</sub>O which includes the same phosphonate ligand.22 Although the chemical composition of the copper compound is quite similar, with two less coordinated water molecules, it shows an infinite ladder-like double chain structure in which the Cu1O<sub>5</sub> square pyramids and Cu2O<sub>6</sub> octahedra are linked by corner-sharing PO<sub>3</sub>C tetrahedra. The chains are stacked in a perpendicular manner, forming a 3D H-bonded network. Both the intrachain arrangement and the stacking mode of the copper compound are completely different from those of Co-10H2O and Ni-10H2O. Thus compounds Co-10H2O and Ni-10H<sub>2</sub>O provide second example of metal phosphonates based on hpdpH4 ligand with a new structural type.

#### Thermal analysis

The two compounds behave in exactly similar way when they are heated at a rate of 5 °C minute<sup>-1</sup> under a dry nitrogen atmosphere (Fig. 2). First they lose six water molecules between 80 and 150 °C with a weight loss of 12.2% (calcd 12.0%) for Co-10H<sub>2</sub>O and 12.0% (calcd 12.0%) for Ni-10H<sub>2</sub>O, which is followed by a second dehydration of four water molecules between 180 and 250 °C with a weight loss of 7.9% (calcd 8.0%) for Co-10H<sub>2</sub>O RSC Advances Paper

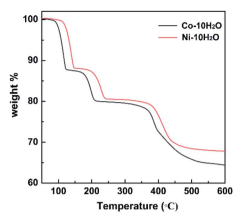


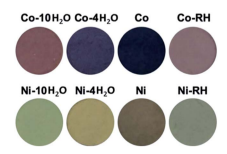
Fig. 2 Thermogravimetric analyses for  $\text{Co-10H}_2\text{O}$  (black) and  $\text{Ni-10H}_2\text{O}$  (red) showing the thermally transformed products.

and 8.0% (calcd 8.0%) for Ni-10H<sub>2</sub>O. Interestingly, the nickel compound exhibits ca. 20 °C higher transformation temperatures than the cobalt one, indicating the stronger binding energies for the former most likely associated with its slightly small ionic radius, as seen in the structural analyses described above (Tables 1, S2 and S3†). A third step follows which corresponds to the decomposition of the piperidine moiety leaving a methyl group. The resulting phase is stable from 500 to 800 °C before it decomposes to metal-oxide and -phosphate.

Since distinct plateau are observed in the thermogravimetry curves, the partially dehydrated  $M_3(hpdpH)_2(H_2O)_4$  (Co-4H<sub>2</sub>O, Ni-4H<sub>2</sub>O) and fully dehydrated  $M_3(hpdpH)_2$  (Co, Ni) were prepared by heating Co-10H<sub>2</sub>O at 140 °C and 230 °C and Ni-10H<sub>2</sub>O at 170 °C and 250 °C, respectively, for 20 minutes for other physical property measurements. The number of remaining coordinated water molecules in Co-4H<sub>2</sub>O and Ni-4H<sub>2</sub>O were verified by further TG analyses (Fig. S5 and S6†).

A strong colour change for the powdered crystals from red to purple to deep blue is observed for Co-10H<sub>2</sub>O upon the two-step dehydration (Fig. 3). While for Ni-10H<sub>2</sub>O, the colour changes from a light green, yellow to brown. On rehydrating the colour of the powders reverts back but with a little bit less intensity.

The crystallinity of the products of the thermal degradation was checked by powder X-ray diffraction because all attempts using selected crystals indicate they are heavily fragmented with very wide mosaic that hampered structural determination. The PXRD of the partially and fully hydrated phases show



**Fig. 3** Photos of the powdered crystals showing the colour changes during the two dehydration processes and after rehydration.

reasonable crystallinity and major changes to the unit cells from the virgin crystals (Fig. 4). However, the quality of the data remains low and has prevented further structural analyses. Interestingly, upon rehydration in water vapor at 70 °C for 3 days the original hydrated forms are reproduced, as confirmed by PXRD and IR measurements (Fig. 4, S7 and S8†).

# **Magnetic properties**

The magnetic properties of the two compounds have been studied by a series of experiments including (a) the susceptibilities on cooling in an applied field of either 100, 1000 or 2000 Oe, (b) the zero-field-cool (ZFC) and field-cool (FC) magnetizations in an applied field of 10 Oe, (c) field dependence of the isothermal magnetization at 2 K after ZFC and the acsusceptibilities in an oscillating field of 2 Oe for three different frequencies (73, 333 and 999 Hz). These data are shown in Fig. 5 for all the samples as clarified in the appropriate captions. The principal results of the analyses of the data are given in Table 2.

The Curie and Weiss constants for the cobalt phases were estimated from least-square fits of the inverse susceptibility data *versus* temperature in the range 150 to 300 K due to the effect of the large spin–orbit coupling. The Curie constants are within the range expected for  $Co(\pi)$  in the octahedral geometry. The large value is due to both spin and orbital contributions. The Weiss temperatures, a measure of the average magnetic coupling between the nearest neighbors, are negative for the three forms. The values for  $Co-10H_2O$  (-14.2 K) and  $Co-4H_2O$  (-15.4 K) are slightly low compared to that expected (-20 K) for an octahedral  $Co(\pi)$  whereas that of Co (-25.2 K) is slightly high, indicating a stronger antiferromagnetic interaction in the fully dehydrated Co compound.

The Curie and Weiss constants for the nickel phases were estimated from similar fits of the susceptibility data in the wider range 50 to 300 K due to the weak effect of single-ion anisotropy. The Curie constants for the three phases correspond to a total of three S=1 Ni( $\pi$ ) in the octahedral geometry. The Weiss temperatures are negative for the three nickel forms with little changes from one another (Table 2), indicating a dominant antiferromagnetic interactions mediated between the Ni( $\pi$ ) centers.

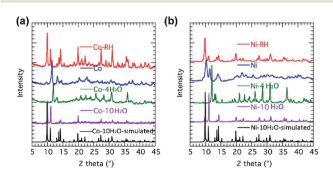


Fig. 4 Powder X-ray diffraction patterns for  $Co-10H_2O$  (a) and  $Ni-10H_2O$  (b) and their partially dehydrated ( $Co-4H_2O$  and  $Ni-4H_2O$ ), fully dehydrated (Co and Ni) and the rehydrated forms (Co-RH and Ni-RH) compared to those simulated from single crystal structural data.

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(a) 3.00F 0.30 0.25 2.00 x (cm² mol¹) 0.20 cm<sup>3</sup> 0.15 × 0.10 0.05 0.00 0.00 100 10 100 Temperature (K) Temperature (K) (d) (c) 3.00 0.30 ---FC 0.25 \_lom χ (cm² mol¹) -ZFC 0.20 x (cm 0.15 0.10 0.05 0.00 100 10 10 100 Temperature (K) Temperature (K) 30.0 · χ<sub>M</sub> →-FC • FC 120 20.0 x (cm³ mol¹) 10.0 x (cm mol1) ZFC ZFC -χ' x 20-80 χ" x 20

Fig. 5 Temperature dependence of the ac- and dc-magnetic susceptibilities for (a)  $Co-10H_2O$ , (b)  $Ni-10H_2O$ , (c)  $Co-4H_2O$ , (d)  $Ni-4H_2O$ , (e) Co and (f) Ni.

100

Temperature (K)

While the susceptibility data at low temperature for Co- $10H_2O$  are featureless, those for Co- $4H_2O$  and Co reveal longrange ordering. For Co- $4H_2O$ , complementary ZFC-FC data in an applied field of 10 Oe indicate a fully compensated antiferromagnetic state at  $T_N = 4.7$  K, with coincidence of the ZFC and FC data, followed by a canting of the moments below 3.5 K, where a bifurcation is observed. The ac-susceptibilities are

independent of frequency and  $\chi'$  follows the ZFC dc-susceptibility while  $\chi''$  only become finite below 3.5 K. This behavior is similar to those observed for  $\text{Co}_2(\text{pyromellitate})$ ,  $\text{Co}_2(\text{OH})_2(\text{terephthalate})$  and  $\text{Co}_4(\text{OH})_2(2\text{-pmb})_2$  [2-pmbH $_3=2$ -(phosphonomethyl)benzoic acid]. For  $\mathbf{Co}$  both ZFC-FC and acsusceptibilities data show a clear bifurcation associated with the anomaly in  $\chi''$  at 12 K.

The isothermal magnetizations were measured at 2 K (Fig. 6). For Co-10H<sub>2</sub>O, a paramagnetic response is observed but it does not reach saturation, which implied some antiferromagnetic coupling between neighboring moment carriers. For Co-4H2O, a complicated behavior emerges with two metamagnetic transformations, one at low field of 30 Oe with a magnetization plateau at 0.15  $\mu_B$  and the other at 1.5 kOe to a plateau of 3  $\mu_B$ , followed by a third gradual increase starting at 25 kOe rising to 6  $\mu_{\rm B}$ . The two low-field transformations are hysteretic, 30 Oe and 150 Oe, respectively. They suggest there are two metamagnetic reversals of different energies from an antiferromagnetic ground state. Given the difference, we associate tentatively the first at 30 Oe to that between layers and the second at 1.5 kOe to that between chains. For Co, there is a single hysteresis loop with a coercive field of 850 Oe and a linear dependence with field up to 70 kOe. This is interpreted as a ferrimagnetic state.26

The nickel series is less complicated. Both Ni-10H<sub>2</sub>O and Ni-4H<sub>2</sub>O behave in similar fashion with long-range antiferromagnetism at  $T_{\rm N}=2.7$  and 3.8 K, respectively and only the real part of the ac-susceptibilities is active and duplicating the desusceptibility. For Ni, all the dc- and ac-susceptibilities have behavior akin that of Co, where the ferrimagnetism sets in at 9.4 K.

The isothermal magnetizations of Ni-10H<sub>2</sub>O and Ni-4H<sub>2</sub>O again behave in similar fashion with a metamagnetic critical field of ca. 10 kOe and saturation approaching 2.2  $\mu_B$ , which would correspond to that of one Ni(II). While the high field may be associated to the spin-flop, the saturation values suggest a ferrimagnetic state results. For Ni, a wide hysteresis with a coercive field of 3 kOe is observed and the saturation in 70 kOe is only 2.5  $\mu_B$ . This is consistent with a ferrimagnetic state.<sup>27</sup>

In the absence of a full structural details of the two less hydrated phases, it is difficult to comment on any structuremagnetic property relationship. However, certain information

 Table 2
 Results of analyses of the magnetization data

	Curie constant (cm <sup>3</sup> K mol <sup>-1</sup> )	Weiss constant (K)	Critical temperature (K)	Critical field (Oe)	Coercive field (Oe)	Saturation moment $(\mu_B)$
Co-10H <sub>2</sub> O	9.62(2)	-14.2(1)	_	_	_	5.9
Co-4H <sub>2</sub> O	9.70(2)	-15.4(3)	$T_{ m N}=4.7$	$H_{c1} = 60$	120	0.15
			$T_{\mathrm{CAF}}=3.5$	$H_{\rm c2}=1500$	150	3
				$H_{\rm c2} = 25~000$	_	6
Co	9.33(2)	-25.2(2)	$T_{ m C}=12$	_	850	4.4
Co-RH	9.34(3)	-16.1(1)	_	_	_	5.8
Ni-10H <sub>2</sub> O	3.75(1)	-13.7(4)	$T_{ m N} = 2.7$	$H_{\rm c} = 10~000$	_	2.1
Ni-4H <sub>2</sub> O	3.80(1)	-16.7(4)	$T_{ m N}=4.0$	$H_{\rm c} = 10~000$	_	2.1
Ni	3.45(1)	-10.8(3)	$T_{ m C} = 9.4$	_	3000	2.5
Ni-RH	3.75(1)	-13.0(3)	$T_{ m N}=\sim 3$	$H_{\rm c} = 10~000$	_	2.1

100

<sup>&</sup>lt;sup>a</sup> CAF = canted antiferromagnet.

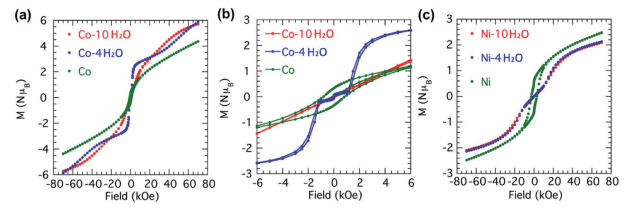


Fig. 6 Isothermal magnetization at 2 K for (a) Co-10H<sub>2</sub>O, Co-4H<sub>2</sub>O, and Co in the field of -70 to 70 kOe; (b) Co-10H<sub>2</sub>O, Co-4H<sub>2</sub>O, and Co in the range of -6 to 6 kOe; (c) Ni-10H<sub>2</sub>O, Ni-4H<sub>2</sub>O and Ni in the range of -70 to 70 kOe.

are justified. One is that the spin states of the different moment carriers within the compounds remain unchanged upon transformation through the two dehydration processes. We can be certain that dehydration of the nickel compound does not form square planar Ni(II) geometry (Table 2) which will be diamagnetic. Whether it adopts a five or six coordination is not certain. We hypothesize that the first-step dehydration process could remove four lattice and two coordination water molecules, leading to the formation of Co-4H2O with a layer structure and thus the inter-chain distance is significantly reduced (Fig. 1d). The clear two steps in the isothermal magnetization observed for Co-4H2O is a direct measure of the weak interactions between the chains within the layer that follows between layers. From the critical fields of the metamagnetism an estimate gives the interchain coupling averaging to ca. 0.1 K and interlayer one of ca. 2 mK (Fig. 6b). Such low coupling energies are what drives these very subtle metamagnetic transformations. Another parameter that can be confirmed is the enhancement of the antiferromagnetic exchange within and between the chains upon total dehydration, which leads to an enhanced ordering temperatures of the ferrimagnetic state in both cases. Interestingly, the rehydrated samples Co-RH and Ni-RH show almost the same magnetic behavior as those of the pristine ones (Fig. S9-S12†). The result indicates that both the structures and magnetic properties of Co-10H2O and Ni-10H2O are reversible upon dehydration and rehydration.

## Conclusions

We report two isostructural compounds,  $M_3^{II}(hpdpH)_2(H_2O)_6$ -  $\cdot 4H_2O$  (M = Co and Ni) based on a polytopic ligand hpdpH<sup>3-</sup>. They contain diamond chains of the metals bridged by O-P-O and connected to one another through H-bonds involving  $H_2O\cdots PO_3$  and  $H_2O\cdots NH_2$  resulting in a 3D-net. The most significant feature is that both compounds undergo a consecutive two-step dehydration process, accompanied by dramatic changes in the magnetic properties – from paramagnet to antiferromagnet/metamagnet to ferrimagnet for M = Co and from antiferromagnet to antiferromagnet to ferrimagnet for M = Ni. Such an interesting magnetic changes must be related to

their structural transformations upon thermal treatment, possibly from chain  $(M-10H_2O)$  to layer  $(M-4H_2O)$  to more dense layer (M). Moreover, the dehydration/rehydration process is reversible with the recovery of both structures and magnetic behaviors. Further work is in progress to explore new metal phosphonates with their magnetic properties responsible to the external stimulus such as heat and light.

# Conflicts of interest

There are no conflicts to declare.

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