Synthesis and Characterization of Magnetoelectric Ba$_7$Mn$_4$O$_{15}$

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**ABSTRACT:** We present the synthesis of a novel binary metal oxide material: Ba$_7$Mn$_4$O$_{15}$. The crystal structure has been investigated by high-resolution powder synchrotron X-ray diffraction in the temperature range of 100–300 K as well as by powder neutron diffraction at 10 and 80 K. This material represents an isostructural barium-substituted analogue of the layered material Sr$_7$Mn$_4$O$_{15}$ that forms its own structural class. However, we find that Ba$_7$Mn$_4$O$_{15}$ adopts a distinct magnetic ordering, resulting in a magnetoelectric ground state below 50 K. The likely magnetoelectric coupling mechanisms have been inferred from a careful symmetry-adapted refinement against the powder neutron diffraction experiments, as well as by making a comparison with the nonmagnetoelectric ground state of Sr$_7$Mn$_4$O$_{15}$.

**INTRODUCTION**

Solid-state phases containing Mn–O systems are of great interest due to the variety of functional properties they may exhibit, such as colossal magnetoresistance, ferromagnetism (FM), and ferroelectricity (FE). Multiferroic materials combine the latter two properties in a single material and have gained popularity recently for both their fundamental chemical novelty and their potential applications in devices. If the onset of magnetic and ferroelectric orderings has different mechanisms, the phase is known as a type I multiferroic. These phases typically have different FM and FE ordering temperatures, and the coupling between the two properties is weak. In a type II multiferroic, the magnetic ordering is induced by the ferroelectric ordering or vice versa, leading to a single ordering temperature and strong coupling between the ferroic states. One of the best-known examples of a Mn–O-containing multiferroic is YMn$_2$O$_5$, an orthorhombic Pbam structure comprising edge-shared chains of Mn$^{4+}$O$_6$ octahedra along the c axis that are linked by Mn$^{4+}$O$_5$ pyramids. The mechanism of the multiferroicity in this phase has been identified as exchange striction, in which the onset of magnetic ordering causes a polar structural distortion. All of the resulting components of the electrical polarization lie along the b axis. Another example of a Mn-containing multiferroic is TbMn$_2$O$_5$, in which the FE polarization is moderated by the Dzyaloshinkii–Moriya (DI) interaction and is thus quite weak. The arrangement of the Mn–O polyhedra in these phases has significant effects on the magnetoelectric coupling, and other multiferroics may be discovered by exploring phases containing unusual Mn–O linkages. A variety of type II multiferroic Mn–O-containing phases may already be found in the literature with ferroelectricity, which is induced by improper mechanisms, magnetic ordering, charge ordering, or orbital ordering.

The Sr$_7$Mn$_4$O$_{15}$ phase was first described by Kriegel et al.; it crystallizes in the P2$_1$/c space group and contains face-sharing Mn$_2$O$_9$ octahedral dimers, which share corners to form strings in the ac plane; these strings stack along a to give a quasi-two-dimensional structure. The face-sharing dimer motif is uncommon compared to solely corner-sharing systems, appearing in 4H-SrMnO$_3$ as well as in perovskites with a broad peak in the DC magnetic susceptibility and divergence between the field-cooled (FC) and zero-field-cooled (ZFC) warming curves. In Sr$_7$Mn$_4$O$_{15}$, these dimer units result in strong antiferromagnetic (AFM) interactions between the neighboring Mn$^{4+}$ sites, a broad peak in the AFM spins are slightly canted. We propose that the behavior could be explained by weak ferromagnetism (wFM) in which the AFM spins are slightly canted. We propose that the space group in which this is allowed, P2$_1$, would also allow local displacements of the oxide anions to produce FE ordering, resulting in a magnetoelectric ground state for the phase.

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general, the low symmetry of this structural type means that a large number of ways exist in which the magnetic exchange interactions may break inversion symmetry and hence allow for the magnetoelectric effect. Motived by this fact and previous reports of limited isovalent substitutions on the Sr site, we have prepared the Ba analogue Ba$_7$Mn$_4$O$_{15}$, which represents a novel binary oxide of Ba and Mn. Our detailed characterization of the magnetic ground state suggests that it may possess magnetoelectric coupling.

### EXPERIMENTAL DETAILS

To synthesize Ba$_7$Mn$_4$O$_{15}$, BaCO$_3$ (99.95%, Alfa Aesar) and MnO$_2$ (99.96%, Alfa Aesar) were ground together in a 7.7:4 ratio (i.e., a 10% molar excess of BaCO$_3$ compared to a stoichiometric reaction) and pressed into a pellet (diameter: 13 mm) under 7.5 metric tonnes of force. The pellet was heated to 900 °C for 48 h, then reground, pressed, and heated to 900 °C again. The grinding, pressing, and heating process was performed five times, and heating was always performed under an atmosphere of flowing N$_2$.

To synthesize Sr$_7$Mn$_4$O$_{15}$, SrCO$_3$ (99.9%, Sigma-Aldrich) and MnO$_2$ (99.96%, Alfa Aesar) were ground together in a 7:4 ratio and pressed into a pellet (diameter: 13 mm) under 7.5 metric tonnes of force. The pellet was heated to 900 °C for 20 h, then reground, pressed, and heated to 1000 °C for 24 h. The grinding, pressing, and heating process was performed six times under air.

High-resolution powder synchrotron X-ray diffraction experiments were performed at Beamline I11 at Diamond Light Source, with diffraction patterns recorded at 300 and 100 K using the multianalyzer crystal (MAC) for both Ba$_7$Mn$_4$O$_{15}$ and Sr$_7$Mn$_4$O$_{15}$, and variable-temperature measurements performed between these temperatures using the Mythen detector for Ba$_7$Mn$_4$O$_{15}$. The beam wavelength was 0.826831 Å for the Ba$_7$Mn$_4$O$_{15}$ diffraction experiment and 0.826341 Å for the Sr$_7$Mn$_4$O$_{15}$ refined using NIST 640c Si standards. Powder neutron diffraction experiments were performed using the GEM instrument at the ISIS Neutron and Muon Source at 80 and 10 K for Ba$_7$Mn$_4$O$_{15}$ and using the D2B instrument at the Institut Laue-Langevin (ILL) at 300, 100, and 1.5 K for Sr$_7$Mn$_4$O$_{15}$.

Rietveld refinements were performed using TOPAS Academic v6. A starting model for the refinement was based on the crystal structure of the related Sr$_7$Mn$_4$O$_{15}$ phase with space group P2$_1$/c. Combined Rietveld refinements were performed using the 100 K dataset from the I11 MAC detector and the 80 K dataset from the GEM experiment. These refinements excluded detector bank 1 and bank 6 from the GEM dataset due to poor signal-to-noise ratios. Refined lattice parameters and atomic coordinates for Ba$_7$Mn$_4$O$_{15}$ at 300 K are summarized in the Supporting Information (Table S1) isotropic displacement parameters were constrained to be equal for a given atom. Refinements of the low-temperature magnetic structure were performed within the symmetry-adapted formalism of the ISODISTORT suite and as implemented through the linear constraints language of TOPAS. Refined values from these refinements are also summarized in the Supporting Information (Table S2).

Magnetization measurements were performed using a Quantum Design MPMS SQUID magnetometer. Magnetic susceptibility versus temperature data were collected between 10 and 325 K in an applied field of 100 Oe under zero-field-cooled warming and field-cooled cooling conditions. Magnetization versus field curves were collected in applied fields of up to 50 kOe at temperatures between 2 and 300 K.

### RESULTS AND DISCUSSION

Sr$_7$Mn$_4$O$_{15}$ has been demonstrated to be flexible to limited substitution by both Ca$^{2+}$ and Ba$^{2+}$ cations at the Sr$^{2+}$ sites. Each substituting cation preferentially occupies different Sr sites within the unit cell, with the smaller Ca$^{2+}$ occupying the Sr1 site and the larger Ba$^{2+}$ occupying the Sr3 and Sr4 sites (see Figure 1). However, substitution levels greater than x = 1 for Sr$_{7-x}$Ba$_x$Mn$_4$O$_{15}$ have not previously been reported. We found that the use of an inert atmosphere and a 10% molar excess of BaCO$_3$ in the synthesis of Ba$_7$Mn$_4$O$_{15}$ was necessary for obtaining the desired phase. Any attempt to synthesize the product in air results in oxidation of the Mn$^{4+}$ cation to Mn$^{5+}$, forming a product which was identified as the Ba$_3$Mn$_2$O$_8$ phase exclusively. Additionally, performing the reaction at a reduced temperature of 900 °C (compared to literature at 1300 °C) produced the purest material. If an inert atmosphere is used without excess BaCO$_3$ in the reaction mixture, the reaction produces a mixture of Ba$_2$Mn$_3$O$_{10}$ and a secondary phase which was identified as Ba$_3$Mn$_2$O$_8$. Since Ba$_7$Mn$_4$O$_{15}$ possesses a greater Ba: Mn ratio than Ba$_2$Mn$_3$O$_{10}$ we found that a 10% excess of BaCO$_3$ produces a near phase-pure product, with just a small quantity of unreacted poorly crystalline BaCO$_3$ in the diffraction pattern. The inert atmosphere, requisite excess of BaCO$_3$ and exceptionally low synthesis temperature required to stabilize this phase are no doubt contributing factors as to why Ba$_7$Mn$_4$O$_{15}$ has hitherto remained unidentified.
collected at 300 K of Ba₇Mn₄O₁₅; the unreacted BaCO₃ appears as a broad peak around 2θ = 13° with a calculated concentration of ~12% by weight. This value may be overestimated due to the low crystallinity of the residual reagent. The unit cell of Ba₇Mn₄O₁₅ is similar to that reported for Sr₇Mn₄O₁₅, containing the same Mn₂O₉ dimer units, but with larger lattice parameters resulting from the greater ionic radius of Ba²⁺ compared to Sr²⁺.

At 300 K, we do not find evidence for the disordering of the Sr(3) and O(6) sites from their high-symmetry positions as described by Vente et al. for Sr₇Mn₄O₁₅.⁰²

The variations in lattice parameters and the angle β with temperature between 300 and 100 K are shown in Figure 3. All parameters vary monotonically with temperature, indicating the absence of any structural phase transitions in this temperature range.

![Figure 3](https://pubs.acs.org/acsicnic/article/61/10/10015)  
**Figure 3.** Lattice parameters of Ba₇Mn₄O₁₅ as a function of temperature.

In the case of Sr₇Mn₄O₁₅, the strings of Mn₂O₉ dimers were predicted to result in strong magnetic exchange interactions in the bc plane, but weak interactions along a.⁰²⁰ Previous reports of magnetic susceptibility experiments on Sr₇Mn₄O₁₅ describe a broad maximum centered around 75–90 K with a small upward tail below around 20 K. The FC and ZFC curves diverge from one another at temperatures below this maximum. This behavior has been explained with two different mechanisms: Vente et al. proposed that it resulted from spin glass-like behavior producing clusters of antiferromagnetically ordered spins, which crystallize into true antiferromagnetic order below ~75 K, whereas we have previously suggested that it might represent a weak FM ordering component, arising from the local symmetry-breaking associated with the aforementioned disorder of the Sr and O sites.⁰²⁰,²¹

We find that the DC susceptibility versus temperature results for Sr₇Mn₄O₁₅ match well with previous descriptions, with the maximum of the broad feature centered around ~84 K and the deviation between the ZFC warming and FC curves below this temperature (Figure 4). In comparison, the DC magnetic susceptibility results for Ba₇Mn₄O₁₅ are relatively featureless. We observe a steady upward trend with decreasing temperature between 300 and 50 K. However, on further cooling a clear divergence between the FC and ZFC warming curves is visible, evidencing a possible long-range magnetic ordering transition. The susceptibility obeys the Curie–Weiss law in the 200–300 K range. A fit to this part of the inverse susceptibility in the ZFC warming data yields a $\mu_{eff} = 3.78(7) \mu_B$ per Mn⁴⁺ site for Ba₇Mn₄O₁₅ that compares favorably against the expected spin-only value of 3.87 $\mu_B$.

To further investigate the magnetic behavior of Ba₇Mn₄O₁₅, we performed powder neutron diffraction at 10 and 80 K (i.e., either side of the divergence) using the time-of-flight powder diffractometer GEM, ISIS. Figure 5 shows the result of a combined Rietveld refinement using the powder synchrotron X-ray diffraction data at 100 K and the powder neutron diffraction data at GEM at 80 K (the same temperature points not having been measured). A fit to data from the same bank at 10 K is also included in the Supporting Information (Figure S1). This refinement of the neutron diffraction data at 80 K is well above the suspected magnetic ordering temperature and shows no significant fit intensity by our nuclear model. Figure 6 shows an enhanced view of the $d = 3.1–3.4$ Å and $d = 4.2–4.9$ Å regions of neutron diffraction data from the same bank at 10 K; magnetic Bragg reflections are evident that index as (0 1 2), (1 0 −2), (1 2 −1), and (2 1 −1).

To fit these magnetic peaks and solve the magnetic structure, the nuclear structure refined using the combined 80 and 100 K data above was used to produce a .cif file, which was used as a
starting model in ISODISTORT for the 10 K neutron data. The nuclear structure was fixed and the only parameters refined in the 10 K models were the components of the magnetic modes. There are two symmetry-inequivalent Mn\(^{4+}\) sites in the asymmetric unit, which share a face in a Mn\(_2\)O\(_9\) dimer. In our refinements, we have constrained the moments of these sites to be antiparallel. This is justified by the strong AFM direct exchange interactions expected for the half-full t\(_2\)g orbitals. We tested relaxing this constraint in our final model, but this led to neither a significant improvement in the fitting statistics nor a substantial deviation from the imposed antiparallel configuration. We tested models considering only a single magnetic propagation vector \(k = [0 0 0]\), as the magnetic intensities can all be indexed on the nuclear cell. At this \(k\)-point there are 4 irreducible representations (irreps), transforming as \(m\Gamma_1^+, m\Gamma_2^+, m\Gamma_1^-, m\Gamma_2^-\), according to the notation used with ISODISTORT. Illustrations of the spin configurations of each of these modes can be found in the Supporting Information (Figure S2); \(m\Gamma_1^+\) has FM components along \([010]\) and AFM components along \([100]\) and \([001]\), \(m\Gamma_2^+\) has FM components along \([100]\) and \([001]\) and AFM components along \([010]\), and both \(m\Gamma_1^-\) and \(m\Gamma_2^-\) have only AFM components along \([100]\), \([010]\), and \([001]\). The calculated components of the magnetic moments along each direction for each of these modes are included in the Supporting Information in the form of “complete modes details” pages found using ISODISTORT.

The results of the models constructed by considering spin orderings that transform as one of these irreps are summarized in Figure 7. The single-irrep models fail to account for all of the magnetic peaks, fitting either the \((012)\) and \((10-2)\) reflections or the \((12-2)\) and \((21-1)\) reflections, but not both. We therefore performed refinements in which a binary combination of modes was constrained to be active in either the \(ac\) plane or along \(b\), following the symmetry requirements of the unit cell. The three basis vectors spanning each irrep that describe the possible spin orderings correspond with moments aligned along the symmetry-unique direction \(b\) or in the \(ac\) plane. We tested all possible combinations exhaustively and report our findings in Table 1. We note that constraining the moments to lie only

Figure 5. Results of combined Rietveld refinement against GEM and I11 diffraction data at 80 and 100 K, respectively, for Ba\(_2\)Mn\(_4\)O\(_{15}\). Visualized is the fit against the data on bank 3 of GEM, though data from all banks was used to generate the model. Blue tick marks indicate reflections for Ba\(_2\)Mn\(_4\)O\(_{15}\); green tick marks indicate reflections for BaCO\(_3\).

Figure 6. Comparison of the powder neutron diffraction data for Ba\(_2\)Mn\(_4\)O\(_{15}\) from GEM bank 3 at 80 and 10 K, showing the development of magnetic Bragg peaks.

Figure 7. Fits to 10 K powder neutron diffraction data from GEM bank 3 at for Ba\(_2\)Mn\(_4\)O\(_{15}\) using single-mode models and dual-mode models, showing the failure of individual modes to capture all magnetic Bragg peak intensity. The underfit intensity in the peak at 4.5 Å is due to the BaCO\(_3\) impurity, marked with a star symbol.
along c made no difference to the fits, despite the moment being unconstrained within the ac plane by symmetry. The binary combinations of irreps produce both solutions where only the direction of the moment is modulated (that we will refer to as a spin-wave solution) and solutions where only the magnitude of the moment is modulated (spin density wave), which are highlighted in blue and red boxes, respectively, in Table 1. For a fixed Mn$^{4+}$ valence state, a spin wave is more physical, so we restrict our discussion to these in what follows. However, the assertions about the magnetoelectric ground state that we present below hold true irrespective of this fact.

The combinations of magnetic modes which resulted in the best fit to the data are as follows: $m\Gamma_1$ with $m\Gamma_1$, both along the c direction, $m\Gamma_1$ along c with $m\Gamma_1$ along b, and $m\Gamma_2$ along b with $m\Gamma_2$ along c. These fits are highlighted in bold in Table 1. Refinements in which the irreps were also allowed to refine in a were also investigated; we find that constraining the irreps along the c and b directions does not negatively impact the quality of the fit to the data, in line with literature predictions of the magnetic structure of Sr$_2$Mn$_2$O$_{15}$. While the components of the magnetic moments were not constrained to be equal along the b and c lattice directions, they consistently refined to approximately equal values as shown in Figure 8.

Despite demonstrating fitting statistics that equal the best models, we discard the first of the combinations ($m\Gamma_1$ with $m\Gamma_1$, both parallel to c) as it results in a spin density wave with unphysical descriptions of the magnitudes of the moments: namely, the calculated magnitude of the moments on half of the Mn sites are equal to 3.5 $\mu_B$ while the other half of the sites have magnitudes of 0.1 $\mu_B$. The remaining combinations of modes result in AFM spin-wave structures with magnetic space groups $Pc$ and $Pc'$. The fits to the data from these models are shown in the right-hand-side column of Figure 7, and the spin configurations within the unit cell are shown in Figure 8. We note that the presence of a small BaCO$_3$ impurity results in slight underfitting of the peak at 4.5 Å. These two models differ only slightly in the arrangement of the magnetic moments, and both produce magnetic moment magnitudes of around 2.3–2.4 $\mu_B$.

This is slightly reduced from the maximum expected value of 3 $\mu_B$, which we attribute to the overlap between $t_2g$ orbitals of the Mn centers within the Mn$_2$O$_3$ dimers causing a loss of spin density.

The fact that these two spin configurations are the best-fitting models to our data at low temperature is of significant interest: the joint action of the two magnetic irreps—one of which conserves inversion symmetry ($\Gamma^+$) and one of which violates inversion symmetry ($\Gamma^-$)—results in a structural space group in which inversion symmetry is globally broken. Indeed, comparing this type of combination (highlighted in Table 1 with a dark box) with combinations where both modes either conserve or do not conserve inversion symmetry, we find that the combinations resulting in a noncentrosymmetric space group fit the data best in all cases. While we are not able to detect an off-centering of any of the high-symmetry positions in Ba$_2$Mn$_2$O$_{15}$ in our powder diffraction data, the crystallographic analysis indicates that this phase possesses a ground state in which the noncentrosymmetric space group ($Pc$) is a direct result of the magnetic ordering, implying that it may be

![Figure 8](https://doi.org/10.1021/acs.inorgchem.2c00889)  
Possible spin configurations for Ba$_2$Mn$_2$O$_{15}$, where the magnetic moments are constrained to be in the bc plane. The total magnetic moment for the $Pc$ model is 2.46 $\mu_B$ (1.6 $\mu_B$ along b, 1.9 $\mu_B$ along c), and the total magnetic moment for the $Pc'$ model is 2.34 $\mu_B$ (1.5 $\mu_B$ along b, 1.8 $\mu_B$ along c). The two symmetry-unique Mn sites in $P2_1/c$ are indicated by blue and red arrows; their coupling is constrained to be AFM as this was found to fit the experimental data best and expected by the strong magnetic direct exchange interactions.
magnetoelectric and/or multiferroic. A table of the space groups which result from the combination of multiple magnetic modes is shown in the Supporting Information (Table S3).

Figure 9 shows magnetization measurements for Ba$_7$Mn$_4$O$_{15}$ as a function of the applied field. A small but clear hysteresis is evident at temperatures less than 25 K, with a maximum moment of 0.083 $\mu_B$ per Mn$^{4+}$ at 2 K and 50 kOe. As this is significantly lower than the effective moment in the paramagnetic region, the ordering is likely to be largely AFM in character, with a small FM component. This can be attributed to a slight canting of the magnetic moments along the c direction, suggesting that there is a small incomplete cancellation of the magnetic moments within the Mn$_4$O$_9$ dimers along this lattice direction. These magnetization results further support the proposed two-irrep magnetic structure and hence also the proposed polar ground state.

Figure 10 shows one of the sets of allowed polar displacements of the O6 anion. This site and the Ba3 site are the only high-symmetry positions in the $P2_1/c$ unit cell, and thus we chose one of these to demonstrate a possible multiferroic coupling mechanism since displacements of these atoms from their average positions will always induce a permanent polarization. The polar displacement transforms as the $\Gamma_2^{-}$ irrep, which is responsible for the displacive distortions that lead to a symmetry reduction from $P2_1/c$ to $Pc$ (basis = {(1,0,0), (0,1,0),(0,0,1)}, origin shift = (0,1/4,0) with respect to the parent cell). Notably, polar distortions transforming as this irrep appear irrespective of which magnetic space group ($Pc$ or $Pc'$) is assigned, thus our result is invariant to the ambiguity of the precise magnetic structure with respect to these two space groups. Since the noncentrosymmetric ground state would be induced by the magnetic ordering, Ba$_7$Mn$_4$O$_{15}$ would be classed as a Type II multiferroic. It is important to emphasize that the displacements indicated in Figure 10 are only one of the possible distortion pathways by which magnetoelectric coupling could be realized. Since our proposed distortions are driven by magnetic ordering, the displacement is expected to be on the order of thousandths of an angstrom. Our experiment is not sensitive to displacements of this magnitude in such a complex structure, and so Figure 10 merely shows the symmetry-allowed character of one such possible displacement, and we can infer nothing more about their magnitude or the relative magnitude of other symmetry-allowed atomic displacements. Additionally, it should be noted that the proposed magnetoelectric ground state arises not only as a result of the canting of the magnetic spins, but is due to the combined action of the two magnetic irreps. While a single magnetic irrep solution can lead to canting in this system, it will never result in a multiferroic ground state.

Finally, we compare the neutron diffraction results for Ba$_7$Mn$_4$O$_{15}$ with those of Sr$_7$Mn$_4$O$_{15}$, for which no direct evidence of a magnetoelectric ground state has previously been reported. A powder neutron diffraction pattern for Sr$_7$Mn$_4$O$_{15}$ is shown in Figure 11, with the magnetic peaks observed at 1.5 K inset. The most plausible fit to the magnetic peaks is achieved using a model containing only the $m\Gamma_2^{-}$ irrep along the b direction to describe the magnetic moments, consistent with previously published models. The spin configuration refined against the data and transforming as the $m\Gamma_2^{-}$ irrep is shown in Figure 12. This produces a calculated magnetic moment of 2.2 $\mu_B$ consistent with the magnitude observed for Ba$_7$Mn$_4$O$_{15}$, giving us further confidence in our proposed spin-wave solutions. The two-irrep fits necessary to model Ba$_7$Mn$_4$O$_{15}$ provide no improvement in the quality of the fit to the magnetic data for Sr$_7$Mn$_4$O$_{15}$, thus the magnetoelectric ground state we report here is specific to Ba$_7$Mn$_4$O$_{15}$. We view the $Pc$ ($m\Gamma_2^{-}$ + $m\Gamma_3^{-}$) structure of Ba$_7$Mn$_4$O$_{15}$ to be the most likely candidate from our refinements, as the $m\Gamma_2^{-}$ mode best describes the magnetic ordering in Sr$_7$Mn$_4$O$_{15}$ and it is to be expected that the...
exchange pathways have a high degree of similarity in these two materials.

It is difficult to assign the difference in magnetic behaviors between Ba$_7$Mn$_4$O$_{15}$ and Sr$_7$Mn$_4$O$_{15}$ to specific structural features due to the low symmetry of the phases. A selection of the nearest-neighbor distances and angles are summarized in Table S3; while the unit cell obviously expands to accommodate the larger Ba$^{2+}$ cations, this expansion does not result in exceptional changes to any of the bond lengths or angles between Mn$^{4+}$ centers. We envisage that substantial future works involving investigating the magnetoelectric ground state of this material, via first-principles calculations, will shed further light on this issue.

CONCLUSIONS

We have successfully synthesized the novel ternary compound Ba$_7$Mn$_4$O$_{15}$. Powder synchrotron X-ray diffraction analysis indicates that this phase remains in the $P2_1/c$ space group in the 100–300 K temperature range. Powder neutron diffraction and SQUID magnetometry indicate that the phase possesses an AFM ground state below 50 K. Careful analysis of this AFM ground state reveals a small FM component and suggests that a pair of magnetic modes, transforming as distinct irreducible representations, act simultaneously, and in doing so couple to a polar distortion. This is the first experimental evidence that this class of materials can support a multiferroic grounds state, and we hope it will stimulate renewed synthetic effort into preparing structural related materials. Further work on Ba$_7$Mn$_4$O$_{15}$ should focus on direct characterization of the nature of the polar displacement, confirming and demonstrating the switchability of the polar state and performing synthetic work with the aim of enhancing the magnetoelectric ordering temperature.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.inorgchem.2c00889.

Supporting data and figures mentioned in the main text (PDF)

Ba$_7$Mn$_4$O$_{15}$ from Rietveld refinement of powder synchrotron X-ray diffraction data at 300 K (CIF)

Ba$_7$Mn$_4$O$_{15}$ from a combined Rietveld refinement of powder synchrotron X-ray diffraction data at 100 K and powder neutron diffraction data at 80 K (CIF)

Components of the magnetic modes derived from ISODISTORT (PDF)

Components of the magnetic modes derived from ISODISTORT (PDF)

Components of the magnetic modes derived from ISODISTORT (PDF)

Components of the magnetic modes derived from ISODISTORT (PDF)

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Figure 11. Rietveld refinement against powder neutron diffraction data collected at D2B for Sr$_7$Mn$_4$O$_{15}$ at 300 K. Blue ticks indicate reflections for the Sr$_7$Mn$_4$O$_{15}$ phase; green ticks indicate reflections for a small SrMnO$_3$ impurity. Inset: magnetic reflections observed at 1.5 K (black crosses) and the same region of $2\theta$ at 300 K (green crosses). The magnetic intensity is well modeled by considering only the $m\Gamma_2^-$ irrep.

Figure 12. Refined $m\Gamma_2^-$ spin configuration for Sr$_7$Mn$_4$O$_{15}$ (space group $P2_1'/c$), where the magnetic moments are constrained to be in the b direction. The magnetic moment is 2.2 $\mu_B$. The two symmetry-unique Mn sites in $P2_1/c$ are indicated by blue and red arrows; their coupling is constrained to be AFM as this was found to fit the experimental data best and expected by the strong magnetic direct exchange interactions.
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Notes
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