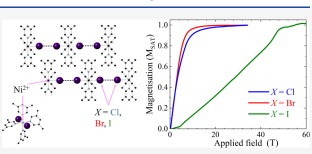
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ABSTRACT: We present the magnetic properties of a new family of S = 1 molecule-based magnets, NiF₂(3,5-lut)₄·2H₂O and NiX₂(3,5 $lut)_4$, where X = HF₂, Cl, Br, or I (lut = lutidine C₇H₉N). Upon creation of isolated Ni-X···X-Ni and Ni-F-H-F···F-H-F-Ni chains separated by bulky and nonbridging lutidine ligands, the effect that halogen substitution has on the magnetic properties of transition-metal-ion complexes can be investigated directly and in isolation from competing processes such as Jahn-Teller distortions. We find that substitution of the larger halide ions turns on increasingly strong antiferromagnetic interactions between adjacent



Ni²⁺ ions via a novel through-space two-halide exchange. In this process, the X···X bond lengths in the Br and I materials are more than double the van der Waals radius of X yet can still mediate significant magnetic interactions. We also find that a simple model based on elongation/compression of the Ni²⁺ octahedra cannot explain the observed single-ion anisotropy in mixed-ligand compounds. We offer an alternative that takes into account the difference in the electronegativity of axial and equatorial ligands.

INTRODUCTION

Theoretical work on low-dimensional magnets¹ and advances in material design² have led to a vast number of experimental investigations into quantum magnetism. This is partly driven by possible applications in data storage, catalysis, and quantum among others. Producing designer magnetic computing,³ systems for physical applications or experimental tests of theoretical predictions is of great interest to physicists and chemists alike.

There have been many experimental studies of the magnetism of low-dimensional $S = \frac{1}{2}$ systems in recent years, such that the influence of structure and composition on the magnetic properties is reasonably well understood.⁷⁻¹¹ This allows experiments on these systems to explore exotic states, such as magnetic Bose-Einstein condensates of magnons¹²⁻¹⁴ or quantum spin liquids.¹⁵ However, investigations into S = 1 magnetic compounds are more rare due to the more complex ground state that these materials exhibit. The S = 1 Hamiltonian is given by

$$\begin{aligned} \hat{\mathcal{H}} &= D \sum_{i} (\hat{S}_{i}^{z})^{2} + E \sum_{i} [(\hat{S}_{i}^{x})^{2} - (\hat{S}_{i}^{y})^{2}] \\ &+ J \sum_{\langle i,j \rangle} \hat{\mathbf{S}}_{i} \cdot \hat{\mathbf{S}}_{j} + J_{\perp} \sum_{\langle i,j' \rangle_{\perp}} \hat{\mathbf{S}}_{i} \cdot \hat{\mathbf{S}}_{j'} + \mu_{\mathrm{b}} \sum_{i} \mathbf{B} \cdot \mathbf{g} \cdot \hat{\mathbf{S}}_{i} \end{aligned}$$
(1)

where J is the magnetic exchange interaction between ions within a chain or plane and $J_{\perp} \ll J$ is the exchange interaction between ions on adjacent chains or planes. For antiferromagnets, we define $J, J_{\perp} > 0$. Angular brackets denote a sum over unique pairs of metal ions, and a primed index indicates an ion in an adjacent chain or plane. D and E are the axial and rhombic single-ion anisotropy (SIA) parameters, respectively. For S = 1 compounds, D > 0 corresponds to easy-plane anisotropy with a singlet ground state (see, for example, the inset of Figure 4a), whereas D < 0 defines easy-axis anisotropy that contains the doublet in the ground state. The final term is the Zeeman splitting. The connection between the g factor and single-ion anisotropy can be found using perturbation theory, which predicts

$$g_z - \frac{g_x + g_y}{2} = \frac{2D}{\lambda}$$
 and $g_x - g_y = \frac{4E}{\lambda}$ (2)

where $\lambda \sim -500$ K is the typical spin–orbit coupling constant for Ni²⁺ in an octahedral environment.¹⁶ The interplay between the different terms in the Hamiltonian leads to a rich and varied phase diagram for S = 1 quantum magnets, which includes the exotic Haldane gapped phase, a rare example of a topologically nontrivial magnetic state.¹⁷ The

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predicted phase diagram of S = 1 antiferromagnetic chains is shown in Figure 1 as a function of J_{\perp}/J and D/J, with the

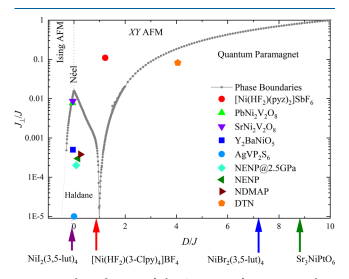


Figure 1. Phase diagram of the S = 1 antiferromagnetic chain recreated from refs 17 and 18. Compounds for which J_{\perp} values are not quantitatively established are represented by arrows along the *x*-axis. From the results in ref 19, it is known that the J_{\perp}/J ratio for NiI₂(3,5-lut)₄ is extremely small. We expect it will be the same for the isostructural NiBr₂(3,5-lut)₄. Reproduced with permission from ref 17. Copyright 2014 American Physical Society. Reproduced with permission from ref 18. Copyright 2014 World Scientific Publishing.

positions of several real materials highlighted. In addition to the Haldane phase, the ordered Néel, Ising, and XY states are also seen, along with the disordered quantum paramagnetic phase at large positive D/J values.¹⁸ Designing bespoke systems to further understand these different phases is currently somewhat hit and miss due to an incomplete understanding of how the Hamiltonian parameters are affected by structure and composition. However, with our family of materials, we are able to span several of these phases and probe them more readily.

Synthesizing high-purity single-phase single crystals is highly resource-intensive, and many compounds are only initially available as powders. The combination of single-ion and exchange effects causes great challenges when characterizing the magnetic properties of polycrystalline samples, especially if $J \approx D$.²⁰ In simple monoligand materials, the link between the distortion of the local octahedral environment and the single-ion anisotropy is clear: an increasingly axially elongated Ni²⁺ octahedron leads to an increase in the easy-plane anisotropy. In contrast, an axially compressed octahedron will promote easy-axis anisotropy. This is also supported by studies of varying distortions in similar mixed-ligand octahedra.²¹ However, predicting how anisotropy energies change between different mixed-ligand materials is necessarily more complex¹⁶ and remains to be fully elucidated.

To shed light on these issues, our study makes use of a family of mixed-ligand magnetic metal-organic complexes. The suitability of these systems for exploring magnetic phase diagrams and their structural and compositional dependences is well established.^{9,14,20,22-31} Using recently published analytical techniques,³¹⁻³³ we present the characterization of the S = 1 chain family NiF₂(3,5-lut)₄·2H₂O and NiX₂(3,5-lut)₄ ($X = HF_2$, Cl, Br, or I). Because of their nonbridging nature, lutidine molecules create an inefficient pathway for inter-

molecular magnetic exchange. Upon creation of isolated Ni-X…X-Ni chains separated by the nonbridging lutidine ligands, the effect that halogen substitution has on the magnetic properties of NiF₂(3,5-lut)₄·2H₂O and NiX₂(3,5-lut)₄ can be detached and investigated in detail. While two-halide exchange interactions have been documented in a few cases of Cu-X…X-Cu pathways for X = Cl and Br,^{34–36} the presence of Jahn-Teller distortions in Cu(II) complexes complicates the structural analysis. The work presented here allows us to span the entire halide series in the absence of Jahn-Teller distortions for the first time by exploiting Ni(II) ions. In addition, the isostructural family that is created provides an ideal test for directly correlating and quantifying the halide radius, the distance between them $(L_{X \dots X})$, and exchange interaction J. We will show that as the radius of the axial halogen ion increases with a corresponding increase in the Ni-X bond distance,²⁰ there is a reduction in the single-ion anisotropy from the strongly easy-plane $Ni(HF_2)_2(3,5-lut)_4$ to the weakly easy-axis $NiI_2(3,5-lut)_4$. Surprisingly, we also find that there are antiferromagnetic interactions along the chains $NiBr_2(3,5-lut)_4$ and $NiI_2(3,5-lut)_4$, despite the distance between halide ions on adjacent NiX2N4 moieties being greater than twice the van der Waals radii of the respective X⁻ ions.

To fully characterize the properties of powder samples of low-dimensional Ni(II) antiferromagnets, we and others have developed a suite of complementary experimental techniques. We use a mixture of bulk (quasi-static and pulsed-field magnetometry) and local probes [electron spin resonance (ESR)] to characterize the magnetic properties of $NiF_2(3,5$ $lut)_4 \cdot 2H_2O$ and $NiX_2(3,5-lut)_4$. Quasi-static magnetometry measurements using a SQUID are relatively quick and economical to perform and have been shown to provide accurate modeling of S = 1 systems exhibiting negligible exchange energies, yielding results in good agreement with those of the more rigorous ESR method.^{21,33} The characterization of low-dimensional antiferromagnetically interacting S = 1 systems cannot rely on susceptibility measurements of powdered samples alone if J and D are similar in magnitude, as there are no models that can be used to reliably fit the data.³² To overcome this, we have successfully used additional information available from low-temperature, high-field magnetization measurements (namely spin-flop, metamagnetic, and saturation fields) to simultaneously extract J and D in the lowtemperature state.^{21,32,33} ESR is highly suited to characterizing the single-ion properties of quantum magnets. In our compounds, the size of the zero-field splitting requires magnetic fields that are higher than can found in commercially available ESR machines. The required equipment is only available at a few specialized facilities worldwide. We therefore reserve ESR measurements for compounds containing both single-ion anisotropy and magnetic interactions, for example, $NiBr_2(3,5-lut)_4$, or in which a complete characterization using magnetometry techniques is not possible, as is the case for $Ni(HF_2)_2(3,5-lut)_4$.

EXPERIMENTAL DETAILS

Synthesis. Chemical reagents were obtained from commercial sources and used as received. All fluoride- and hydrogen difluoride-containing compounds were manipulated and stored in plasticware. The desired products of each compound were obtained in good yield (>70%), and the reaction conditions were not optimized any further.

Synthesis of NiF₂(3,5-lut)₄·2H₂O. Ni(BF₄)₂·6H₂O (0.5023 g, 1.48 mmol) was dissolved in 2 mL of H₂O to produce a green solution. Ammonium fluoride (0.2721 g, 7.35 mmol) was dissolved in 2 mL of H₂O, and the mixture slowly added to the Ni solution. Lastly, 4 mL of neat 3,5-lutidine was slowly pipetted into the aqueous solution of in situ NiF₂·6H₂O, separating upon contact into a clear layer and a blue layer. The beaker was covered with perforated Parafilm and allowed to slowly evaporate at room temperature. After the mixture had stood for several months, 0.2005 g of small medium-blue crystals was collected via vacuum filtration and carefully washed with a minimal amount of fresh H₂O.

Synthesis of Ni(HF₂)₂(3,5-lut)₄. Ni(NO₃)₂·6H₂O (0.3050 g, 1.05 mmol) and 4 equiv of NH₄HF₂ (0.2376 g, 4.17 mmol) were dissolved together in 5 mL of H₂O to produce a light green solution. A 2 mL volume of neat 3,5-lutidine was dissolved in 5 mL of acetonitrile and slowly added to the Ni solution. The resulting solution changed color to dark teal upon contact. All steps were performed in plastic. The beaker was covered with perforated Parafilm, and its contents were allowed to slowly evaporate at room temperature. Upon standing for 3 days, the solution produced medium-blue block-shaped crystals.

Synthesis of NiCl₂(3,5-lut)₄. NiCl₂·6H₂O (0.3500 g, 1.47 mmol) was dissolved in 30 mL of H₂O at room temperature, and the solution slowly mixed with neat 3,5-lutidine (0.9467 g, 8.83 mmol). After a few hours, small crystals were observed growing on the walls of the Erlenmeyer flask. After 3 days, a larger mass of teal-colored crystals had grown, and they were collected via vacuum filtration, washed with fresh 2 × 5 mL aliquots of a 1:1 H₂O/3,5-lutidine mixture, and lightly dried to avoid possible decomposition.

Synthesis of $NiBr_2(3,5-lut)_4$. Preparation of the $NiBr_2(3,5-lut)_4$ compound was accomplished in the same manner as for $NiCl_2(3,5-lut)_4$ but substituting $NiBr_2 \cdot 4H_2O$ for $NiCl_2 \cdot 6H_2O$.

Synthesis of $Nil_2(3,5-lut)_4$. The preparation of this compound has been described by Williams et al.¹⁹

X-ray Structure Determination. Single-crystal X-ray diffraction measurements on $\rm NiF_2(3,5\text{-}lut)_4\text{-}2H_2O$ and $\rm Ni(HF_2)_2(3,5\text{-}lut)_4$ were performed using a Bruker D8 Venture CPAD DUO microsource diffractometer using θ and ω scans and graphite monochromatic Mo $K\alpha$ (λ = 0.71073 Å) radiation. Suitable single crystals were selected from the bulk, mounted on a cryoloop, and cooled to 150(2) K using an LN2 cryostream. Data sets were corrected for Lorentz and polarization effects as well as absorption. The criterion for observed reflections was $I > 2\sigma(I)$. Lattice parameters were determined from least-squares analysis of reflection data. Empirical absorption corrections were applied using SADABS. Structures were determined by direct methods and refined by full-matrix least-squares analysis on F^2 using the SHELXT functionality within X-SEED. Non-hydrogen atoms were refined anisotropically using the SHELXL program. Hydrogen atoms were located by difference Fourier synthesis but placed in idealized geometric positions with $U_{iso} = 1.2U_{eq}$ of the atom to which they were attached; $U_{iso} = 1.5U_{eq}$ for methyl groups. The O atoms of the H₂O molecules are 0.125 occupied owing to 16-fold positional disorder; however, the associated H atoms were not located or modeled. Methyl groups were equally modeled over two positions as suggested by final residual difference maps. Tables S1 and S6 list structural and refinement parameters.

Low-temperature, single-crystal X-ray diffraction measurements of NiX₂(3,5-lut)₄ (X = F, Cl, Br, or I) were conducted on the ChemMatCARS 15-ID-B microcrystal beamline of the Advanced Photon Source (APS) at Argonne National Laboratory (ANL). Microcrystals measuring 10 μ m × 10 μ m × 2 μ m were selected from the bulk samples using a cryo-loop and mounted on a Huber three-circle X-ray diffractometer equipped with an APEX II CCD area detector. The sample was cooled to 100(2) K using an LN2 cryojet. Further details of the beam profile and data collection and analysis, alongside the full details of the resulting structural refinement, bond lengths, and angles, are given in the Supporting Information. The atom labeling scheme is shown in Figure S1.

Magnetometry. Pulsed-field magnetization measurements of powdered NiCl₂(3,5-lut)₄ and NiBr₂(3,5-lut)₄ and single crystals of NiF₂(3,5-lut)₄·2H₂O and Ni(HF₂)₂(3,5-lut)₄ were performed at the

National High Magnetic Field Laboratory (NHMFL) in Los Alamos. Fields of \leq 40 T with typical rise times of \approx 10 ms were used. Samples were mounted in 1.3 mm diameter PCTFE ampules (inner diameter of 1.0 mm) that can be moved into and out of a 1500-turn, 1.5 mm bore, 1.5 mm long compensated-coil susceptometer, constructed from 50 gauge high-purity copper wire.⁹ When the sample is within the coil and the field pulsed the voltage induced in the coil is proportional to the rate of change of magnetization with time, dM/dt. Accurate values of the magnetization are obtained by numerical integration of the signal with respect to time, followed by subtraction of the integrated signal recorded using an empty coil under the same conditions.⁹ The magnetic field is measured via the signal induced within a coaxial 10turn coil and calibrated via observation of de Haas-van Alphen oscillations arising from the copper coils of the susceptometer.⁹ The susceptometer is placed inside a ³He cryostat, which can attain temperatures as low as 500 mK.

Pulsed-field data were calibrated using magnetization measurements made in a Quantum Design MPMS 7 T SQUID magnetometer. Powdered samples of a known mass were loaded into gelatin capsules and fixed to a stainless steel rod. The samples were cooled in zero field to the measured temperature, and then the field was increased in increments of 0.1 T from 0 to 7 T, with the field stabilized before each measurement was taken. The pulsed-field measurements were then scaled onto the SQUID data. This experimental setup was also used to obtain dc susceptibility measurements, with the sample zero-field cooled to 1.8 K. The field was then set to 0.1 T, and temperature-dependent susceptibility data were taken on warming to 300 K.

ESR. High-field, high-frequency ESR spectra of powdered Ni- $(HF_2)_2(3,5-lut)_4$ and NiBr₂ $(3,5-lut)_4$ were recorded on a home-built spectrometer at the EMR facility, NHMFL, Tallahassee, FL. Microwave frequencies (ν) in the range of 160–516.9 GHz at temperatures ranging from 3 to 30 K were used in the measurement. The instrument was a transmission-type device and used no resonant cavity. Powdered samples were loaded into thin Teflon vessels and lowered into the cryostat. The microwaves were generated by a phase-locked Virginia Diodes source, generating frequencies (ν) in the range of 12–18 GHz, and equipped with a cascade of frequency multipliers to generate higher harmonic frequencies. The resultant signal was detected using a cold bolometer. A superconducting magnet capable of reaching fields of \leq 15 T was employed.

In a polycrystalline S = 1 sample with axial single-ion anisotropy (*E* = 0), multiple ESR transitions are expected between the split triplet energy levels. At a given frequency, for the field applied parallel to the local *i*-axis (i = a, b, or c), there are two transitions possible that obey the ESR selection rule $\Delta m_s = \pm 1$: one at low field and one at high field, which we label β_i and γ_i , respectively (inset of Figure 4a). In addition, it is also possible to see an excitation with $\Delta m_s = \pm 2$; this socalled half-field transition is labeled α_i . Formally, such transitions are forbidden, but when the Zeeman energy is comparable to the zerofield splitting, strong mixing between m_s states occurs and the selection rule is relaxed. Double-quantum transitions may also occur when two microwave packets enter the sample at the same time. Additional lines may also be observed at positions that do not correspond to one of the Cartesian axes. These off-axis resonances may be present at the half-field transitions and have been known to dominate the polycrystalline spectra.^{37–39}

Polarized Neutron Diffraction. Neutron diffraction studies of a single crystal of Ni(HF₂)₂(3,5-lut)₄ were performed at the Orphée 14MW reactor of the Laboratory Léon Brillouin, CEA Saclay. The diffraction patterns were collected on a model 5C1 diffractometer, equipped with a position-sensitive detector, which has a cylindrical geometry covering 80° and 25° in the horizontal and vertical directions, respectively. Incident neutrons of wavelength $\lambda = 0.84$ Å were obtained with a Heusler-alloy monochromator, and the beam polarization was 0.88.

RESULTS

X-ray Crystallography. $NiX_2(3,5-lut)_4$ compounds (X = HF₂, Cl, Br, or I¹⁹) crystallize in tetragonal space group P4/ nnc. The nickel octahedron contains four equatorial nitrogen atoms donated from the lutidine molecules and two axial halogen ions (Figure 2a). Because of the tetragonal structure of

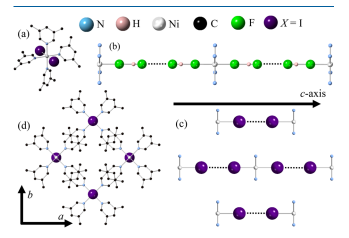


Figure 2. Low-temperature structure of Ni $X_2(3,5-lut)_4$. (a) Layout of the local environment around each Ni²⁺ ion (silver). (b) Ni-HF₂... HF₂-Ni chain showing that the F-H-F molecule axis is oriented along the *c* direction in Ni(HF₂)₂(3,5-lut)₄. (c) Ni-I...I-Ni chains in NiI₂(3,5-lut)₄ (isostructural with X = Cl and Br), where Ni²⁺ ions in adjacent chains are offset in the *c* direction. (d) Lutidine molecules keep Ni-X...X-Ni chains well separated in the *a*-*b* plane. Lutidine hydrogen atoms have been omitted for the sake of clarity.

each compound, the rhombic anisotropy parameter *E* is determined to be zero.⁴⁰ Ni²⁺ ions are linearly bridged along the *c*-axis via two HF₂ molecules (Figure 2b) or two halogen ions (Figure 2c), where F–H–F bonds are linearly oriented along the *c*-axis in Ni(HF₂)₂(3,5-lut)₄. The X…X linkages arise via halogen bonds, which are noncovalent through-space interactions shown to be capable of mediating magnetic exchange.^{41–43} Nonbridging lutidine molecules keep the chains well separated (Figure 2d), and nickel ions in adjacent chains are offset from each other along the *c* direction. There are no clear exchange pathways between adjacent nickel ions within the *a*–*b* plane, and therefore, it is likely that any significant magnetic interaction will be limited to the nickel–halogen chains.

Within NiF₂(3,5-lut)₄·2H₂O, which is isostructural to the Co analogue CoF₂(3,5-lut)₄·2.6H₂O,⁴⁴ adjacent intrachain octahedra are rotated by 45° about the crystallographic *c*-axis as shown in panels a and b of Figure 3. Despite the difference in space group (*P*4/*mcc*), adjacent Ni²⁺ sites remain well separated with an unchanged local environment (Figure 3c). Positionally disordered water molecules sit between fluorine ions that are coordinated to adjacent Ni²⁺ ions. In NiF₂(3,5-lut)₄·2H₂O, the local environment has tetragonal symmetry and therefore E = 0 as for the other compounds.

The relevant structural and bond information for NiX₂(3,5-lut)₄ is given in Tables 1 and 2. As the Ni–N bond lengths for all NiX₂(3,5-lut)₄ compounds are all very similar, this gives an excellent starting point for understanding how the single-ion anisotropy changes due to the substitution of the halogen ion. The nearest-neighbor Ni–Ni distance along the chain decreases as $X = Cl \rightarrow I$ in NiX₂(3,5-lut)₄ despite the increase in the Ni–X bond length. This contraction of L_{Ni-Ni} is driven

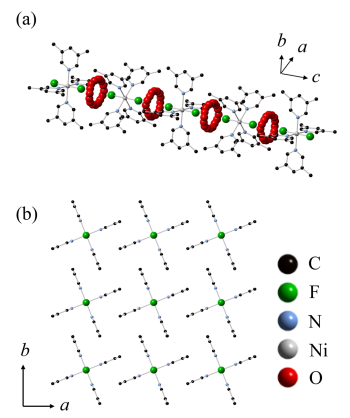


Figure 3. 150 K structure of NiF₂(3,5-lut)₄·2H₂O. (a) Ni-F···F-Ni chain with lutidine paddle wheels rotated by 45° on adjacent sites. Water molecules located between F⁻ ions along Ni-F···F-Ni bond pathways are positionally disordered (water hydrogen atoms have been omitted for the sake of clarity). (b) View down the Ni²⁺ chains highlighting the rotation of the lutidine paddle wheels along the *c*-axis. Lutidine hydrogen atoms in panels a and b have been omitted for the sake of clarity.

Table 1. Unit Cell Parameters for $NiX_2(3,5\text{-lut})_4$ and $NiF_2(3,5\text{-lut})_4\text{-}2H_2O$

	a, b (Å)	c (Å)	space group
$NiF_2(3,5-lut)_4 \cdot 2H_2O$	9.1773(2)	17.2386(4)	P4/mcc
$Ni(HF_2)_2(3,5-lut)_4$	11.1287(3)	11.7600(4)	P4/nnc
$NiCl_2(3,5-lut)_4$	11.4902(9)	10.5512(11)	P4/nnc
$NiBr_2(3,5-lut)_4$	11.7232(5)	10.1153(4)	P4/nnc
$NiI_2(3,5-lut)_4^{19}$	12.0048(5)	9.9568(4)	P4/nnc

by the decrease in the $L_{X\cdots X}$ distance. Coincident with this change is the near 1 Å increase in lattice parameters *a* and *b*, wherein the lutidine torsion angle (C1–N1–Ni–X) decreases from 45.52° for X = Cl to 40.29° for X = I. The structure of NiF₂(3,5-lut)₄·2H₂O does not follow this trend owing to the different space group and interstitial H₂O molecule, resulting in a much shorter intrachain nearest-neighbor Ni²⁺ distance.

Magnetic Properties. $NiF_2(3,5-lut)_4 \cdot 2H_2O$. Single-crystal magnetic susceptibility $\chi(T)$ data for the field parallel and perpendicular to the crystallographic *c*-axis are shown in Figure 4a. For the field applied parallel to *c*, $\chi(T)$ shows a broad hump centered around $T \approx 5$ K, while with the field perpendicular to *c*, $\chi(T)$ continues to increase as the temperature is decreased. These data hint at the beginning of a plateau as the temperature approaches the lowest experimentally accessible value of T = 1.8 K. The $\chi(T)$ curves show that the anisotropy in NiF₂(3,5-lut)₄·2H₂O is easy-plane,⁴⁶ with the hard axis

Table 2. Bond	Lengths i	n NiX ₂ (3,5-lut) ₄	and NiF ₂ ($(3,5-lut)_4 \cdot 2H_2O$
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	$L_{\rm Ni-X}$ (Å)	$L_{\rm Ni-Ni}^{a}$ (Å)	$L_{\mathrm{X}\cdots\mathrm{X}}$ (Å)	van der Waals radius of X^{b} (Å)	$L_{\rm Ni-N}$ (Å)
$NiF_2(3,5-lut)_4 \cdot 2H_2O$	2.036(2)	8.619(2)	4.547(6)	1.35	2.124(2)
$Ni(HF_2)_2(3,5-lut)_4$	2.0120(15)	11.7600(4)	3.137	1.35	2.0945(16)
$NiCl_2(3,5-lut)_4$	2.4502(9)	10.5512(11)	5.651(2)	1.8	2.122(2)
$NiBr_2(3,5-lut)_4$	2.6170(4)	10.1153(4)	4.881(2)	1.95	2.121(2)
$NiI_2(3,5-lut)_4^{19}$	2.8292(4)	9.9568(4)	4.298(2)	2.1	2.123(3)

 ${}^{a}L_{\text{Ni-Ni}}$ corresponds to the nearest-neighbor Ni–Ni distance along the Ni–X···X–Ni chains. ^bvan der Waals radius of the halide ion in NiF₂(3,5-lut)₄.²H₂O and NiX₂(3,5-lut)₄.⁴⁵

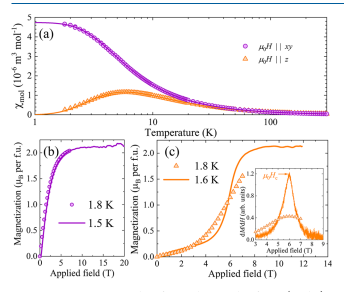


Figure 4. Magnetometry data for single crystals of NiF₂(3,5-lut)₄· 2H₂O. (a) Zero-field-cooled susceptibility (χ_{mol}) measured at $\mu_0 H =$ 0.1 T for the field applied along the *z*-axis (orange triangles) and within the *x*-*y* plane (purple circles). Solid lines are fits to the models described in the text. (b) Pulsed-field magnetization (line) calibrated using similar temperature dc-field SQUID (circles) measurements with the field applied within the *x*-*y* plane. (c) Pulsed-field magnetization (line) calibrated using g_z obtained from modeling $\chi(T)$ with the field applied parallel to the *z*-axis. SQUID data (triangles) are also shown. The differences between the data sets are caused by a magnetocaloric cooling in the pulsed-field measurements. The inset shows a peak in the pulsed-field dM/dH data (line) centered at $\mu_0 H_c$ is consistent with the feature in the 1.8 K SQUID dM/dH data (triangles).

along the crystallographic *c* direction. Therefore, we define the *z*-axis to be along the *c* direction, with the x-y plane analogous to the a-b plane. We use a model containing SIA and paramagnetic contributions³³ and perform a global fit to the data for the field applied parallel and perpendicular to the *z*-axis, shown by solid lines in Figure 4a. The fitting returns the following values: $g_{xy} = 2.12(1)$, $g_z = 2.13(2)$, and D = +8.7(3) K. We note that the fitted g_z and g_{xy} are the same within errors. In this case, the simple perturbation theory of eq 2 suggests that the magnitude of the spin—orbit coupling could be very large.

Pulsed-field magnetization for the field applied within the x-y plane rises smoothly to a broad saturation at 2.13(4) $\mu_{\rm B}$ per formula unit shown in Figure 4b (data calibrated using MPMS SQUID measurements). For the field applied along the *z*-axis (Figure 4c), it is not possible to use SQUID measurements to calibrate the pulsed-field data due to some magnetocaloric cooling in the latter experiment. Therefore, we use the value of g_z found from the measurement of $\chi_z(T)$ to calibrate the saturation at $M_{\rm sat} = 2.13 \ \mu_{\rm b}$ per formula unit. In this orientation, the magnetization shows an initial slow rise before a sharp upturn and eventual rounding to a plateau at saturation.

The midpoint of the upturn occurs at $\mu_0 H_c$, where $g_c \mu_B \mu_0 H_c$ = D, and is the point at which the applied field closes the gap between the $m_s = 0$ and $m_s = -1$ states (see the inset of Figure 5a). Using the peak in dM/dH (inset of Figure 4a), we find that $\mu_0 H_c = 6.0(2)$ T and D = +8.7(4) K, in excellent agreement with the susceptibility measurements. We note that the same critical field can be seen in the SQUID data, but the feature is narrower and sharper in the pulsed-field measure-

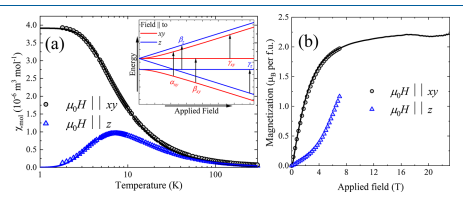


Figure 5. Magnetometry data for an aligned single crystal of Ni(HF₂)₂(3,5-lut)₄. (a) Zero-field-cooled susceptibility plotted vs temperature for the field applied parallel (blue triangles) and perpendicular (black circles) to the crystallographic *c*-axis. Solid lines correspond to a fit to the model described in the text. The inset is a representative sketch of an S = 1 energy-level diagram, showing the labeling scheme for transitions observed in ESR measurements of Ni(HF₂)₂(3,5-lut)₄ and NiBr₂(3,5-lut)₄. (b) Pulsed-field magnetization (solid black line) calibrated to similar temperature DC-field SQUID data (black circles) with the field perpendicular to the *c*-axis. SQUID magnetometry data with the field parallel to the *c*-axis (blue triangles) are also shown.

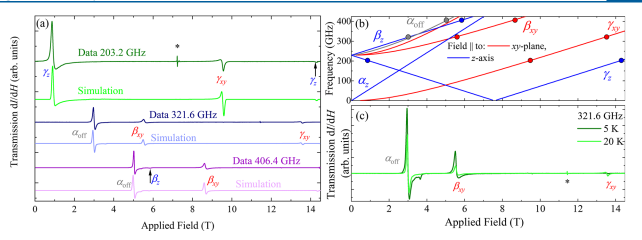


Figure 6. (a) ESR spectra of Ni(HF₂)₂(3,5-lut)₄ recorded at 20 K and frequencies of 203.2, 321.6, and 406.4 GHz. Large resonances are observed in all three spectra, which were fitted using a model with single-ion anisotropy *D*. The obtained parameters were then simulated (Sim) in good agreement with the data. (b) Field frequency positions of the observed resonances (circles) and the result of the fit (lines) with the following extracted parameters: $g_{xy} = 2.22(1)$, $g_z = 2.16(2)$, and D = +11.0(1) K. (c) Temperature dependence of the 321.6 GHz Ni(HF₂)₂(3,5-lut)₄ ESR spectra.

ment due to a magnetocaloric cooling associated with the high dB/dt in the pulsed-field measurement.¹⁴

 $Ni(HF_2)_2(3,5-lut)_4$. Single-crystal susceptibility data for $Ni(HF_2)_2(3,5-lut)_4$ (Figure 5a) show a rise to a broad hump centered around $T \approx 8$ K for the magnetic field parallel to the *c*-axis. For the field perpendicular to the *c*-axis, the data continue to rise and appear to begin to plateau as the temperature approaches 1.8 K. The $\chi(T)$ measurements are consistent with a singlet ground state and D > 0. Following the treatment given to NiF₂(3,5-lut)₄·2H₂O, we define the *z*-axis and x-y plane to be along the *c*-axis and a-b plane, respectively. Fitting the data to a model containing contributions from SIA and paramagnetism⁴⁶ gives the following values: $g_{xy} = 2.20(1)$, $g_z = 2.13(3)$, and D = +11.0(6) K. Muon spin rotation measurements also show no evidence of long-range order at temperatures as low as 20 mK,⁴⁷ consistent with $J \approx 0$ for Ni(HF₂)₂(3,5-lut)₄.

The calibrated pulsed-field magnetization data with the field applied perpendicular to the *z* direction (Figure 5b) rises smoothly to a broad saturation at 2.18(1) $\mu_{\rm B}$ per formula unit. For the field applied parallel to the *z*-axis, SQUID magnetization displays an upturn due to the field-induced ground-state energy-level crossing in the *S* = 1 triplet (inset of Figure 4a). This is consistent with easy-plane anisotropy in Ni- $(HF_2)_2(3,5-lut)_4$. The application of a magnetic field above 7 T in the *c* direction consistently caused shattering of crystals, regardless of the method used to fix the samples in position. As a result, M(H) data cannot be used to quantify *D* for Ni $(HF_2)_2(3,5-lut)_4$.

To confirm the Hamiltonian parameters derived from magnetometry, high-field ESR measurements of polycrystalline Ni(HF₂)₂(3,5-lut)₄ were performed and show resonances corresponding to γ_{xyy} $\gamma_{z'}$, β_{xyy} and β_z transitions as shown in Figure 6a. A feature corresponding to the off-axis resonance is also present in the 321.6 and 406.4 GHz sweeps. The resonance marked with an asterisk is attributed to an impurity. The field frequency positions of the resonances were fitted (Figure 6a), with the resulting parameters extracted: $g_{xy} = 2.22(1)$, $g_z = 2.16(1)$, and D = +11.0(1) K. Simulating spectra using a model containing only SIA at the experimentally measured frequencies and temperatures provides results in excellent agreement with the data. A temperature-dependent

study at 321.6 GHz was performed (Figure 6c) to confirm the sign of *D*. The data show an increase in the intensity of the β_{xy} transitions as the temperature decreases. This is due to an increased occupation of the $m_s = -1$ state at lower temperatures. At the same time, the intensity of the γ_{xy} resonance decreases as the temperature decreases due to a reduction in the occupation of the $m_s = 0$ state. Both of these trends are consistent with Ni(HF₂)₂(3,5-lut)₄ exhibiting easyplane anisotropy. This is further supported by the fact that $g_{xy} > g_{zy}$ and the resulting value of $\lambda \approx -400$ K is consistent with that expected for Ni²⁺ ions in octahedral environments.¹⁶ The value of *D* extracted from ESR measurements is in excellent agreement with that found from fitting the susceptibility data above.

To provide further evidence of the easy-plane nature of $Ni(HF_2)_2(3,5-lut)_4$, polarized neutron diffraction measurements were performed on a single crystal of Ni(HF₂)₂(3,5 $lut)_4$. Data were collected at temperatures of 1.8 and 10 K in a range of magnetic fields between 1.5 and 6 T applied parallel to the y- and z-axes. The magnetization density was extracted from flipping ratio R, the ratio of the intensity of neutrons scattered when polarized parallel and antiparallel to the external magnetic field. Initial data refinement indicated a strong anisotropy between the y and z directions, which are aligned along the crystallographic b- and c-axes, respectively. This anisotropy has been taken into account using the susceptibility approach described by Gukasov and Brown.⁴⁸ In the orthorhombic axes, the symmetry constraints for the magnetic Ni atom occupying the "2b" site in the P4/nnc space group imply $\chi_{xx} = \chi_{yy}$ and $\chi_{xy} = \chi_{xz} = \chi_{yz} = 0$. Thus, only two independent susceptibility parameters, χ_{xx} and χ_{zz} need to be determined to fully characterize the magnetic behavior of the Ni²⁺ ion.

The susceptibility component χ_{zz} can be accurately determined on the basis of a set of flipping ratios collected in a magnetic field applied along the z-axis. At the same time, susceptibility component χ_{yy} has a significant effect on the flipping ratios measured in the magnetic field applied along the y-axis. Thus, a joint refinement procedure, in which two data sets with the field oriented along the y- and z-axes are fitted simultaneously, is required to simultaneously calculate the χ_{yy} and χ_{zz} components. The joint refinement of the data at 10 K shows no evidence of anisotropy, whereas at 1.8 K, the joint refinement of the lower-field (H//y = 1.5 T, and H//z = 3.3 T) data sets and higher-field (H//y = 6 T, and H//z = 4 T) data indicates that $\chi_{yy} \gg \chi_{zz}$. The susceptibility components are combined to produce magnetization ellipsoids (Figure 7),

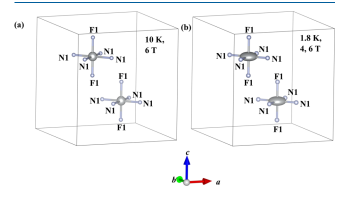


Figure 7. Illustration of magnetization ellipsoids for Ni atoms in a single crystal of Ni(HF₂)₂(3,5-lut)₄ as determined by refinement of polarized neutron diffraction data (see the text): (a) high-temperature, high-field data and (b) low-temperature, high-field data. The ellipsoids in panel b confirm the *xy* orientation of the Ni²⁺ moment in the paramagnetic state attributed to D > 0.

showing that $Ni(HF_2)_2(3,5-lut)_4$ exhibits easy-plane anisotropy, with the hard axis oriented along the z-axis. These results are in excellent agreement with the magnetometry and ESR measurements.

 $NiCl_2(3,5-lut)_4$. Single crystals of sufficient size to perform bulk magnetometry measurements of $NiCl_2(3,5-lut)_4$ and $NiBr_2(3,5-lut)_4$ are not currently available. Therefore, we use the methods of extracting the Hamiltonian parameters of polycrystalline samples described by Blackmore et al.³³

Susceptibility data of powdered NiCl₂(3,5-lut)₄ increase as the temperature is decreased before appearing to plateau at ≈ 2 K (Figure 8a), consistent with negligible magnetic interactions and the presence of single-ion anisotropy with D > 0. The data are fitted using a model incorporating single-ion anisotropy, paramagnetism,⁴⁶ and a small temperature-independent contribution $\chi_0 = -1.46(7) \times 10^{-9}$ m³ mol⁻¹ that is obtained from a Curie–Weiss fit to $\chi^{-1}(T)$ (inset of Figure 8a). The parameters extracted are D = +10.1(1) K and powder-averaged g factor $\tilde{g} = 2.16(1)$.

Pulsed-field magnetization measurements of polycrystalline NiCl₂(3,5-lut)₄ show a smooth, seemingly featureless, concave rise to a broad saturation point (Figure 8b), while differentiating the data reveals a weak bumplike feature (inset). The observations are consistent with the simulations for an S = 1 system with negligible exchange interactions.^{31,33} The weak bump in dM/dH at $\mu_0H_c = 6.6(5)$ T is caused by the ground-state energy-level crossing, which for E = 0 occurs for only easy-plane systems $(D > 0)^{33}$ and yields a D value of +9.6(7) K, in agreement with the susceptibility result. As NiCl₂(3,5-lut)₄, we determine that the z-axis is located along the *c*-axis and the x-y plane is located within the a-b plane in NiCl₂(3,5-lut)₄.

 $NiBr_2(3,5-lut)_4$. DC susceptibility data of powdered $NiBr_2(3,5-lut)_4$ (suspended in Vaseline to prevent grain movement) (Figure 9a) show an increase in $\chi(T)$ as the temperature is decreased, before leveling off as T approaches 1 K. The data are fitted to a model that includes a small mean-

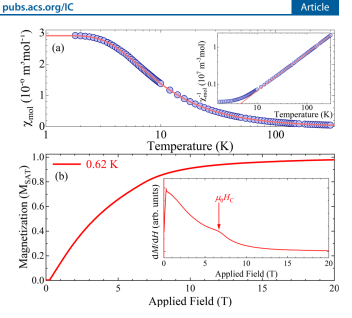


Figure 8. (a) Temperature-dependent susceptibility measurements of polycrystalline NiCl₂(3,5-lut)₄ performed at an applied field of 0.1 T and fitted to a model including a single-ion anisotropy contribution (see the text). The inset of panel b shows inverse susceptibility data of NiCl₂(3,5-lut)₄ fitted to a Curie–Weiss model and a small temperature-independent contribution in the range 36–300 K. (b) Magnetization and differential susceptibility (inset) of polycrystalline NiCl₂(3,5-lut)₄. A feature indicating a ground-state energy-level crossing is apparent in the dM/dH data at $\mu_0H_c = 6.6(5)$ T. The sharp rise below 0.3 T is due to an artifact of the measurement.

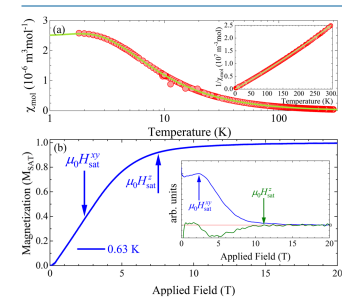


Figure 9. (a) Temperature-dependent susceptibility measurements of powdered NiBr₂(3,5-lut)₄ suspended in Vaseline performed at an applied field of 0.1 T. The data are fitted to a model containing both single-ion and magnetic exchange terms [green (see the text)].⁴⁹ The inset shows inverse susceptibility data fitted to a Curie model in the range of 50–300 K. (b) Pulsed-field magnetization and differential susceptibility (inset) of NiBr₂(3,5-lut)₄. Critical fields are observed in the d*M*/d*H* data indicating saturation of moments for the field parallel to the easy plane at $\mu_0 H_{sat}^{sy} = 2.3(2)$ T and the easy axis at $\mu_0 H_{sat}^z = 11.1(5)$ T.

field perturbation caused by magnetic exchange on a single-ion anisotropy model, as described by Carlin and de Jongh.⁴⁹ The parameters extracted from the resultant fit are $\tilde{g} = 2.11(1)$, D =

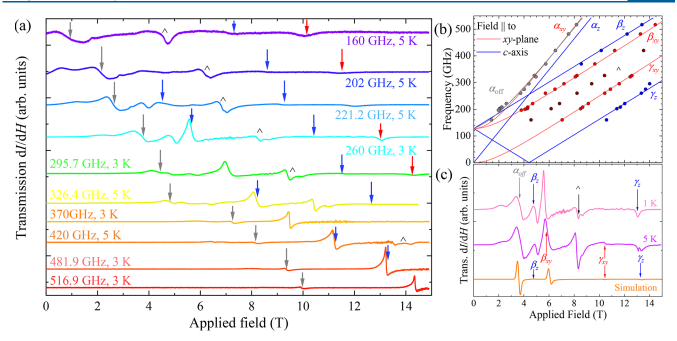


Figure 10. (a) ESR spectra of polycrystalline NiBr₂(3,5-lut)₄ collected in derivative mode at low temperatures and high frequencies. Red arrows show the position of the γ_z transition, and blue arrows show the positions of the γ_{xy} (at high field) and β_{xy} (at low field) transitions. Gray arrows and \wedge mark the off-axis and double-quantum resonances, respectively. (b) Plot of frequency vs field showing the peak positions (circles). The lines are the expected locations of resonances from the fit to the experimental data described in the text, with the dark gray line and circles corresponding to the off-axis resonances and the brown circles referring to the double-quantum resonances. (c) Temperature dependence of the 260 GHz spectra. The orange line is a simulation at 5 K using the parameters obtained from the fit in panel b.

+6.84(9) K, $\chi_0 = -5(1) \times 10^{-9}$ m³ mol⁻¹, and nJ = 1.09(4) K, where n = 2 is the number of magnetically interacting nearest neighbors. The model recreates the data well, and the parameters are consistent with the weak exchange and stronger single-ion anisotropy that we find in the analysis of M(H) and ESR data below. This strongly indicates the presence of weak magnetic interactions along the Ni–Br…Br–Ni chains. The inverse susceptibility data show paramagnetic behavior down to very low temperatures and were fitted to a Curie model (inset of Figure 9a). The parameters obtained are $\tilde{g} = 2.09(1)$ with a small temperature-independent contribution $\chi_0 = -6.05(4) \times 10^{-9}$ m³ mol⁻¹.

Pulsed-field (Figure 9b) measurements of polycrystalline $NiBr_2(3,5-lut)_4$ show that the magnetization rises quasi-linearly at first before approaching a broad saturation. The differential susceptibility (inset of Figure 9b) similarly shows a flattish region that ends in a short rise to a peak before dropping to saturation with no indication of a spin flop. This is consistent with an S = 1 compound with easy-plane single-ion anisotropy and modest magnetic interactions between adjacent Ni²⁺ ions.³³ In general, two critical fields can be identified in polycrystalline M(H) data of Ni²⁺ antiferromagnets that are attributed to moments saturating when the field is parallel to the easy-plane $g_{xy}\mu_B\mu_0H_{sat}^{xy} = 2SnJ$ and along the hard-axis $g_z \mu_B \mu_0 H_{sat}^z = 2(nf + D)S$, where *n* is the number of nearest neighbors. The lower saturation field occurs at $\mu_0 H_{sat}^{xy} = 2.3(2)$ T. The upper saturation field is estimated as the point at which d^2M/dH^2 first crosses zero, which occurs at $\mu_0 H_{sat}^z = 11.1(5)$ T. Using $g_{xy} = 2.20(1)$ and $g_z = 2.08(1)$ obtained from ESR measurements (see below), values of J = 0.85(7) K and D =+6.1(5) K are extracted, consistent with the fit to $\chi(T)$ presented above.

Low-temperature ESR measurements of polycrystalline $NiBr_2(3,5-lut)_4$ were recorded in the first-derivative mode at

frequencies between 160 and 516.9 GHz (Figure 10a). No evidence for long-range magnetic order has been observed in this system, nor is it to be expected as the temperature range (3) $K \leq T \leq 5$ K) at which these data are taken is larger than energy scale of the exchange interaction estimated from magnetometry. For this reason, it is possible to identify features in the spectra that arise from transitions between anisotropy-split S = 1 energy levels only, neglecting the effects of magnetic exchange. The low-frequency data show resonances corresponding to γ_z and γ_{xy} transitions at higher fields, which move out of the field window above 326.4 GHz. The β_{xv} transition is also observed at $\nu \geq 196.8$ GHz, and the β_z resonance develops as the frequency increases from 370 GHz. A feature corresponding to $\alpha_{\rm off}$ is also present in all field sweeps. Resonances marked with a " \wedge " are ascribed to the double-quantum transition. The resonances caused by transitions between magnetic energy levels of the Ni²⁺ ion are plotted in Figure 10b and are fitted to a Hamiltonian containing a single-ion anisotropy term and no exchange. The extracted parameters are $g_{xy} = 2.20(1)$, $g_z = 2.08(1)$, and |D| =6.2(1) K. A simulation at 260 GHz using these parameters is plotted in Figure 10c and successfully replicates the position and amplitudes of the $\alpha_{\rm off}$, β_{xy} , γ_{xy} , and γ_z transitions. The position of the simulated β_z transition is in good agreement with the data but is recreated at a much smaller amplitude.

The sign of D can be extracted from observing the temperature dependence of the β and γ peaks. The 260 GHz data (Figure 10c) show the intensity of the γ_{xy} transition increases as the temperature increases, corresponding to a transition between two excited states. The β_{xy} and γ_z resonances are present at very low temperatures, indicating that these are caused by transitions between the ground state and an excited state. The temperature dependence of these transitions implies that D > 0 for NiBr₂(3,5-lut)₄, consistent

with the observation that $g_{xy} > g_z$.¹⁶ The calculated value of $\lambda \approx -100$ K is of the order expected for Ni²⁺ in such an environment. The value of D = 6.2(1) K obtained here is in excellent agreement with that extracted from magnetization measurements and consistent with that obtained from the $\chi(T)$ data.

 $Nil_2(3,5-lut)_4$. Nil $_2(3,5-lut)_4$ has been previously characterized by Williams et al.¹⁹ and exhibits easy-axis single-ion anisotropy with D = -1.2(3) K, which was extracted from inelastic neutron scattering measurements. Pulsed-field magnetization shows a very high saturation field relative to the other compounds reported here, indicating strong antiferromagnetic interactions of J = 17.5(1) K along the Ni–I···I–Ni chains. Nil $_2(3,5-lut)_4$ was also shown to lie in the Haldane phase with anisotropic energy gaps of $\Delta^{\parallel} = 5.3(1)$ and $\Delta^{\perp} =$ 7.7(1) K. The structural and Hamiltonian parameters of Nil $_2(3,5-lut)_4$ will be used in the discussion below.

DISCUSSION

The magnetic parameters of $NiF_2(3,5-lut)_4 \cdot 2H_2O$ and $NiX_2(3,5-lut)_4$ (X = HF₂, Cl, Br, or I) are summarized in Table 3. Magnetometry measurements show that $NiF_2(3,5-lut)_4$ (X = HF₂, Cl, Br, or I) are summarized in Table 3.

Table 3. Experimentally Derived Parameters for NiF₂(3,5-lut)₄·2H₂O and NiX₂(3,5-lut)₄^{*a*}

Х	J (K)	D (K)	g factor
F	≈ 0	8.7(4)	$g_{xy} = 2.12(1)$
			$g_z = 2.13(1)$
HF_2	≈ 0	11.0(1)*	$g_{xy} = 2.22(1)^*$
			$g_z = 2.16(1)^*$
Cl	≈ 0	9.6(7)	$\tilde{g} = 2.16(1)$
Br	0.85(7)	6.1(5)	$g_{xy} = 2.20(1)*$
			$g_z = 2.08(1)^*$
Ι	17.5(1)	-1.2(3)	$g_{xy} = 2.19(1)*$
			$g_z = 2.13(1)^*$

^{*a*}All values are determined from magnetometry data, except those marked with an asterisk, which are derived from ESR measurements.

 $lut)_4{\cdot}2H_2O$ and $NiCl_2(3,5{\cdot}lut)_4$ exhibit no magnetic interactions between adjacent Ni^{2+} ions, and modeling the susceptibility and magnetization data give the same D parameters within errors. In the case of $Ni(HF_2)_2(3,5-lut)_4$, the D parameter extracted from ESR measurements is in excellent agreement with that found from the susceptibility data. The magnetization data do not provide any additional information in this particular case as the crystal shatters upon application of a significant field parallel to the hard direction, an effect that has also been observed previously.⁵⁰ The dM/dHcurves of NiBr₂(3,5-lut)₄ are significantly different from that of $NiCl_2(3,5-lut)_4$. There is a peak indicating the presence of magnetic interactions, which is supported by fitting the susceptibility data to a a mean-field correction to account for the exchange term. High-frequency ESR measurements confirm the value of single-ion anisotropy of $NiBr_2(3,5-lut)_4$, which is in excellent agreement with that extracted from the magnetization data.

Here we will discuss the trends linking the structure and composition (Tables 1 and 2) to the magnetic exchange and the anisotropy.

Exchange Interactions. Starting with X = Cl, Br, and I, as the size of bridging halogens increases, adjacent Ni²⁺ ions move closer together along the *z*-axis and strong magnetic

interactions are turned on from $X = Cl \rightarrow I$ as indicated by the difference in shape of the dM/dH data for these compounds. There is also a corresponding decrease in the X…X distance $(L_{X \dots X})$, with X = I having the smallest separation. In each case, this separation exceeds twice the van der Waals radii for the halides (1.80, 1.95, and 2.10 Å for Cl, Br, and I, respectively) (Table 2). The difference is smallest for X = I (~0.1 Å) and increases for NiBr₂(3,5-lut)₄ (~1 Å) and NiCl₂(3,5-lut)₄. This is surprising given the measured antiferromagnetic interaction energies in both the X = Br and I materials and implies an efficient through-space mechanism for mediating these interactions. The crystal structures are modeled by treating all atoms and/or ions as spherical shapes. We believe that the heavier halides such as Br⁻ and I⁻ become increasingly aspherical or polarized along the c-axis. This will cause their respective shapes to elongate and enhance their potential for magnetic interactions.

NiF₂(3,5-lut)₄·2H₂O and Ni(HF₂)₂(3,5-lut)₄ follow a separate trend from the other materials due to their structural differences. The smallest nickel–nickel distance (L_{Ni-Ni}) is found in NiF₂(3,5-lut)₄·2H₂O, while Ni(HF₂)₂(3,5-lut)₄ has the largest L_{Ni-Ni} . The F…F distance linking F atoms on different Ni–HF₂ moieties is also ~0.1 Å more than twice the van der Waals radius of fluorine. This length scale of the halide…halide distance is similar to that found for NiI₂(3,5-lut)₄. However, the magnetometry data show that there are no antiferromagnetic interactions in either NiF₂(3,5-lut)₄·2H₂O and Ni(HF₂)₂(3,5-lut)₄.

Single-Ion Anisotropy. Another trend linking structure and composition to magnetic parameters in our compounds is the increase in the axial Ni–halogen bond (L_{Ni-X}) relative to the equatorial Ni²⁺—nitrogen bond (L_{Ni-N}) in the NiX₂N₄ octahedron. This is accompanied by a corresponding decrease in *D*, such that easy-plane anisotropy is suppressed at the extreme end of the family, and the axially elongated NiI₂N₄ octahedron in NiI₂(3,5-lut)₄ actually gives rise to easy-axis anisotropy. Furthermore, the NiF₂N₄ octahedra in NiF₂(3,5-lut)₄·2H₂O and Ni(HF₂)₂(3,5-lut)₄ are both axially compressed, but we find that both materials display strong easy-plane anisotropy. All of this behavior contradicts the simplest predictions that increasingly elongated Ni²⁺ octahedra should correspond to an increasingly more positive *D* parameter.^{16,52,53}

The variation in anisotropy on changing the axial ligands involved in the NiX₂N₄ octahedra in our family of materials fits with the picture¹⁶ that for mixed-ligand complexes the relative values of the axial and equatorial crystal-field strength and covalency are additional factors in establishing the size and sign of D. To explore these issues further, Figure 11 plots the value of D as a function of both the axial/equatorial Ni-L bond length ratio and the bonded atom electronegativity ratio⁵⁴ for a number of different materials with NiX₂N₄ local environments. Although the relationship between covalency (or orbital reduction factor) and electronegativity is not straightforward and the electronegativity of the bonded atom may not be entirely representative of the effect of the coordinated ligand as a whole, certain trends are highlighted by the figure. For materials in which the halogen remains unchanged, specifically those with F and Cl axial ligands, the electronegativity ratio is fixed. In this case, the simple picture is upheld and D becomes more positive as the axial bond weakens and the octahedron elongates. However, when the coordinated halide changes, the effect of reducing the electronegativity competes with the

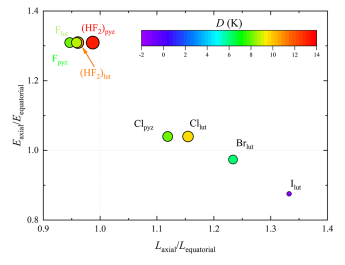


Figure 11. Bond length and bonded atom electronegativity dependence of *D* for NiX₂(3,5-lut)₄ (X = HF₂, Cl, Br, or I), NiF₂(3,5-lut)₄·2H₂O, NiCl₂(pyz)₂)^{20,51} NiF₂(pyz)₂·3H₂O,²¹ and [Ni-(pyz)₂(HF₂)₂)(SbF₆).³² The compounds presented in this paper are labeled with X_{lut} with their NiX₂(pyz)₂ relatives labeled with X_{pyz} . The larger red circles are strongly easy-plane, and the smaller bluer circles correspond to an easy-axis *D*.

lengthening of the axial bond. For compounds in which there is a significant difference between the electronegativity of the axial and equatorial ligands, the sign of D can be estimated with confidence. The Ni-F axial bond, for example, has by far the most ionic character and yields a consistently easy-plane magnet in all cases here and for several other materials with NiF_2N_4 octahedra.²¹ At the other end of the scale, we would expect that a NiI₂N₄ environment would consistently yield an easy-axis anisotropy, as is the case here for $NiI_2(3,5-lut)_4$. The situation for an electronegativity ratio closer to unity is less clear, with $NiCl_2(3,5-lut)_4$ exhibiting greater axial elongation and a D that is slightly more positive than that of $NiF_2(3,5$ lut)₄·2H₂O, despite the decreased electronegativity ratio. Furthermore, while $NiBr_2(3,5-lut)_4$ exhibits a slightly easyplane D, recent neutron diffraction measurements are indicative of easy-axis behavior for $NiBr_2(pyz)_{2,j}$ which contains the same NiBr₂N₄ octahedra, ⁵¹ suggesting that, in this case at least, the coordinated molecule must be considered as a whole.

Gapped Ground States. The ground state of an S = 1AFM chain depends on the size of the J_{\perp}/J and D/J ratios. The complete phase diagram for these systems displays Ising and XY-ordered states along with two gapped phases: the Haldane and quantum paramagnetic states (see Figure 1).18 It has already been shown that NiI₂(3,5-lut)₄ is a highly isotropic [D/J = -0.07(1)] example of a Haldane chain.¹⁹ In addition, because it displays no magnetic order even when the Haldane gap has been closed by the application of an applied magnetic field, the value of J_{\perp} in this material is believed to be extremely small. The similarity between the interchain structure in our family of samples suggests that J_{\perp} will also be small for NiBr₂(3,5-lut)₄. This, together with the measured D/J ratio, predicts that this material would lie in the quantum paramagnetic state (see Figure 1). While we see no direct evidence of the anticipated spin gap in the data, it seems that the gap is observed clearly only in magnetometry measurements of single crystals (see, for example, Sr₃NiPtO₆⁵⁵ and DTN)¹³ but is not apparent in powdered samples like ours (see the data for NENC,⁵⁶ which has a D/J ratio of 15 and is

known to reside in the QPM phase from the results of heat capacity⁵⁶ and EPR measurements).⁵⁷

The ability to populate experimentally underexplored phase diagrams via chemically simple substitution is a benefit to studying molecule-based magnets. In addition, their often low interaction energies mean their novel quantum critical behavior lies well within an experimentally accessible regime. Measurements are underway on the newly synthesized doped compound Ni(Br_xI_{1-x})₂(3,5-lut)₄ to further probe these gapped phases using chemical pressure.

SUMMARY

We have successfully characterized four new related Ni²⁺-chain compounds. This enabled us to study the effect that different halide ions have on single-ion anisotropy and two-halide exchange in isolation. NiF₂(3,5-lut)₄·2H₂O, Ni(HF₂)₂(3,5 $lut)_4$, NiCl₂(3,5-lut)₄, and NiBr₂(3,5-lut)₄ have all been shown to exhibit easy-plane single-ion anisotropy. Magnetometry data suggest that the compounds with axial F, HF₂, and Cl ligands contain negligible magnetic exchange with D = 8.7(1), 11.0(1),and 9.6(7) K, respectively. Magnetization measurements indicate that $NiBr_2(3,5-lut)_4$ contains magnetic exchange along the Ni²⁺ chains with J = 0.85(7) K and a value of D =6.1(5) K, which is confirmed by ESR. $NiI_2(3,5-lut)_4$ was found to be weakly easy-axis, with D = -1.2(3) K with strong magnetic exchange interactions J = 17.5(1) K. We have shown that the different halides drastically alter the magnetic properties. The single-ion anisotropy is strongly influenced not only by the distortion of the local octahedra but also by the relative size of the electronegativity of the axial and equatorial ligands. As a result, in general for systems with Ni²⁺ octahedra, we would expect D > 0 for strongly electronegative axial ligands (such fluorine) even when the octahedron is axially compressed but D < 0 in the case of weakly electronegative axial ligands (for example iodine) even when the octahedron is axially compressed. At the same time, upon substitution of larger halide ions into the magnetic exchange pathway, the $L_{X...X}$ distance decreases and J significantly increases. The size of the $L_{X...X}$ distances implies a through-space pathway for the exchange in the X = Br and I systems. It is expected that there will be extreme sensitivity to tuning $L_{X \cdots X}$ on J via the application of strain and pressure in these types of materials. While further study is required to fully understand this through-space mechanism, it is clear that this is a novel kind of exchange pathway that can be exploited in the design of bespoke magnetic materials.

ASSOCIATED CONTENT

1 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.inorgchem.1c02483.

Additional data (ZIP)

Further details of the crystal structure data and refinement, polarized neutron diffraction measurement analysis, and a description of the thermal stability of the pulsed-field magnetization measurements (PDF)

Accession Codes

CCDC 2074899–2074902 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

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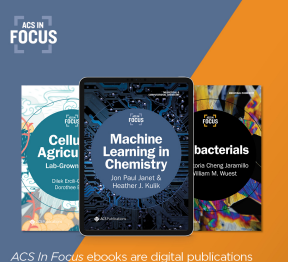
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