

$Mn_2(Fe_{0.8}Mo_{0.2})MoO_6$: A Double Perovskite with Multiple Transition Metal Sublattice Magnetic Effects

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Supporting Information

ABSTRACT: Transition-metal-only perovskite oxides can introduce additional magnetic functionality with robust magnetoelectric properties but are rare. In this work we prepared a new transition-metal-only perovskite $Mn_2(Fe_{0.8}Mo_{0.2})MoO_6$ at high pressure and temperature. Uniquely, $Mn_2(Fe_{0.8}Mo_{0.2})MoO_6$ was discovered as a line phase upon composition modulation that was motivated from the above-room-temperature multiferroic Mn₂FeMoO₆ corundum phase. It exhibits ferrimagnetic Fe–Mo sublattice ($T_{\rm C}$ = 194 K) and Mn sublattice antiferromagnetic $(T_{\rm m} \sim 45 \text{ K})$ transitions. Below $T_{\rm m}$ the two sublattice orderings are coupled and give rise to canted components in both. A first-order field-



induced transition is also observed below 45 K. Mn₂(Fe_{0.8}Mo_{0.2})MoO₆ is a Mott variable range hopping semiconductor. These findings for the first time show that either an exotic perovskite or a corundum phase can be achieved by composition modulation besides the pressure effect.

1. INTRODUCTION

Exotic perovskite oxides with transition-metal ions at both Aand B-sites are expected to introduce additional magnetic functionality with possible robust magneto-electric properties.^{1–10} The A-site ordered $AA'_{3}B_{4}O_{12}$ quadruple perovskites usually have transition-metal ions in 3/4 of the A-sites (A') as in $LaCu_3Fe_4O_{12}$ and $CaCu_3Fe_2Re_2O_{12}$, which show aboveroom-temperature reversible metal-insulator transition and spin-polarized half-metallic state, respectively.^{3,4} Although $A_2BB'O_6$ -type double perovskites Mn_2BSbO_6 (B = Cr, Fe) possess Mn^{2+} (high-spin d⁵) at the A-sites, they are less promising (antiferromagnetic (AFM) order ~ 55 K) compared with the above-mentioned quadruple perovskites,^{11,12} likely due to the nonmagnetic B'-site Sb⁵⁺. Thus, the design and synthesis of transition-metal-only perovskites with enhanced magnetic interactions are desirable. To the best of our knowledge, so far

only six transition-metal-only perovskite oxides have been prepared, including MnVO₃, ACu₃V₄O₁₂ (A = Mn, Cu), ζ - Mn_2O_3 , and Mn_2BReO_6 (B = Mn, Fe). Metallic MnVO₃ is the only ABO₃ simple perovskite.¹ Quadruple perovskites $ACu_3V_4O_{12}$ (A = Mn, Cu) demonstrate interesting cationic rattling of the 12-coordinated Mn²⁺ and Cu²⁺ at the A-sites.^{13,14} The ζ -Mn₂O₃ in a distorted quadruple perovskite structure $(Mn^{2+}(Mn^{3+})_3(Mn^{3.25+})_4O_{12})$ is a semiconductor with direct narrow bandgap and switchable p-n electrical conduction.^{5,6} $Mn_2FeReO_6^{-7,8}$ and $Mn_2MnReO_6^{-9,10}$ are the only two $A_2BB'O_6$ double perovskites: Mn₂FeReO₆ is a ferrimagnetic (FiM) halfmetal (Curie temperature $T_{\rm C} \sim 520$ K) that shows giant

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positive magnetoresistance of 220% at 5 K and 8 T,^{7,8} while Mn_2MnReO_6 is a one-dimensional variable-range hopping semiconductor with AFM transition at ~ 110 K.^{9,10}

In this work, we prepared a new transition-metal-only double perovskite $Mn_2Fe_{0.8}Mo_{1.2}O_6$ (MFMO) under high pressure via the B-site composition modulation motivated by the recent synthesis at high pressure (8 GPa) of Mn_2FeMoO_6 and the findings of exotic properties of this new phase.^{15,16}

2. EXPERIMENTAL DETAILS

Synthesis. Polycrystalline $Mn_2Fe_{0.8}Mo_{1.2}O_6$ was prepared from a stoichiometric mixture of MnO (99.99%, Alfa Aesar), Fe (99.998%, Alfa Aesar), Fe₂O₃ (99.998%, Alfa Aesar), and MoO₃ (99.8%, Alfa Aesar) at 1623 K for 1 h under 8 GPa in a Walker-type Multi Anvil Press and then quenched to room temperature by turning off the voltage supply to the resistance furnace as reported in our previous work.^{7,10,16–22} The pressure is maintained during the temperature quenching and then decompressed slowly in 8–12 h.

Powder X-ray and Neutron Diffraction. Room-temperature powder X-ray diffraction (PXD) data were collected on a Bruker D8 Advance Diffractometer (Cu $K_{\alpha\nu} \lambda = 1.5418$ Å) with a SOL-X solid state detector. Room-temperature synchrotron powder X-ray diffraction (SPXD) data were recorded on beamline X-16C ($\lambda = 0.70019$ Å) at the Brookhaven National Synchrotron Light Source (NSLS) in the U.S. Powder neutron diffraction (PND) data were collected on an about 500 mg polycrystalline sample at 10, 60, and 300 K on the POWGEN instrument at the Spallation Neutron Source at Oak Ridge National Laboratory (U.S.). The magnetic structure symmetry analysis was performed with ISODISTORT software.²³ The EXPGUI interface of the GSAS program²⁴ was used for Rietveld refinement²⁵ of the atomic and magnetic spin structures. Representations of the crystal and spin structures were made with VESTA-3.²⁶

X-ray Absorption Near-Edge Spectroscopy. Mn– and Fe–K X-ray absorption near-edge spectroscopy (XANES) data were collected in both the transmission and fluorescence mode with simultaneous standards. The Mo XANES was collected in fluorescence mode in a He-atmosphere-chamber with standards run in temporal proximity. All of the spectra were fit to linear pre- and postedge backgrounds and normalized to the unity absorption step across the edge.^{7,27–33} All of the XANES was performed on beamline X-19A at the Brookhaven NSLS with a Si-111 double-crystal monochromator.

Magnetism and Magnetotransport. Magnetization measurements were carried out with a Quantum Design superconducting quantum interference device (SQUID) magnetometer. The magnetic susceptibility (χ) was measured in zero field cooled (ZFC) and field cooled (FC) conditions under 0.1, 1, and 5 T applied magnetic field (H) for temperatures ranging from 5 to 400 K. Isothermal magnetization curves were obtained at 5, 10, 100, 170, 200, 220, 300, and 400 K under an applied magnetic field varying from -5 to 5 T. The magnetotransport properties were measured on a pellet sample with the standard four-probe technique in a physical property measurement system (PPMS) from Quantum Design at 0 and 7 T, respectively. The magnetic-field-dependent conductivity variation was measured between -14 and 14 T at 5 K.

3. RESULTS AND DISCUSSION

3.1. Line Phase and Crystal Structure. The Ni₃TeO₆type multiferroic Mn₂FeMoO₆ is a highly spin-polarized aboveroom-temperature FiM ($T_{\rm C} \sim 340$ K) semiconductor; it shows an unexpected low-temperature (150–300 °C) cationic rearrangement at ambient pressure, accompanied by a Ni₃TeO₆-to-ordered-ilmenite structural transition and dramatic physical property changes.^{15,16} These findings inspired further composition modulations in Mn₂FeMoO₆ for optimal physical properties as in Sr₂FeMoO₆ and Sr₂Fe_{1+x}Mo_{1-x}O₆.^{32,34} However, attempts to prepare Mn₂Fe_{1+x}Mo_{1-x}O₆ (-0.3 $\leq x$ \leq 0.33) solid solutions up to 8 GPa and 1623 K were unsuccessful. PXD data show that only a double perovskiterelated line phase MFMO can be achieved at x = -0.2 besides the Ni₃TeO₆-type Mn₂FeMoO₆ (x = 0) (Figure S1 in Supporting Information (SI)). Detailed analysis of the SPXD and PND data of MFMO collected at 300 K indicates a distorted monoclinic $P2_1/n$ phase (a = 5.222(1) Å, b =5.382(1) Å, c = 7.605(1) Å, $\beta = 90.04(1)^\circ$, V = 213.72(1) Å³), which yielded a good fit ($R_p/R_{wp} = 6.81/3.19\%$) in a combined SPXD and PND refinement with the structural model of Mn₂MReO₆ (M = Mn, Fe)⁷⁻¹⁰ and Mn₂MSbO₆ (M = Sc, Fe, Cr) perovskites^{11,12,35} as shown in Figures 1A and S2. The final



Figure 1. (A) SPXD plots of MFMO from combined refinements in monoclinic $P2_1/n$ structure at room temperature. The asterisks (*) denote the peaks of diamond internal standard. The inset presents the crystal structure of MFMO showing rock-salt ordering of (Fe_{0.73(1)}Mo_{0.27(1)})O₆ and MoO₆ octahedral connections. Mn atoms, blue spheres; O, green spheres; (Fe_{0.73(1)}Mo_{0.27(1)})O₆ octahedra, brown; MoO₆ octahedra, yellow. (B) A superimposed comparison of the Mo-L₃ edge of MFMO to those of the d^0/Mo^{6+} -SrFe_{0.75}Mo_{0.25}O_{3- δ}, d^1/Mo^{5+} -Sr₂Cr_{1.2}Mo_{0.8}O_{5.8}, and ~ 4 d^2 -Mo⁴⁺-Sm₂Mo₂O₇ standards, indicating Mo^{4+/5+} in MFMO. The double-headed arrows denote the t_{2g}^- and e_g -related A and B features respectively based upon the octahedral crystal electric field splitting.

crystallographic data are shown in Table S1. There are no oxygen defects observed during the refinements within the standard uncertainties. The A- and B'- sites are fully occupied by Mn and Mo, respectively, while the B-site is mixed with 73(1)% Fe and 27(1)% Mo, giving the structural formula of $Mn_2^{A}(Fe_{0.73(1)}Mo_{0.27(1)})^{B}Mo^{B'}O_{6'}$ in reasonable agreement with the starting stoichiometry.

The crystal structure of MFMO is presented in the inset of Figure 1A, where Mn is in 8-fold coordination, $(Fe_{0.73(1)}Mo_{0.27(1)})O_6$, and MoO_6 octahedra are in a corner-shared rock-salt arrangement. The average $\langle Mn-O \rangle$ distance of 2.388 Å is in line with those in isostructural $Mn^{2+}_2BB'O_6$ (B = Sc, Cr, Mn, Fe; B' = Sb, Re) between 2.379 and 2.405 Å, giving

bond valence sums (BVS) of 1.93 as listed in Table S2.^{10–12,35} The $\langle (Fe_{0.73(1)}Mo_{0.27(1)}-O \rangle$ of 2.021 Å is nearly identical with that of the $\langle Fe-O \rangle$ in $Mn_2Fe^{3+}Re^{5+}O_6$ (2.022 Å)^{7,8} and $Mn_2Fe^{3+}Sb^{5+}O_6$ (2.017 Å),¹¹ which suggests Fe^{3+}/Mo^{4+} at the Fe/Mo site given their almost identical ionic radius values (0.645 and 0.65 Å for Fe^{3+} and Mo^{4+} , respectively, in octahedral coordination).³⁶ This gives an average charge of 3.27 at the Bsite, comparable with the BVS value of 3.18. Assuming Mn²⁺ at the A-site and $(Fe^{3+}_{0.73(1)}Mo^{4+}_{0.27(1)})$ at the B-site, the B'-site Mo therefore has an average charge of 4.73 (compared to its BVS of 4.58) from a mix of 73% Mo^{5+} and 27% Mo^{4+} according to charge balance. This gives overall 2.0-Mn²⁺, 0.73-Fe³⁺, 0.54- $\begin{array}{l} Mo^{4+}, \text{ and } 0.73\text{-}Mo^{5+}, \text{ namely, } Mn^{2+A}(Fe^{3+}_{0.73(1)}Mo^{4+}_{0.27(1)})^B \\ (Mo^{4+}_{0.27(1)}Mo^{5+}_{0.73(1)})^{B'}O_6, \text{ which has been corroborated by} \end{array}$ the XANES results shown in Figures 1B and S3-4. The low energy chemical shift and pre-edge features further clearly indicate Mn²⁺ and Fe³⁺ valence states in the Mn- and Fe-K edge XANES in Figures S3 and S4. The Mo-L₃ edge XANES in Figure S5 illustrates both the systematic increase of the A $(t_{2g}$ -related) feature intensity, relative to the B $(e_g$ -related) features, and the increasing chemical shift of the combined A-B feature in the configuration sequence from $\sim 4d^2/Mo^{\sim 4+}$ to $4d^0/Mo^{6+}$. The Mo-L₃ spectra in Figure 1B support the mixed Mo^{4+/5+} valence by both the relative A-feature intensity and A-B feature chemical shift of the MFMO spectrum lying intermediate between those of the $\sim d^{1}/Mo^{5}$ $(Sr_2Cr_{1,2}Mo_{0,8}O_{5,8})$ and $\sim 4d^2/Mo^{4+}$ $(Sm_2Mo_2O_7)$ standard spectra.

3.2. Magnetic Properties at Low Field. Figures 2A and 2B, respectively, present constant field magnetization, M(T),



Figure 2. (A) Temperature-dependent magnetization (*M*) evolution of MFMO between 5 and 400 K at 0.1, 1, and 5 T, respectively, in both ZFC and FC modes. (B) Isothermal M(H) loops of MFMO at 5, 10, 100, 220, and 400 K between 5 and -5 T. Note the hysteretic field-induced transitions upon moving up ("u") and down ("d") in field in the T = 5 and 10 K loops.

and isothermal magnetization, M(H), measurements on MFMO. For clarity/simplicity the low-field and low-temperature magnetic measurement results will be discussed first to motivate the neutron scattering spin structure measurements. The discussion of the more complex higher-field magnetic behavior will then be returned once the H = 0 spin structure is solidly established.

In Figure 2A, the temperature-dependent ZFC and FC magnetizations of MFMO, for H = 0.1 T, clearly exhibit at least two transitions: an apparently FiM transition at $T_{\rm C} = 194$ K and a lower temperature AF-type transition at $T_{\rm m} \sim 45 \pm 3$ K. The nonlinear FiM-like M(H) results at T = 100 K and the finite field-induced transition in the M(H) results at T = 5 and 10 K (Figure 2B) support the two-transition-temperature hypothesis. From the above magnetic results the temperatures T = 10 K (below $T_{\rm m}$) and T = 60 K ($T_{\rm m} < T < T_{\rm C}$) were chosen for H = 0 PND spin structure determination which is discussed in the next section.

3.3. Spin Structures. The magnetic spin structures of MFMO at 10 and 60 K were determined by PND data refinements (Figures 3A, 3B, and S2C, respectively). The



Figure 3. Rietveld refinement plots of the PND data for MFMO of (A) 1.333 Å bank at 10 K and (B) 3.731 Å bank at 60 K. Tick marks show the position of allowed nuclear and magnetic reflections of the perovskite phase.

MFMO magnetic reflections were indexed with the $k = (0\ 0\ 0)$ propagation vector and magnetic space group (MSG) $P2_1/n$. Other $k = (0\ 0\ 0)$ MSGs $P2'_1/n$, $P2'_1/n'$, and $P2_1/n'$ were discounted as they did not permit all the observed magnetic reflections. The magnetic structures of MFMO at 10 and 60 K are shown in Figure 4. At 60 K antiparallel order of the B and B' site moments (going forward they will also be referred to as Fe³⁺ and Mo^{4.7+} sites) observed is consistent with spin



Figure 4. Magnetic structure of MFMO at (A) 10 K and (B) 60 K. Brown = Fe/Mo at B-site, yellow = Mo at B'-site, blue = Mn at A-site.

structures reported for other FiM double perovskites (Figure 4B). The net ferromagnetic component is 1.4 $\mu_{\rm B}/{\rm f.u.}$ Above $T_{\rm m}$ \sim 45 K (the magnetic reorientation transition temperature) the A-site Mn²⁺ spins have not yet ordered as they are partially frustrated by the dominant FiM order of the B/B' sites.⁸ At 10 K the refined spin structure (Figure 4A) has Mn²⁺ moments antiferromagnetically ordered in the m_{xz} plane, and the Fe³⁺ moments (which above $T_{\rm m}$ were oriented perpendicular along m_{ν}) now exhibit a degree of canting with new alternating $\pm m_{x}$ component (AFM type, $\theta_{\text{FeFe}} = 20(2)^{\circ}$). Canting of the B/B' site moments below T_m is also observed for the other closely related Mn²⁺ A-site FiM double perovskites Mn₂FeReO₆ ($\theta_{\text{FeFe}} = 15^{\circ}$) and Mn₂MnReO₆ ($\theta_{\text{BsiteBsite}} = 15^{\circ}$).^{7–10} In our PND refinements, the precise magnitude and direction of canted Mo^{4.7+} moments could not be fully determined at 10 K, and therefore only the m_{ν} ordered spin component was refined. In view of the canting of both the FiM-Fe/Mo and AFM-Mn sublattices in the low-temperature PND-determined phase, this low-field phase will be referred to as the cFiM/AFM in subsequent discussions.

3.4. Magnetic Properties at Finite Field. The neutron diffraction (H = 0) spin structure determinations now provide a sound basis for discussion of the higher-field magnetic results. A tentative H-T magnetic phase diagram is displayed in the SI because it must be regarded only as tentative. Starting from high temperature, the onset of the FiM phase is manifested by the inflection points in the M(T) curves as indicated by the dashed "i" lines in Figure 2A. These i-features are emphasized by the negative peaks in the dM/dT curves in Figure S6A. Here it should be emphasized that model calculations show that the FiM phase onset is a true second-order phase transition only in the H = 0 limit and that the onset is continuous, albeit nonlinear, in the presence of a finite symmetry-breaking magnetic field (see SI for more details).^{37,38} The nonlinear Sshaped M(H) loops at T = 100 K (in Figure 2B) and that at T = 170 and 200 K (Figure S6B) are consistent with the FiM transition at $T_{\rm C}$.

The spin structure determination below $T_{\rm m}$ clearly indicated that this transition involved the AFM-type ordering of the Mn sites into a canted-AFM (cAFM) order that is coupled to a reorientation of the Fe–Mo sublattice order into a canted FiM state (cFiM). Moreover, the fact that the Fe-site-canted component has an alternating AFM-like character (see Spin Structures section) further underscores the coupling of the two sublattices.

In dual magnetic site ABO₃ perovskites, magnetic ordering of the B-site at a higher temperature, followed by A-site magnetic ordering at a lower temperature, is not uncommon.^{7,8,39–42} Moreover, a complex A/B site magnetic coupling, involving moment competition/frustration/canting, is also a leitmotif in such materials. Understandably, the application of a magnetic field leads to a still more complex interplay of the sites and moment orientations in such multiple-magnetic-sublattice systems. The line "ii" in Figure 2A indicates the local maximum in the M(T) curves at higher magnetic fields. This phenomenon is inferred to reflect the presence of Mn-site AFM correlation coupling to the Fe/Mo site dominated by net magnetization. Indeed, the maximum observed magnetization at H = 5 T occurs near T = 100 K in both the M(H) and M(T) results.

As previously noted, the onset of the coupled cFiM/AFMtype order at $T_{\rm m}$ is marked by a sharp structure in the H = 0.1T M(T) curves in Figure 2A. Broader downturn features occur in the magnetization curves for $H \ge 1$ T, at temperatures somewhat less than $T_{\rm m}$ (see the line "iii" in Figures 2A and S8), and are inferred to indicate the onset of a high-field phase involving cFiM and AFM order and are denoted as HF-cFiM/ AFM hereafter (see Figure S8). Field-induced transitions into this HF-cFiM/AFM are discussed below.

For $T < T_{\rm m}$, the presence of a first-order, field-induced transition from the low-H cFiM/AFM phase into a higher magnetization phase (HF-cFiM/AFM) is emphasized by the 5 and 10 K M(H) loops in Figure 2B and the M(H) curves in Figure S6B. The increasing field metastability limits for the cFiM/AFM to HF-cFiM/AFM transition are identified by "u" in M(H) curves in Figures 2B and S6B. The decreasing field metastability limits for the HF-cFiM/AFM to cFiM/AFM transition are identified by "d" in M(H) curves in Figures 2B and S6B. M(H) and dM/dH(H) curves at 5 K in Figure S7 illustrate the hysteretic field-induced transition around ± 1.7 T. It is worth noting that the field-dependent magnetoresistance (Figure 5B) also shows a sharp peak that agrees well with the increasing field cFiM/AFM to HF-cFiM/AFM transition as illustrated in Figure S7. The HF-cFiM/AFM phase extends to the highest field studied at temperatures below \sim 30–45 K. The line labeled "iii" in Figure 2B and in the schematic phase diagram in Figure S8 indicates an estimate for the hightemperature border of the HF-cFiM/AFM phase.

Returning to the M(T) curves in Figure 2A the sharp disparities between the FC and ZFC curves, for both the H =0.1 and 1 T (labeled by "iv" in Figure 2A), should be noted. The temperatures and fields for the step-like changes in these two ZFC M(T) curves, collected by heating from the lowest temperatures after ZFC (i.e., in the cFiM/AFM phase), are included in the phase diagram in Figure S8, and importantly they coincide closely with the magnetic-field-induced metastability limit for the cFiM/AFM to HF-cFiM/AFM transition. This motivates the association of the "iv" features, in the two lowest-field ZFC M(T) curves, with crossing in temperature a metastability limit for the cFiM/AFM to HF-cFiM/AFM transition upon heating.

It should be emphasized that field-induced transition states discussed here will differ depending on the detailed specific crystal-axis orientation of the applied field so that the polycrystalline results in this paper represent an average over such details. Transitions and phase mixtures will also differ for differing crystal axis magnetic field orientations and for differing field/temperature experimental H-T paths. Accordingly, extensive single-crystal studies would be required for a further



Figure 5. (A) Temperature (*T*)-dependent resistivity (ρ) plots of MFMO at 0 and 7 T, respectively. Inset shows the linear fit to the plot of ln ρ versus $T^{-1/4}$ above 225 K, indicating Mott's VRH conduction mechanism. (B) Isothermal magnetoresistance at T = 5 K with field paths identified in the sequence: ZFC-prepared, H = 0 to +15 T (path 1, dotted red); +15 to -15 T (path 2, dashed green); and -15 to 15 T (path 3, solid blue). Note that the peaks of the magnetoresistance correspond closely in field to the field-induced transitions in the T = 5 K M(H) and dM/dH results shown in Figure S7.

clarification of the low-temperature H-T phase diagram of this material.

3.5. Magnetotransport and Magnetoresistance. Resistivity (ρ) and magnetoresistance measurements on a polycrystalline MFMO pellet are shown in Figure 5. The resistivity is about 1.15 × 10³ and 18.9 Ω ·mm without applied magnetic field at 5 and 300 K, respectively, which are almost identical with the values measured at 7 T. The overall resistivity shows characteristic semiconducting behavior between 5 and 300 K (Figure 5A). The temperature dependence of ρ was determined by trial fitting the relations $(1/T)^p$, which follows the relation with p = 1/4 between 40 and 260 K and indicates a Mott variable range hopping mechanism $\rho = \rho_0 \exp(T_0/T)^{1/4}$, where $T_0 = 281.24$ K and $\rho_0 = 0.933$ Ω ·mm.

The magnetic-field-dependent magnetoresistance of MFMO at 5 K gives a peak value of magnetoresistance ratio of \sim 5% (Figure 5B). Comparing Figure 5B to the dM/dH field-cycling curves in Figure S7 indicates that the sharp peaks in the magnetoresistance, at $H = \pm 1.7$ T, occur at the peeks of the dM/dH curves that determine the field-induced transition. Thus, the maximal magnetoresistance in this regime is directly coupled to the maximal phase inhomogeneities that occur at the field-induced cFiM/AFM to HF-cFiM/AFM transition. The reverse HF-cFiM/AFM to cFiM/AFM transition shows up as a much smaller shoulder on the magnetoresistance hysteresis loop. This behavior is similar to what was observed in halfmetallic Sr₂FeReO₆ where the peaks in dM/dH and the magnetoresistance occur at the same fields.⁴³ Sr₂FeReO₆ is however simply a FiM material with no moment on the perovskite A-site.

Insight into the wider temperature range magnetoresistance can be gleaned by calculating the resistivity ratio $\rho_0/\rho_{\rm H}$ (H = 7 T) using the data in Figure 5A. The results for $\rho_0/\rho_{\rm H}$ are shown in Figure S9. The utility of resistivity ratio lies in the assumption that the very high-field resistivity ($\rho_{\rm H}$) represents a more homogeneous, less resistive state and that the magnetoresistance is dominated by the field suppression of inhomogeneities in the H = 0 resistivity (ρ_0).⁴⁴ In Figure S9 the field-induced value of $\rho_{0}/\rho_{\rm H}$ first increases with decreasing temperature, reaching a maximum near the FiM $T_{\rm C}$. This is presumably due to fluctuation-induced inhomogeneities increasing in magnitude upon approaching the ordering temperature from above. Below $T_{\rm C}$, as the magnetic domains grow, $\rho_{0}/\rho_{\rm H}$ decreases. In an ~ 40 K range around $T_{\rm m}$ the $\rho_{0}/\rho_{\rm H}$ becomes less than 1.0 indicating that the magnetic field increases, rather than decreases, inhomogeneities. Finally, at low temperatures $\rho_{0/}\rho_{\rm H}$ becomes greater than 1.0 again and is dominated by transiting the field-induced transition and entering the more homogeneous high-field phase. The value of $\rho_{0}/\rho_{\rm H}$ from the magnetoresistance curves in Figure 5B, included in Figure S9, supports this analysis. Interpretation of the degree of interdomain/intergrain tunneling versus intergrain affects in the magnetoresistance would be speculative at this stage.

4. CONCLUSIONS

In conclusion, we have prepared the third transition-metal-only double perovskite $Mn_2(Fe_{0.8}Mo_{0.2})MoO_6$ at high pressure via composition modulation. $Mn_2(Fe_{0.8}Mo_{0.2})MoO_6$ is isostructural with Mn_2BReO_6 (B = Mn, Fe). The Fe–Mo sublattice of this compound exhibits a ferrimagnetic transition with $T_{\rm C}$ = 194 K. The Mn sublattice orders antiferromagnetically at $T_{\rm m} \sim 45$ K. The concomitant canting of the coupled ferrimagnetic (Fe-Mo) and antiferromagnetic (Mn) sublattices is underscored by dubbing T_m as a spin "reorientation" transition. The canted coupling of the two sublattices is still more complex in the presence of magnetic field as emphasized by hysteretic firstorder, field-induced phase transition at low temperatures. $Mn_2(Fe_{0.8}Mo_{0.2})MoO_6$ is a Mott variable range hopping semiconductor and exhibits a magnetoresistance coupled to the nonlinear magnetization at low temperature. These transition-metal-rich exotic perovskites can be stabilized at higher pressure over competition with the corundum phases^{11,35,45} as observed in the corundum-perovskite transitions in Mn_2BSbO_6 (B = Sc, Fe).^{11,35} In contrast, the high-pressure made perovskite phase can also transform into ilmenite structure upon heating at ambient pressure as reported in Mn₂CrSbO₆.¹² So far, only the perovskite phases have been obtained for Mn_2FeReO_6 and Mn_2MnReO_6 .⁷⁻¹⁰ This finding may be the consequence of the specific spin structures and possible spin-orbit coupling (Re) in Mn₂MnReO₆ and Mn₂FeReO₆ that stabilize a lower-energy perovskite relative to the corundum structure. In the work presented here, the presence of only two line phases, namely, Mn₂FeMoO₆ corundum and Mn₂(Fe_{0.8}Mo_{0.2})MoO₆ perovskite in $Mn_2Fe_{1+x}Mo_{1-x}O_{6}$, raises a new possibility. The Ni₃TeO₆type Mn₂FeMoO₆ is known to be stabilized by the spin structure,¹⁵ while the absence of solid solutions and the formation of a new perovskite line phase Mn₂(Fe_{0.8}Mo_{0.2})-MoO₆ could be a synergy of thermodynamics, crystal, and spin structure (mixed-valence $Mo^{4+/5+}$) based energetic effect. Further theoretical calculations are necessary to better understand these correlations and guide the design of new polar and magnetic corundum phases and transition-metal-only double perovskites, which are important for spintronic applications.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemma-ter.8b00250.

Line phase mining, Mn- and Fe-K edge and pre-edge XANES figures, detailed magnetism and magnetic phase diagram analysis (PDF)

Crystallographic information file for Mn2Fe.8-Mo1.2O6_300K (CIF)

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Notes

The authors declare no competing financial interest.

The supporting crystallographic information file may also be obtained from FIZ Karlsruhe, 76344 Eggenstein-Leopoldshafen, Germany (e-mail: crysdata@fiz-karlsruhe.de), on quoting the deposition number CSD-433446.

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