A multifunctional magnetic material under pressure

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[Fe II(Metz) 6](Fe III Br4)2 (Metz = 1-methyltetrazole) is one of the rare systems combining spin-crossover and long-range magnetic ordering. The spin-crossover phenomenon of the cationic sub-lattice [Fe II(Metz) 6]2+ coexists in this multifunctional material with an antiferromagnetic order of the anionic sub-lattice (Fe IIIBr4)–. A combined study involving neutron and X-ray diffraction and bulk and single-crystal magnetometry allows determining the collinear antiferromagnetic structure of this system, and shows that the magnetic ordering is favoured by the application of pressure, with an increase of the Néel temperature from 2.4 K at ambient pressure, to 3.9 K at 0.95 GPa. Applied pressure also enables a full high-spin to low-spin switch at ambient temperature.

Introduction

The molecular approach for the design of multifunctional materials makes use of the flexibility offered by molecular systems for combining different functional properties in the same compound. Several recent examples illustrate the success of this approach, featuring e.g., magnetic and optical properties,1 porosity and magnetism,2 conductivity and magnetism,3 superconductivity and magnetism,4 or even a multiferroic character.5 The spin crossover (SCO) phenomenon6 is in itself a source of multifunctionality,7 with drastic variations in magnetic, optical, dielectric and structural properties potentially taking place concomitantly with the change of spin state between high-spin (HS) and low-spin (LS) electronic configurations.8 Conveniently, this molecular bistability can be addressed by an external perturbation9 like temperature,10 magnetic field,11 light irradiation,12 chemical stimuli13 or pressure.14 Very recent reports have in addition shown that the light-induced HS form of an Fe(II) SCO material can behave as a Single-Molecule Magnet (SMM) and that photomagnetic chains can be designed by linking SCO and SMM building blocks.15 On the other hand, although magnetic ordering is one of the most pursued functional properties due to its potential use in a variety of applications,16 the molecular-based systems combining it with SCO are still scarce. This combination has been achieved in hybrid crystalline solids that integrate FeIII SCO centres together with MnII-CrIII-oxalate-based magnetic networks.17 In FeII SCO systems, which are particularly interesting since the involved LS state is diamagnetic, the combination of SCO and long range magnetic ordering remains more elusive. An exception is a recent Fe–Nb metal–organic framework featuring a light-induced magnetic ordering.18 [FeII(Metz)6](FeIIIBr4)2 (Metz = 1-methyltetrazole)19 (I) is in this respect an exciting new example combining SCO and magnetic order. The trigonal structure of I has one [Fe(Metz)6]2+ cation and two FeBr4– anions in the unit cell, building hexagonal layers extended in the ab plane (Figure 1). These layers are pillared along the c axis, in a perfectly eclipsed conformation. According to magnetic and calorimetric measurements, [Fe(Metz)6]2+ cations present a gradual complete thermal SCO centred at 165 K, coexisting with a long range antiferromagnetic order below $T_N = 2.4$ K of the FeBr4–.19 We report here the magnetic structure of this original material and present a detailed study of how applied pressure affects both the SCO and the magnetic order.

Figure 1. View along the b axis of the structure of 1 at 10 K – [Fe(N)6] polyhedra in purple, FeBr4– anions in yellow, H···Br short contacts as blue sticks.
Results and discussion

The compound [FeIII(Metz)6][FeBr4]2 was readily prepared as shiny red polycrystalline material, as previously reported. Slower evaporation of the ethanolic solution containing the mixture of FeBr2, FeBr3 and Metz allows the formation of larger hexagonal rod-shape single crystals, suitable for single-crystal magnetic measurements (ESI). Bulk calorimetric and magnetic data clearly indicate the presence of an antiferromagnetic order of the FeBr4− anionic sublattice in I below 2.4 K, in addition to the SCO of the [Fe(Metz)6]2+ cation observed at higher temperatures. We have now used powder neutron diffraction (PND) and single-crystal magnetometry for determining the magnetic structure of I. A first high-resolution PND data collection aimed at refining the nuclear structure at low temperature has been performed at 10 K – with the cationic sublattice in the LS diamagnetic state and the anionic one in a paramagnetic state – using a wavelength of 1.59 Å (see ESI for full experimental and crystallographic details). The diffraction pattern essentially matches the reported structure determined by X-ray at 104 K, although the better sensitivity of neutrons to hydrogen compared with X-rays reveals a rotation of the methyl groups of Metz that slightly changes the described hydrogen-bonding network, with shortened H···Br distances that create double-layered planes parallel to ab (Figure 1, Tables 1 and S2). The Rietveld refinement (Figure S1, Tables 1 and S1) shows also a small decrease of volume of ca. 2%, which can be assigned to thermal compression of the cell parameters.

Table 1. Cell parameters and relevant bond distances (Å) and angles (º) as well as short contacts (Å) in the structure of I at 10 K / ambient pressure and 298 K / 2.3 GPa. See Figure 5 for labelling. Data from the structures at ambient pressure and 298 K and 104 K are recalled for comparison.

<table>
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<tr>
<th>Parameter</th>
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<th>298(2)</th>
<th>296(2)</th>
<th>104(2)</th>
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<tr>
<td>T, K</td>
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<td>298(2)</td>
<td>296(2)</td>
<td>104(2)</td>
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<td>p, GPa</td>
<td>amb.</td>
<td>2.3</td>
<td>amb.</td>
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<td>0.71073</td>
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<td>crystal system</td>
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<td></td>
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<td>space group</td>
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<tr>
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<td>12.2039(14)</td>
<td>12.9487(1)</td>
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<td>c, Å</td>
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<td>6.4241(7)</td>
<td>6.9577(1)</td>
<td>6.7890(14)</td>
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<td>V, Å³</td>
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<td>826.6(2)</td>
<td>1010.30(2)</td>
<td>947.0(3)</td>
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<td>1.952(3)</td>
<td>2.187(2)</td>
<td>2.003(2)</td>
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<td>7.018(6)</td>
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<td>2.683(1)</td>
<td>2.911(1)</td>
<td>2.807(1)</td>
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</table>

Symmetry operations: #1: x+1, y+1, z+1; #2: x, y, 1+z

Figure 2 shows the PND patterns collected at 1.5 and 4.5 K in the high-flux diffractometer D1B using a wavelength of 2.52 Å. At the lowest temperature, an increase in intensity of some nuclear Bragg reflections – (100), (101) / (011) – is noticed, indicating the appearance of long-range magnetic order. The observed magnetic contribution can be indexed with a magnetic propagation vector \( \mathbf{k} = (0,0,0) \), that is, the magnetic unit cell coincides with the nuclear one.

The representational analysis technique described by Bertaut has been used to determine the possible magnetic structures compatible with the P–3 space group of I. This gives four possible magnetic configurations considering the two magnetic FeIII ions in the unit cell, namely (i) ferromagnetic with moments along c, (ii) antiferromagnetic with moments along c, (iii) ferromagnetic with moments in the ab plane, and (iv) antiferromagnetic with moments in the ab plane (it should be noted that PND is not sensitive, in the present symmetry, to the absolute orientation angle of the magnetic moments contained in the ab plane). In order to determine the magnetic configuration, we consider as a first approximation a fixed magnetic moment of 3.5 \( \mu_B \) in each FeIII site, in agreement with the expected value at 1.5 K for a Heisenberg 3D ordering of \( S = \frac{5}{2} \) spins and \( T_N = 2.4 \) K (see below) and we calculate the four possible magnetic structures to be compared against the experimental data. This comparison immediately reveals (see Figure S3) that the only configuration compatible with the data is antiferromagnetic with moments in the ab plane (configuration iv, Figure 3). In order to verify this result overcoming the symmetry restrictions, a simulated annealing method was applied, using a spherical coordinate system to describe the orientation of the moment. With the magnetic moment of the FeIII ions constrained, as previously, to a fixed value of 3.5 \( \mu_B \), but the spherical angles left free, the procedure always converges to a magnetic structure corresponding to moments almost exactly antiparallel and lying in the ab plane (Figure 3), thus confirming the previous result. Using this model, the Rietveld refinement with unconstrained magnetic moment (Figure 2) gives a value of 3.44 ± 0.25 \( \mu_B \) (\( R_{mag} = 12.30 \% \)). Single-crystal magnetic measurements (Figures S4–S6) fully agree with the c axis being a hard axis of magnetization and the ab plane an easy plane, as evidenced by \( M \) vs. \( H \) (Figure S5) and \( \chi \) vs. \( T \) data (Figures 4 and S6). The latter also confirms the value of \( T_N = 2.4 \) K obtained on the bulk.
Figure 3. View of the magnetic structure of $[\text{Fe(Metz)}_6](\text{FeBr}_4)_2$. Same colour code as in Fig. 1. The moments’ orientation in the $ab$ plane is arbitrary (see text).

The neutron data are also in good agreement with this value of $T_N$ since the intensity of the magnetic reflections decreases with temperature, vanishing between 2.3 and 2.45 K. The refined value of the magnetic moment, as shown in Figure 4, decreases with temperature, vanishing between 2.3 and 2.45 K. The refined value of the magnetic moment correspondingly decreases (Figure 4), showing a temperature dependence that approximately follows a power-law $C|T - T_N|^\beta$ corresponding to a Heisenberg 3D ordering of $S = 5/2$ spins (with $C = 5$, $t = |T - T_N|$, $T_N = 2.4$ K, and $\beta = 0.364$), as expected for Fe$^{III}$ ions. A 3D antiferromagnetic order in the same temperature range has indeed recently been reported in the plastic crystal choline$[\text{FeCl}_4]$, although the absence of structural data impedes magneto-structural correlation in this case. The magnetic order in I finds its origin in two types of interactions among these Fe(III) spins. First, the 1D symmetric stacks of FeBr$_4$– anions gives rise to a direct d-d ferromagnetic exchange interaction, that we previously evaluated on basis of the Fe···Fe distance along the stacks at 104 K. The shorter distance observed here in the 10 K structure provides a more accurate larger value of $J/k_B = 0.32$ K, which is in excellent agreement with that derived from a Curie-Weiss fit (Figure S7). The value of the critical temperature, $T_N = 2.4$ K, is a measure of the energy per spin involved in the ordering mechanism. Therefore, neglecting crystal-field interactions, since the intrachain energy is $|E_d| = JS^2 = 2.0$ K, there is a remaining 0.4 K that we associate to the inter-stack coupling, i.e. $J'/k_B$ of the order of $-0.09$ K. This weak though noticeable interaction could arise from exchange coupling through direct Br···Br contacts of the FeBr$_4$– ions within the $ab$ plane (Table S2, Figure S8), but the fact that the magnetic moments are oriented perpendicular to the ferromagnetic chains would also be consistent with a magnetostatic (dipolar) origin for the antiferromagnetic order. The estimation of the dipolar ground state yields $|E_{dp}| = 0.5$ K, thus making the latter hypothesis more plausible.

As mentioned before, pressure is one of the external stimuli used for switching the spin state in SCO based multifunctional materials. In the present case, besides the well-established stabilization of the LS state, pressure would likely affect the low temperature magnetic properties as well, and could be a suitable external parameter for addressing the magnetic ordering of the complex. The crystal structure of I was thus determined at RT at a pressure of 2.3 GPa, using a Diamond Anvil Cell and synchrotron radiation (see ESI for details). An ellipsoid representation of the structure is provided in Figure 5, showing the $[\text{Fe(Metz)}_6]^2+$ cation and one FeBr$_4$– anion. Significant compression of the cell is observed here in the 10 K structure provides a more accurate larger value of $J/k_B = 0.32$ K, which is in excellent agreement with that derived from a Curie-Weiss fit (Figure S7). The value of the critical temperature, $T_N = 2.4$ K, is a measure of the energy per spin involved in the ordering mechanism. Therefore, neglecting crystal-field interactions, since the intrachain energy is $|E_d| = JS^2 = 2.0$ K, there is a remaining 0.4 K that we associate to the inter-stack coupling, i.e. $J'/k_B$ of the order of $-0.09$ K. This weak though noticeable interaction could arise from exchange coupling through direct Br···Br contacts of the FeBr$_4$– ions within the $ab$ plane (Table S2, Figure S8), but the fact that the magnetic moments are oriented perpendicular to the ferromagnetic chains would also be consistent with a magnetostatic (dipolar) origin for the antiferromagnetic order. The estimation of the dipolar ground state yields $|E_{dp}| = 0.5$ K, thus making the latter hypothesis more plausible.

Figure 4. Left: Single-crystal molar magnetic susceptibility ($\chi$) vs. $T$ with 0.1 T dc field perpendicular (green) and parallel (orange) to the crystallographic c axis. The average $\chi$ (blue) compares well with bulk data (black). Right: temperature dependence of the magnetic moment obtained from PND at the indicated pressures. Solid lines are the power-law $C|T - T_N|^\beta$ with $C = 5$, $\beta = 0.364$, and $T_N = 2.4$ K (black), 2.7 K (green) and 3.1 K (orange).
Magnetic susceptibility ($\chi$) measurements under pressure (Figure 6, ESI for details) confirm the structural observations, the SCO centred at 165 K at ambient pressure being shifted to 325 K at 0.8 GPa. This indicates that the spin-state of the Fe$^{II}$ ion in 1 is in fact switched to LS at room temperature by only applying ca. 1 GPa. Moreover, as can be seen in Figure 6, the application of such pressures also affects the long-range magnetic ordering of 1. The low temperature data at various applied pressures indeed evidence a shift towards higher temperatures and broadening of the maxima corresponding to the antiferromagnetic ordering of the FeBr$_4^-$ ions. The maxima in the curves mark the ordering temperature, $T_N$, that increases with pressure following a linear dependence (Figure 6). PND under applied pressure confirms the enhancement of the magnetic interaction revealed by this $T_N$ increase. PND patterns collected at different temperatures between 1.5 and 10 K, and at applied pressures of 0.19 and 0.50 GPa are compatible with the same magnetic structure model described previously for ambient pressure (Figure S9). The increase of $T_N$ is deduced from the fact that a significant magnetic moment can still be refined at 2.5 K, with values of 2.32 ± 0.86 $\mu_B$ ($R_{\text{mag}} = 16.80 \%$) and 2.88 ± 0.60 $\mu_B$ ($R_{\text{mag}} = 14.23 \%$) for 0.19 and 0.50 GPa respectively. As shown in Figure 4, these values are in agreement with the same power-law dependence proposed for ambient pressure, $Ct^\beta$, ($C = 5$, $\beta = 0.364$) just scaling to the appropriate $T_N$ given by the susceptibility measurements (2.7 and 3.1 K for 0.19 and 0.50 GPa respectively). The increase of $T_N$ with applied pressure is likely a joint effect of the structural modifications of the FeBr$_4^-$ ions packing. Indeed, both the ferromagnetic exchange interaction within the FeBr$_4^-$ ions along $c$ and the antiferromagnetic dipolar interaction between these stacks in the $ab$ plane are expected to increase with reduced Fe$^{III}$$\cdots$Fe$^{III}$ distances.$^{21,23}$ The effect of the former should however be dominant since the corresponding Fe$^{III}$$\cdots$Fe$^{III}$ distances along the $c$ axis suffer a stronger reduction under pressure than those distances contained in the $ab$ plane (−7.7% vs. −5.6%, Tables 1 and S2), and the exchange interaction is much more sensitive to distance variations (∼$r^{-10}$) than the dipolar interaction (∼$r^{-3}$).$^{30}$

Conclusions

In this work, we have successfully applied neutron diffraction and single-crystal magnetic measurements to determine the magnetic structure of [Fe$^{II}$(Metz)$_6$(Fe$^{III}$Br$_4$)$_2$, a unique multifunctional material presenting a magnetically-ordered phase at low temperatures as well as the spin-crossover phenomenon. The effect of applied pressure on the magnetic order has been determined and correlated with aspects of the X-ray structure determined at higher pressures. The latter also demonstrated the efficient spin-crossover induced by pressure at ambient temperature, which was corroborated by magnetic measurements. Altogether our results highlight the great potential of simple two-network hybrid spin crossover systems to provide original multifunctional materials.

Acknowledgements

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Notes and references


In a mean-field approach and using the $-JS_i S_j$ formalism, the Weiss constant is defined as
$$\theta = \frac{zJS(S+1)}{3k_B},$$
where $z$ is the number of nearest neighbours for each spin $S$. For $z = 2$, as proper for the 1D stacks of FeBr$_4^-$ ions, the fit provides $\theta = 1.90$ K, thus yielding $J/k_B = 0.33$ K for the dominant intrachain exchange.