Dalton Transactions



PAPER

View Article Online



Cite this: *Dalton Trans.*, 2015, **44**, 20441

Received 4th September 2015, Accepted 12th October 2015 DOI: 10.1039/c5dt03445k

www.rsc.org/dalton

High pressure synthesis of polar and non-polar cation-ordered polymorphs of Mn₂ScSbO₆†

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Two new cation-ordered polymorphs of Mn_2ScSbO_6 have been synthesised at high-pressure. At 5.5 GPa and 1523 K Mn_2ScSbO_6 crystallizes in the Ni_3TeO_6 -type structure with the polar R3 space group and cell parameters a=5.3419 (5) Å and c=14.0603 (2) Å. Below $T_C=42.0$ K it exhibits ferrimagnetic order with a net magnetization of $0.6\mu_B$ arising from unusual site-selective Mn/Sc disorder and is thus a potential multiferroic material. A double perovskite phase obtained at 12 GPa and 1473 K crystallizes in the non-polar $P2_1/n$ monoclinic space group with cell parameters a=5.2909 (3) Å, b=5.4698 (3) Å, c=7.7349 (5) Å and $\beta=90.165$ (6) °. Magnetization and neutron diffraction experiments reveal antiferromagnetic order below $T_N=22.3$ K with the spins lying in the ac plane.

Introduction

Material properties are dependent upon atomic arrangement and the degree of order can be tailored by controlling the pressure–temperature conditions and bulk composition. At a given pressure *P* and temperature *T*, there are different possible atomic arrangements that correspond to local minima of free energy, the lowest-energy conformation being the thermodynamically-stable phase. However, high pressure and high temperature (HPHT) synthesis conditions may favour higherenergy minima and metastable phases, making them kinetically stable under ambient conditions and therefore recoverable.

Amongst transition metal oxides, HPHT helps to stabilize unusual oxidation states and environments that result in useful properties, e.g. the room temperature ferromagnetic metal CrO_2 . Moreover, HPHT promotes interesting structural mechanisms. The ilmenite (IL) FeTiO₃, for example, crystallizes in the space group $R\bar{3}$ with Fe and Ti stacked into alternate layers along the c-axis. It transforms into an unquenchable distorted perovskite (Pv) structure (space group Pbnm) at 16 GPa and converts back into a polar LiNbO₃-type

(LN, space group R3c) with Fe and Ti cations being ordered in the same layers.^{3,4}

Moreover, ABO₃ oxides with Mn²⁺ on the A site are of fundamental interest due to the electronic and magnetic phenomena that emerge from the coupling of spin, charge and orbital degrees of freedom. HPHT conditions are often needed to stabilize MnBO₃ materials, e.g. perovskite-type MnVO₃ with an incommensurate magnetic structure and metallic conductivity,5 and LiNbO3-type MnTiO3-II with a weak ferromagnetism through anisotropic exchange interactions. The use of HPHT on ordered quaternary systems (AA'B2O6 or A2BB'O6) is a relatively less explored area with only a few reports. 7,8 Amongst these, Mn₂FeB'O₆ (B' = Ta and Nb) with a LN-type structure⁹ or (B' = Mo, W) with the Ni_3TeO_6 -type (NTO)^{10,11} order shows polar and magnetic properties, while Mn₂BSbO₆(B = Fe and Cr) shows polymorphism between the IL- and double perovskite-types (DPv) depending on whether they are synthesized at moderate (3-5 GPa) or higher pressures (5.5-8 GPa). 12,13

In this work, we present the order–disorder effects on the high pressure polymorphs of Mn₂ScSbO₆ oxide. Mn and Sc cations are randomly distributed in a corundum-related type structure when this material is synthesized under ambient conditions and it shows no long-range magnetic ordering.¹⁴ The double perovskite (DPv) can be achieved with pressures higher than 10 GPa, but below 5.5 GPa the polar NTO-structure is obtained. NTO_Mn₂ScSbO₆ shows an unusual ferrimagnetism due to partial substitution of non-magnetic Sc³⁺ at just one of the two Mn²⁺ sites, whereas DPv_Mn₂ScSbO₆ is antiferromagnetic. Combined X-ray and powder neutron diffraction refinements and electron micro-diffraction experiments confirm the non-centrosymmetry of the NTO_Mn₂ScSbO₆ polymorph allowing a structural polarization that is predicted to

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[†] Electronic supplementary information (ESI) available. CCDC 1428909–1428911. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5dt03445k

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be 28.3 μ C cm⁻² at room temperature. These results demonstrate that cation ordering can be induced by high pressure synthesis providing access to new metastable phases with unusual properties including multiferroicity. We also show that site selective disorder provides an unusual way to induce ferrimagnetism.

II. Experimental section

 $\rm Mn_2ScSbO_6$ was previously synthesized at ambient pressure by Kosse $\it et al.^{15,16}$ and subsequently studied by Ivanov $\it et al.^{14}$ NPD studies showed a random distribution of Mn and Sc crystallizing in a corundum-related structure ($\it R\bar{3}$) with no long-range magnetic ordering. We prepared a sample at 1373 K and Supplementary Fig. 1 (SF1†) shows the Rietveld fit to the XRD pattern that agrees with a random cation distribution.

Both double perovskite and Ni_3TeO_6 -type polymorphs were synthesized under high pressure and high temperature conditions. The precursor, prepared by grinding stoichiometric amounts of Mn_2O_3 , Sc_2O_3 and Sb_2O_3 oxides, was treated at 12 GPa and 1473 K for 20 minutes in a Walker-type multianvil apparatus for the preparation of the perovskite phase. The application of 5.5 GPa and 1523 K for 35 minutes in a belt-type press led to the formation of the Ni_3TeO_6 polymorph. In both cases, the sample was quenched by rapid cooling and the pressure was progressively released down to ambient conditions.

The crystal structures were first characterized by Rietveld refinement of powder X-ray diffraction collected on a Phillips X'Pert Pro Alfa 1 diffractometer using Cu $K\alpha_1$ radiation, equipped with a Ge (111) monochromator working in Bragg–Brentano geometry. The diffractograms were collected between 10 and 120 degrees with a step size of 0.017°.

For transmission electron microscopy (TEM) studies, samples were ground in *n*-butyl alcohol and ultrasonically dispersed. A few drops of the resulting suspension were deposited on a carbon-coated grid. The study of reciprocal space by selected area electron diffraction (SAED) and microdiffraction was carried out using a JEOL JEM2100 microscope operating at 200 kV with a double tilt (±42°) goniometer. High resolution transmission electron microscopy (HRTEM) and electron energy loss spectroscopy (EELS) experiments were performed with a JEOL JEM 3000F microscope operating at 300 kV (double tilt (±25°) point resolution 0.17 nm), fitted with an energy-dispersive X-ray spectroscopy (XEDS) microanalysis system (OXFORD INCA) and an ENFINA spectrometer with an energy resolution of 1.3 eV.

Simulations of the HRTEM images were performed with the software MacTempas X_1^{17} using the refined structures from neutron diffraction data. The oxidation state of Mn in both materials was determined by EELS with the relationship between the white-line intensity ratio (L_3/L_2) and the oxidation state. ¹⁸

Neutron powder diffraction (NPD) patterns were collected at 300 K on a high resolution D2B diffractometer (Institut Laue-Langevin, Grenoble) between 0 and 160° with a step-width of

0.05°, using a neutron wavelength $\lambda = 1.594$ Å and a standard He cryostat. The NTO polymorph was also cooled down to 4 K, where a long scan was measured under the same conditions. The evolution of the magnetic structures was studied using the sequential patterns collected at each degree from 3 K up to 55 K on the high intensity D1B instrument. Each pattern was measured in the angular range $0^{\circ} \le 2\theta \le 130^{\circ}$ with a step-width of 0.1° using $\lambda = 2.520$ Å. Long scans were also taken at 3 K.

The nuclear structures were refined from room temperature D2B data, using Rietveld analysis through the FullProf software package¹⁹ and considering a Thomson–Cox–Hastings function to optimize the shape of the peaks. The magnetic symmetry analysis of the low temperature D1B data was performed by means of the program BasIreps.²⁰

Magnetic susceptibility measurements were performed on a Quantum Design XL-MPMS SQUID magnetometer under DC Zero-Field-Cooling (ZFC) and Field-Cooling (FC) conditions in the temperature range 2 K < T < 300 K under a magnetic field of 3 kOe. Magnetization dependence on the magnetic field strength was studied at 5, 20, 40 and 90 K for the perovskite polymorph up to 5 T and at 2, 20 and 40 K for the NTO-type oxide up to 7 T.

III. Results and discussion

(A) NTO_Mn₂ScSbO₆

Structural characterization. The lower pressure polymorph of Mn_2ScSbO_6 can be obtained at 5.5 GPa and 1523 K. Fig. 1 (bottom) shows the XRD diffraction pattern of this material. It can be indexed in a hexagonal cell with a = 5.34186 (5) Å and c = 14.0603 (2) Å cell parameters. This cell is indicative of a corundum-type related structure, as observed in other $Mn_2BB'O_6$ oxides. $^{9-13}$ The presence of the (003) and (101) reflections (inset Fig. 1-bottom) demonstrates the absence of a c-glide plane and therefore excludes the corundum- ($R\bar{3}c$) and LN-type (R3c) structures, suggesting an ilmenite-type ordering ($R\bar{3}$). Rietveld refinement was performed with the IL- Mn_2FeSbO_6 structure as a starting model. The refinement converged to $R_{wp} = 11.3\%$ and $R_p = 6.24\%$ agreement factors.

However, NPD experiments on the same compound revealed no intensity for the (003) and (101) reflections (Fig. 1-top) and the IL-type model obtained from XRD refinement showed a poor fit to the neutron data. The data can be fitted assuming a LN-type ordering (R3c) with $R_{\rm wp}$ = 4.40% and $R_{\rm p}$ = 3.41% agreement factors.

In order to clarify this situation, electron microscopy experiments were performed. The SAED patterns exhibit the reflection conditions: hkil: -h + k + l = 3n, hki0: -h + k = 3n, 000l: l = 3n, h - h00: h = 3n, h - h0l: h + l = 3n and the only feasible space groups are $R\bar{3}$ and $R\bar{3}$ (Fig. 2a–c).

Fig. 2d and e show the microdiffraction of the $[2-1-1\ 0]$ zone axis. The no periodicity difference between the zero and first order Laue zones (ZOLZ and FOLZ) indicates the absence of a c-glide plane, in agreement with XRD. Moreover, the analysis of the whole pattern (WP) of the $[2-1-1\ 0]$ and $[0\ 1-1\ 0]$

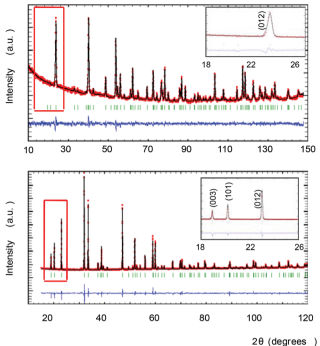


Fig. 1 Rietveld refinements of the nuclear structure of NTO_Mn_2SCSbO_6 from room temperature NPD profile collected using a D2B diffractometer (top) and XRD data (bottom). The insets show an enlargement of the low angle range identified with the red rectangles, where reflections (003) and (101) are observed in the XRD pattern although they are not seen in the NPD profile, as discussed in the text.

zone axes (Fig. 2f) unequivocally distinguishes between $R\bar{3}$ (two-fold axis) and $R\bar{3}$ (no symmetry).²¹

A model with *R*3 symmetry was then tested. In order to break the inversion centre of the IL-type model, we assumed a complete ordering within the (00z)-layers between Mn1/Sc and Mn2/Sb; this cation arrangement is adopted in the Ni₃TeO₆-type structure with the polar *R*3 space group. The NTO-type structure has also been observed in the related compounds Mn₂FeMoO₆ and Ni₂ScSbO₆. The refinement converged smoothly; Fig. 1 shows the Rietveld refinements of the NPD (top) and XRD (bottom) data for the NTO_Mn₂ScSbO₆ model and Table 1 summarizes the structural details and agreement factors resulting from this combined refinement.

The final model clarifies the apparent contradiction between the XRD and NPD results. The cation arrangement of the R3 NTO-type structure is illustrated in Fig. 3 (left). The (00z) layers alternate between Mn1/Sc and Mn2/Sb composition. For XRD, Sc (Z=21) and Mn (Z=25) look similar when compared with the consecutive layer of Mn and Sb (Z=50). Therefore, IL-type ordering is apparent and the (003) and (101) reflections are observable. On the other hand, the difference in neutron scattering lengths for Mn, Sc and Sb (-3.73, 12.29 and 5.57 fm respectively) breaks the apparent inversion centre, as the (012) planes are alternatively constituted by Mn or by Sc/Sb, which makes the scattering distribution resemble that of an LN-type. For clarity, both IL- and LN-type orders are high-

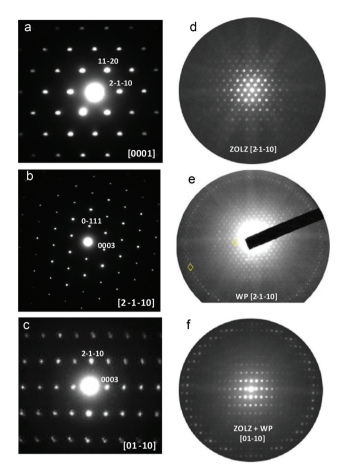


Fig. 2 SAED patterns for NTO_Mn₂ScSbO₆ along the $[0\ 0\ 0\ 1], [2\ -1\ -1\ 0]$ and $[0\ 1\ -1\ 0]$ zone axes (a-c, respectively) and microdiffraction patterns (d-f) along the $[2\ -1\ -1\ 0]$ and $[0\ 1\ -1\ 0]$ zone axes.

lighted in Fig. 3 (left) with black and red dashed lines respectively. In conclusion, the combination of SAED, NPD and XRD techniques unequivocally leads to the true NTO-type structure of this compound.

EELS analysis performed on several crystallites yields a +2 oxidation state for Mn (see ESI Fig. SF2a†). A structural HRTEM image along the $[2-1-1\ 0]$ zone axis (SF2b†) shows a well ordered material and the absence of extended defects.

The structure of NTO_Mn₂ScSbO₆ can be described as a corundum derivative with (00z) layers alternatively occupied by [Mn1O₆]/[ScO₆] and [Mn2O₆]/[SbO₆] edge sharing octahedra. The order is such that the polyhedra sharing faces along the c axis are always Mn1 against Sb and Sc against Mn2 (see Fig. 3 left). All site occupancies were tested by allowing cation exchange among the different sites, but the only significant result was found between Sc and Mn2 positions and the final refinement converged to a value of 12.3 (2)% of anti-site mixing. The insertion of this small amount of Sc³⁺ in the Mn2 site decreases its octahedral distortion from 5.5 × 10⁻³ observed for Mn1 down to 1.7 × 10⁻³, while the observed Δ [ScO₆] value remains 1.1 × 10⁻⁴ as that observed for [SbO₆].

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Table 1 Crystallographic parameters, selected interatomic distances (Å) and angles (°) for NTO_Mn₂ScSbO₆ from the combined refinement of room temperature NPD and XRD data in the R3 space group^a

a = 5.3419 (5) (Å), c = 14.0603 (2) (Å)							
Atom	Site	x	y	z	Occ		
Mn1	3a	1/3	2/3	0.1619(8)	1		
Mn2/Sc	3a	2/3	1/3	0.3423(8)	0.877(2)/0.123(2)		
Sc/Mn2	3a	0	0	0.2224(8)	0.877(2)/0.123(2)		
Sb	3a	1/3	2/3	0.3823(8)	1		
O1	9b	0.641(1)	0.695(2)	0.2926(7)	1		
O2	9b	0.998(2)	0.622(2)	0.4555(5)	1		
$(Mn1-O_1) \times 3$	2.42(1)	$(Mn2-O_1)\times 3$	2.12(1)	$\Delta[Mn1O_6]$	5.5×10^{-3}		
$(Mn \ 1-O_2) \times 3$	2.09(1)	$(Mn2-O_2) \times 3$	2.31(1)	$\Delta[SbO_6]$	1.1×10^{-4}		
$(Sb-O_1) \times 3$	2.02(1)	$(Sc-O_1)\times 3$	2.04(1)	$\Delta[Mn2O_6]$	1.7×10^{-3}		
$(Sb-O_2) \times 3$	1.98(1)	$(Sc-O_2) \times 3$	2.18 (1)	$\Delta[ScO_6]$	1.1×10^{-4}		
<mn1-o1-mn2></mn1-o1-mn2>	121.0 (4)		<mn1-o2-mn2></mn1-o2-mn2>		118.4 (3)		
BVS(Mn1) b = 1.89	BVS (Sb) = 5.18		BVS (Mn2) = 2.01		BVS (Sc) = 2.93		

^a Fitting residuals: $R_{\rm p} = 3.04\%$, $R_{\rm wp} = 3.86\%$, $R_{\rm B} = 6.18\%$ and $R_{\rm F} = 5.04\%$, $^bV_i = \sum_j S_{ij}i = \exp(r_0 - r_{ij}/0.37)$. Values calculated using $r_{ij} = 1.79$ Å for Mn²⁺, 1.849 Å for Sc³⁺ and 1.942 Å for Sb⁵⁺. Polyhedral distortions calculated from $\Delta = 1/n \times \sum [(d_i - d_{\rm av})/d_{\rm av}]^2$.

The displacement of the different cations from their ideal sites with reference to the oxygen octahedra are labeled in Fig. 3(left). Mn cations show larger displacements in comparison with the d⁰ Sc or d¹⁰ Sb, and in accordance with the greater distortion found for their polyhedra (see Table 1).

The theoretical value of polarization along the z axis has been calculated from $P = \sum_{i} q_{i}u_{j}^{z}$, where q_{j} stands for the charge over a generic j ion and u_i^z for its displacement along the z axis, resulting in $P_{\text{expected}} = 28.3 \, \mu\text{C cm}^{-2}$. This estimate neglects the effects of Mn/Sc disorder. Attempts were made to measure the experimental polarization, using a modified Sawyer-Tower circuit.²³ Due to the low resistivity, sample polarization is masked by conducting effects and typical inconclusive P vs. E ellipse-shaped cycles were obtained.

Magnetic behaviour and magnetic structure. The magnetic susceptibility of NTO_Mn2ScSbO6, represented in Fig. 4a with its inverse, follows a Curie-Weiss behaviour above 75 K with a

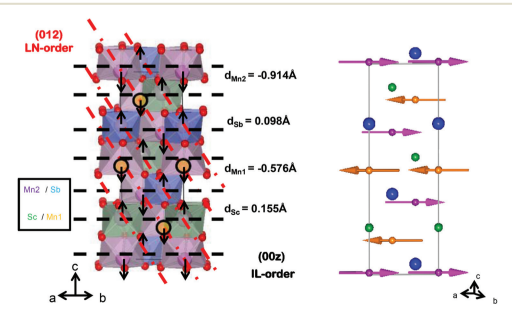


Fig. 3 Unit cell of the nuclear (left) and magnetic (right) structures of the NTO polymorph of Mn₂ScSbO₆. In both, Sc is depicted in green, Sb in blue and the two different sites for Mn cations are identified with purple and orange spheres. [MO₆] octahedral coordination can be observed in the left, where both Mn octahedra are purple to facilitate the visualization of LN-type order. (00z) and (012) layers showing the apparent IL and LN-type orders, respectively, are labeled as black and red dashed lines. The black arrows show the cationic displacements (d_M) along the c axis. Purple and orange arrows in the right part of the figure indicate the modulus and direction of Mn²⁺ spins located in Mn1 and Mn2 sites respectively.

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calculated effective moment of $\mu_{NTO} = 6.1 \ \mu_{B}$ per Mn²⁺ and $\theta_{\rm NTO}$ = -138 K. A ferro- or ferri-magnetic transition is observed at $T_{\rm C}$ = 42 K. There is no significant divergence between ZFC and FC branches, and field dependent magnetization measurements (inset in Fig. 4a and SF3† top) show that NTO_Mn₂ScSbO₆ is a very soft ferrimagnet with a saturated magnetization of 0.6 $\mu_{\rm B}$ per f.u. at 2 K.

From the structural point of view, Mn²⁺ cations are located in different octahedral sites within (00z) layers and are connected to each other through sharing corners. This allows cooperative AFM Mn-O-Mn superexchange interactions that are governed by <Mn1-O1-Mn2> = 121.0 (4)° and <Mn1-O2- $Mn2 > = 118.4 (3)^{\circ}$ angles.

The 4 K NPD data collected for NTO_Mn2ScSbO6 at D2B instrument are shown in Fig. 4b. The magnetic structure of this polymorph (Fig. 3 right) was determined from the fit to these data, where 27°-30° and 46°-48° angular ranges are excluded due to the presence of peaks arising from the sample holder. Additional magnetic neutron diffraction peaks are

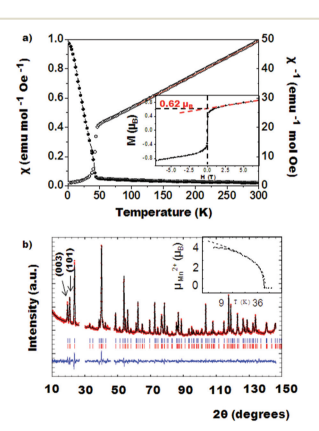


Fig. 4 (a) NTO_Mn₂ScSbO₆ magnetic susceptibility as a function of temperature under an applied magnetic field of 3 kOe (black circles). Reciprocal susceptibility (open circles) is indexed on the right axis and fitted (red line) to the Curie-Weiss law. Inset in (a) shows the magnetization loop measured at 2 K showing the saturated magnetization per formula unit. (b) Rietveld refinement of the magnetic structure from the 4 K NPD pattern. The main magnetic peaks (003) and (101) are indexed. Inset shows the temperature dependence of the magnetic moment of Mn^{2+} with a fit (dashed line) to a critical law $\mu(T) = \mu(0) \times [1 - (T/T_C)]^{\beta}$, as described in the text.

observed below $T_{\rm C}$ = 42 K, the most intense being (003) and (101), which can be indexed with the propagation vector k =[0 0 0]. Fits to the magnetic peaks confirm a FM coupling of Mn1 spins within 00z layers, where spins are confined to the ab plane, but antiferromagnetically coupled with the Mn2 cations of the adjacent layers. The refined moment per Mn²⁺ cation is 4.62 (3) $\mu_{\rm B}$. The spin order and Mn site occupancies observed by NPD reveal the origin of the ferromagnetic behavior of NTO_Mn2ScSbO6. Antiparallel Mn1 and Mn2 spins are symmetry-inequivalent and so a small uncompensated ferromagnetic moment is expected, but this is greatly enhanced by site-selective disorder as Mn1 sites are fully occupied by Mn2+ spins, but 12.3% of Mn2 sites are substituted by non-magnetic Sc^{3+} . The predicted uncompensated moment (12.3% \times 4.62 μ_{B} = 0.57 $\mu_{\rm B}$) is in excellent agreement with the observed magnetization of 0.6 $\mu_{\rm B}$. Hence, the ferromagnetic component arises from an unusual site-selective substitution of non-magnetic Sc³⁺ at just one of the two Mn²⁺ sites. A similar ferrimagnetism due to selective disorder has been recently observed in DPvtype La₃Ni₂SbO₉.²⁴ Magnetic diffraction from the ferromagnetic component in NTO_Mn2ScSbO6 is expected to be very weak and is not observed in the difference pattern between NPD data sets collected at 50 K and 3 K on the D1B instrument (SF4† lower panel).

The evolution of the magnetic moment below the Curie temperature (inset on Fig. 4b), was fitted to a critical law $\mu(T)$ = $\mu(0) \times [1 - (T/T_C)]^{\beta}$ in the $(T_C/2) < T < T_C$ temperature range between 21 K and 41 K. It led to the parameters $T_{\rm C}$ = 42.0 K and $\beta = 0.37$, consistent with 3D Heisenberg behaviour for which β = 0.36 is predicted.

Overall, NTO_Mn2ScSbO6 is notable as a potential multiferroic material, with ferroelectricity arising from the acentric cation order in the NTO-type arrangement, and ferrimagnetism from the uncompensated antiparallel order of Mn²⁺ spins enhanced by site-selective Mn/Sc disorder. This material seems to be a type I multiferroic as the two orders are decoupled. The magnetoelectric coupling between the electrical and magnetic polarizations may be small as they are orthogonal, being respectively parallel and perpendicular to the c-axis, although this could lead to an unusual switching mechanism as predicted for LiNbO3-type MnTiO3-II.6

(B) DPv_Mn₂ScSbO₆

Structural characterization. As in other related Mn₂BSbO₆ compounds (B = Fe, V, Cr, Ga, Al),²⁵ high pressure synthesis was needed to stabilize the small Mn2+ cation in the highly coordinated A site of the perovskite structure.

Fig. 5a (top) shows the Rietveld refinement of NPD data of DPv_Mn₂ScSbO₆ collected at room temperature. It was fitted using the Mn_2BSbO_6 (B = Fe, Cr)^{12,13} structure as the starting model and crystallizes in the $P2_1/n$ monoclinic space group with lattice parameters a = 5.2909 (3) Å, b = 5.4698 (3) Å, c =7.7349 (5) Å and β = 90.165 (6)°. The resulting crystallographic parameters, interatomic distances and bond angles are summarized on Table 2. Anti-site disorder was found between Sc and Mn, resulting in 9.1% of the A-site occupied by Sc³⁺. The

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a) (inc) (in

Fig. 5 (a) Rietveld refinements of room temperature (top) and 3 K (bottom) NPD patterns. 20.1% of the NTO polymorph is included as the secondary phase. First and second rows of Bragg positions in the lower patterns are the nuclear and (indexed) magnetic phases of the perovskite polymorph and the third and fourth rows are the nuclear and magnetic structures of the NTO_Mn₂ScSbO₆ secondary phase. The inset in the lower panel shows the temperature dependence of the magnetic moment of Mn²⁺ fitted (dashed line) to the critical law described in the text. (b) Nuclear and magnetic structures of DPv_Mn₂ScSbO₆. [ScO₆] octahedra are depicted in red and [SbO₆] in green; Mn²⁺ cations, are shown as purple spheres with arrows indicating their magnetic moments. (c) Direct and reciprocal susceptibility (dark and empty circles respectively) as a function of temperature.

100

200

Temperature (K)

300

secondary phase included in the refinement of DPv_Mn₂ScSbO₆ is the NTO polymorph formed at lower pressures during the synthesis reaction. A 20.1% of this phase has been found to crystallize within this high pressure compound. The presence of such a high proportion of NTO_Mn₂ScSbO₆ is due to the high stability of the NTO-type structure at high temperatures. The effect of this secondary phase on the magnetic behaviour of DPv Mn₂ScSbO₆ is discussed below.

Fig. 5b depicts the nuclear (and magnetic) structure of DPv_Mn_2ScSbO₆; Sb and Sc are six-fold coordinated and ordered in a rock-salt configuration. Mn cations occupy the highly distorted cuboctahedral voids. The tilt angle, calculated as $\Phi = (180^{\circ} - \theta)/2$ from <B-O-B'> bond angles (see Table 2) is $\Phi = 21.0$ (3)°. This value is indicative of the high degree of distortion attained in this polymorph. The polyhedral distortions (Δ in Table 2) are also abnormally large (Δ [ScO₆] = 2.9×10^{-4} and Δ [SbO₆] = 1.6×10^{-5}), close to those observed in other perovskites containing small-sized cations as Mn²⁺ and Sc³⁺ at the A site, as MnVO₃ or ScVO₃ with Φ = 18.02° and 26° and

The microstructure of this double perovskite is clearly observed in the HRTEM image recorded along the [010] zone axis (see SF2c†). The image and its corresponding FFT indicate the absence of extended defects and a good agreement with the calculated image is found. EELS experiments performed on the Mn L-edge in several crystals of DPv_Mn₂ScSbO₆ revealed an average oxidation state of +2 for Mn, in correspondence with BVS calculations.

Magnetic behaviour and magnetic structure. The ZFC and FC (black and red circles, respectively) molar magnetic susceptibilities for the DPv_Mn₂ScSbO₆ polymorph and the reciprocal susceptibility (open circles) as a function of temperature under an applied field of 3 kOe are depicted in Fig. 5c. It follows the Curie–Weiss behaviour with an additional 0.0579 emu⁻¹ mol Oe temperature independent paramagnetic (TIP) contribution

Table 2 Crystallographic parameters, selected interatomic distances (Å) and angles (°) for DPv_Mn₂ScSbO₆ from the room temperature NPD Rietveld refinement in the $P2_1/n$ space group^a

$a = 5.2909 (3) (Å), b = 5.4698 (3) (Å), c = 7.7349 (5) (Å), \beta = 90.165 (6) (°)$							
Atom	Site	x	у	Z	Occ		
Mn/Sc	4e	-0.009(3)	0.038(2)	0.749(4)	0.909(4)/0.091(4)		
Sc/Mn	2d	1/2	0	0	0.909(4)/0.091(4)		
Sb	2c	Õ	$\frac{1}{2}$	0	1		
01	4e	0.313(2)	0.321(2)	0.940(1)	1		
O2	4e	0.327(2)	0.307(2)	0.570(1)	1		
O3	4e	0.878(1)	0.427(1)	0.760(2)	1		
$(B-O_1) \times 2$	2.07(1)	$(Sb-O_1)\times 2$	1.982(9)				
$(B-O_2) \times 2$	2.10(1)	$(Sb-O_2) \times 2$	1.99(1)	$\Delta[BO_6]$	2.9×10^{-4}		
$(B-O_3) \times 2$	2.15(1)	$(Sb-O_3) \times 2$	2.00(1)	$\Delta[SbO_6]$	1.6×10^{-5}		
<a-o1-a></a-o1-a>	113 (1)	<a-o2-a></a-o2-a>	104 (1)	<a-o3-a></a-o3-a>	123 (1)		
<b-o1-b'></b-o1-b'>	140.9 (3)	<b-o2-b'></b-o2-b'>	137.1 (3)	<b-o3-b'></b-o3-b'>	136.8 (3)		
$BVS(Mn)^b$	1.7	BVS (Sc)	2.9	BVS (Sb)	5.3		

^a Fitting residuals: $R_{\rm p} = 2.00\%$, $R_{\rm wp} = 2.51\%$, $R_{\rm B} = 6.04\%$ and $R_{\rm F} = 4.36\%$. $^bV_i = \sum_j S_{ij}i = \exp(r_0 - r_{ij}/0.37)$. Values calculated using $r_{ij} = 1.79$ Å for Mn²⁺, 1.849 Å for Sc³⁺ and 1.942 Å for Sb⁵⁺. Polyhedral distortions calculated from $\Delta = 1/n \times \sum [(d_i - d_{\rm av})/d_{\rm av}]^2$.

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above 100 K. The Weiss temperature is $\theta_{DPv} = -94$ K and the effective paramagnetic moment $\mu_{DPv} = 5.5 \mu_B$ per Mn²⁺ is slightly lower than the expected value for $S = 5/2 \text{ Mn}^{2+}$ (5.92 $\mu_{\rm B}$). The magnetic susceptibility increases abruptly at ~42 K due to the presence of the NTO polymorph as a secondary phase. This obscures the antiferromagnetic transition at 22.3 K, determined below for this polymorph from NPD. The absence of magnetic cations neither in B nor in B' - sites yields the only possible magnetic interactions in this compound to be those operating between the nearest neighboring Mn²⁺ cations. The long Mn-Mn distance and the high coordination of the A - site, favors the development of cooperative superexchange interactions through oxygen anions. The nature and strength of these interactions is defined by the Mn-O-Mn angles, which take the values of 104 (1)°, 113 (1)° and 123 (1)°. According to the Goodenough-Kanamori rules²⁷ these angles rule the antiferromagnetic interactions among d⁵ cations, as observed in the magnetization curves (see SF3† bottom). From the Rietveld refinement, it has been found that 9.1% of A-sites are occupied by diamagnetic Sc3+ cations, resulting in a magnetic dilution effect on the overall Mn²⁺ sublattice.

The magnetic structure of DPv_Mn₂ScSbO₆ was determined by NPD using data from the D1B instrument. Additional magnetic neutron diffraction peaks are observed below 22 K.

Fig. 5a (bottom) shows the fit to the 3 K data. The magnetic reflections can be indexed with the $k = [0 \ 0 \ 0]$ propagation vector in the $P2_1/n$ space group. Irreducible representations and their basis vectors were obtained by using the program BASIREPS. Fig. 5b shows the resulting magnetic structure. It can be described as an antiparallel orientation of Mn²⁺ spins located in the ac-plane, with a total moment of 4.54 (1) $\mu_{\rm B}$ at 3 K. The deviation of the spins with respect to the c axis has been calculated to be $\gamma = 20.34$ (1)°, resulting in 1.58 (1) μ_B and 4.26 (1) $\mu_{\rm B}$ components along the a- and c-axes respectively. The inset in Fig. 5a shows the thermal evolution of the ordered magnetic moment, fitted to the critical law. The fitting results in T_N = 22.3 K and β = 0.37, where β agrees again with 3D Heisenberg antiferromagnetic behaviour.

IV. Conclusions

Our results demonstrate that two new phases of Mn₂ScSbO₆ are accessible by high pressure-high temperature synthesis and their structural and microstructural details and magnetic properties have been investigated.

Although cations are completely randomly located within the room pressure polymorph, frustrating any long range magnetic order, the synthesis of this oxide under high pressure conditions induces different cationic arrangements leading to interesting properties. The NTO-type moderate pressure modification crystallizes in the rhombohedral R3 polar space group, which allows a predicted room temperature polarization of 28.3 μC cm⁻². It has ferrimagnetic order below 42 K, with the spins lying in the ab plane. A ferromagnetic component of $0.6 \mu_{\rm B}$ has been determined to arise from unusual site-selective

Mn/Sc disorder. NTO_Mn2ScSbO6 is thus a potential multiferroic material. The high pressure phase has a double perovskite structure crystallizing in the $P2_1/n$ space group, which exhibits an antiferromagnetic order below 22.3 K. Its magnetic structure has antiparallel Mn^{2+} spins in the *ac* plane.

Acknowledgements

The authors thank MINECO and Comunidad de Madrid for foundation through projects MAT2013-44964 and S-2013/ MIT-12753, respectively. EPSRC and the Royal Society are also acknowledged for support, and Dr Gallardo-Amores and Dr Romero-de Paz for HPHT synthesis and magnetic measurements.

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