Magnetic Transitions in Exotic Perovskites Stabilized by Chemical and Physical Pressure

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ABSRACT

Exotic Perovskites significantly enrich materials in multiferroic and magnetoelectric applications. However, their design and synthesis are the challenge due to the mostly required extreme conditions at high pressure. Herein, we have presented the $Ca_{2-x}Mn_xMnTaO_6$ ($0 \le x \le 1$) 1.0) solid solutions stabilized by chemical pressure assisted with intermediate physical pressure up to 7 GPa. The incorporation of Mn²⁺ into the A-site neither drives any cationic ordering nor modifies the orthorhombic *Pbnm* structure, namely written as $(Ca_{1-x/2}Mn_{x/2})(Mn_{1/2}Ta_{1/2})O_3$ with disordered A and B site cationic arrangements. The increment of x is accompanied by a ferromagnetic to antiferromagnetic transition around x = 0.2, which is rooted in the super-exchange interactions between A-site Mn²⁺ and B-site Mn³⁺. Partial charge disproportionation of the B-site Mn^{3+} into Mn^{2+} and Mn^{4+} occurs for x above 0.8 samples as manifested by X-ray spectrum and magnetic behaviors. The coexistence of B-site Mn³⁺ (Jahn-Teller distortion ion) and B'-site Ta⁵⁺ (second-order Jahn-Teller distortion ion) could be energetically responsible for the absence of A-site columnar ordering as observed in other quadruple perovskites with half of the A-sites occupied by small transition-metal cations. These exceptional findings indicate that exotic perovskites can be successfully stabilized chemical and intermediate physical pressure, and the presence of John-teller distortion cations at the same lattice should be avoid to enable cationic ordering.

INTRODUCTION

Perovskite oxides have attracted great interest because of their enriched structural, magnetic and electronic properties.¹⁻⁴ The stability and crystal system of ABO₃ perovskites can be depicted by the octahedral factor $\mu = r_B/r_O$ and tolerance factor $t = (r_A + r_O)/\sqrt{2}(r_B + r_O)$ proposed by Goldschmidt (r_i stands for the ionic radius of ion i).^{5, 6} The radii of ions and tilting of octahedron have a significant impact on stabilizing perovskite structure in light of their chemical, octahedral, stretch, and tilt limits. 7-12 In contrast to conventional perovskite with large A-site cations, the exotic perovskite can adaptively incorporate small cations (especially transition-metal ions) into the A-site. $^{13-16}$ A-site columnar-ordered quadruple perovskites $A_2A'A''B_4O_{12}$ with 50% of the small A-sites (denoted as square-planar coordinated A' and tetrahedrally coordinated A'', respectively) was firstly discovered in $Ca_2Fe'Fe''Ti_4O_{12}$ (also known as $CaFeTi_2O_6$ with A' = A''= Fe)¹⁷ and then $Ca_2Mn'Mn''Ti_4O_{12}$ (known as $CaMnTi_2O_6$ with A' = A'' = Mn).¹⁸ Ca₂Fe'Fe"Ti₄O₁₂ crystallizes in centrosymmetric P4₂/nmc (No. 137), while Ca₂Mn'Mn"Ti₄O₁₂ adopts a polar P42mc (No. 105) structure in that Mn' displaces off from the Mn'O4 square-plane (Figure 1a) other than exactly stays in the oxygen plane as observed for Fe' in Fe'O₄ in Ca₂Fe'Fe"Ti₄O₁₂. Pock-salt *B*-sites ordering in *A*₂*A'A*"*B*₂*B'*₂O₁₂ (**Figure 1b**) provides higher compositional freedom as reported in the $P4_2/n$ (No. 86) type Ca₂Mn'Mn" B_2 Re₂O₁₂ (B = Mn, ²⁰ Fe^{20} Co^{21} Ni^{21}), $Ca_2Mn^ACu^A(Fe_2)^B(Re_2)^BO_{12}$, and $R_2Mn^2Mn^2Mn_2Sb_2O_{12}$ (R = La, Pr, Nd, Pr, NSm). 22 The A'O₄ (square-planar), A"O₄ (tetrahedral), BO₆ (octahedral), and B'O₆ (octahedral) sites in $A_2A'A''B_2B'_2O_{12}$ are highly adaptable, ²³ and can be occupied by the same element as recently observed in RMn_3O_6 (R = Y, Gd, Dy, Ho, Er, Tm; A' = A'' = B = B' = Mn), ²⁴⁻²⁶ which crystallize in *Pmmn* (No. 59, Figure 1c) and can be structurally written as $R^{3+}_{2}(Mn^{2+})^{A'}(Mn^{3+})^{A''}(Mn^{3+})_{2}^{B'}(Mn^{3.5+})_{2}^{B'}O_{12}$ with layered charge ordering over the *B*-sites. When

75% of the A-sites are occupied by small cations, $AA'_{3}B_{4}O_{12}$ -type quadruple perovskites can be formed with square-planar coordinated A'-site, such as $A' = Mn^{3+}$, Co^{2+} , Cu^{2+} , Pd^{2+} in $LaM{n_3}{V_4}{O_{12}},^{27}\ LaC{u_3}F{e_4}{O_{12}},^{28}\ CaC{o_3}{V_4}{O_{12}},^{29,\ 30}\ CaP{d_3}{Ti_4}{O_{12}},^{31,\ 32}\ respectively.\ Most\ known$ $AA'_{3}B_{4}O_{12}$ quadruple perovskites are in cubic Im-3 (No. 204) symmetry (Figure 1d) unless the B-site charge-ordering induced distortion occurs as in rhombohedral manganites A²⁺Mn₃Mn₄O₁₂ (R-3, No. 148, A = Ca, Cd, Pb, Pr, Sr, Figure 1e). 33-35 1322-type B-site ordering of quadruple perovskites AA'₃B₂B'₂O₁₂ leads to a symmetry evolution from Im-3 to Pn-3 (No. 201, Figure 1f) such as reported in CaCu₃Fe₂Re₂O₁₂, ³⁶ CaCu₃Fe₂Nb₂O₁₂, ³⁷ and NaCu₃Fe₂Os₂O₁₂, ³⁸ In A³⁺Mn₇O₁₂ (A = La, Pr, Nd) series the trivalent A-site ion drives more Jahn-Teller distorted Mn³⁺ component on the B-site, which, together with charge ordering, results in monoclinic I2/m (No. 12) structure (**Figure 1g**) as in $Pr^{3+}Mn^{3+}{}_{3}(Mn^{2.99+})^{8}{}_{2}(Mn^{3.01+})^{8}{}_{2}O_{12}$. The spatial effect of $6s^{2}$ -lone pair electrons of Bi³⁺ in BiMn₇O₁₂ further leads to complex temperature-dependent symmetry evolution of Im-3 (above 608 K) - I2/m (460-608 K) - Im (290-460 K) - P1 (below 290 K) upon cooling. 41 Full occupation of the A-site with small cations usually draws pressure-dependent polymorph competition, and mostly the perovskite phases need to be stabilized from higher pressure synthesis. $^{16, 42-44}$ GdFeO₃-based *Pnma* (No. 62, **Figure 1h**) and $P2_1/n$ (No. 14) structures (Figure 1i) have been successively discovered in ABO₃-type (such as MnVO₃ and ScCrO₃)⁴⁵⁻⁴⁷ and $A_2BB'O_6$ -type (such as Mn₂BSbO₆ (B = Fe, ⁴³ Sc, ⁴⁸V⁴⁹), Mn₂BReO₆ (B = Mn, ^{14, 50} Fe, ^{15, 51} Co^{52}), and $\mathrm{Mn_2(Fe_{0.8}Mo_{0.2})MoO_6)^{13}}$) perovskites, respectively. $AA'_3B_4O_{12}$ -type quadruple perovskites can also be prepared in this catalog if the pressure is high enough, such as $ACu_3V_4O_{12}$ (Im-3, A = Cu, Mn), and ζ -Mn₂O₃ (structurally written as Mn²⁺(Mn³⁺)₃(Mn^{3.25+})₄O₁₂, Figure 1j, P-1 (No. 2)). 53-55 Post-perovskite structural compounds (Figure 1k, Cmcm, No. 63) such as δ -Mn₂O₃ and MgSiO₃ can exist at extremely high pressure but unquenchable to ambient

conditions. 56, 57

Partial or full occupation of small cations on the A-site in exotic perovskite not only depicts an enriched image of structural chemistry, but also provides a continuous impetus to search for emergent physical properties, in that the small A-site cations, especially transition-metal ions, enhance the quantum degree of freedom such as lattice, spin, charge, and orbital. The reduced t can cause structural distortion and thus large spontaneous electrical polarization (P_S) in noncentrosymmetric materials, 58, 59 while the transition-metal-rich lattices lead to robust magnetic interplays and thus magnetoelectric effect. ^{15, 28, 51} For examples, CaMnTi₂O₆ is the only lone-pair-electron free switchable ferroelectric double perovskite ($P_S \sim 24 \, \mu \text{C/cm}^2$), ¹⁸ demonstrating potential piezoelectric ferroelectric-photovoltaic applications; and $\text{Ca}_2\text{Mn}^A\text{'Cu}^A\text{''}(\text{Fe}_2)^B(\text{Re}_2)^B\text{'O}_{12}$ is an above room-temperature ferromagnet (Curie temperature T_C of 560 K) with large room-temperature magnetizations and low-temperature switchable magnetoresistance;²⁰ LaCu₃Fe₄O₁₂ undergoes a temperature-dependent charge transfer around 400 K, accompanied by a metal-insulator transition behavior; ²⁸ CaCu₃Fe₂Re₂O₁₂ is a ferrimagnetic ($T_{\rm C} \sim 560$ K) half metal with large saturated magnetization of 8.7 $\mu_{\rm B}$; ³⁶ Mn_2FeReO_6 displays giant positive magnetoresistance up to 220%; ^{15, 51} ζ - Mn_2O_3 is the hardest direct narrow bandgap semiconductor, showing switchable p-n electronical conduction and spin-induced multiferroicity. 53-55 This list can be even longer. However, these exotic perovskites are thermodynamically metastable and need to be prepared at high-pressure (usually above 6 GPa) and temperature (HPT). This costly procedure and small-scale sample amount (usually milligram level) significantly limit the applications of these materials. Therefore, synthesis at ambient or much lower pressure, which remains a challenge, is highly desired.

Recently, Zhou *et al.* successfully stabilized gram-level high-pressure $Ca_{2-x}Mn_xTi_2O_6$ ($x \le 0.6$) phase under a very modest pressure (below 0.1 GPa). Ca_{1.4}Mn_{0.6}Ti₂O₆ is isostructural with CaMnTi₂O₆ ($P4_2mc$) and shows similarly high ferroelectric transition temperature. These findings suggest that it is possible to achieve large-scale high-pressure product driven by chemical potential (pressure)⁶⁰ and very modest physical pressure. Understandably, Ca^{2+} and Mn^{2+} have the same charge and comparable ionic radii (at eight coordination, VIII $_7$ (Ca^{2+}) = 1.12 Å, VIII $_7$ (Mn^{2+}) = 0.96 Å),⁶¹ so that the chemical pressure and geometric homogeneity can assist to underpin Mn^{2+} in the Ca^{2+} matrix. In this work we reported the perovskite solid solutions of $Ca_{2-x}Mn_xMnTaO_6$ ($0 \le x \le 1.0$) stabilized by the combination of chemical and physical pressure, and intensively studied the composition-dependent evolution of the crystal structure, formal oxidation states of cations, and magnetic properties.

EXPERIMENTAL SECTION

Synthesis. The Ca₂MnTaO₆ (CMTO) precursor was prepared by solid-state reactions with appropriate stoichiometric amount of CaCO₃ (MACKLIN, 99.99%), Mn₂O₃ (Sigma-Aldrich, 99.99%), and Ta₂O₅ (Alfa Aesar, 99.993%). The mixture was ground and pressed into pellets, calcined at 1275 K for 8 h to decompose the carbonate. Then, the product was reground and pressed into pellets, sintered in air for two periods of 48 h at 1653 K with intermediate grinding and pelletizing. Ca_{2-x}Mn_xMnTaO₆ samples with x = 0.2, 0.4, 0.6, 0.8, 1.0, 1.5, 1.6 and 2.0 were synthesized from the stoichiometric mixtures of the as-prepared CMTO, MnO (Alfa Aesar, 99.99%), Mn₂O₃, and Ta₂O₅. Samples with x = 0.2 were heated at 1625 K for 24 h under ambient pressure (AP), whereas x = 0.4 - 2.0 were prepared in a Walker-type multi-anvil apparatus at

1523-1625 K under 5-8 GPa for 30 min in Pt capsules and then quenched to room temperature, followed by a gradual release of the pressure.

X-ray Powder Diffraction and Energy Dispersive X-ray Spectroscopy. X-ray powder diffraction (XRD) data were collected at room temperature on a RIGAKU RINT-2000 diffractometer by using Cu-K α radiation ($\lambda = 1.5418$ Å). Here the 2θ range between 10° and 120° with a step size of 0.02° was measured, using a counting time of 0.24 s per step at 40 kV and 26 mA. The TOPAS 4.2 software package⁶² was applied to perform diffraction data analysis and Rietveld refinement. Cross sectional scanning electron microscopy (SEM) with energy-dispersive X-ray spectroscopy (EDS) images were recorded using a FEI Quanta 400F with an Oxford-instruments INCA 400 EDS detector operating at an accelerating voltage of 20 kV. EDS elemental composition could be expressed quantitatively as weight percentage or atomic percentage.

X-ray Absorption Near Edge and Photoelectron Spectroscopy. Mn-K and Ta-L_{2,3} X-ray absorption near edge spectroscopy (XANES) data were collected in both the fluorescence and the transmission modes with simultaneous standards. All of the spectra were fit to linear pre- and post-edge backgrounds and normalized to the unity absorption step across the edge. $^{13, 63, 64}$ The Ca_{1,8}Mn_{0,2}MnTaO₆ XANES measurements was performed on at the Brookhaven NSLS-II on the QAS 7-BM beamline using a Si-111 channel cut monochromator. Most of the standard spectra were performed at X-19A at NSLS-I with a Si-111 double crystal monochromator. X-ray photoelectron spectroscopy (XPS) was performed on a Nexsa XPS system equipped with a monochromatic Al $K\alpha$ X-ray source ($h\nu = 1486.6$ eV) operated at 720 W, and background

pressure was kept about 2×10^{-9} mbar. All binding energies were calibrated using surface contaminant carbon (C 1s = 284.8 eV) as a standard to scale.

Magnetic Measurements. Magnetic measurements were implemented with a Physical Properties Measurement System (PPMS, Quantum Design). The temperature-dependent magnetization was measured in zero field cooled (ZFC) and field cooled (FC) modes in a temperature scope of 5-400 K under 0.1 T magnetic field (*H*). Field dependence of isothermal magnetization was measured under an applied magnetic field varying from -5 to 5 T between 5 and 300 K.

RESULTS AND DISCUSSION

Synthesis and Structural Characterization.

The degree of cationic ordering and octahedral tilting in double perovskite oxides kinetically depends on synthetic conditions as in Ca_2MnTaO_6 , which was reported to be monoclinic $P2_1/m$ (No. 11, ordered Mn and Ta at the B-sites)⁶⁵ or the competitive orthorhombic Pbnm (disordering arrangement of Mn and Ta)⁶⁶ by different researchers. In our cases, all the $Ca_{2-x}Mn_xMnTaO_6$ ($0 \le x \le 2.0$) samples synthesized under AP and HP display black color. Both the x = 0 and 0.2 phases can be stabilized by chemical pressure only and synthesized under AP with a better explanation in orthorhombic Pbnm from the XRD patterns (**Figure 2a**). The chemical pressure alone is not enough to stabilize the x = 0.4 and 0.6 samples, since impurity peaks are observed beside the target phase as shown in **Figure S1a** and **c**. Attempts to purify x = 0.4 and 0.6 series with additional physical pressures between 2 and 5 GPa were unsuccessful, since the impurity phases can be suppressed at higher pressure but are still distinct in XRD patterns of the 5 GPa products

(Figure S1b and d). Therefore, higher physical pressure is required to stabilize and drive pure phase with x above 0.4. Figure 2a clearly shows that the pure orthorhombic phase can be obtained for $0.4 \le x \le 1.0$ at 7 GPa. The chemical composition of selected (x = 0.4-1.0) specimen were confirmed by EDS analyses (Figure S2 and Table S1). No single phase was achieved for x above 1.5 up to 8 GPa as shown in Figure S1e-i, where MnO and Mn₃Ta₂O₈ related phases are dominated in the x = 2.0 (Mn₂MnTaO₆) trial (Figure S1h and i). Conclusively, chemical pressure can adequately entangle Mn²⁺ at the Ca²⁺ site in Ca_{2-x}Mn_xMnTaO₆ for x around 0.2, and addition physical pressure of 7 GPa can assist to pump the phase boundary (x) around 1.0 (CaMnMnTaO₆) but less than 1.5 (Ca_{0.5}Mn_{1.5}MnTaO₆). Single phase in different structure type(s) may exist at higher pressure for x > 1.0 in Ca_{2-x}Mn_xMnTaO₆, which is, however, not the theme of this work.

The XRD peak evolution of $Ca_{2-x}Mn_xMnTaO_6$ is highlighted in the enlarged 2θ area between 21°-25° and 31°-34° in Figure 2b, respectively. The peaks continuously shift toward higher degree (right) with increasing x, suggesting successful incorporation of Mn^{2+} into the Ca^{2+} sites regarding their ionic radius difference. The main peak between 32-33° is somewhat broadened in the x = 0.6 case compared with others, suggesting symmetry degrading or the coexistence of two phases with very similar unit cell parameters. The structure refinements were conducted for all x = 0-1.0 samples as shown in Figure 3, and the corresponding crystal structures are present in Figure 4. The crystal structures, besides x = 0.6, can be well refined in orthorhombic cell (*Pbnm*) starting from the model of La₂MgTiO₆. The XRD patterns of x = 0.6sample can be well fitted by two *Pbnm* phases with similar cell parameters, which appeared to be around 0.51(1) (Ca_{1.49(1)}Mn_{0.51(1)}MnTaO₆, 57(5)% by weight) 0.32(1)(Ca_{1.68(1)}Mn_{0.32(1)}MnTaO₆, 43(7)% by weight) phases if extrapolated from the x-dependent cell parameter evolution diagram in **Figure 5a**. In the initial structural model, sites of La and Mg/Ti ions in La₂MgTiO₆ were replaced by Ca/Mn and Mn/Ta ions, respectively, with fixed occupancies according to the suggested chemical formula. The refined structural parameters and the coordinates of atoms are shown in **Table 1**, and main bond lengths are listed in **Table 2**. The unit-cell dimension evolution tolerably follows Vegard's law (**Figure 5a**), 68 and the deviation may be attributed to the effect of physical pressure. The average bond lengths of (Ca1/Mn2)-O1, (Ca1/Mn2)-O2, (Ta1/Mn1)-O1, and (Ta1/Mn1)-O2 show small fluctuations (**Figure 5b**), which is unobvious to state shortened ionic bond length under chemical pressure. The bond valence sums (BVS) calculations suggest that, the *A*-site (Ca1/Mn2) is under bonded with increasing *x*, while the *B*-site (Mn1/Ta1) is over bonded. To further confirm the formal oxidation state of cations in Ca_{2-x}Mn_xMnTaO₆, XANES and XPS measurements were conducted for selected samples.

XANES and XPS Analyses

The Mn-K main edge of Ca_{1.8}Mn_{0.2}MnTaO₆ is shown in **Figure 6a** along with a series of standard spectra for comparison⁶⁹⁻⁷¹. The Sr₂Mn²⁺ReO₆, and Ca₂Mn³⁺TaO₆ and CaMn⁴⁺O₃ spectra illustrate the systematic "chemical shift" of the Mn-K edge, for Mn on the perovskite *B*-site, to higher energy with increasing formal Mn-valence.⁶⁹⁻⁷¹ The Mn²⁺₂FeReO₆ spectrum, on the other hand, illustrates the much lower energy onset and peak typical for Mn²⁺ on the perovskite *A*-site.^{63, 72} Close inspection of the Ca_{1.8}Mn_{0.2}MnTaO₆ spectrum indicates: a dominant perovskite-*B*-Mn³⁺ component based on the proximity of the main peak to that of Ca₂Mn³⁺TaO₆; and a much smaller perovskite-*A*-Mn²⁺ component based on the excess spectral weight below the peak in the energy range where the Mn²⁺₂FeReO₆ spectrum peaks lie. To emphasize this

difference, spectrum (Diff. Spect. in the **Figure 6a**-top) was calculated by first subtracting the appropriately weighted $Ca_2Mn^{3+}TaO_6$ spectrum from the $Ca_{1.8}Mn_{0.2}MnTaO_6$ spectrum and the results were normalized to the standard unity absorption across the edge. Despite the crudeness of this approximation the difference spectrum manifests a very clear spectral peak at precisely the peak energy of the perovskite-*A*-site standard $Mn^{2+}_2FeReO_6$. Thus, there is a strong conclusion that the Mn-K edge results support the $\sim Mn^{3+}$ (perovskite-*B*-site) and $\sim Mn^{2+}$ (perovskite-*A*-site) assignments respectively for the Mn and $Mn_{0.2}$ components in the $Ca_{1.8}Mn_{0.2}MnTaO_6$ compound formula.

The prominent bimodal A/B peak features in 5d-row $L_{2,3}$ edge features, in octahedrally coordinated oxides, have been useful probes of d-configuration/valence by virtue of the systematic decrease in the A (t_{2g} -hole related) feature intensity, relative to that of the B (e_g -hole related) feature. The systematic A-feature spectral weight decrease with increasing d-count (decreasing t_{2g} -hole count) is clearly illustrated in **Figure 6b-c** for standard 5d-row compounds between d^0 and d^5 . 69,70,73,74 The Ta- $L_{2,3}$ edge spectra of $Ca_{1.8}Mn_{0.2}MnTaO_6$ is plotted as a solid red line in **Figure 6b-c** and its large A-feature intensity very clearly supports the d^0 /Ta⁵⁺ configuration/valence assignment for this compound. In summary the XANES results for $Ca_{1.8}Mn_{0.2}MnTaO_6$, manifest an A-site $\sim Mn^{2+}$ state, a B-site $\sim Mn^{3+}$ state, and a B'-site d^0 /Ta⁵⁺ state.

As shown in **Figure S3**, the XPS spectrum for Mn $2p_{3/2}$ region of CaMnMnTaO₆ (x = 1.0) was recorded and fitted by using XPS standard software. The spectrum exhibits three main peaks at about 640.5, 641.9, and 643.6 eV, respectively. The binding energy values are in good agreement with Mn²⁺, Mn³⁺, and Mn⁴⁺ oxidation states as reported in the literature.^{75,76} It should be pointed out that Mn²⁺ is attributed to *A*-site Mn ions and Mn³⁺ corresponds to *B*-site Mn ions. The

appearance of Mn^{4+} could derive from charge disproportionation of Mn^{3+} into Mn^{2+} and Mn^{4+} at B-sites. 77, 78

Magnetic Characterization.

Figure 7 shows the temperature-dependent ZFC and FC magnetic susceptibilities (χ) for x=0-1.0 samples measured at 0.1 T. In Figure 7a, the ferromagnetic (FM) transition temperature (Tc) can be determined as 19.6 K for x = 0 by means of a Curie-Weiss (CW) law fit ($\gamma = C/(T - 1)$ θ)), which implies that magnetic transition is due to spin-only Mn³⁺ (S = 2) and Mn³⁺-O²⁻-Mn³⁺ superexchange interactions. As shown in Figure 7b, on account of the spin magnetization of Mn^{3+} at B site and the doping of Mn^{2+} at A site, two magnetic transitions appeared with x=0.2. The effective magnetic moment ($\mu_{eff} = 5.37 \mu_B$) could be obtained by CW law fitting, which is close to the theoretical value (5.57 μ_B) corresponding to spin-only A site Mn²⁺ (HS d^5) and B site Mn^{3+} (HS d^4) moments as evidenced by the crystal structure and XANES results. With the increasing of dopant, the samples of x = 0.4 and 0.6 phases display an antiferromagnetic (AFM) state as shown in Figure 7c-d. The substitution of Ca²⁺ ions by Mn²⁺ can be defined as the "chemical pressure", which can be attributed to the decrease of magnetic transition temperature. 79,80 Their effective magnetic moment are greater than the calculated values due to magnetic interactions between Mn²⁺ and Mn³⁺. As exhibited in Figure 7e-f, another interesting finding is that a negative ZFC $\chi(T)$ is observed in the x = 0.8 and 1.0 samples. This has been proposed in the literature that misplaced B-site cations can result in the antiphase boundary coming into being, which accompanied by a short-ranged FM coupling between Mn²⁺ and Mn⁴⁺ ions at B sites as evidenced by XPS results, and an AFM coupling occurs in the antiphase boundary. In the ZFC mode, the magnetic field of 0.1 T is insufficient to align all frozen clusters

and domain spins in the direction of field. By this sense, as the temperature decreases, the spin anti-parallel or tilt state is stabilized, resulting in the residual magnetization tends to be negative. A similar phenomenon occurred in R_2NiMnO_6 (R = Pr, Nd, Y, and Ho) and $La_{2-x}Bi_xCoMnO_6$ (x = 0 and 0.1). 81,82

The isothermal magnetization M(H) was measured at different temperatures from 5 to 300 K under magnetic field between -5 and 5 T. The M(H) results presented in **Figure 8** support this conclusion with the magnetic hysteresis loop below T = 10 K, most dramatically evidencing a first-order increasing-field-induced transition out of the AFM state. Samples of x = 0 and 0.4 had S-type shape curves in 10 K, which reflected typical spin-glass-like magnetic properties, displayed in **Figure 8a** and **c**, respectively. The other samples showed a clear hysteresis loop given in **Figure 8b**, **d-f**. Their FM properties appeared with decreasing temperature owing to the concentration of induced magnetic moments on Mn at the A site increasing. The magnetic parameters of $Ca_{2-x}Mn_xMnTaO_6$ system with $0 \le x \le 1.0$ were summarized in **Table 4**, and have been used to map the magnetic phase diagram as shown in **Figure 9**. Samples of x = 0 and 0.2 were FM phases with $T_C = 19.6$ and 44 K, respectively. It can be found in $Ca_{2-x}Mn_xMnTaO_6$ for $0.4 \le x \le 1.0$ that Mn^{2+} at the A site involved in the long-range magnetic order and enhances the AFM order Mn-O-Mn super-exchange with increment of x, and T_N gradually becomes larger and tends to be gentle.

Unlike the other exotic perovskite with 50% of the *A*-site occupied by transition metal cations, such as the *A*-site columnar-ordered $P4_2/n$ -type quadruple perovskites CaMn*B*ReO₆ (B = Mn, 20 Fe, 20 Co, 21 Ni²¹), CaMn_{0.5}Cu_{0.5}FeReO₆, 20 and *R*MnMnSbO₆ (R = La, Pr, Nd, Sm), 22 and $P4_2mc$ structural Ca_{1.4}Mn_{0.6}Ti₂O₆ in Ca_{2-x}Mn_xTi₂O₆ ($x \le 0.6$), 19 either *A*- or *B*-site cationic ordering is absent in Ca_{2-x}Mn_xMnTaO₆, and the structural formula can be written as

 $(Ca_{1-x/2}Mn_{x/2})(Mn_{0.5}Ta_{0.5})O_3$, namely $(Ca^{2+}_{0.5}Mn^{2+}_{0.5})(Mn^{3+}_{0.5}Ta^{5+}_{0.5})O_3$ other than CaMnMnTaO₆ for x = 1.0, which is, to the best of our knowledge, for the first time observed in exotic perovskite with half of the A-site occupied by small transition metal ions. In $Ca_{2-x}Mn_xTi_2O_6$ prepared under chemical and soft physical pressure, similar A-site Ca/Mn disordering was observed for x below 0.4. However, A-site columnar-ordered quadruple perovskite $A_2A'A''B_4O_{12}$ is obtained for $Ca_{2-x}Mn_xTi_2O_6$ with $0.4 \le x \le 0.6$. When Ti^{4+} was replaced by Mn/Ta in $Ca_{2-x}Mn_xTi_2O_6$, the A-site columnar-ordered vanished in this work. Probably, the coexistence of B-site Mn^{3+} (Jahn-Teller distortion ion) and B'-site Ta^{5+} (second-order Jahn-Teller distortion ion) does not favor the formation of A-site columnar ordered perovskite structure $(Ca)^A_2(Mn)^{A'}(Mn)^{A''}(Mn)^{B_2}$ ($Ta)^{B'}_2O_{12}$ (x = 1.0 case), and the cationic disordering at both the A- and B-sites, together with charge disproportionation of Mn^{3+} into Mn^{2+} and Mn^{4+} at the B-sites, synergically renders a lower energy ground state in Pbmm structure in $Ca_{2-x}Mn_xMnTaO_6$.

Conclusions

In conclusion, we have prepared new exotic perovskite oxides $Ca_{2-x}Mn_xMnTaO_6$ ($0 \le x \le 1.0$) by means of combining chemical and physical pressure technique. The lower-Mn compounds ($x \le 0.2$) can be solely stabilized by chemical pressure, while the target phase dominated samples can be achieved at pressure of 2-5 GPa for $0.4 \le x \le 0.6$ and pressure around 7 GPa at $0.8 \le x \le 1.0$. The crystal structure remains Pbnm as in Ca_2MnTaO_6 , and can be structurally written as $(Ca_{1-x/2}Mn_{x/2})(Mn_{1/2}Ta_{1/2})O_3$ without expected A-site columnar-ordering. Both the X-ray absorption near edge and photoelectric spectroscopy data suggested $Ca_{2-x}Mn^{2+}_xMn^{3+}Ta^{5+}O_6$ ($0 \le x \le 0.6$) and $Ca_{2-x}Mn^{2+}_x(Mn^{3+,2+/4+})Ta^{5+}O_6$ ($0.8 \le x \le 1.0$). The spin-only A site Mn^{2+} (high spin d^5) and B site Mn^{3+} (high spin d^4) moments enhance a ferromagnetic ($x \le 0.2$) to

antiferromagnetic (0.4 $\leq x \leq$ 1.0) transition around the boundary of x = 0.2. The partial charge

disproportionation of the B-site Mn^{3+} into Mn^{2+} and Mn^{4+} x = 0.8 and 1.0 introduces negative

ZFC magnetization stemming from the formation of spin antiparallel or ferromagnetic clusters

and domains separating by the antiphase boundaries. This charge disproportionation over the

B-sites could have synergically contributuion to stabilize the highly disordered Pbnm structure in

Ca_{2-x}Mn_xMnTaO₆. The present findings update the fundamental understanding of fixing of

high-pressure phase in an ambient-pressure phase matrix, and imply that the coexistence of

Jahn-Teller distorted ions at the B- and B'-sites should be avoid in A₂A'A"B₂B'₂O₁₂ to ensure

cationic ordering and strong magnetic interactions.

ASSOCIATED CONTENT

Supporting Information

XRD patterns for unsynthesized phases; Mn 2p_{3/2} XPS binding energy; EDS spectra and analysis

Crystallographic information file for Ca_{1.8}Mn_{0.2}MnTaO₆ (CIF)

Crystallographic information file for Ca_{1.6}Mn_{0.4}MnTaO₆ (CIF)

Crystallographic information file for Ca_{1.2}Mn_{0.8}MnTaO₆ (CIF)

Crystallographic information file for CaMnMnTaO₆ (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: x = 0.2 (CSD1972421), x = 0.4 (CSD 1972420), x = 0.8 (CSD1972419), x = 1.0 (CSD 1972422).

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Figures and Captions

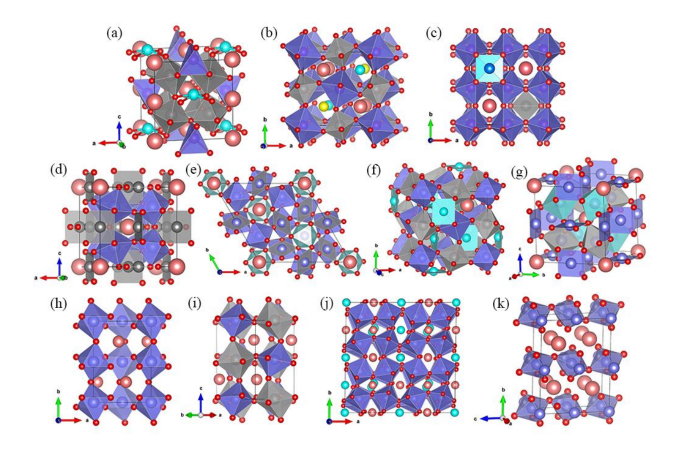


Figure 1. Crystal structures of exotic perovskite oxides with small A-site cations. A, pink spheres; O, red spheres; BO₆ octahedra, violet; B'O₆ octahedra, silver gray. A₂A'A"B₄O₁₂ with 50% of the small cationic A-sites (A' and A"): (a) A-site columnar-ordered quadruple perovskites $A_2A'A''B_4O_{12}$ (tetragonal, $P4_2mc$). $A'O_4$ square-planar, cyan; $A''O_4$ tetrahedral, violet; BO_6 octahedra, silver gray. (b) Rock-salt B-sites ordering in $A_2A'A''B_2B'_2O_{12}$ (tetragonal, $P4_2/n$). A', cyan spheres; A", yellow spheres. (c) RMn₃O₆ with layered charge ordering over the B-sites (orthorhombic, Pmmn). Mn1O₄ square-planar, cyan; Mn2O₄ tetrahedral, silver gray; MnO₆ octahedra, violet. $AA'_3B_4O_{12}$ with 75% of small cationic A-sites (A'): (d) Most known $AA'_3B_4O_{12}$ quadruple perovskites (cubic, Im-3). A'O₄ square-planar, silver gray. (e) B-site charge-ordering of A^{2+} Mn₃Mn₄O₁₂ (rhombohedral, R-3). Mn₁O₄ square-planar, silver gray; Mn₂O₆ octahedra, violet; Mn3O₆ octahedra, cyan. (f) 1322-type B-site ordering of AA'₃B₂B'₂O₁₂ (cubic, Pn-3). A'O₄ square-planar, cyan. (g) Trivalent A-site driving Jahn-Teller distorted Mn³⁺ component on the B-site of A³⁺Mn₇O₁₂ (monoclinic, I2/m), MnO₄ square-planar, violet; Mn4O₆ octahedra, silver gray; Mn5O₆ octahedra, cyan. Exotic perovskite with 100% of small cationic A-sites: (h) GdFeO₃-based ABO_3 -type (orthorhombic, Pnma); (i) $A_2BB'O_6$ -type (monoclinic, $P2_1/n$), (j) ζ -Mn₂O₃-type (triclinic, F-1), and (**k**) Post-perovskite (orthorhombic, Cmcm).

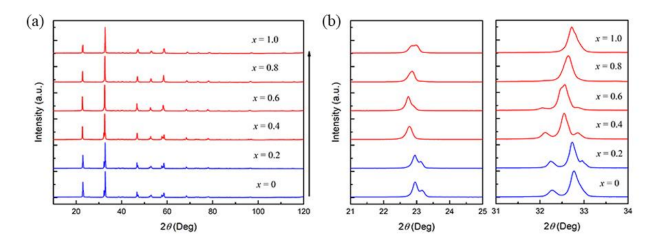


Figure 2. (a) XRD patterns of $Ca_{2-x}Mn_xMnTaO_6$ with x = 0, 0.2, 0.4, 0.6, 0.8 and 1.0, where blue and red lines denote the phase prepared at AP and HP, respectively. (b) The enlarged XRD patterns between $21^{\circ}-25^{\circ}$ (left) and $31^{\circ}-34^{\circ}$ (right) ranges.

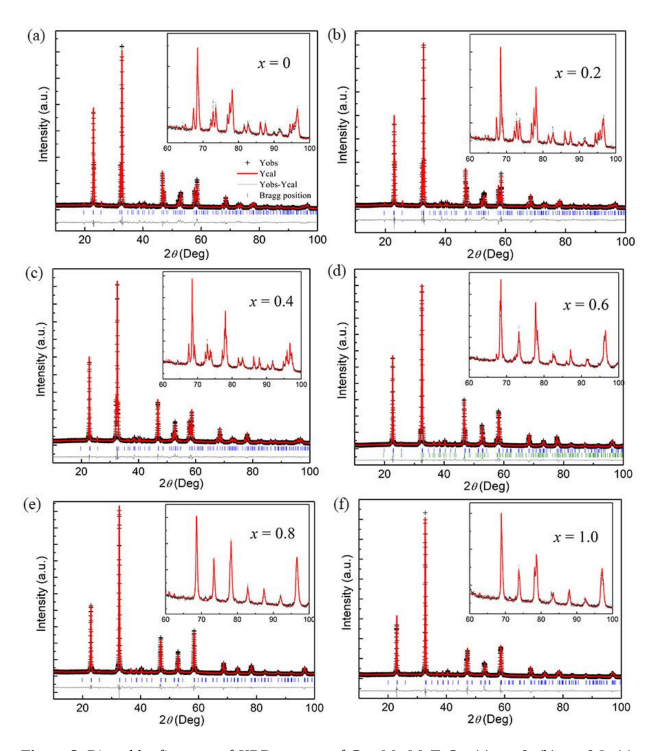


Figure 3. Rietveld refinement of XRD patterns of $Ca_{2-x}Mn_xMnTaO_6$: (a) x = 0; (b) x = 0.2; (c) x = 0.4; (d) x = 0.6; (e) x = 0.8; (f) x = 1.0. The insets (60-100°) show enlarged views.

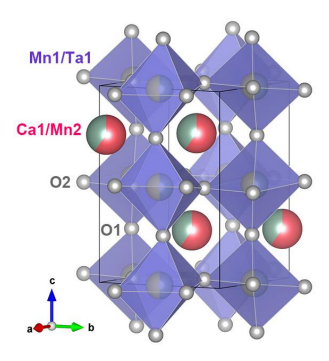


Figure 4. Crystal structure of $Ca_{2-x}Mn_xMnTaO_6$. Ca1/Mn2 are located at the *A* site and Mn1/Ta1 are disordered over the *B* site, the violet octahedra are (Mn1,Ta)O₆.

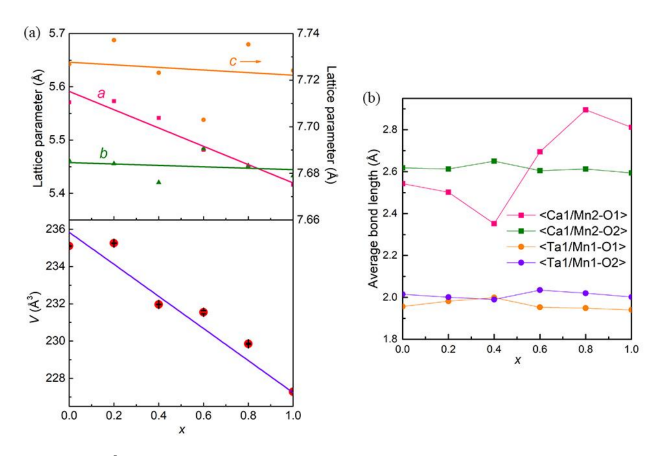


Figure 5. Mn²⁺ ion concentration dependence of (a) the orthorhombic lattice parameter a(Å), b(Å), c(Å), the unit cell volume $V(\text{Å}^3)$, and (b) average bond lengths in Ca_{2-x}Mn_xMnTaO₆ refined from XRD data.

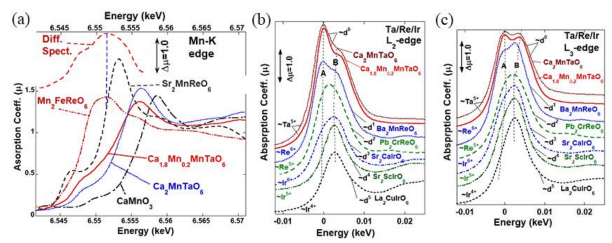


Figure 6. (a) The Mn–K edge spectra for $Ca_{1.8}Mn_{0.2}MnTaO_6$, compared with those of a series of standard compound spectra: the *A*-site perovskite based $Mn^{2+}_{2}FeReO_6$; the *B*-site perovskite based $Sr_2Mn^{2+}ReO_6$, $Ca_2Mn^{3+}TaO_6$ and $CaMn^{4+}O_3$. The spectrum labeled as "Diff. Spect." is a weighted difference spectrum (with normalization) to estimate the *A* site Mn spectrum in $Ca_{1.8}Mn_{0.2}MnTaO_6$. (b) A superimposed comparison of the $Ta-L_2$ and L_3 edge of $Ca_{1.8}Mn_{0.2}MnTaO_6$ to those of the d^0/Ta^{5+} - Ca_2MnTaO_6 , d^1/Re^{6+} - Ba_2MnReO_6 , d^2/Re^{5+} - Pb_2CrReO_6 , d^3/Ir^{6+} - Sr_2CaIrO_6 , d^4/Ir^{5+} - Sr_2ScIrO_6 and d^5/Ir^{4+} - La_2CuIrO_6 standards, indicating Ta^{5+} in $Ca_{1.8}Mn_{0.2}MnTaO_6$.

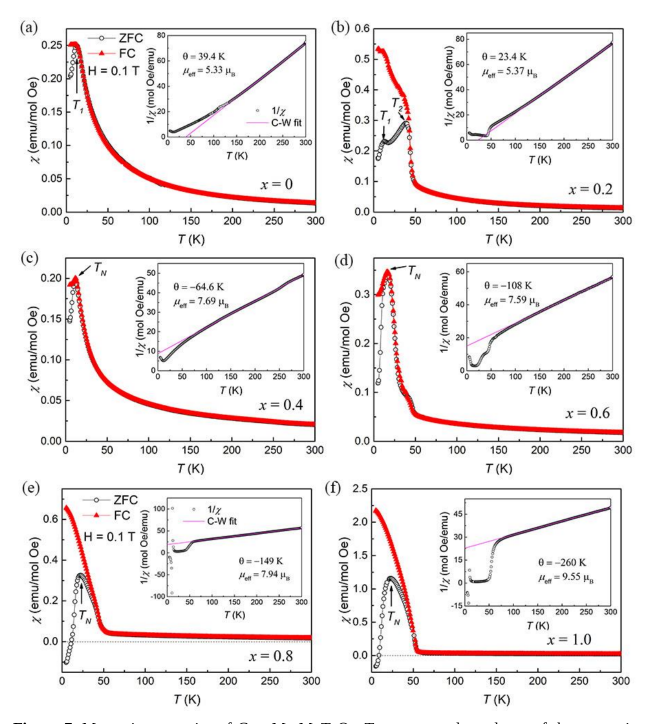


Figure 7. Magnetic properties of $Ca_{2-x}Mn_xMnTaO_6$. Temperature dependence of the magnetic susceptibilities measured at 0.1 T under zero-field-cooled (ZFC, empty circles) and field-cooled (FC, filled triangles) conditions between 5 and 300 K. (a) x = 0; (b) x = 0.2; (c) x = 0.4; (d) x = 0.6; (e) x = 0.8; (f) x = 1. The insets show the ZFC $\chi^{-1}vs$ T curves at 0.1 T with the Curie–Weiss fits.

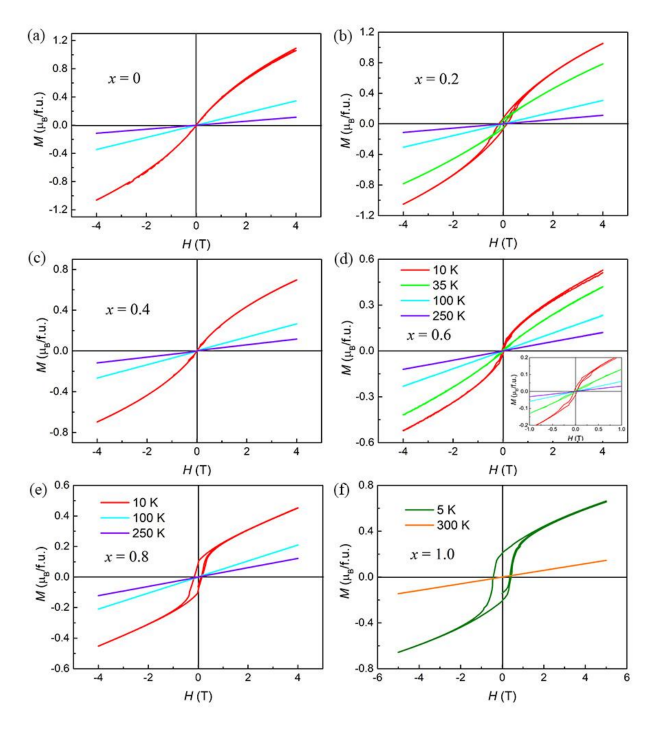


Figure 8. Isothermal magnetization curves of $Ca_{2-x}Mn_xMnTaO_6$ at 10, 35, 100, and 250 K between -4 and 4 T. (a) x = 0; (b) x = 0.2; (c) x = 0.4; (d) x = 0.6; (e) x = 0.8; (f) x = 1. The inset in (d) shows the expanded region between -1.0 and 1.0 T, showing clear hysteresis loops at different temperatures.

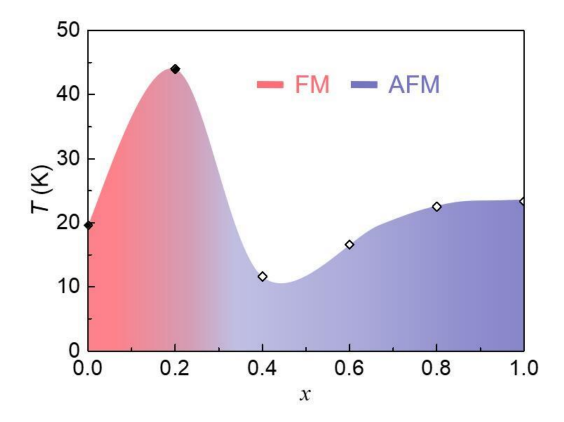


Figure 9. Magnetic phase diagram of $Ca_{2-x}Mn_xMnTaO_6$. The symbols are transition temperatures of $Ca_{2-x}Mn_xMnTaO_6$ (filled diamonds for T_C $0 \le x \le 0.2$, open diamonds for T_N of $0.4 \le x \le 1.0$). Pink area is for ferromagnetic (FM) phase, blue area for antiferromagnetic (AFM) phase.

Tables and Captions

Table 1. Structure refinements results of fractional atomic coordinates, occupancies and isotropic displacement parameters ($Å^2$) of Ca_{2-x}Mn_xMnTaO₆ with x = 0, 0.2, 0.4, 0.6, 0.8 and 1.0

	x = 0	x = 0.2	x = 0.4	x = 0.6		x = 0.8	x = 1.0	
				x = 0.51	x = 0.32			
a/Å	5.5707 (2)	5.5731 (1)	5.5414 (1)	5.4816 (5)	5.5512 (2)	5.450 (1)	5.4160 (2)	
b/Å	5.4620 (2)	5.4558 (1)	5.4203 (1)	5.4838 (5)	5.4197 (3)	5.4519 (9)	5.4330 (3)	
$c/ ext{Å}$	7.7268 (2)	7.7371 (2)	7.7231 (2)	7.7030 (2)	7.7478 (5)	7.7353 (3)	7.7241 (3)	
V/Å ³	235.10(1)	235.25 (1)	231.966 (8)	231.55 (3)	233.10 (2)	229.85 (6)	227.28 (2)	
Ca1/Mn2 4c (x, y, 1/4)								
X	-0.0166 (8)	-0.0262 (8)	-0.0409 (5)	-0.0327 (8)	-0.0377 (13)	0.0325 (10)	0.0315 (10)	
у	0.0316 (8)	0.0227 (12)	0.0247 (7)	-0.0248 (13)	0.0489 (12)	-0.0379 (8)	-0.0357 (8)	
Occ.	1	0.9/0.1	0.8/0.2	0.75/0.25	0.84/0.16	0.6/0.4	0.5/0.5	
$B_{ m iso}$	2.00 (15)	2.00 (16)	2.0(2)	2.0(2)	2.0 (9)	2.0(2)	2.0(3)	
Mn1/Ta1 4a (1/2, 0, 0)								
$B_{ m iso}$	2.44 (14)	2.21 (15)	2.1 (2)	2.5 (2)	2.1 (8)	2.3 (2)	2.7 (3)	
O1 4c (x, y, 1/4)								
X	0.0525 (18)	0.050(2)	0.001(3)	0.051 (4)	0.024 (6)	-0.044 (6)	-0.034 (5)	
у	0.481 (3)	0.439 (3)	0.4048 (16)	0.470 (6)	0.469 (4)	0.505 (3)	0.493 (3)	
$B_{ m iso}$	4.0 (4)	4.0 (5)	4.0 (4)	4.0 (6)	4.0 (11)	3.5 (7)	3.6 (6)	
O2 8d (x, y, z)								
X	0.7965 (15)	0.7971 (17)	0.7966 (11)	0.683 (4)	0.794 (3)	0.832 (6)	0.810(3)	
у	0.212 (2)	0.210(2)	0.2120 (14)	0.302 (5)	0.198 (3)	0.180(7)	0.175 (4)	
z	0.0496 (8)	0.0385 (13)	-0.0397 (9)	-0.0505 (13)	-0.015 (4)	0.0177 (19)	0.0305 (18)	
$B_{ m iso}$	1.9 (4)	2.6 (4)	1.9 (3)	3.0 (4)	1.5 (11)	2.1 (4)	4.0 (5)	
R_{wp} (%)	7.93	8.96	5.77	5.	42	6.69	8.09	
R_p (%)	5.77	6.53	4.25	4.	06	4.84	5.78	
χ^2	2.35	2.72	2.25	2.	12	2.54	3.29	
R_B (%)	4.13	4.81	2.72	2.87	2.33	2.64	2.67	

Table 2. Selected bond lengths (Å) and BVS of $Ca_{2-x}Mn_xMnTaO_6$ with x = 0, 0.2, 0.4, 0.6, 0.8 and 1.0

	x = 0	x = 0.2	x = 0.4	$x = 0.51^a$	x = 0.8	<i>x</i> = 1.0
(Ca1/Mn2)-O1	2.485 (17)	2.311 (17)	2.073 (10)	2.75 (3)	2.989 (17)	2.894 (17)
(Ca1/Mn2)-O1	2.600 (11)	2.693 (12)	2.631 (16)	2.64 (2)	2.80(3)	2.73 (3)
(Ca1/Mn2)-O2 (×2)	2.110 (9)	2.166 (11)	2.473 (7)	2.29 (2)	2.33 (2)	2.372 (17)
(Ca1/Mn2)-O2 (×2)	2.807 (10)	2.805 (11)	2.617 (7)	2.634 (15)	2.42 (3)	2.450 (15)
(Ca1/Mn2)-O2 (×2)	2.938 (8)	2.868 (11)	2.859 (7)	2.89 (2)	3.09 (3)	2.961 (18)
<(Ca1/Mn2)-O1>	2.543 (14)	2.502 (15)	2.352 (13)	2.695 (3)	2.895 (10)	2.812 (10)
<(Ca1/Mn2)-O2>	2.618 (9)	2.613 (11)	2.650 (7)	2.605 (6)	2.613 (3)	2.594 (17)
<(Ca1/Mn2)-O>	2.599 (10)	2.585 (12)	2.575 (9)	2.627 (5)	2.684 (5)	2.649 (15)
BVS(Ca1)	2.14	2.09	1.95	1.61	1.60	1.56
BVS(Mn2)	-	1.29	1.21	1.00	0.99	0.96
BVS(Ca1/Mn2)	2.14	2.01	1.80	1.43	1.36	1.26
(Mn1/Ta1)-O1 (×2)	1.9565 (17)	1.982 (3)	1.999 (2)	1.953 (4)	1.949 (4)	1.940(3)
(Mn1/Ta1)-O2 (×2)	1.977 (10)	1.967 (10)	1.950 (7)	1.98 (3)	1.98 (4)	1.944 (18)
(Mn1/Ta1)-O2 (×2)	2.053 (9)	2.035 (10)	2.029 (7)	2.09 (2)	2.06(3)	2.06(2)
<(Mn1/Ta1)-O1>	1.9565 (17)	1.982 (3)	1.999 (2)	1.953 (4)	1.949 (4)	1.940 (3)
<(Mn1/Ta1)-O2>	2.015 (10)	2.001 (10)	1.990 (7)	2.035 (3)	2.020 (4)	2.002 (10)
<(Mn1/Ta1)-O>	1.996 (12)	1.995 (8)	1.993 (5)	2.008 (3)	1.996 (4)	1.981 (8)
BVS(Mn1)	3.19	3.19	3.21	3.11	3.19	3.33

^aThe data for x = 0.51 phase is shown only in the x = 0.6 case.

Table 3. Magnetic parameters for $Ca_{2-x}Mn_xMnTaO_6$ (x = 0-1.0) measured in 0.1 T magnetic field

Compound	$\theta_{\mathrm{W}}\left(\mathrm{K}\right)$	$T_{\rm C}/T_{\rm N}$ (K)	$\mu_{\rm eff}$ ($\mu_{\rm B}/{\rm f.u.}$)	$\mu_{\rm cal}$ ($\mu_{\rm B}/{\rm f.u.}$)
x = 0	39.4	19.6	5.33	4.90
x = 0.2	23.4	44.0	5.37	5.57
x = 0.4	-64.6	11.6	7.69	6.14
x = 0.6	-108	16.6	7.59	6.71
x = 0.8	-149	22.5	7.94	7.21
x = 1.0	-260	23.3	9.55	7.68

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