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Magnetic frustration in the high-pressure Mn₂MnTeO₆ (Mn₃TeO₆–II) double perovskite.

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A new double perovskite Mn_2MnTeO_6 has been obtained by high pressure phase transformation of a corundum-related precursor. It is antiferromagnetic below 36 K and develops a magnetic structure with magnetic moments of 4.8 μ_B and 3.8 μ_B for Mn^{2+} at the A and B sites respectively. This new polymorph accounts for a recently reported decrease in the bandgap of Mn_3TeO_6 under pressure that may lead to useful light-harvesting properties.

The ABO₃ perovskite structure has proven to be very versatile, since it can accommodate several cations at both A and B sites. The consequent accessibility to a wide range of magnetic and electrical properties have attracted much attention. Perovskites show increasing distortion with decreasing ionic radius of the A-site cation, down to tolerance factors t = $(r_A+r_O)/(\sqrt{2(r_B+r_O)})$ of 0.75, where r_A , r_B and r_O are the ionic radii of the A, B and O ions¹. However, the use of high pressures enables small cations such as Mn^{2+} (r = 0.96 Å)² to be accommodated at the A-sites, allowing complex electronic and magnetic interactions. For example Mn²⁺V⁴⁺O₃ perovskite shows the coexistence of localized $d^5 \ Mn^{2+}$ and itinerant d^1 electron V⁴⁺ cations³. Further variety is achieved in high pressure double perovskites (DPvs) Mn₂BB'O₆ with ordering of two cations on the B sites; Mn_2BSbO_6 (B = Sc, Cr, and Fe)⁴⁻⁶, Mn_2BReO_6 (B = Mn, Fe and Co)⁷⁻⁹ and $Mn_2(Fe_{0.8}Mo_{0.2})MoO_6$.¹⁰ Tellurium based oxides M₂MTeO₆ (M = Mn, Co and Ni) have structures based on the corundum arrangement at ambient pressure (see Supplementary Figure 1) and show a variety of complex magnetic orders. Mn₃TeO₆-I crystallises in the Mg₃TeO₆-type structure.¹¹ Below 24 K it shows the coexistence of an elliptical helix and a sinusoidal spin density wave, both being incommensurate with the crystal structure. This phase has shown a dielectric response below 21 K and so is a type-II multiferroic 12 . Co $_3$ TeO $_6$ has 5 independent Co sites, providing a rich magnetic phase diagram 13 and Ni $_3$ TeO $_6$ is a non-hysteretic colossal magnetoelectric material with a collinear antiferromagnetic (AFM) structure stabilised on stacked honeycomb layers 14 .

A recent study of light-harvesting properties of Mn_3TeO_6-I discovered an irreversible 39% bandgap reduction during insitu measurements under pressure¹⁵. We have explored the possibility of transforming ambient pressure Mn_3TeO_6-I phase into a recoverable double perovskite using high pressure and high temperature conditions, and we report here the new double perovskite Mn_2MnTeO_6 , which is likely to be responsible for the reported bandgap engineering.

 Mn_3TeO_6 -I, prepared at 1270 K by the solid state synthesis method of reference 11 was packed into a Pt capsule, pressed at 8 GPa and heated at 1173 K using a Walker-type module in a multianvil press. After 20 minutes, the temperature was quenched and the pressure was slowly released. Synchrotron X-ray diffraction (SXRD) powder data were collected at room temperature using the BL04 beamline at ALBA, Barcelona. Neutron powder diffraction (NPD) data were taken on 6 combined high pressure samples (\approx 120 mg) at 50 K, 1.5 K and several intermediate temperatures using the D1B beamline at the ILL, Grenoble. Zero-field cooled and field cooled (ZFC and FC) magnetic susceptibilities and a magnetisation-field loop at 2 K were measured using a PPMS Dynacool from Quantum Design

Preliminary laboratory X-ray diffraction evidenced the transition from $Mn_3TeO_6\text{-I}$ into a new double perovskite, $Mn_3TeO_6\text{-II}$. This was found to have a monoclinic $P2_1/n$ rocksalt B-site ordered DPv structure from Rieveld fitting of 300 K SXRD data (Fig. 1a). A few % of $\delta\text{-M}n_3O_4$ (CaMn $_2O_4\text{-type}$) and Pt from the capsule are included in the refinements. The SXRD results are summarised in Table 1 and the structure is shown in Fig. 1b.

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[†] Data that support the findings of this study have been deposited at https://datashare.is.ed.ac.uk/handle/10283/838.

 $[\]ddagger$ Electronic Supplementary Information (ESI) available: supporting figures and tables. See DOI: 10.1039/x0xx00000x

COMMUNICATION Journal Name

Antisite disorder between Mn_B and Te positions was checked but no mixing was found within error showing that Mn_3TeO_6 -II is fully cation-ordered. The refined structural model shows that the Mn_BO_6 octahedra are distorted while TeO_6 octahedra are more regular (Table 1). Octahedral tilts ~20° are comparable to those for other $Mn_2BB'O_6$ double perovskites and reflect the significant distortion in agreement with the small t-value (t = 0.8) for this high pressure polymorph.

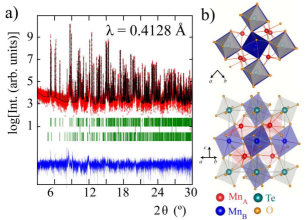


Figure 1. Rietveld fit (a) of the DPv structure of Mn_2MnTeO_6 (b) from 300 K SXRD data. Second and third rows of Bragg tick marks are minor amounts of Pt from the capsule (2.2(1) %) and δ -Mn₃O₄ (1.9(1) %). Table 1. Atomic positions and main interatomic distances and bond angles from the Rietveld fit of 300 K SXRD data. S.G. $P2_1/n$, a=5.29370(1) Å, b=5.45203(1) Å, c=7.80894(1) Å, $\beta=89.62514(5)$ 9.

Riso

Site	х	х		У	z	BISO	Occ	
						(Ų)		
Te (2a)	0		0		0	0.43(1)	0.5	
$Mn_B(2b)$	0.5		0.5		0	0.56(2)	0.5	
Mn_A (4e)	0.4887(2)		0.0475(1)		0.2397(1)	0.66(2)	1	
O1 (4e)	0.1677(8)		0.2784(7)		0.0808(6)	0.31(5)	1	
O2 (4e)	0.7106	0.7106(8)		45(8)	-0.0520(5)	0.31(5)	1	
O3 (4e)	0.8794	1(7)	0.9244(7)		0.2215(5)	0.31(5)	1	
	$d_{M-O} (\mathring{A})$ < $M-O-M>(0)$							
2x (Mn _B -C	2x (Mn _B -O1)		2.308(4)		∕ln _B -O1-Te	136.2(3)		
2x (Mn _B -O2)		2.13	2.133(5)		/ln _B -O2-Te	144.9(3)		
2x (Mn _B -O3)		2.30	2.308(4)		∕ln _B -O3-Te	137.0(3)	
<Mn _B $-$ O $>$		2.24	19(6)					
2x (Te-O1)		1.87	70(5) N		n _A -O1-Mn _B	96.5(1	L)	
2x (Te-O	2)	1.85	1.851(5)		n _A -O2-Mn _B	109.6(2)	
2x (Te-O	2x (Te-O3)		1.885(4)		n _A -O3-Mn _B	98.7(1)		
<te-o></te-o>		1.86	8(5)					
2x Mn _A -O1		2.188(5) 2.458(5)			ф	21.50		
3x Mn _A -O2		2.179(5)			•	19.70		
		2.648(5) 2.660(5)			θ			
2 14 02		2.178(5)			A[N4= O]	2.20*10-2		
2x Mn _A -O3		2.19	92(4)		$\Delta[Mn_BO_6]$	2.30*10 ⁻³		
<mn<sub>A-O</mn<sub>	<mn<sub>A-O></mn<sub>		58(4)		Δ[TeO ₆]	1.04*10-4		

Octahedral distortions calculated from $\Delta [MO_6]$ =(1/n)* $\Sigma (d_{i^-}d_{av})^2/d_{av}$. Tilt angles along [001] φ and [110] θ are calculated from (180-<M-O-M>)/2. R_p = 16.8%, R_{wp} = 21.9%, R_f = 5.34%, R_B = 7.50%, χ^2 = 2.22 %.

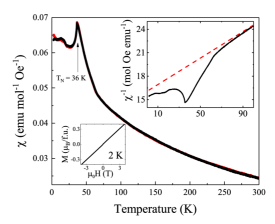


Figure 2. ZFC (black) and FC (red) magnetic susceptibility of Mn₂MnTeO₆ Insets show the reciprocal susceptibility with C-W from high temperature (red dashed line) (right) and the field dependent magnetisation at 2K (left).

ZFC and FC magnetic susceptibilities were measured under an external field of 500 Oe (Figure 2). They show a single antiferromagnetic transition at $T_{\rm N}$ = 36(0.5) K. A Curie-Weiss fit to the inverse susceptibility in the 100 - 300 K temperature range gives $\mu_{\rm eff}$ = 5.6 $\mu_{\rm B}/{\rm Mn^{2+}}$, in good agreement with the theoretical value of 5.92 $\mu_{\rm B}$ for spin only Mn²+ (S = 5/2). The large negative Weiss constant θ = -183(1) K indicates predominant antiferromagnetic interactions, and the linear dependence of magnetisation on applied field at 2 K (left inset in Figure 2), demonstrates the absence of a ferromagnetic component to the low temperature order. The upturn of the susceptibility observed below 25 K may be due to traces of the Mn³TeO₆-I phase (T_N = 24 K).

Rietveld fits to the NPD data collected at D1B at 50 K and 1.5 K are shown in Figure 3a. The large difference between neutron scattering lengths for Mn (-3.73 fm) and Te (5.80 fm) appear ideal to test for antisite disorder, thus the refinements confirmed the well-ordered structural model obtained from SXRD. Oxidation states estimated from BVS calculations on the 50 K NPD model are 2.1, 1.7 and 6.0 for Mn_A, Mn_B and Te sites respectively, demonstrating that the charge distribution is $Mn_2^{2+}Mn_2^{2+}Te^{6+}O_6$.

Prominent magnetic peaks observed below T_N can be indexed with the $k = [\frac{1}{2} \ 0 \ \frac{1}{2}]$ propagation vector. The magnetic symmetry analysis was performed using the BasIreps tool implemented in the Rietveld refinement program FullProf Suite¹⁶. The resulting irreducible representations (Ireps) are summarised in Table S1 in the ESI and all four simple possible models were tested. The best fit is when both A- and B-site Mn spins follow Irep Γ_4 and can be refined independently ($R_B = 6.60\%$, $R_f = 8.54\%$, $R_{Mag} = 10.1\%$, $\chi^2 = 1.233$ for T = 1.5 K). The 1.5 K - 50 K difference pattern containing only magnetic intensity is shown as an inset in the lower panel of Figure 3a and magnetic refinement details are shown in ESI and Table S2.

Journal Name COMMUNICATION

The magnetic structure of Mn₂MnTeO₆-II, depicted in Figure 3b, can be described as *ab* layers of Mn_A spins AFM coupled with Mn_B spins perpendicular to the *a* direction in the magnetic unit cell. These layers are separated by diamagnetic TeO₆ octahedra, stacked AFM along the *c* direction and with the spins confined in the *ac* plane. The 90° superexchange Mn_A-O-Mn_B d⁵-d⁵ interactions are assumed to be the dominant ones (see Table 1), and satisfy the Goodenough-Kanamori-Anderson rules (GKA).¹⁷ The refined ordered magnetic moments at 1.5 K converge to μ_A = 4.8(6) μ_B and μ_B = 3.9(1) μ_B for A- and B-site Mn²⁺ respectively. The refined value for Mn_A agrees well with the expected ordered magnetic moment of 5 μ_B for Mn²⁺, while the reduced value obtained for the Mn_B sublattice reflects some frustration present in the system, the later also supported by the ratio f = $|\theta|/T_N \sim 5$.

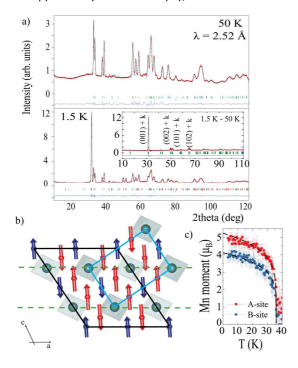


Figure 3. a) Rietveld fits of the nuclear and magnetic structures to the 50 K (top) and 1.5 K (bottom) NPD data. The right inset shows the fit of the magnetic model to the 1.5 K – 50 K difference pattern. b) ac plane of the magnetic structure of Mn_2MnTeO_6 with Mn_A and Mn_B spins as red and blue arrows respectively. Magnetic unit cell t is delimited with a black line and related to the crystallographic cyan one via the transformation matrix: [(1 0 -1), (0 -1 0), (-2 0 0)] with an origin shift of (0, 0, 1/2) and magnetic space group P_c2_3/n (14.80). Green dashed lines show the AFM planes described in the text. c) Thermal evolution of the Mn^{2+} magnetic moments and fit to the critical law (dashed black line) as described.

The magnetic model was used to fit the NPD scans at intermediate temperatures giving the evolution of the magnetic moments at both sites, as shown in Fig. 3c. Their fit by the critical law $\mu(T)=\mu(0)^*[1-(T/T_N)]^\beta$ in the $(T_N\ /\ 2) < T < T_N$ temperature range led to the parameters $T_N=35.8(1)$ K and $\beta \approx 0.19(1)$, confirming a simultaneous order of both sublattices, in agreement with magnetisation measurements. The small β value suggests a reduced dimensionality, being close to the

expected value of 0.23 for a 2D-XY model 18 and agreeing with the refined magnetic structure with spin directions confined into the ac plane.

In this context, the comparison with related high-pressure double perovskites Mn₂BB'O₆ is remarkable. Table 2 summarises the main features of their magnetic behaviours, which suggests the essential role of the diamagnetic B' cation on the frustrated B-site magnetic moments of these compounds as detailed below. Mn₂MnTeO₆, Mn₂MnReO₆ and Mn₂CoReO₆ are the only Mn₂BB'O₆ compounds reported to stabilise a DPv structure with divalent B and hexavalent B' cations. The highest magnetic frustration is observed in Mn₂MnTeO₆ and is released in Mn₂BReO₆ by the presence of the paramagnetic cation Re⁶⁺, which enables further superexchange pathways within the B sublattice and consequently increases T_N. Both Mn₂CoReO₆ and Mn₂MnReO₆ become ordered around 100 K into almost perpendicular AFM A and B sublattices with $k = [\frac{1}{2} \frac{1}{2} 0]$ and with reduced moments in B.9,7 The isoelectronic Mn2+ cations at A and B sites in Mn₂MnReO₆ give rise to magnetoelastic effects that prompt Re magnetic ordering at lower temperatures than the Mn (f = 1.4). However, in Mn_2CoReO_6 with Co^{2+} in a high spin state, some GKA paths are blocked and it recovers some frustration increasing f up to 3.6. In Mn₂FeReO₆ the strong coupling between Fe3+ (d5) and Re5+ (d2) leads to a ferrimagnetic ordering on the B/B'-sites at high temperatures into a simple $k = [0\ 0\ 0]$ magnetic structure (f = 0.9).8 Likewise, $Mn_2Fe_{0.8}Mo_{0.2}MoO_6$ shows almost no frustration (f = 0.3) due to the B-sublattice disordering of cations and the presence of Mo in a 4+ and 5+ oxidation states.

Our comparison has to include compounds of type Mn₂BSbO₆ (B = Sc, Cr and Fe) due to the presence of a diamagnetic B' cation as in Mn₂MnTeO₆. All of them achieve high degrees of magnetic frustration with transitions lower than 60 K. B = Sc, where only Mn²⁺ at the A sites are paramagnetic, has a strong frustration (f = 4.3) with AFM order at 22 K into a simple collinear magnetic structure with k = [0 0 0].4 B = Cr3+ shows the closest magnetic structure to that of Mn₂MnTeO₆, with k = [½ 0 ½] and AFM d5-d3 A-O-B superexchange interactions at $90^{\circ}.^{5Erreur\,!~Signet~non~d\acute{e}fini.}$ The weaker nature of these interactions compared to the d5-d5 Mn2+A-O-Mn2+B of Mn₂MnTeO₆, justifies the slightly lower frustration of Mn₂CrSbO₆. Finally, Mn₂FeSbO₆ is isoelectronic to Mn₂MnTeO₆ with both Mn²⁺ and Fe³⁺ d⁵ cations. Mn₂FeSbO₆ presents an incommensurate helical magnetic structure with k = [0 0.426 0].6 The origin of this difference is that the exchange integral for Fe3+ is much larger than that for Mn2+,19 inducing strong competitions between AFM d5-d5 A-O-B and A-O-A interactions, thus frustrating the compound into a helical magnetic structure. In the case of Mn²⁺ the larger difference between these interactions, prevents the rotation of the spins and the release of the strong magnetic frustration, see Supporting Information for the interplays between A-A, B-B and A-B with frustrated topologies.

Finally, Mn_3TeO_6 has been recently reported as a promising light-harvesting material due to an irreversible pressure-

COMMUNICATION Journal Name

induced phase transition occurring at 16-18 GPa, where an energy bandgap reduction of $\Delta E_{gap}\approx 1$ eV is observed. ^15 Our experimental plots of the Kubelka-Munk function of the UV-VIS reflectance spectra in Figure 4 show a $\Delta E_{gap}\approx 0.8$ eV reduction from the Mn_3TeO_6 precursor (E_gap = 2.6 eV) to the Mn_2MnTeO_6 perovskite (E_gap = 1.8 eV) and we conclude that this is very probably the phase observed in the high pressure measurements. The change in connectivity from edge- to corner-shared octahedra between both polymorphs explains the decrease in the bandgap in the double perovskite.

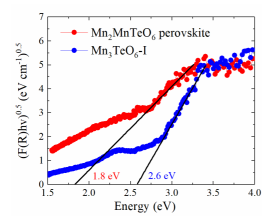


Figure 4. Kubelka-Munk plots (for an indirect bandgap) - for Mn₃TeO₆-l and the Mn₂MnTeO₆ high-pressure double perovskite polymorph, both measured at ambient pressure and temperature.

Table 2. Main features of the magnetic behaviour for $Mn_2BB'O_6$ double perovskites, including, electronic configuration of B and B' cations, transition temperature, Weiss constant (0), frustration index (f = $|\theta|/T_N$) and propagation vector (k).

В	B'	t _{2g}	e _g	d _{B′}	T _N /T _C (K)	θ	f	k
Sc	Sb	0	0	10	22	94	4.3	0 0 04
Mn	Te	3	2	10	36	183	5.1	½ 0 ½*
Cr	Sb	3	0	10	55	220	4.0	½ 0 ½ ⁵
Fe	Sb	3	2	10	60	180	3.0	$0 k_y 0^6$
Co	Re	5	2	1	94	26	3.6	½ ½ 0 ⁹
Mn	Re	3	2	1	109	147	1.4	½ ½ 0 ⁷
$Fe_{0.8}Mo_{0.2}$	Mo	3	2	1	194	50	0.3	$0\ 0\ 0^{10}$
Fe	Re	3	2	2	520	452	0.9	0 0 08

*This work

In conclusion, a new DPv Mn_2MnTeO_6 has been prepared by high-pressure high-temperature transformation of the rhombohedral polymorph. Due to the large charge and size difference between Mn^{2+} and Te^{6+} , Mn_2MnTeO_6 shows a fully ordered B/B' rock-salt type DPv structure. It is AFM below $T_N=36$ K and it develops a complex magnetic structure with ordered magnetic moments of 4.8 (1) μ_B and 3.8 (1) μ_B at the A and B sites respectively. The magnetic structure describes AFM ab layers of A- and B-site spins, AFM coupled along the c axis, with the spins confined in the ac plane. Mn_2MnTeO_6 shows the highest frustration index among the A-site manganites with

DPv structure. Replacement by non-magnetic B' cations proves to be a way to enhance the frustration and may be used as tuning parameter in partial B' substitutions. Formation of this DPv polymorph accounts for the previously reported bandgap reduction under pressure.

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Conflicts of interest. There are no conflicts to declare.

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