Phase stability of the layered oxide, Ca$_2$Mn$_3$O$_8$; probing interlayer shearing at high pressure†

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We have performed high-pressure neutron diffraction studies on the layered oxide, Ca$_2$Mn$_3$O$_8$. Studies up to approximately 6 GPa at temperatures of 120 and 290 K demonstrate that there are no structural phase transitions within this pressure range. Fits of the unit-cell volume to a Birch-Murnaghan equation of state gives values for the bulk modulus of 137(2) GPa and 130(2) GPa at temperatures of 290 K and 120 K respectively possibly suggesting that Ca$_2$Mn$_3$O$_8$ is more compressible at lower temperature. Furthermore, compression along the principal axes are anisotropic on the local scale. Comparison of individual bond lengths and bond angle environments further demonstrate that compression is complex and likely results in a shearing of the layers.

1 Introduction

Layered metal oxides continue to attract extensive attention due to their large compositional flexibility and wide variety of potential applications. For example, applications in renewable energy$^{1-7}$ and catalysis$^{8-10}$ have been reported. The prototypical layered transition metal materials are the delafossite family typified by the formulae ABO$_2$ (where A is a monovalent cation such as Na$^+$ or Cu$^+$ and B is a trivalent transition metal such as Fe$^{3+}$ or Mn$^{3+}$). The structure can be described by BO$_6$ layers with the A$^+$ cations situated between these layers. When B is a magnetic ion with unpaired electrons the triangular connectivity between these species often results in the realisation of exotic magnetic states.$^{11-16}$

More recently, interest is growing in the layered transition metal oxide, Ca$_2$Mn$_3$O$_8$, first reported by Horowitz et al. in 1978.$^{17}$ They showed that Ca$_2$Mn$_3$O$_8$ crystallises with a monoclinic $C2/m$ layered structure described by Mn$_6$O$_8^{4-}$ sheets, formed from edge sharing MnO$_6$ octahedra and separated by trigonal bipyramidal CaO$_6$ sites as shown in Fig. 1.$^{17-19}$ In contrast with delafossites, these materials have a nominal formulae of Ca$_{0.5}$Mn$_{0.75}$O$_2$ as a result the B-site cation layers are incomplete (when compared with the triangular lattice exhibited by delafossite materials) with a quarter of the triangular lattice Mn$^{4+}$ octahedral) sites vacant. This leads to a network of ordered voids within the 2D layers realising a ‘bow-tie’ connectivity of the Mn$^{4+}$ ions (Fig. 1).$^{17-19}$ The location of this void alternates between layers giving rise to ABAB stacking in the lattice $a$-direction.$^{17-19}$ White et al. later characterised the magnetic behaviour of the material as antiferromagnetic from SQUID magnetometry.$^{20}$ As with delafossite materials, Ca$_2$Mn$_3$O$_8$ has been investigated for the catalytic splitting of water$^{21-27}$ and as a potential battery electrode.$^{28}$ However, the understanding of the structure–property relationships in Ca$_2$Mn$_3$O$_8$ has largely been limited by the low crystallinity of the as synthesised materials.$^{29}$

Previously, we have demonstrated that synthetic route can have a large effect on the morphology observed.$^{29}$ Solid-state methods give large spherical particles comprised of small crystallites, sol–gel methods produce nano-sized particles and hydrothermal methods yield flake-like geometries with large surface areas.$^{29}$ However, all synthetic routes led to materials with low crystallinity making further analysis difficult. Recently, we developed a molten salt synthetic route which allowed us to synthesise highly crystalline Ca$_2$Mn$_3$O$_8$ bulk materials.$^{30}$ The high crystallinity of these materials allowed us to perform more comprehensive studies of the thermal stability and magnetic behaviour of Ca$_2$Mn$_3$O$_8$.$^{30,31}$ Low temperature studies (1.8–300 K) showed that Ca$_2$Mn$_3$O$_8$ orders antiferromagnetically with an $\uparrow \uparrow \downarrow \downarrow$ arrangement of the magnetic spins at a Néel temperature ($T_N$) of approximately 58 K.$^{30}$ Additionally, an invariant behaviour below 130 K is seen in the lattice parameter, $b$ and the monoclinic $\beta$ angle. This occurs at the same temperature as a deviation from Curie–Weiss behaviour.
observed in magnetic susceptibility measurements and is consistent with short-range spin correlations. Data collected between 300 and 1273 K show that the monoclinic structure is remarkably stable up to a temperature of approximately 1200 K before it decomposes into the perovskite, CaMnO$_3$ and marokite, CaMn$_2$O$_4$ phases. Moreover, electrical measurements confirm that Ca$_2$Mn$_3$O$_8$ is an electronic conductor in the temperature range, 400–700 K with an activation energy of approximately 0.50(1) eV.31

Given the interesting nature of Ca$_2$Mn$_3$O$_8$ and the phase stability of the C2/m structure as a function of temperature it is of importance to determine the distortion of this structure on the application of hydrostatic pressure. Applied pressure is an effective way to change co-ordination environments and tune property behaviour driving very different distortion behaviour upon increasing pressure. For example, in the perovskite family of materials there are examples of both decreasing and increasing structural distortions which are controlled by the relative compressibilities of the two types of polyhedra present within the structure. A number of compressibility studies (both experimental and computational) have been performed on CuFeO$_2$ and related rhombohedral (R3m) delafossites. Structural studies show that the compression of the lattice is anisotropic with the compressibility within the lattice c direction (between the layers) being far larger than that of the lattice a direction (within the layers). In CuFeO$_2$ the R3m structure has been shown to be stable up to pressure of approximately 18 GPa before transformation to a monoclinic, C2/c phase. At approximately 23 GPa there is a second partial transformation of the structure to a trigonal P3m phase resulting from an interatomic valence change from Cu$^{2+}$/Fe$^{3+}$ sites in the low pressure phases to Cu$^{2+}$/Fe$^{3+}$ above 23 GPa. In contrast in CuGaO$_2$ and CuAlO$_2$ the R3m structure is stable up to a pressures of 28 GPa and 35 GPa respectively. Above these pressures there is an irreversible transition (in both CuGaO$_2$ and CuAlO$_2$) to an unidentified phase which seems to be driven primarily by the Cu site. The pressure dependent magnetism has also been extensively studied in CuFeO$_2$. At ambient pressure CuFeO$_2$ exhibits two distinct antiferromagnetically ordered states. The first magnetic transition occurs At 14 K ($T_N1$) with the spins ordering with an incommensurately modulated collinear spin density wave. Below 11 K ($T_N2$) a second magnetic phase transition occurs with the magnetic spins now adopting a collinear commensurate four sub-lattice (4SL) $\uparrow \uparrow \downarrow \downarrow$ ordering. At $T_N2$ there is also a lowering of the symmetry from rhombohedral to monoclinic. With applied pressure the transition temperature of the spin density wave ($T_N1$) increases. In contrast the behaviour below $T_N2$ is far more complex. With increasing pressure the transition temperature falls slightly and a number of pressure induced magnetic phase transitions occur. Firstly, a transition to an incommensurately modulated proper screw ordered state is observed at approximately 3 GPa. This is followed by a transition to an incommensurate state which has signatures of both colloidal and proper screw configurations at approximately 4 GPa. Finally, at 6 GPa only the high temperature incommensurate spin density wave is observed (i.e. there is only a single magnetically ordered state). More recently, more complex magnetic correlations as a function of both applied pressure and magnetic field have been reported.

In this paper we have explored the pressure–temperature phase diagram for Ca$_2$Mn$_3$O$_8$ using neutron powder diffraction. We demonstrate that the C2/m structure is shown to be robust up to pressures of approximately 6 GPa at temperatures of both 290 and 120 K. Compression behaviour is shown to be complex and largely results from a shearing of the layers.

2 Materials and method

2.1 Sample synthesis and characterisation

Ca$_2$Mn$_3$O$_8$ was synthesised using a molten salt method as previously reported by us. Briefly, stoichiometric ratios of
CaCO₃ (Sigma Aldrich, ≥ 99%), MnCO₃ (Sigma Aldrich, ≥ 99%) and a eutectic mixture of KCl/NaCl (350% w/w) were mixed and reacted for 48 hours at 973 K followed by a second heating cycle of 24 hours at 973 K. After both heating cycles, the material was quenched to room temperature. Between heating cycles the material was washed and dried under ambient conditions and an additional 350% (w/w) eutectic mixture of KCl/NaCl was added for the second heating.

Phase purity was confirmed using a Rigaku Miniflex 600 powder X-ray diffractometer (40 kV and 15 mA, with λ = 1.5406 Å) over a 2θ range of 10° to 70° (data not shown here).

### 2.3 High-pressure diffraction

Neutron powder diffraction (NPD) data were collected under applied pressures of 0–6 GPa at 290 K (room temperature) and 0–5 GPa at 120 K, using the PEARL diffractometer at the ISIS Neutron and Muon Facility (UK). A powdered sample of Ca₂Mn₃O₈ was loaded into toroidal profile anvils machined from zirconia toughened alumina, encapsulated in a null scattering TiZr gasket and sealed in the Paris-Edinburgh press with an applied load of 6 tonnes. Time-of-flight neutron powder diffraction patterns were collected in ~1 GPa increments for ~4 hours each. Data were focused, normalised and intensity corrected for anvil attenuation using in-house software. Rietveld refinements were performed using the GSAS suite of programs (full details are provided in the ESI†). The data collection at 120 K was performed in the same manner described above except we used a modified Paris-Edinburgh press which was cooled with liquid N₂ cooling rings and is described in detail elsewhere. In both experiments a small piece of lead was included in the gasket chamber and used as a pressure marker, and a 4:1 (by volume) mix of perdeuterated methanol: ethanol was used to ensure compression was made under hydrostatic conditions.

### 3 Results and discussion

Fig. 2 shows the room temperature time-of-flight diffraction pattern of Ca₂Mn₃O₈ as collected on the PEARL instrument at the sealing load of 6 tonnes in the Paris-Edinburgh press. The associated Rietveld refinement is also shown. Table 1 shows the results of the Rietveld refinement compared to those published previously. The Rietveld fit to the data clearly indicates the phase, Pb pressure marker, the Al₂O₃ and ZrO₂ phases of the anvils.

#### Table 1

| Parameter | This study | Previous study\(^{18}\) |
|-----------|------------|------------------------|
| a (Å)     | 11.021(6)  | 11.014(4)              |
| b (Å)     | 5.846(3)   | 5.851(3)               |
| c (Å)     | 4.9428(3)  | 4.942(2)               |
| β (°)     | 109.785(4) | 109.73(5)              |
| Unit cell volume (Å³) | 299.707(3) | 299.78 |
| Ca x      | 0.722(6)   | 0.72442(3)             |
| Ca z      | 0.6613(12) | 0.66593(7)             |
| Mn(2) y   | 0.25100(15)| 0.25914(4)             |
| O(1) x    | 0.0980(4)  | 0.10059(7)             |
| O(1) y    | 0.2220(6)  | 0.22158(15)            |
| O(1) z    | 0.3894(8)  | 0.39171(17)            |
| O(2) x    | 0.3955(6)  | 0.59726(11)            |
| O(2) z    | 0.08918(12)| 0.90202(25)            |
| O(3) x    | 0.6042(3)  | 0.60385(11)            |
| O(3) z    | 0.9585(10)| 0.96258(26)            |
| w₀        | 2.43       | 3                      |
| χ²        | 1.027      | —                      |

Fig. 2 Rietveld refinement of the room temperature data collected for Ca₂Mn₃O₈ as loaded in the Paris-Edinburgh high pressure cell. The black circles represent the data collected, the red line the calculated model and the blue line the difference between the two. The vertical tick marks indicate the predicted peak positions for (top to bottom) the Ca₂Mn₃O₈ phase, Pb pressure marker, the Al₂O₃ and ZrO₂ phases of the anvils.

CaCO₃ (Sigma Aldrich, ≥ 99%), MnCO₃ (Sigma Aldrich, ≥ 99%) and a eutectic mixture of KCl/NaCl (350% w/w) were mixed and reacted for 48 hours at 973 K followed by a second heating cycle of 24 hours at 973 K. After both heating cycles, the material was quenched to room temperature. Between heating cycles the material was washed and dried under ambient conditions and an additional 350% (w/w) eutectic mixture of KCl/NaCl was added for the second heating.

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Upon increasing pressure at temperatures of both 290 and 120 K there is no change in the diffraction pattern beyond that expected with increasing pressure. Rietveld refinement of the structural model permits the lattice parameters of the monoclinic cell to be determined and with increasing pressure they show a continuous decrease as shown in Fig. 3. We find no evidence for a change in symmetry (at either 120 or 290 K) and thus conclude that the C₂/m crystal symmetry is maintained up to the highest pressure studied (~ 6 GPa) confirming the robust nature of the C₂/m phase. This is consistent with studies on related delafossite materials which are also seen to be stable over this pressure range.
in the behaviour of the monoclinic unit-cell angle ($\beta$). The beta angle decreases from initial values of 109.785(4)° and 109.745(4)° at 290 and 120 K respectively at ambient pressure to 109.720(5)° and 109.671(13)° at 290 and 120 K at pressures of 5.6 GPa and 4.6 GPa respectively possibly suggesting that the lattice is more compressible at low temperature. Overall, the structure is becoming less distorted with increasing pressure as suggested by the decrease in monoclinic cell angle ($\beta$) consistent with what has been observed in other layered delafossite materials.40

Fig. 4 shows the variation of the unit-cell volume with increasing pressure at 290 and 120 K. At both temperatures the cell volume decreases smoothly with increasing pressure indicating no structural changes consistent with lattice parameter behaviour. The variation of the volume has been fitted with a Birch-Murnaghan equation of state. At 290 K the determined bulk modulus ($B_0$) is 136.8(20) GPa with a pressure derivative ($B'$) of 4.5(5) (Table 2). Such a bulk modulus is comparable with other layered oxides such as CuFeO$_2$ (156 GPa and 2.6), PbCoO$_2$ (225 GPa and 0.7) and CuGaO$_2$ (202 GPa and 3.9).40 In Ca$_2$Mn$_3$O$_8$ there is a small decrease in the bulk modulus upon cooling to 120 K (Table 2).

We have established that upon compression at 290 and 120 K there is no change in crystal symmetry of Ca$_2$Mn$_3$O$_8$ with increasing pressure. However, compression behaviour can only be understood directly from unit cell dimensions when they are related to short range magnetic correlations which are known to propagate at temperatures below approximately 130 K at ambient pressure.30 In order to understand this further the behaviour of the individual bond lengths and polyhedra need to be understood upon compression.
Fig. 6 shows the variation in the average Mn–O bond lengths in the two crystallographically distinct MnO₆ octahedra. Both octahedra (Mn(1) and Mn(2)) display a decrease in the average bond length and individually the bond lengths within the octahedra show the same behaviour (see ESI†). The average Mn(2)O₆ bond lengths show a greater compressibility than those of the Mn(1)O₆ (Fig. 6). This behaviour is also reflected in the rate of change of the polyhedral volume with pressure (Fig. 6); the volume of the Mn(1)O₆ polyhedra decreases at a lower rate compared to that of the larger Mn(2)O₆ polyhedra. However, for both polyhedra no measurable change in distortion is seen. Within the Mn(1)O₆ octahedra there is a clear contraction of the Mn(1)–O(1) bond lengths which lie in the a–b plane (see ESI†). In contrast for the most part there is little change in the Mn(1)–O(2) bond lengths, which predominantly lie in the c-lattice plane within this octahedral unit (Fig. 1). There is a decrease in the Mn(2)–O(1) bond length with increasing pressure, however, we see very little change in the Mn(2)–O(3) bond lengths (see ESI†). We note that Mn(2)–O(1) and Mn(2)–O(3) bonds predominantly lie in the lattice a–b plane. Likewise, we see a change in the Mn(2)–O(2) bond lengths (lying predominantly along the c-lattice direction) with increasing pressure. There is very little change in the Mn–O–Mn bond angles, suggesting there is no rotation of the MnO₆ octahedra or buckling of the layers. Additionally, whilst both octahedral sites contribute to lattice contraction in the a and b directions only the Mn(2)O₆ octahedra drives the contraction of the layers in the c-lattice direction. This is perhaps not surprising given that the MnO₆ octahedra are edge-shared within the layers consistent with the relatively small change in dimensions seen in the c lattice parameter when compared to the a and b lattice parameters.

As described previously, in Ca₂Mn₃O₈, the calcium cation is sited between the MnO₆ layers within a six coordinate trigonal bipyramidal site as shown in Fig. 1. The contraction of these bonds is a little more complex with all three bond lengths (Ca–O(1), Ca–O(2) and Ca–O(3)) contributing to lattice contraction in all three crystallographic directions. The average Ca–O bond length (shown in Fig. 6) decreases with increasing pressure. The rate of decrease is greater than the Mn–O bond contraction. Also Fig. 6 shows a significant decrease in CaO₆ polyhedral volume with increasing pressure.

Looking at the individual Ca–O bonds within the polyhedra, there is only a small change in the Ca–O(1) bond length with increasing pressure whilst both the Ca–O(2) and Ca–O(3) bond lengths show a clear contraction. This would seem to suggest...
that the CaO₆ polyhedra become increasingly distorted with increasing pressure. Previous work using the difference in the absolute bond valence sum (|ΔV|) has suggested that a value of >0.1 is a result of strain within a polyhedron.³² |ΔV| for the CaO₆ polyhedra shows a clear increase in value with increasing pressure for which R₀ = 1.967 Å, B = 0.37 and formal expected charge of the calcium ion as 2+. The individual bond distances within the octahedra compress at different rates as described in the ESI,† and gives rise to an increase in geometric distortion of the octahedra as suggested by |ΔV| (Fig. 6). Indeed for the CaO₆ polyhedra the initial value is 0.2 and this doubles upon application of a pressure of ~6 GPa. This is in contrast with the behaviour of the MnO₆ polyhedra where the Mn(2)O₆ shows no real strain or changes with increasing pressure and only the Mn(1)O₆ polyhedra show an increase in strain suggested by the bond valence difference from 0.28 to 0.4 at 5.6 GPa (for both octahedra R₀ = 1.753 Å, B = 0.37) with a formal charge on the manganese ion of 4+. Again within each individual octahedra the individual Mn–O bonds compress at different rates and this leads to the difference in overall behaviour suggested by |ΔV|.

In contrast with the Mn–O–Mn bond angles, there are systematic changes in the bond angles associated with the calcium site (as shown in the ESI†). The Ca–O(1)–Ca bond angle shows a small increase with increasing pressure. This effectively acts to pull the layers closer together as shown in Fig. 7. The interlayer stacking in Ca₂Mn₃O₈ is ABAB with the intralayer voids formed within the MnO₆ framework shifted with respect to each other in adjacent layers.¹⁷–²⁰,²⁹–³¹ This means the Ca–O–Mn bond environments are different above and below the Ca²⁺ cation as shown in Fig. 7. These environments also alternate as you move between Ca²⁺ cations within the layer. The Ca–O(1)–Mn₁ and Ca–O(1)–Mn(2) bond angles are each described by two different values depending on if the angle describes bond angles linked to the ‘top’ or ‘bottom’ layer (as is depicted in the arrangement in Fig. 7). In fact all bond angles associated with the top layer (Ca–O(1)–Mn₁, Ca–O(1)–Mn(2) and Ca–O(3)–Mn(2)) all show little change with increasing pressure. In contrast those bond angles associated with the bottom layer show consistent changes with increasing pressure. In addition to the decreasing bond angle trends described above for Ca–O(1)–Mn(1) and Ca–O(1)–Mn(2), the Ca–O(2)–Mn(2) and Ca–O(2)–Mn(1) (non-bonding) bond angles both subtly increase with increasing pressure. This suggests that a shearing (consistent with the decrease of the β angle) of the layers contributes to the overall contraction in the a-lattice direction. This is consistent with the larger contractions seen in the interlayer dimensions in the delafossites (in this case the c-lattice direction).⁴⁰–⁴³,⁴⁵ From these data we can also suggest that the shearing effect occurs uniformly across the material, as we see no evidence to support buckling or superstructure formation. However, considering the ‘average’ nature of these experiments we cannot rule out local or modulation effects. Further work is necessary to elucidate the local behaviours in these materials.

### 4 Conclusions

In summary we have investigated the temperature-pressure phase diagram of the layered oxide, Ca₂Mn₃O₈ at temperatures of 290 K and 120 K and pressures up to 6 GPa. These studies demonstrate that there is a smooth decrease in the lattice parameters and unit cell volume with increasing pressure. Compression of the structure is largely anisotropic and predominantly occurs in the interlayer directions as opposed to the intralayer directions consistent with data reported for the compression of similar layered oxides.⁴⁰–⁴⁵ On the local scale compression behaviour is far more complex. Bond lengths and bond angle values suggest that there is little tilting or strain associated with the MnO₆ octahedra. In contrast the CaO₆ trigonal bipyramidal site becomes increasingly strained with increasing pressure. Moreover, contraction appears to be driven by one side of the CaO₆ polyhedra with the other side remaining largely unchanged. The nature of the structure means that this behaviour alternates between neighbouring Ca²⁺ ions which results in a shearing of the layers. These studies provide insight into the compression behaviour of Ca₂Mn₃O₈ which may be useful for optimising dopant (chemical pressure) studies, as well as providing a deeper understanding which will allow optimisation of this material for application.

### Conflicts of interest

There are no conflicts to declare.
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