Chloride reduction of Mn³⁺ in the mild hydrothermal synthesis of a charge ordered defect pyrochlore, CsMn²⁺Mn³⁺F₆, a canted antiferromagnet with a hard ferromagnetic component

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Abstract

Geometrically frustrated systems play an important role in studying new physical phenomena and unconventional thermodynamics. Charge ordered defect pyrochlores $AM^{2+}M^{3+}F_6$ offer a convenient platform for probing the interplay between electron distribution over M^{2+} and M^{3+} sites and structural distortions; however, they are limited to compounds with $M^{2+/3+} = V$, Fe, Ni, and Cu due to difficulties in the simultaneous stabilization of other 3d elements in the +2 and +3 oxidation states. Herein, we employ Cl⁻ anions under hydrothermal conditions for the mild reduction of Mn₂O₃ in concentrated HF to obtain the CsMn²⁺Mn³⁺F₆ composition as a phase pure sample and study its properties. The magnetism of CsMn₂F₆ was characterized by measuring the magnetic susceptibility and isothermal magnetization data, and a magnetic transition to a canted antiferromagnet state was found at 24.1 K. We determined the magnetic structure of CsMn₂F₆ using powder neutron diffraction, which revealed successive long-range ordering of the Mn²⁺ and Mn³⁺ sites that is accompanied by a second transition. The role and strength of magnetic exchange interactions were characterized using DFT calculations.

Introduction

The pyrochlore structure-type has long been used for the design of new materials with a number of potential applications encompassing magnetism, ^{1–5} oxygen evolution reactions and storage, ^{6–8} energy storage materials, ^{9,10} optical materials, ¹¹ and others. ¹² The pyrochlore structure, $A_2B_2O_7$, consists of two interpenetrating vertex-sharing tetrahedral frameworks with *A* and *B* metal cations, which offers a unique atomic arrangement that promotes frustrated magnetism.^{3,13–16} This arrangement has resulted in the development of a number of rare-earth titanate and zirconate compounds exhibiting spin ice and spin liquid behavior.^{17–20} A spin ice does not have a single ground state with a minimum energy; on the contrary, magnetic frustration results in multiple ground states with a non-zero residual entropy, making a spin ice material closely related to water ice and offering a unique magnetic system to study quantum effects and abnormal thermodynamics in extended solids. To express the degree of frustration in a material and show how the magnetic ordering temperature deviates from the magnetic exchange interactions, one can use the frustration index *f* = $|\Theta_{CW}|/T_C$, where Θ_{CW} is the Curie-Weiss temperature and *T*_C is magnetic ordering temperature. Large values of the frustration index, *f*, indicate possible candidates for spin liquid materials and many highly frustrated materials based on pyrochlore structure have been reported to date.

Although the parent pyrochlore structure type is based on an oxide material, other classes of inorganic materials can readily adopt the same structure type, paving a path toward the design of new geometrically frustrated materials. One of the recent advances in this direction has been oxygen replacement with fluorine to create a new class of frustrated materials based on 3d metals with a NaAB₂F₇ (A = Ca or Sr; B = Fe, Co, and Ni) composition, which have been grown as large single crystals *via* the floating zone method.²¹⁻²⁴ Geometric frustration in these pyrochlore antiferromagnets with magnetically coupled 3d cations that occupy one of the two tetrahedrally connected sublattices results in a low temperature spin freezing point and high frustration indexes f, from 19 to 58. Another intriguing class of geometrically frustrated fluorides with composition $AM^{2+}M^{3+}F_6$ (A = alkali cations and NH₄⁺, M^{2+} and M^{3+} are 3d transition metal cations such as $Mn^{2+/3+}$, $Fe^{2+/3+}$, Co^{2+} , Ni^{2+}) exhibits a modified pyrochlore structure, in which the A site is half-occupied and one fluorine atom is missing for charge balance.^{25,26} $AM^{2+}M^{3+}F_6$ defect pyrochlores can adopt either a disordered cubic structure with both M^{2+} and M^{3+} cations occupying the same site or an orthorhombic structure, in which the 3d metal cations are charge ordered between the two sites. One notable compound that belongs to the cubic crystal systems is CsNiCrF₆, which has been extensively studied with respect to its magnetic properties and, despite strong antiferromagnetic interactions between the metal cations, does not exhibit magnetic long-range order at any temperature, with a spin-glass anomaly at about 2.3 K.^{27–29} Of all possible M^{2+} and M^{3+} combinations, charge ordered

 $AM^{2+}M^{3+}F_6$ phases are less numerous and include, for example, $AFe^{2+}Fe^{3+}F_6$ ($A = NH_4^+$, Rb^+ , Cs^+), $AV^{2+}V^{3+}F_6$ (A = Rb⁺ and Cs⁺), $ANi^{2+}Ni^{3+}F_6$ (A = Rb⁺ and Cs⁺), CsCu₂F₆, CsAgFeF₆, NH₄ZnAlF₆, and NH₄CoAlF₆ compositions.^{30–39} RbFe₂F₆, which has been obtained hydrothermally by reacting respective metal fluorides with trifluoroacetic acid, orders antiferromagnetically at 16 K with two orthogonally oriented magnetic moments on the Fe sites. This compound has a significant frustration index f of 17 and exhibits anharmonic magnon excitations found by neutron spectroscopy.⁴⁰ Complete understanding of what drives the formation of charge ordered $AM^{2+}M^{3+}F_6$ orthorhombic phases over the cubic ones is yet to be attained, but some orthorhombic phases, e.g. CsFe₂F₆ and CsV₂F₆, undergo a phase transition to the cubic phase upon heating, indicating possible transitions between the two structure types. Since almost all charge ordered defect pyrochlores contain the same 3d element on the M sites, there are still gaps in this family of highly frustrated defect pyrochlores. One group includes the compositions containing Mn on both M sites, and has a reverse, as compared to AFe_2F_6 , spin arrangement on the M^{2+} and M^{3+} sites, S = 5/2 and 2, respectively, for the Mn compound. Given the overall tetrahedral framework arrangement of the M sites in the AM_2F_6 structure, one would not expect a significant difference in the magnetism upon replacement of Fe by Mn; however, there are distortions in the M coordination environments that can cause a difference in their properties. The purpose of this work was to study the effect of such Fe to Mn replacement and to probe its influence on the properties of resulting compound belonging to charge ordered defect pyrochlore family.

In this manuscript, we report for the first time the hydrothermal synthesis of $CsMn_2F_6$ by Mn_2O_3 reduction with Cl^- and its full magnetic and structural characterization with respect to its nuclear and magnetic structures (Figure 1). We examined the influence of ionic radii of the constituent elements in the structures of $12 NaAB_2F_7$ and $90 AM^{2+}M^{3+}F_6$ compounds, revealing that smaller *M* cations favor the formation of defect pyrochlore structure. The magnetic properties of $CsMn_2F_6$ were characterized by measuring magnetic susceptibility vs. temperature and magnetization vs. field plots. After finding a magnetic transition at around 24.1 K, we studied the magnetic structure of $CsMn_2F_6$ below this temperature and found a second transition between 10 and 14 K. The magnetic structures support the presence of weak spontaneous magnetization, in contrast to the purely antiferromagnetic RbFe₂F₆ analogue. $CsMn_2F_6$ also demonstrates a surprising transition of its ferromagnetic component from an extremely soft, highly susceptible magnet at around 24 K to a hard magnet with coercive field significantly exceeding 5 T at 2 K, suggesting a potential use of such materials for reliable data storage in high magnetic field environments or magnetic field recording.



Figure 1. (top) Synthesis of $CsMn^{2+}Mn^{3+}F_6$ via hydrothermal synthesis using CsCl as a reducing agent. $CsMn^{2+}Mn^{3+}F_6$ exhibits a magnetic transition at 24.1 K to a canted antiferromagnetic state, in which (middle) coercive field grows rapidly as temperature decreases. (bottom) The calculated magnetic structure is in a very good agreement with the experimental magnetic structure and highlights the important role of spin canting in its stabilization.

Results and discussion

Synthesis and crystal structure. Our successful application of the mild hydrothermal synthesis method has enabled us to prepare a large number of new fluoride compositions, which motivated us to employ this method for obtaining $CsMn_2F_6$.^{41–48} After several trial syntheses and reaction conditions optimization, we found that the target phase forms readily by partially reducing Mn_2O_3 with CsCl at 200 °C $(E^0(Mn^{3+}/Mn^{2+}) = 1.54 \text{ V}, E^0(Cl_2/2Cl^-) = 1.36 \text{ V})$ in concentrated HF according to the following equation:

 $2 Mn_2O_3 + 2 CsCl + 12HF = 2 CsMn^{2+}Mn^{3+}F_6 + 6H_2O + Cl_2$

We found that the product of the reaction is strongly dependent on the Mn_2O_3 to CsCl ratio. A phase pure sample of CsMn₂F₆ can be obtained using a 1: 2 ratio, while even a slight deviation from this ratio or from the quantity of HF used results in the formation of side products, e.g. CsMnF₄·2H₂O, CsMnF₄, CsMnCl₃·2H₂O, or MnF₂.⁴⁹ To illustrate the role of the chloride ion as a reducing agent, we performed a series of reactions where CsCl was gradually replaced by CsF. Upon decreasing of the CsCl content in the reaction, the amount of Mn³⁺-containing side products, CsMnF₄·2H₂O and CsMnF₄, increases (Figures 2 and S1, Table S1). Ultimately, a complete substitution of CsCl by CsF results in the formation of only Mn^{3+} -containing products $CsMnF_4$ and $CsMnF_4 \cdot 2H_2O$. At the same time, additional sources of Cl^- (e.g. HCl) in the reaction lead to the formation of Mn^{2+} -containing products, such as $CsMnCl_3 \cdot 2H_2O$ and MnF_2 .



Figure 2. (left) PXRD of the product for the reaction between Mn_2O_3 , CsCl, and CsF as a function of the CsCl/CsF molar ratio in 2 θ range 26.0–29.5 deg. Red, blue, and green areas highlight the diffraction peaks for CsMn²⁺Mn³⁺F₆, CsMn³⁺F₄, and CsMn³⁺F₄·2H₂O, respectively. Red, blue, and green percentages are the content of CsMn²⁺Mn³⁺F₆, CsMn³⁺F₄, and CsMn³⁺F₄·2H₂O, respectively, estimated using whole pattern fitting. (right) CsMn₂F₆ percentage in the product as a function of Cl⁻ content in the starting reaction mixture.

Although a number of related fluorides with formula $AM^{2+}M^{3+}F_6$ have been obtained using flux methods and large (mm³ to cm³ size scale) single crystals were grown,²⁵ our own attempts to grow large crystals of CsMn₂F₆ using fluxes were not successful and resulted in the formation of known manganese fluorides. Larger single crystals (~0.1 mm³) were grown *via* partial replacement of CsCl with CsF and using more diluted HF, creating less reducing conditions and slowing down the crystallization process. The purity of the product was confirmed by powder X-ray diffraction (PXRD) and neutron diffraction; moreover, no magnetic transitions that can be associated with other manganese phases were found in the magnetic susceptibility data.

We determined the structure of $CsMn_2F_6$ using single crystal X-ray diffraction and found that it crystallizes in a distorted defect pyrochlore structure (*Pnma* space group, Tables S2 and S3) with unit cell parameters a = 7.3013(2) Å, b = 7.7141(2) Å, and c = 10.1926(3) Å. The unit cell parameters agree well with previously reported values found from PXRD data, but the crystal structure of this compound has not been reported.⁵⁰ The structure consists of distorted MnF₆ octahedra that are connected through the vertices to build up a framework with pores that are occupied by Cs cations (Figure 3). The Mn–F bond distances

allow for unambiguous distinction between the Mn^{2+} and Mn^{3+} cations: the $Mn^{2+}F_6$ octahedra have four longer, 2.1535(12)–2.1612(13) Å, and two shorter, 2.0205(18) and 2.0416(18) Å, bonds, while $Mn^{3+}F_6$ octahedra exhibit four shorter, 1.8594(12)–1.8682(12) Å, and two longer, 2.0414(6) Å, bonds (Table S4). The bond-valence sum calculations result in values of 1.97 and 3.02 for Mn^{2+} and Mn^{3+} , respectively, indicating a fully charge ordered distribution of the Mn atoms in the structure. In contrast to the $RbFe^{2+}Fe^{3+}F_6$ analog, in which Fe^{3+} atoms are located in a nearly regular octahedral environment with Fe^{3+} –F distances varying over the range of 1.91–1.95 Å, the geometry of the $Mn^{3+}F_6$ octahedron is strongly affected by a Jahn-Teller distortion.⁵⁰ Interestingly, although the other Mn site should not be affected by a Jahn-Teller distortion, as it contains a $3d^5$ cation, there is nonetheless a strong distortion of the $Mn^{2+}F_6$ octahedron, in which one of the F–Mn–F angles is 111.15(7)° (Table S5), which can be attributed to either a structural distortion inherent to the structure type or to a cooperative Jahn-Teller distortion.³⁸ In the Mn sublattice, each Mn is separated from six other surrounding Mn by 3.63–3.86 Å, creating a nearly uniformly spaced net of magnetically interacting Mn atoms.



Figure 3. (left) A view down the x-axis of the structure of $CsMn_2F_6 \equiv CsMn^{2+}Mn^{3+}F_6$ and (right) its Mn-sublattice, which consists of corner sharing $Mn^{2+}_2Mn^{3+}_2$ tetrahedral units.

As the structure is influenced by the size of its constituent elements, we made an attempt to identify the size parameters that favor the formation of defect pyrochlores as well as their preference for either the ordered or the disordered structure type. Lui and coworkers introduced a tolerance factor for the pyrochlore structure type based on $A_2M_2O_7$ series and defined it as $\tau = \frac{3(R(A) + R(O))}{\sqrt{17}((R(M) + R(O)))}$, where τ falls into a small range of 0.826–0.943 for pyrochlore oxides.⁵¹ By using the same equation for over 90 defect pyrochlore fluoride structures of the type $AM^{2+}M^{3+}F_6$, we found that all of them have a tolerance factor ranging from 1.032 to 1.191 (Table S9), whereas for 12 pyrochlore fluorides of the type $AA'M^{2+}_2F_7$, the tolerance factor ranges between 0.925 and 0.974 (Table S10). Given that there is a gap between the two ranges for fluoride structures, one can surmise that a regular pyrochlore structure will likely form for $\tau <$

1, whereas a $\tau > 1$ indicates a higher probability of defect pyrochlore formation. We found it less straightforward to predict which variation of the defect pyrochlore structure, ordered orthorhombic or disordered cubic, will form based on the size of the constituent ions. Since a larger difference between M^{2+} and M^{3+} ionic radii should promote the formation of the charge ordered phase as the large size difference hinders two elements from sharing the same site,³⁹ we plotted the size difference as a function of the A cation ionic radius (Figure 4). Although a large M cation size difference, >0.25 Å, clearly favors the formation of the ordered structure, both structure types are observed when the size difference falls below 0.25 Å. Molokeev et al. showed that another possible reason for symmetry lowering from cubic to orthorhombic is a mismatch between the A- and average M-site sizes, in which a small A cation size can cause a structural distortion that reduces the volume that the A cation occupies.³⁴ As can be seen from Figure 4, there is a large number of cubic phases with $A = Cs^+$ or Rb^+ when the average size of the framework-forming M cations is below 0.72 Å. For even smaller A-cations ($A = NH_4^+$ and K^+) both the cubic and the orthorhombic phases form for an average *M*-site size range of 0.60–0.70 Å. The importance of the A-site cation size can also be illustrated by an orthorhombic to cubic phase transition at high temperatures observed in several orthorhombic AM_2F_6 phases. Although one would expect all thermal motion in an AM_2F_6 structure to change simultaneously with the temperature, neutron diffraction studies on $RbFe^{2+}Fe^{3+}F_6$ have revealed that the thermal ellipsoids, and thus the effective radius, of the A-site cation increases disproportionally faster than the other sites in the structure as the temperature increases, reducing the size difference between the A- and M-sites.³³ Due to this, compositions such as RbV_2F_6 , CsV₂F₆, and CsFe₂F₆ adopt a charge ordered orthorhombic structure type at room temperature and undergo a transition to the cubic structure type, accompanied by charge disordering, at high temperature.



Figure 4. Radius of the A cation (top) vs. the size difference between M^{2+} and M^{3+} cations and (bottom) average *B* site size in the $AM^{2+}M^{3+}F_6$ defect pyrochlore family. Black and blue circles correspond to disordered cubic and charge ordered orthorhombic pyrochlores, respectively. Compounds that crystalize in both charge ordered and disordered structures or exhibit a high temperature transition from orthorhombic to cubic phase are shown with red circles. Gray and blue areas schematically represent the stability range for the cubic and orthorhombic pyrochlore phases, respectively. Ionic radii data were taken from Shannon⁵² and NH₄⁺ ionic radii data were taken from ref.⁵³ where for *A* and *M* cations coordination numbers of XI and VI were used, respectively.

A reversible transition from orthorhombic to cubic defect pyrochlore is facilitated for compositions where both M^{2+} and M^{3+} sites are occupied by the same element in the +2 and +3 oxidation states, resulting in an abundance of orthorhombic phases with ternary *vs.* quaternary compositions. In fact, all ternary AM_2F_6 defect pyrochlores are orthorhombic because, for them, the cubic to orthorhombic transition can readily be achieved by a simple charge (i.e. electron) redistribution. In contrast, the transition from cubic to orthorhombic in quaternary $AM^{2+}M'^{3+}F_6$ phases has to be accompanied with the redistribution of the actual cations. For that reason, the disordered cubic phase can be obtained under either high temperature conditions in flux reactions with $T > 1000 \, ^\circ C$,²⁵ where the effective size of the *A* cations is high due to

disproportionally high increase in their thermal motion in the resulting crystal structure, or by entangling the mobility of the M cations (Figure 5). A good illustration of the latter is the recently obtained (NH₄)MnFeF₆ defected pyrochlore, which can be synthesized as a cubic phase via ball milling of the fluoride precursors.⁵⁴ Another example is the decomposition of [Co³⁺(NH₃)₆]AlF₆ at 280 °C, which results in the formation of cubic NH₄Co²⁺AlF₆, whereas a reaction between CoF₂, AlF₃, and NH₄F under hydrothermal conditions at 400 °C yields an ordered orthorhombic phase.³⁹ Moreover, the cubic phase of NH₄CoAlF₆ can be transformed to the orthorhombic one by annealing it at 350 °C for one week, which enables the mobility of the M cations in the solid state, illustrating the metastable nature of the cubic phase. One would expect that the phases in the intermediate size difference regime in Figure 4 could also undergo a similar cubic to orthorhombic ordering transition through prolonged heating at elevated temperatures; although, as of yet, there are no reports in the literature of this process. The overall conclusion is that the formation of the orthorhombic phases is favorable for the ternary phases and can be promoted by 1) a larger difference between the M^{2+} and M^{3+} cation sizes, 2) a smaller size of the A cation, and 3) mid-temperature reaction conditions, whereas the formation of the cubic phases is unlikely for the ternary phases and can be promoted by either 1) high temperature flux synthesis or 2) the decomposition of a precursor at rather low, below 300 °C, temperatures.



Figure 5. A schematic representation of defect pyrochlore structure stability as a function of temperature. Low diffusion rates at low temperature stabilize the cubic phases with M^{2+} and M^{3+} statistically distributed over the M site. Intermediate temperature range with fast M cation diffusion promotes the formation of the orthorhombic phases. At high temperatures, the cubic phase is stabilized due to the A cation large effective size, which increases disproportionately relative to the size of other cations.

We performed thermogravimetric analysis of $C_{s}Mn_{2}F_{6}$ under nitrogen flow to probe if it undergoes a phase transition to the tetragonal or cubic crystal system, as was observed in the case of the $CsFe_2F_6$ analog. The differential thermal analysis (DTA) data showed an endothermal effect with an onset temperature of 410 °C, which is not associated with any weight loss (Figure S2). Such an effect could be indicative of a phase transition and high temperature PXRD (HT PXRD) was employed to check this possibility. PXRD patterns of a powder sample of CsMn₂F₆ were collected under vacuum up to 500 °C with 50 °C step intervals (Figure S3). The PXRD patterns from room temperature to 400 °C indicate no changes in the sample symmetry, followed by sample decomposition at a temperature between 400 and 450 °C, in a good agreement with the DTA endothermic peak observed at 410 °C. As the decomposition products could not be identified from the HT PXRD pattern, a fresh sample of CsMn₂F₆, which was placed into a carbon crucible and sealed under vacuum inside a silica tube, was heated to 450 °C overnight and an ex-situ room temperature PXRD pattern was collected. In the silica tube reaction, two major products formed during the decomposition, MnF₂ and Cs₂MnF₆, along with a Mn₂O₃ impurity due to a reaction between $CsMn_2F_6$ and residual moisture and air in the tube, indicating the disproportionation of Mn³⁺ in the fluoride starting compound. Although MnF₂ and Cs₂MnF₆ do not match with the HT PXRD pattern at 450 °C, both methods confirm the thermal decomposition of CsMn₂F₆ at a temperature between 400 and 450 °C and a lack of a structural transition to the tetragonal or cubic crystal system.

Magnetism. The magnetic properties of $CsMn_2F_6$ were characterized by collecting DC magnetic susceptibility vs. temperature data over the 2–300 K range using polycrystalline powder and also a multidomain single crystal (Figures 6 and S4, respectively). The magnetic susceptibility plot can be divided into two temperature regions: a low-temperature region below 100K in which a magnetic transition is visible at 24.1 K and high-temperature region (100 to 300K) exhibiting Curie-Weiss behavior. The inverse magnetic susceptibility data were fitted with a linear function over the 150–300 K range (Figure 6), and a Weiss temperature of –93.7 K was derived from it, indicating antiferromagnetic correlations in the sample. Dividing the Weiss temperature by the ordering temperature results in a frustration index *f* of 3.9, which indicates low frustration in the system. The measured magnetic moment per formula unit of 8.01 μ_B is above the theoretical value of 7.68 μ_B that was calculated using the spin-only magnetic moments for Mn²⁺ and Mn³⁺ cations, 5.92 and 4.90 μ_B , respectively. Although higher experimental magnetic moments for Mn³⁺ have been previously reported and can be one possible explanation of higher than expected paramagnetic moment of CsMn₂F₆,^{55,56} the moment can also be adjusted to lower values by including a small temperature independent paramagnetic (TIP) component. While fitting the data to a modified Curie-Weiss law results in a negligibly small negative χ_0 ($\chi_0 \sim$ -0.00009 emu/mol), using a small positive χ_0 of 0.0011 emu/mol results in a magnetic moment of 7.66 μ_B (Figure S5), which is in good agreement with the expected theoretical value of 7.68 μ_B , and insignificantly decreases the quality of the data fit, see Figure S5. In the low temperature region, below 100 K, the susceptibility plot deviates from Curie-Weiss behavior and ultimately undergoes a transition to an ordered magnetic state. The d*M*/*H*d*T* vs. *T* plot indicates a clear transition at 24.1 K (Figure 6); there are no other apparent transitions at lower temperatures. A small difference in the magnetization between the ordered and paramagnetic states, ~0.012 vs. ~0.005 emu, indicates either a ferri- or canted antiferromagnetic ordering in the sample. The AC data exhibit a single sharp peak at a temperature of 24.1 K, indicating a single transition to a long range ordered magnetic state. The sample is unlikely to exhibit spin liquid behavior based on the overlapping AC susceptibility plots collected at different frequencies (Figure 6).



Figure 6. (top) DC molar magnetic susceptibility (χ_{mol}) and inverse susceptibility (χ_{mol}^{-1}) vs. temperature plots of polycrystalline CsMn₂F₆. (middle) Enlarged 0–30 K region showing zero-field cooled χ_{mol} vs. *T* plot along with dM/HdT vs. *T* plot, which exhibits only one peak corresponding to a magnetic transition at 24.1 K and (bottom) AC susceptibilities as function of temperature and frequency.

To better understand the nature of the low temperature magnetic transition, a series of magnetization vs. field (MvH) data were collected in the temperature range of 2 to 25 K. Above 24 K, the MvH plots exhibit a proportional change of the magnetization as a function of field (Figure 7), which is typical for paramagnetic samples that do not reach their saturation magnetization in the applied field. Right below the transition temperature, at 23.5 K, the magnetization changes rapidly when a weak non-zero field is applied in either direction, indicating a weak metamagnetic effect as the applied field H approaches zero. At temperatures between 23.5 and 10 K, a hysteresis loop appears and extends to stronger fields as the temperature decreases. The low remnant magnetization of the sample and its linear behavior in higher applied fields indicates a small ferromagnetic component in a predominately antiferromagnetic system, which is characteristic of ferrimagnets or canted antiferromagnets. It is worth noting that all hysteresis curves start with a non-zero magnetization, despite demagnetization of the sample by warming it to room temperature after each MvH data collection and zero-field cooling to the next measurement temperature. The non-zero magnetization is likely due to the metamagnetic transition at 23.5–24.5 K: when crossing this temperature region, even a small applied field, such as a residual field in a SQUID instrument, can magnetize the sample. Our attempts to minimize the residual field by using scans with the Pd standard were not successful, and even a small, nearly undetectable field well below 10^{-4} T resulted in sample magnetization upon cooling. The MvH plots below 10 K show gradual "closing" of the hysteresis loop until it turns to a direct proportional dependence of the applied field on the magnetization, with an offset magnetization value. It is noteworthy that at even a 5 T applied field does not change the magnetization of the sample and shows that the ferromagnetic component in the sample turns from an extremely soft magnet that is sensitive to the applied field at around 24 K to a very hard one at 2 K.

Since the small ferromagnetic component in the magnetization vs. field plots are obscured by a significant antiferromagnetic component, we derived the ferromagnetic components' M vs. H plots by subtracting linear antiferromagnetic contributions from the full magnetizations (Figure 8). The derived plots show a clear magnetic transition at 23.5 K with a small saturation moment of $0.8 \cdot 10^{-2} \mu_B/f.u$. As the temperature decreases to 20 K, one can observe a hysteresis loop with a coercive field $H_c \approx 0.1$ T, which then nonlinearly increases as temperature decreases, to ≈ 0.2 and 0.5 T at 17.5 and 15 K, respectively. When the temperature approaches 10 K, H_c reaches a high value of ≈ 1.7 T. A further temperature decrease hardens the ferromagnetic component, and even a 5 T maximum magnetic field is insufficient to fully

revert the magnetization and to establish the coercive field of the sample at 5 and especially 2 K, where no change of remnant magnetization occurs after a full $-5 \rightarrow 5$ T field sweep of the sample. The "locked" remnant magnetization suggests that the coercive field in the sample is significantly higher than the maximum applied field, 5 T, and provides evidence of a dramatic change of coercive field in a narrow temperature range, from $H_c \approx 0$ at a transition temperature of 24.1 K to $H_c >> 5$ T at 2 K. Such an effect, if used in a low temperature environment, such as space, or if brought to a more practical temperature range, can have potential applications in long term information storage in situations where reliable data preservation is needed when the data storage device is exposed to high magnetic fields. Further characterization of the sample and the understanding of the remnant magnetization origins required magnetic structure characterization of the sample.



Figure 7. Magnetization as a function of field M(H) plots of CsMn₂F₆ measured at 24.5 (above $T_c \approx 24.1$ K), 23.5 (slightly below T_c), 15, 5 and 2 K. All samples were measured with -5 to 5 T field sweeps, but only -1 to 1 T regions are shown. There is a weak metamagnetic transition at 23.5 K corresponding to an easy magnetization reversal when a weak magnetic field is applied. As temperature decreases, the coercive field of the ferromagnetic component increases until it becomes a very hard magnetization even when a full 5 to -5 T sweep is made, indicating a very hard ferromagnetic component in the sample. The orientation of initial magnetization depends on the direction of a residual magnetic field in the instrument.



Figure 8. Magnetization as a function of field plots for ferromagnetic components of $CsMn_2F_6$ between 23.5 and 10 K. The plots were derived by fitting linear antiferromagnetic part and subtracting it from the magnetization vs. field plots shown in Figure 7.

Neutron diffraction and magnetic structures. To determine the magnetic structure of $CsMn_2F_6$ and to further characterize the "locked" remnant magnetization effect at low temperatures, we collected temperature dependent neutron diffraction data. Several diffraction peaks appeared below the transition temperature of ~24.1 K (Figures 9 and S6–S24) that indicate long-range magnetic order and that correspond to a propagation vector $\mathbf{k} = (0,0,0)$. We found a good fit to the experimental data below the transition point by refining the magnetic structure in the *Pnm'a'* magnetic space group. The magnetic structure refinement revealed a large magnetic moment on the Mn³⁺ atoms, $3.25(2) \mu_B$ at 18 K, which aligned in a canted antiferromagnetic fashion with an angle of 167.2° between two neighboring magnetic vectors (Figure 10). The other Mn-sublattice exhibits a low magnetic moment of $0.59(5) \mu_B$, which is inconsistent with the $3d^5$ configuration of the Mn²⁺ atoms and indicates only a partial magnetic ordering of the sites. Like the Mn³⁺-sublattice, the magnetic moment of the unit cell, $0.83 \mu_B$ at 18 K, is aligned along the *a* direction, and mostly consists of the magnetic moment on the Mn²⁺ atoms, $0.636 \mu_B$ along the *a* axis (Table S7). The magnetic moments along the other two directions cancel each other out due to symmetry constraints on both Mn sites.



Figure 9. (top) Neutron diffraction patterns of $CsMn_2F_6$ at 26, 18, and 1.5 K. The compound undergoes a transition first from the *Pnma* to the *Pnm'a'* magnetic space group and then to the $P2_1/c$ space group. Diffraction peaks due to magnetic ordering are highlighted with blue and orange for the first and the second transition, respectively. (bottom) Rietveld refinement plot of neutron diffraction data collected at 18 K.



Figure 10. Magnetic structure of $CsMn_2F_6$ at (top) 18 K and (middle) 1.5 K. (bottom) Magnetic moments on the manganese atoms in the magnetic structure of $CsMn_2F_6$ at 1.5, 6, 10, and 14 - 23 K. The dashed lines are guides to the eye.

At a temperature of 10 K and below, the magnetic structure refinement resulted in unaccounted for intensities on the magnetic peaks along with the emergence of additional magnetic peaks, indicating that the sample undergoes a second magnetic transition, which was not detected in the magnetic susceptibility data. A search through the subgroups of the *Pnm'a'* group revealed $P2_1/c$ as a potential lower symmetry magnetic space group, which was confirmed by the successful Rietveld refinement of the neutron diffraction data (Figures S22–S24). In the low temperature magnetic structure at 1.5 K, the Mn³⁺ atomic positions, which already acquired a high moment after the first transition, split into two crystallographically independent positions, Mn1 1 and Mn1 2, and slightly increase their magnetic moments from 3.25(2) to 3.40(3) and 3.52(4) μ_B , while the Mn²⁺ cations significantly increase their magnetic moments from 0.59(5) to 4.13(3) μ_B , in perfect agreement with the higher unpaired electron count on the Mn²⁺ cations. The Mn²⁺ sites show significant increase of their magnetic moments along the b axis, which are prohibited by symmetry in the *Pnm'a*' space group. The net magnetic moment in the unit cell is symmetry constrained along the *a* axis of the original cell (corresponds to the *b* axis in the standard setting of the $P2_1/c$ space group) and is equal to 0.69 μ_B (Table S8). Unlike in the higher temperature magnetic structure, the net moment is dominated by the contribution from Mn1 1 and Mn1 2 atoms, while the moment on the Mn2 sites is oriented in an antiparallel fashion, reducing the overall magnetic moment of the cell. The small value of the net magnetization, which corresponds to a weak canting of predominantly antiferromagnetically ordered spins and which is almost unchanged during the second phase transition (0.64 vs 0.69 μ_B), agrees well with the absence of any significant changes in the susceptibility data and the presence of the remnant magnetization observed in the sample in M vs. H plots (Figure 7).

Neutron diffraction data that were collected in a temperature range between 1.5 and 24 K allowed us to examine how the magnetic moments on both lattices vary as the temperature changes (Figure 10). The moment on the Mn1 rapidly saturates as the temperature decreases, reaching $\approx 3.3-3.4 \mu_B$ at 17 K. Transition to the low temperature modification splits the site into two, but the magnetic moment on both sites stays at approximately the same saturation value. The magnetic moment on the Mn2 sites displays a different behavior: at first it slowly increases with an increasing rate as the temperature decrease, then rapidly increases at around the second transition temperature, and finally reaches a maximum at 1.5 K. Although there are not enough datapoints at low temperatures, one can surmise that the moment of 4.13(3) μ_B found for Mn2 site at 1.5 K is very close to its saturation limit.

There are a number of compounds with $AM^{2+}M^{3+}F_6$ composition, but only a few of them exhibit charge ordering and long range magnetic ordering of the *M* sites.^{33–35,40,57,58} Among them is a series of isostructural iron compounds $AFe^{2+}Fe^{3+}F_6$ ($A = NH_4^+$, Rb^+ , Cs^+) that undergo a one-step antiferromagnetic transition at temperatures of 19, 16, and 13.7 K, respectively.^{31,33,34,57} Although only the magnetic structures of the NH₄⁺ and Rb⁺ analogs have been reported, similar magnetization behavior of CsFe₂F₆ suggests that it might have a similar magnetic structure. As in CsMn₂F₆, the nuclear and magnetic unit cell in NH₄Fe₂F₆ and RbFe₂F₆ have identical metrics; however, the symmetry of the CsMn₂F₆ magnetic unit cell after the first transition is different from the symmetry of the Fe-based analogs. Both Fe containing compounds undergo a transition to the *Pnma* magnetic space group (#62.441), in which the net unit cell magnetization is constrained to zero by symmetry. The Fe1 sites can have magnetic components in all three M_x , M_y , and M_z directions and the Fe2 site moment is restrained to have zero M_x and M_z components. Although there are three degrees of freedom of the magnetic moment on the Fe1 site, it aligns along the *a* direction with $M_x = -4.29 \ \mu_B$ and $M_y = M_z = 0 \ \mu_B$ in RbFe₂F₆. The Fe₂ site acquires magnetization along the only symmetry allowed direction, the b direction, with $M_y = 3.99 \mu_B$, resulting in an orthogonal arrangement of the magnetic moments on the two different metal sublattices. In contrast to the Fe analogs, $CsMn_2F_6$ exhibits a magnetic structure with *Pnm'a'* (#62.447) symmetry, for which the net magnetization of the unit cell can be non-zero along the a axis. The Mn1 site orders predominately along the b direction, $M_x = 0.05(10)$, $M_y = -3.234(14)$, $M_z = 0.36(6) \mu_B$, while the Mn2 site magnetization is perpendicular to the b axis due to symmetry constraints, $M_x = 0.16(8)$, $M_z = 0.57(4) \mu_B$. Given that the Mn2 site does not exhibit its full magnetic moment and stays in a mixed ordered/paramagnetic state, one can surmise that there is exchange between Mn1 and Mn2 site that prevents the latter from complete ordering. As temperature decreases and the symmetry of $C_{s}Mn_{2}F_{6}$ reduces to $P2_{1}/c$, the symmetry constraint on the Mn2 M_y component lifts, and it readily acquires magnetization along the b axis, resulting in an overall nearly parallel alignment of the magnetic moments, in stark contrast to orthogonal moments in RbeFe₂F₆. To better understand such magnetic structure transitions in CsMn₂F₆, we performed Density Functional Theory (DFT) calculations to shed light on exchange interactions occurring in this compound.

Theoretical calculations of magnetic exchange interactions in CsMn₂F₆

To clarify the complex magnetic behavior of $CsMn_2F_6$, we have conducted first principles DFT calculations. Five different collinear magnetic structures were selected for calculations: a ferromagnetic case (FM); a ferrimagnetic structure where different magnetic moments on the Mn1 and Mn2 sites orient antiparallel (FI1); a state with each of these fourfold sites further subdivided into two antialigned twofold sites (FI2); and two antiferromagnetic states – the groundstate AF1, which most closely resembles the experimental magnetic structure, and AF2, which corresponds to the first excited state (Figure 11).



Figure 11. A depiction of the calculated AF1 groundstate. Atoms labelled Mn1 and Mn2 have a spin moment of 3.49 μ_B , Atoms Mn3 and Mn4 have $-3.49 \mu_B$, while atoms Mn5 and Mn6 have $4.22 \mu_B$ spin moment and Mn7 and Mn8 have $-4.22 \mu_B$ spin moment. In the first excited state AF2 Mn5 and Mn6 have $-4.22 \mu_B$ moment and Mn7 and Mn8 have $4.22 \mu_B$ spin moment, with the other moments unchanged. Atoms Mn1–4 occupy the crystallographic Mn1 site and atoms Mn5-8 the crystallographic Mn2 site.

The results of DFT calculations are summarized in Table 1 and agree very well with the experimental observations. Despite significantly varying energies of the four most energetically favorble optimized structures, up to 59 meV/Mn atom, the magnetic moments on the Mn sites are consistent across the different structure, falling in the narrow ranges of $3.47-3.53 \ \mu_B$ and $4.22-4.23 \ \mu_B$ for Mn1 and Mn2, respectively, indicating their local character. The groundstate structure AF1 confirms charge ordering in CsMn²⁺Mn³⁺F₆ structure and is in excellent agreement with the experimental values obtained in the neutron diffraction experiment, $3.49 \ \mu_B$ vs 3.40(3) and $3.52(4) \ \mu_B$ for Mn1 sites and $4.22 \ \mu_B$ vs $4.13(3) \ \mu_B$ for Mn2 site. A good match between the calculated and experimental values indicates high accuracy of the selected calculation approach.

State	Energy above	Mn1 moment	Mn2 moment	Total moment
	AF1 (meV/Mn)	(µ _B)	(µ _B)	$(\mu_B/f.u.)$
FM	44.2	3.53	4.22	9.00
FI1	58.9	3.47	-4.23	-1.00
FI2	304.3	+3.50/-1.75	+/4.22	+1.00
AF1	0	+/-3.49	+/4.22	0
AF2	29.6	+/-3.49	+/4.22	0

Table 1. DFT-calculated energies of magnetic structures and magnetic moments on the Mn sites. Positive and negative magnetic moments are parallel and antiparallel to the b axis, respectively

Previous reports show that the energy difference and first excited state may serve to probe the transition temperature as it depends on energies of competing states, and can be calculated as three times the energy difference.^{59–61} Applying this approach to $CsMn_2F_6$ one can expect a transition temperature at around 100 K, which is around for times higher than the temperature of the first transition. Such discrepancy highlights the significance on the spin non-collinearity as well as the presence of two phase transitions. Interestingly, both ground AF1 and first excited AF2 states have the same arrangements of the moments on the Mn1 site, which orders magnetically first and therefore can be compared to the next excited state, FM. The energy difference between AF2 and FM levels, 14.6 eV/Mn atom, indicates a phase transition at 44 K, which is closer to the experimental value of 25 K, but deviates from it due to non-collinearity of the magnetic moments.

Calculations of the spin-density in the observed magnetic ground state (Figure S25) find a significant spatial extent of the Mn-based orbitals, with significant spin density associated with the F atoms, consistent with their exhibiting a slight (as high as $0.026 \ \mu_B$) spin moment in the calculations. This is an induced moment and derives from relatively close distances of F to the Mn atoms (as close as 1.86 Å).

Finally, we comment briefly on the experimental finding of substantial hard magnetic behavior at the the lowest temperatures, with coercive fields exceeding 5 T. Our main point here is that unlike in the case of ferromagnetic permanent magnet materials,^{62–64} small net magnetizations M_s generated by canted antiferromagnets allow generation of quite substantial anisotropy fields H_A and hence coercivities H_c , even if the effective first anisotropy constant K_1 is rather small. The reason is the inverse relationship between these quantities, which can be expressed in the equation $H_A = 2K_1/M_s$.⁶⁵ Quantitatively, if M_s is 0.008 µ_B per Mn, as in Figure 8, M_s is approximately 0.015 Tesla, and for a K₁ assumed to be just 0.1 MJ/m³ (nearly an order of magnitude smaller than in hcp cobalt, for example), one can expect an anisotropy field of nearly 17 Tesla, which could well allow a coercivity as high as 5 T. This suggests that canted antiferromagnets, even whose magnetic elements are only the usually magnetically soft 3d elements, can produce highly coercive behavior potentially of interest for spintronic and other applications.

Conclusions

We obtained a new charge ordered defect pyrochlore fluoride $CsMn^{2+}Mn^{3+}F_6$ via a hydrothermal route using CsCl to partially reduce Mn_2O_3 oxide in the presence of hydrofluoric acid. Crystallochemical

analysis of the new and reported fluoride defect pyrochlore structures showed that their formation is promoted by smaller M site cations. Moreover, the present structural and synthetic data indicate that the formation of the ordered orthorhombic defect pyrochlore structure is favored by a large size difference between the M^{2+} and M^{3+} cations, small A cation sizes, and a moderate, ca. 300–500 °C temperature regimes during crystallization.

Magnetic susceptibility data collected using a powder sample of the new phase CsMn₂F₆ exhibits a transition to a long-range spin ordered state at 24 K, which was confirmed by neutron diffraction data that also revealed a second transition between 10 and 14 K. The first transition to the *Pnm'a'* magnetic space group is caused by spin ordering on the Mn³⁺ sublattice and partial ordering on the Mn²⁺ sublattice, while a complete ordering of the latter results in further symmetry lowering to the *P*₂₁/*c* magnetic space group during the second transition. The canted antiferromagnetic ordering resulted in a 0.69 μ_B net magnetization in the unit cell along the *a* axis, which is consistent with a hysteresis loop and weak remnant magnetization observed in *M* vs. *H* plots. Using DFT calculations, we found two low energy antiferromagnetic states, which differ by spin arrangement on the Mn²⁺ sites. The ground state structure matches the experimentally observed spin arrangement and the excited state is 29.6 meV/Mn atom above the groundstate. Given the discrepancy between predicted and experimental values of transition temperatures (100 vs 24.1 K, respectively), we conclude that canting is crucial for CsMn₂F₆ magnetic structure.

A notable feature of this material is that the coercive field of its ferromagnetic component is highly dependent on the temperature and changes from nearly $H_c \approx 0$ T at around the transition temperature $T_{c,1} = 24.1$ K to $H_c > 5$ T below 10 K. Such effect results in an interesting phenomenon, in which the residual field in the SQUID instrument is sufficient to induce magnetization in the CsMn₂F₆ sample upon zero-field cooling, which then becomes an extremely hard magnet at 2 K, and the magnetization of the sample stays unaffected upon exposure to a 5 T field. Such an effect can be useful in a situation when there is a need to record data in a weak field and then to protect the data from fields that are orders of magnitude stronger than the recording field. Surprisingly, such behavior contrasts with purely antiferromagnetic AFe_2F_6 analogs, suggesting that distortion of the M^{2+} site plays an important role in the properties of charge ordered defect pyrochlore fluorides $AM^{2+}M^{3+}F_6$.

Experimental

Caution: Hydrofluoric acid is toxic and corrosive and must be handled with extreme caution and the appropriate protective gear! If contact with the liquid or vapor occurs, proper treatment procedures should

immediately be followed.

Caution: Chlorine gas is released during the reactions described below. Chlorine gas causes acute damage in the upper and lower respiratory tract and must be handled in a well-ventilated hood with appropriate protective gear.

General considerations. All reagents were obtained from commercial sources and were used without further purification. Phase purity was confirmed by powder X-ray diffraction using a Bruker D2 Phaser instrument (CuK α , $\lambda = 1.54184$ Å).

Synthesis. Phase pure (by PXRD) powder of $CsMn_2F_6$ was obtained hydrothermally by reacting 0.70 g of Mn_2O_3 and 1.50 g of CsCl in a 1:2 molar ratio with 3.00 mL of 49% HF. The starting materials were mixed in a PTFE liner and sealed in a steel autoclave. The autoclave was heated to 200 °C for 12 h in a convection oven and cooled to room temperature by turning off the oven. After finishing the reaction, the autoclave was opened, releasing greenish, chlorine smelling gas. The product was filtered on a paper filter using a Buchner funnel and washed on the filter with about 50 mL of acetone. $CsMn_2F_6$ is stable in air for several months, but rapidly dissolves in water.

Large $(0.1-0.5 \text{ mm}^3)$ dark brown single crystals of CsMn₂F₆ along with a pale pink MnF₂ side product were obtained by reacting 0.350 g of Mn₂O₃, 0.56 g CsCl, and 0.17 g of CsF with 4.00 mL of 49% HF and 2.67 mL of H₂O. The reagents were sealed in a PTFE-lined steel autoclave and heated to 200 °C at a rate of 0.1 °C/min, dwelled at this temperature for 12 h and cooled to room temperature by shutting off the convection oven. We found that the crystal size is very sensitive to the PTFE liner that was used, with generally larger crystal size for the liners with a coarse surface, which presumably promotes nucleation sites for the product. Quantitative elemental analysis was performed on product single crystals using a Tescan Vega-3 SEM instrument equipped with a Thermo EDS attachment (Table S6 and Figure S26).

Phase pure (by PXRD) powder of CsMnF₄ was obtained by dehydrating a mixture of CsMnF₄ and CsMnF₄·2H₂O under N₂ flow at 120 °C, well above the decomposition temperature of CsMnF₄·2H₂O, 69–70 °C, which was determined using thermal gravimetric analysis (TGA) (Figures S27 and S28). The purity of CsMnF₄ was confirmed by PXRD (Figure S28). The mixture of CsMnF₄ and CsMnF₄·2H₂O was obtained by reacting 0.70 g of Mn₂O₃ and 1.35 g of CsF in a 1:2 molar ratio with 3.00 mL of 49% HF. The starting materials were mixed in a PTFE liner and sealed in a steel autoclave. The autoclave was heated to 200 °C for 12 h in a convection oven and cooled to room temperature by turning off the oven. The product was filtered on a paper filter using a Buchner funnel and washed on the filter with about 50 mL of acetone.

Crystal structure. Single-crystal X-ray diffraction data were collected at 300(2) K on a Bruker D8 QUEST diffractometer equipped with an Incoatec IµS 3.0 microfocus radiation source (MoK α , λ = 0.71073 Å) and a PHOTON II area detector. The crystals were mounted on a microloop using immersion oil. The raw data reduction and absorption corrections were performed using SAINT and SADABS programs.^{66,67} Initial structure solutions were obtained with SHELXS-2017 using direct methods and Olex2 GUI.⁶⁸ Full-matrix least-square refinements against F² were performed with SHELXL software.⁶⁹The crystallographic data and results of the diffraction experiments are summarized in Tables S2 and S3. Crystal data for CsMn₂F₆: space group *Pnma* (no. 62), *a* = 7.3013(2) Å, *b* = 7.7141(2) Å, *c* = 10.1926(3) Å, *V* = 574.08(3) Å³, *Z* = 4, *T* = 301.46 K, µ(MoK α) = 10.681 mm⁻¹, *D_{calc}* = 4.128 g/cm³, 6789 reflections measured (6.624° ≤ 2 θ ≤ 54.966°), 708 unique (*R*_{int} = 0.0313, R_{sigma} = 0.0146) which were used in all calculations. The final *R*₁ was 0.0142 (I > 2 σ (I)) and *wR*₂ was 0.0289 (all data).

Magnetism. Magnetic property measurements were performed using a Quantum Design MPMS 3 SQUID magnetometer. For DC measurements, 32.71 mg of thoroughly ground sample was sealed in a plastic VSM powder sample holder and placed in a brass sample holder. For single crystal DC measurements (Figure S4), a multi-domain non-merohedrally twinned crystal of about $1 \times 1 \times 0.2$ mm³ size was glued to a quartz paddle in a random orientation. AC measurements were carried out using 57.27 mg of powder sample sealed in a gelatin capsule. The zero-field cooled and field cooled measurements were carried out in the SQUID DC mode in the 2–300 K temperature range and in an applied field of 0.1 T. AC measurements for magnetic transition detection were performed at applied field $H_{DC} = 0$ and $h_{ac} = 5$ Oe with a frequency *f* of 5, 7.57, 15, 50, 100, 150, and 500 Hz. MvH plots were measured at applied field from –5 to 5 T at temperatures of 2, 5, 10, 15, 17.5, 20, 23.5, 24.5, and 25 K. Before collecting MvH data, the sample was heated to 300 K, and dwelled at this temperature for 10 minutes to ensure removal of remnant magnetization. For magnetic moment calculation, the DC magnetic susceptibility was corrected for sample shape and radial offset.⁷⁰

To calculate the magnetic moment of $CsMn_2F_6$, several magnetic susceptibility vs. temperature datasets from different reaction batches were collected, yielding consistent results with a magnetic moment of around 8.0 μ_B , which is higher than the theoretical value of 7.68 μ_B that was calculated based on spin only moments of 5.92 and 4.90 μ_B for Mn²⁺ and Mn³⁺. For comparison, the magnetic susceptibility data for anhydrous CsMnF₄ were collected, resulting in the moment of 4.96 μ_B (Figure S29), which is consistent with the theoretical value of 4.90 μ_B for Mn³⁺. The latter result precludes an instrumental error, while reproducibility of the results and the phase purity of the studied material based on PXRD and neutron diffraction, indicates that the higher than expected magnetic moment of CsMn₂F₆ is likely an intrinsic property of the material. **Neutron diffraction.** Neutron powder diffraction measurements were conducted at Oak Ridge National Laboratory (ORNL). Preliminary data at 40 and 10 K were obtained using the POWGEN automatic sample changer *via* the mail-in program of the POWGEN diffractometer at the Spallation Neutron Source (SNS) (Figure S6) and temperature-dependent neutron diffraction data were collected on the instrument HB-2A at the High Flux Isotope Reactor (HFIR). For the experiment at HB-2A, a constant wavelength of 2.4067 Å was selected from the (113) reflection of a Ge monochromator. A PG filter was used to remove any scattering from higher order reflections. The sample was loaded in to a vanadium can and neutron diffraction data were collected at 1.5, 6, 10, 14–24 (with a 1 K step), 26, 40, and 290 K, well above the transition temperature (Figures S7–S24). The sample temperature was obtained using a top-loading ⁴He cryostat with a 3-sample changer stick. No apparent impurities were detected in the neutron data. The data were collected using 44 ³He detectors, with 12' collimation before each detector, in a Debye-Scherrer geometry that covered a 2θ range of 5–130 degrees. Data were collected in 0.05 degree steps. Figures S7–24 show neutron powder diffraction data along with the Rietveld refinements.

The magnetic structures were solved and refined using the Bilbao Crystallographic Server and FullProf Rietveld refinement suite.^{71,72} The nuclear structure and refinement variables were refined against the diffraction data at 26 K and the model was used to solve the magnetic structure at 18 K. The propagation vector \mathbf{k} for the magnetic structure was found to be (0,0,0) by searching additional peaks in the difference diffraction pattern. The crystal structure of $CsMn_2F_6$ along with the propagation vector were uploaded to the MAGNEXT program on the Bilbao server and several possible magnetic space groups were found.^{73,74} We found the best fitting magnetic structure in the *Pnm'a'* group by attempting trial structure refinements in different possible magnetic space groups. The obtained magnetic model was then refined against the other datasets at the temperatures between 24 and 10 K. Refinement of the same Pnm'a' model against the 1.5 K data revealed large residual peaks, indicating the presence of a second magnetic transition at lower temperatures. The symmetry of the low temperature magnetic phase was determined by removing symmetry elements from the *Pnm'a*' supergroup using MAGNEXT and refining the model against the 1.5 K data. The only reasonable solution was obtained using the $P2_1/n11$ space group, resulting in a good model with reasonable magnetic moments on the Mn atoms. The original cell can be transformed to the standard $P2_1/c$ setting by the (0 0 -1 1 0 0 0 -1 1) transformation matrix. The magnetic model was used for refinement against the 6 K data.

First-principle calculations were carried out using the all-electron code WIEN2K,^{75,76} working within the generalized gradient approximation (GGA),⁷⁷ as well as calculations using its extension, the GGA+U approach, in which an additional potential is applied to the relevant atomic orbitals. All calculations were performed using the experimental structure as a starting model. Ordinarily our approach is to employ the

experimental lattice parameters and then relax non-symmetry-dictated internal coordinates within an assumed ferromagnetic configuration, to account for potential magnetoelastic effects.^{78–80} However, this procedure produced a structure far from the experimental one, including in particular Mn1-F distances as much as 0.1 A larger than experiment, and in addition tended to equalize the moments on the two Mn sites, again in contradiction to the experimental data. Structures optimized within antiferromagnetic arrangements yielded only slight structural improvement. This departure of optimized structures from experimental ones is reminiscent of the situation with the iron-based superconductors,⁸¹ where relaxing the As height can yield a structure far from experiment. Here the physics is of large Mn local moments coupling strongly to the lattice, and it is likely that the non-collinearity (not studied here) plays a significant role in the structure. For this reason all calculations were performed in the experimental structure. Muffin-tin radii of 1.59, 1.76 and 2.50 Bohr were used, respectively for the F, Mn and Cs spheres, and an RK_{max} value (the product of the smallest radius and the largest plane-wave vector) of 7.0 was used; spin-orbit coupling was not employed. We report here results from the GGA+U approach with a U value of 5.25 eV applied to the Mn d orbitals.⁸² Given the highly electronegative nature of fluorine a correlated approach is likely more accurate than the straight GGA, and in fact yields moment magnitudes in better agreement with the neutron results.

Supporting Information

Crystalographic data, magnetic moments, a list of tolerance factors for $AM^{2+}M^{3+}F_6$ and $AA'M_2^{+2}F_7$ compounds, DTA and TGA results, high temperature PXRD, neutron diffraction data and fitting, EDS results.

Accession Codes

CCDC 2079680 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via <u>www.ccdc.cam.ac.uk/data_request/cif</u>, or by emailing <u>data_request@ccdc.cam.ac.uk</u>, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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