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Structural refinement of Ca₃Mn₂O₇ using electron diffraction

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Abstract

The digital large-angle convergent-beam electron diffraction (D-LACBED) technique is applied to $Ca_3Mn_2O_7$ for a range of temperatures. We use Bloch-wave simulations to examine the effects that changes in different parameters have on the intensity in D-LACBED patterns and refine atomic coordinates, thermal atomic displacement parameters and apparent occupancy to achieve a good fit between simulation and experiment. The sensitivity of the technique to subtle changes in structure is demonstrated. Refined structures are in good agreement with previous determinations of $Ca_3Mn_2O_7$ and show the decay of anti-phase oxygen octahedral tilts perpendicular to the c-axis of the the $A2_1am$ unit cell with increasing temperature, as well as the robustness of oxygen octahedral tilts about the c-axis up to $\sim 400^{\circ}C$. The technique samples only

the zero-order Laue zone and is therefore insensitive to atom displacements along the electron beam direction. For this reason it is not possible to distinguish between inphase and anti-phase oxygen octahedral tilting about the c-axis using the [110] data collected in this study.

1. Introduction

Convergent beam electron diffraction (CBED) has a long history of application to symmetry determination (Buxton et al., 1976) and the accurate measurement of individual structure factors (Spence et al., 1989; Zuo, 1998; Zuo & Spence, 2017). However, despite its demonstrated sensitivity to many interesting and useful parameters, it is rarely used for structural refinement (Ogata et al., 2004). Recently, we have demonstrated that atomic coordinates can be extracted from 'digital' large angle convergent beam electron diffraction (D-LACBED) patterns (Beanland et al., 2013) with sub-picometre precision and accuracy using Al₂O₃ as a model material (Hubert et al., 2019). D-LACBED patterns, formed by combining hundreds or thousands of CBED patterns each with a slightly different incident beam orientation, have the advantage of a much larger angular range and many more data points than conventional CBED. They also can be obtained from a region the size of the convergent beam and a single measurement accesses many diffracted beams, unlike conventional LACBED (Tanaka et al., 1980). The interference between different Bloch waves as they propagate through a crystalline specimen produces regions in D-LACBED patterns that are particularly sensitive to small changes in phase and give a unique fingerprint for different parameters. Here, we apply the D-LACBED technique to the Ruddlesden-Popper phase Ca₃Mn₂O₇, which has attracted interest as a multiferroic (Benedek & Fennie, 2011; Fawcett et al., 1998) with the emergence of ferroelectric-ferromagnetic coupling occurring through the coupling of two non-polar modes with different symmetries. This effect has become known as hybrid improper ferroelectricity (Benedek & Fennie, 2011) and is currently being studied in a variety of related materials (Senn et al., 2015; Bansal et al., 2019; Xu et al., 2019; Chen et al., 2020)].

Ruddlesden-Popper phases in general have the formula $A_{n+1}B_nO_{3n+1}$ and a structure that consists of a rock-salt AO bilayer every $n ABO_3$ perovskite unit cells along [001]. In $Ca_3Mn_2O_7$ n=2 and the oxygen atoms form corner-sharing MnO_6 octahedra. The prototype (undistorted) structure has space group I4/mmm (No. 139) at high temperature. Just as in ABO_3 perovskites, these octahedra can tilt and distort, here lowering the symmetry from the prototype structure and giving rise to macroscopic ferroic behaviour. An orthorhombic structure for $La_{2-2x}Ca_{1+2x}Mn_2O_7$ (0.8 < x < 1) with space group $Cmc2_1$ (No. 36) or Ama2 (No. 40) was proposed by Bendersky et al. based on a TEM and electron diffraction study. (Bendersky et al., 2001; Bendersky et al., 2003) Guiblin et al. then performed structural refinement at room temperature (RT) using single crystal X-ray diffraction, only finding an acceptable fit for $Cmc2_1$. They used the alternative (non-standard) setting $A2_1am$, which allows the long axis of the unit cell to always be the c axis, although there is a $\sim 45^{\circ}$ difference in the a and b axes between I4/mmm and $A2_1am$ due to a doubling of periodicity caused by the oxygen octahedral tilting. We use the same setting here, i.e. perform refinements in the space group $A2_1am$. A similar structural determination was obtained by Lobanov et al. using powder neutron diffraction, which also gave magnetic data, although a minority additional phase was detected at RT, using very high resolution X-ray measurements, that was assigned to be I4/mmm. The $A2_1am$ RT structure is shown in Figures 1a-d. The oxygen octahedra are both anti-phase tilted about the [110] axis and in-phase twisted about the [001] axis, corresponding to the X_3^- and X_2^+ irreps respectively (Benedek & Fennie, 2011). At intermediate temperatures, the evolution of distortion is more complicated. While the X_2^+ and X_3^- distortions may be coupled at RT, it is less clear that they should have the same behaviour as a function of temperature. In fact, as the material is heated the disappearance of one or other of the distortions before a final transition to the prototype structure might seem inevitable. If the X_2^+ distortion disappears leaving only the anti-phase X_3^- octahedral tilt about [110] (Fig. 1f), the structure has space group Amam (No. 63). Conversely, the presence of a X_2^+ distortion without X_3^- (Fig. 1e) produces an Acam (No. 64) structure (Senn $et \ al.$, 2015).

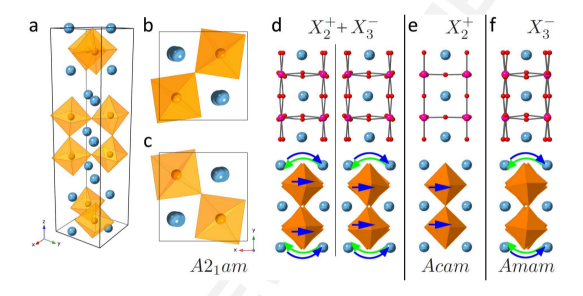


Fig. 1. The room-temperature $A2_1am$ structure of Ca₃Mn₂O₇ (Guiblin et al., 2002) and possible intermediate phases. a) Perspective view of the $A2_1am$ unit cell, Ca atoms blue and MnO₆ octahedra orange. b) View along [001] showing the upper and lower oxygen octahedra, which twist in-phase about the [001] axis. c) View along [001] showing the central octahedra, which also twist in-phase about [001]. d) Projections of the $A2_1am$ structure, as a ball-and-stick model for the upper perovskite block (Mn atoms magenta, oxygen atoms red, Ca atoms blue) and showing oxygen octahedra for the lower perovskite block. The RT structure has X_2^+ and X_3^- distortions, and appears differently when viewed along [$\bar{1}10$] (left) and [110] (right). Note the lack of mirror symmetry; the anti-phase octahedral tilts about [110] and in-phase tilts about [110] split the visible oxygen sites and displace the oxygen atoms adjacent to the Mn atom asymmetrically. e) A structure with the X_3^- distortion only, with symmetry Acam. f) A structure with the X_2^+ distortion only, with symmetry Amam. For both e) and f) the [110] projection is identical to the [$1\bar{1}0$] projection.

However, the study of Senn et al., which examined both $Ca_3Mn_2O_7$ and $Ca_3Ti_2O_7$ as a function of temperature using high-resolution X-ray and neutron powder diffraction, gave the unexpected result that a Acaa (No. 68) was in fact the best fit structure to $Ca_3Mn_2O_7$ at 500K (227°C). This is surprising because the Acaa space group corresponds to a X_1^- irrep, with anti-phase tilting along the c-axis. Although Acaa is also a subgroup of the I4/mmm prototype, it is not an intermediate between $A2_1am$ and I4/mmm, and can only be in competition with the Acam or Amam structures. The transition from $A2_1am$ to Acaa must be of first order, either with or without passing through an intermediate Acam or Amam phase. Their data indicated a coexistence of $A2_1am$ and Acaa structures between 200K and 320K, with single phase $A2_1am$ below 200K (-73°C) and pure Acaa above 320K (\sim 50°C). This proposal was given further credence by an observed contraction of the c-axis with increasing temperature, consistent with a 'trapped' X_3^- soft phonon mode that can only occur in the Acaa structure, and explained the additional RT phase detected by Lobanov et al.

Here we consider the question whether electron diffraction - specifically, D-LACBED - has the accuracy and precision required to provide a refinement of the structure of Ca₃Mn₂O₇. As noted by Senn *et al.*, the A2₁am structure requires 19 free parameters to describe the fractional atomic coordinates of the seven atoms that make up the basis (the unit cell contains 48 atoms in total). In addition, each atom in the basis has an occupancy and thermal atomic displacement parameters and there are the three lattice parameters of the orthorhombic unit cell. We use the term 'structural parameter' for all of these variables in what follows. Finally, for electron diffraction four parameters that relate to experiment, rather than the material being examined, also need to be determined - the crystal thickness, the accelerating voltage of the microscope, the angular range covered in the patterns and the point spread function of the camera. Considering only isotropic thermal atomic displacement parameters, this makes forty

parameters in total that could in principle be refined by best match of simulation to experiment for the $A2_1am$ structure. For the I4/mmm structure, the basis consists of six atoms with only four free parameters describing atomic coordinates, giving 18 structural parameters. While it is obvious that a typical D-LACBED data set, containing $>10^7$ pixels, should have sufficient information to refine this comparatively small number of unknown values, whether this is possible in practice is less clear. Furthermore, it is possible that uncontrolled parameters or effects - such as distortions in the experimental patterns, the use of an independent atom model (IAM) for Bloch wave simulations, (Kirkland, 2010), the lack of any energy filtering or unsophisticated background subtraction may compromise the measurement and make it unreliable. Finally one must consider the utility of the experiment and the relative difficulty in comparison with X-ray or neutron diffraction; does the D-LACBED method produce information that is different or more easily obtained than well-established methods?

2. Experiment

Ca₃Mn₂O₇ ceramics were synthesized by solid-state reaction using mechanochemical activation before calcination. CaCO₃ and MnO₂ powders (Sigma-Aldrich, 99% purity) were mixed in stoichiometric ratio, then milled intensively in a Fritsch Pulverisette 7 planetary ball micromill for 135 min in a dry environment followed by 23 min in suspension with n-heptane. ZrO₂ grinding bowls (25 ml) and balls (12mm diameter, acceleration 14 g) were used. The suspension was dried under an infrared lamp and the dried powder was pressed in a uniaxial press (330 MPa, 3 min) into pellets of diameter 13 mm. The pellets were calcined for 24 h at 1200°C in pure O₂ to prevent the reduction of Mn4⁺, then milled and pressed by the same procedure as above and sintered at 1300°C for 24 h in pure O₂. The composition of calcined pellets from XRD was 97% of Ca₃Mn₂O₇ and 3% of Ca₂MnO₄. Traces of Ca₂MnO₄ were

also observed in sintered pellets. The pellets were then ground and flattened using a dimpler before being ion milled to electron transparency using ${\rm Ar^+}$ ions at 6kV. The resulting specimens were examined in a standard JEOL 2100 LaB₆ transmission electron microscope operating at 200 kV.

Room temperature in our TEM was measured as 29°C. D-LACBED data was constructed from 3,721 CBED patterns using an incident beam convergence semi-angle of 0.56 mrad and a probe size of approx. 8nm full width at half maximum on the specimen. A $Ca_3Mn_2O_7$ crystal was aligned to the $[1\bar{1}0]$ $A2_1am$ zone axis. Patterns were imaged using a Gatan Orius SC600 camera with binning 4. An angular raster of 61×61 incident beam orientations at a rate of approximately 10 patterns/second covered an angular range of 35 mrad. Data was collected from roughly the same area at nominal temperatures of 100°C, 200°C, 300°C and 400°C with heating provided by a Gatan 652 double-tilt heating holder. Due to the distance between the electron transparent area and the heating element, actual temperatures are likely to be slightly lower than nominal ones. Irreversible changes were observed in the specimen at nominal temperatures of 500°C and higher, presumably due to loss of oxygen; these results were discarded as being unreliable. For temperatures above 100° C only 2,601 (51×51) CBED patterns were collected covering 29.3 mrad, since the sample was slightly less stable and subject to drift of more than a few nm. Diffuse background was subtracted from the CBED discs by 2D cubic-spline interpolation. Patterns were rotated to give h00 horizontal using bicubic interpolation and cropped to 400×400 pixels, reducing the angular range to 20 mrad. Where symmetry was evident, averaging was used to improve signal to noise ratio and correct linear distortions due to intermediate lens aberrations. Linear image distortions that could not be corrected this way were removed by comparison with simulations.

Refinement of parameters to optimise the fit between experiment and simulation

was performed using the Bloch-wave simulation program felix (Beanland et~al., 2019). Here we refer to simulations as LACBED patterns, while retaining the term D-LACBED pattern for experimental data. Experiment and simulation were compared using a zero-mean normalised cross-correlation fit index R for pixel intensities $I_{p,n}$ in LACBED patterns with mean intensities \bar{I}_n and standard deviation σ_n

$$R = 1 - \sum_{n=1}^{N} \sum_{p=1}^{P} \frac{1}{N} \left(\frac{I_{p,n}^{e} - \bar{I}_{n}^{e}}{\sigma_{n}^{e}} \cdot \frac{I_{p,n}^{s} - \bar{I}_{n}^{s}}{\sigma_{n}^{s}} \right), \tag{1}$$

where the sum p is performed over all P pixels in each pattern and the sum n is performed over all N experimental D-LACBED patterns. Superscripts e and s refer to experimental and simulated patterns respectively. This comparison ignores the absolute intensity of each LACBED pattern and concentrates instead on the changes in the detail of each pattern individually. Equation (1) gives a value of zero for a perfect fit and a value of unity for completely uncorrelated experiment and simulation. Data was binned by four to 100×100 to reduce simulation times and symmetryrelated patterns were not included to reduce memory utilisation. The simulation uses isotropic Debye-Waller factors and atomic Born scattering factors for neutral atoms from (Kirkland, 2010). Absorption is calculated using the numerical integration package QUADPACK, (Piessens et al., 1983) following the Bird and King model (Bird & King, 1990) also using Kirkland scattering factors. Simulations on a cluster used 400 zero-order Laue zone reflections chosen on a pixel-by-pixel basis from a pool of 1200, which typically completed in 108 seconds using 168 cores. Changes in intensity in the set of LACBED patterns produced by any structural parameter are both complex and distinctive, which allows a best fit to experimental D-LACBED patterns to be optimised for each parameter independently using a downhill gradient optimisation routine. A global minimum was always obtained, independent of starting position. Errors were determined by the change in structural parameter that produced an increase of 10^{-5} in R, corresponding to the reproducibility of the solution obtained from IUCr macros version 2.1.11: 2020/03/03

a multitude of starting points.

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3. Results

3.1. Room temperature imaging and diffraction

A typical TEM image at RT is shown in Fig. 2a. Grains, often with (001) planar boundaries, were up to 2 μ m in size. Imaging perpendicular to the [001] direction reveals a significant density of (001) stacking faults, with a spacing between 2 and 100 nm, i.e. between 1 and 50 unit cells. These faults are interruptions in the 2:1 perovskite:rock-salt n=2 Ruddlesden-Popper stacking sequence and were found to always consist of additional perovskite layers, indicating a slight excess ($\sim 1\%$ Mn) above the nominal 2:3 Mn:Ca ratio. Bands of brighter and darker contrast are observed in dark field images as can be seen in the upper part of Fig. 2a. As will be shown later, these bands are merohedral twins, as first proposed by Bendersky et al. and also observed by Gao et al.. D-LACBED data were collected from the centre of stacking fault-free regions, typically between 30 and 100 nm in size, using an electron probe of 10nm FWHM.

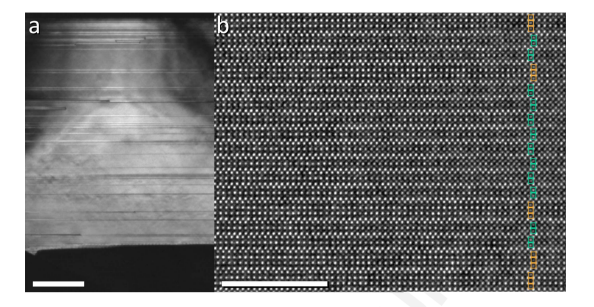
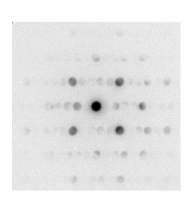


Fig. 2. Transmission electron micrographs of $Ca_3Mn_2O_7$ at the [110] $A2_1am$ zone axis. a) Dark field 220 image showing planar defects and contrast bands in a single grain (scale bar 200 nm). b) High resolution [110] lattice image showing the occurrence of n=3 perovskite blocks forming planar defects. Scale bar 10 nm.

3.2. Room temperature D-LACBED

Figure 3a shows an individual RT $Ca_3Mn_2O_7$ CBED pattern at an incident beam orientation very close to the $[1\bar{1}0]$ zone axis, while Fig. 3b shows the average CBED pattern, obtained by aligning the 000 beam of all 3,721 CBED patterns prior to summation. Due to the relatively large angular range covered by the incident beam, the average CBED pattern shows a more even distribution of intensities and extends further out in reciprocal space, a similar effect to that seen in precession electron diffraction (PED) (Vincent & Midgley, 1994). Here, we only reconstruct D-LACBED patterns that have their pattern centre within the angular range covered, and choose the one hundred and nine discs inside the marked area in Fig. 3b.





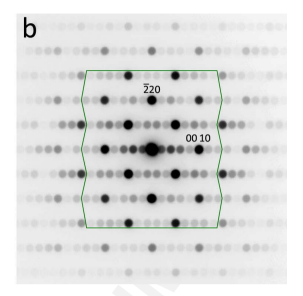


Fig. 3. a) Individual RT CBED pattern obtained at the Ca₃Mn₂O₇ [110] A2₁am zone axis. b) Average RT CBED pattern, the sum of 3,721 CBED patterns. The 109 spots used to reconstruct D-LACBED patterns lie inside the green boundary.

The corresponding RT D-LACBED patterns are shown in Fig. 4. As we have noted before (Beanland et al., 2013), determination of symmetry using D-LACBED patterns is straightforward in comparison with other methods, particularly for materials with large lattice parameter that restrict the convergence angle of the incident beam. In the current case, the assembly of D-LACBED patterns in the upper part of Fig. 4 exhibits a (001) mirror, but not a (110) mirror, as expected from the [110] projection of the RT structure in Fig. 1d. Since electron diffraction does not obey Friedel's law, the lack of a centre of symmetry is visible from the difference between $\pm \mathbf{g}$ patterns, or equivalently by comparison of Bijvoet pairs $h\bar{h}l$ $\bar{h}hl$ related by the broken mirror, as shown for selected patterns in the lower part of Fig. 4. The D-LACBED patterns on the left show that symmetry breaking is quite striking in some cases (e.g. $\bar{2}22$ -, $\bar{1}17$ -, $\bar{2}28$ -type patterns), but hardly visible in others (e.g. 22 10-type patterns). The presence of (001) mirror symmetry and absence of (110) mirror symmetry is also visible in the D-LACBED patterns on the 00h systematic row. As is clear from Fig. 1d, the symmetry

is broken primarily by oxygen atom displacements of a few picometres, demonstrating the sensitivity of D-LACBED to the coordinates of relatively light elements. It is impossible to observe the symmetry in the conventional CBED pattern (Fig. 3a) - in fact since the discs must be so small to avoid overlap and are essentially featureless, it is hardly possible to align the crystal exactly on the zone axis. Furthermore, while the average CBED pattern contains significant intensities that cover a larger angular range (Fig. 3b), the absence of the (110) mirror is still not evident, presumably because the intensity differences only affect a small part of the the D-LACBED patterns and so have a small impact on the total intensity. We expect this problem may also affect PED data, which may not sample the particular incident beam orientations that produce strong intensity differences between Bijvoet pairs.

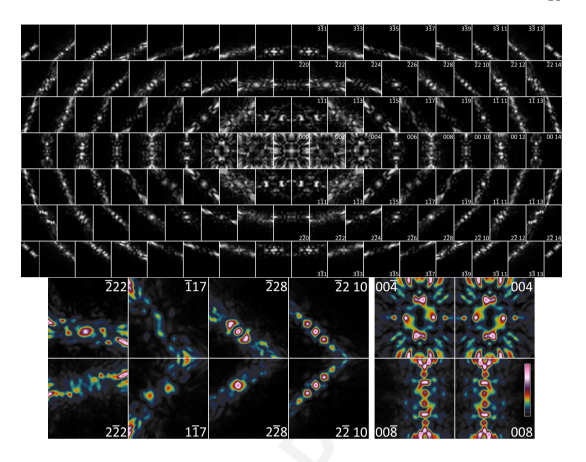


Fig. 4. Top: the 109 experimental RT (29°C) D-LACBED patterns at the Ca₃Mn₂O₇ [1 $\bar{1}$ 0] zone axis corresponding to the spots indicated in Fig. 3b. All patterns have an angular range of 20 mrad and are normalised to the full intensity range for display. A mirror is present on (001) but not (110), which can be seen in 00h patterns and by comparing Bijvoet pairs $\bar{h}hl$ and $h\bar{h}l$ (false colour patterns, bottom).

The difference in $\pm \mathbf{g}$ D-LACBED patterns in Fig. 4 provides a simple method to demonstrate that the dark and light bands seen in dark field TEM images are caused by merohedral twinning. Four twin variants are expected from the ratio of the order of the point groups of the prototype and RT structure, 4/mmm (order 16) and mm2 (order 4) respectively, i.e. $4/mmm = mm2 \times 2/m$, where \times indicates a direct group product. In the case of $A2_1am$ Ca₃Mn₂O₇, one may choose $2/m = \{1, 2_{[110]}, m_{(110)}, \bar{1}\}$. However, since D-LACBED patterns are from the zero order Laue zone (ZOLZ) they do not contain any three-dimensional information from higher-order

Laue zones (HOLZ). They correspond to a projection of the crystal structure and it is not possible to distinguish between all four of these different twinning operations; a twin generated by $2_{[110]}$ is invisible and that generated by $m_{(110)}$ appears identical to that generated by $\bar{1}$. This explains the observation of only two types of contrast, dark and light, even though there are four distinct twin variants. Fig. 5 shows selected D-LACBED patterns, 002 and 222-type, which are sensitive to the broken mirror symmetry obtained from bands of different contrast. Reflections that appear equivalent in conventional (CBED, SAED) diffraction patterns are labelled with the same indices, using the subscript T to denote twinned (darker) material. The relationship between the different bands is readily apparent, e.g. the $2\bar{2}2_T$ pattern from the dark band is equivalent to the $\bar{2}2\bar{2}$ pattern from the light band.

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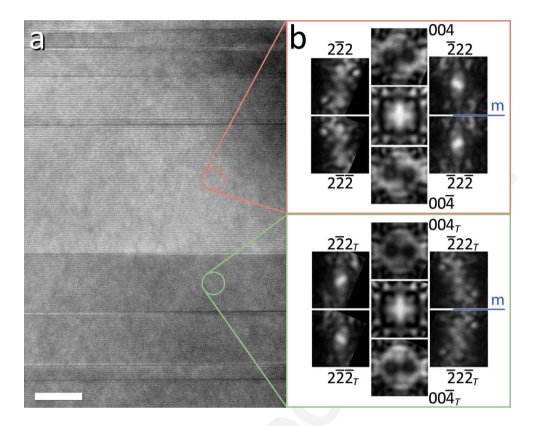


Fig. 5. a) contrast bands in a TEM image of $Ca_3Mn_2O_7$ close to the $[1\bar{1}0]$ axis. b) Selected D-LACBED patterns taken from different bands showing that the structures are related by a twinning operation such as a (110) mirror or an inversion, although it is not possible to distinguish between them.

3.3. Structural refinement

We first consider atomic coordinates, with 19 structural parameters. Refinement was performed by minimising differences between simulations and experimental RT data as described in section 2, starting from the I4/mmm structure given by Bendersky et al. in the space group $A2_1am$ with some small initial displacements from I4/mmm to ensure that the sense of octahedral rotations were consistent with the model of Lobanov et al.. The initial fit was $R \sim 10\%$, which refined to R=3.7%. The refined RT structure is given in Table 1. Column δs gives the difference in picometres between this refinement and the structure determined by Lobanov et al. using

synchrotron X-ray diffraction. Average disagreement for all coordinates is 1.7 pm, with the largest difference (9 pm) in the position of the Ca atoms; our refinement is closer to that of Lobanov $et\ al$. than is the X-ray study of Guiblin $et\ al$. This result clearly demonstrates the ability of D-LACBED to reliably determine structure with a precision comparable to other diffraction methods. Differences in electron and X-ray/neutron refinements may be due to the inevitable inclusion of stacking faults and small amounts of other phases in the X-ray/neutron data, (Lobanov $et\ al$., 2004) which are avoided by the local measurement in defect-free material that is possible with D-LACBED. The RT structure refined from our D-LACBED data shows octahedral tilting about x and z axes, as expected.

Site		Coordinate			$\delta s \; (\mathrm{pm})$	$B(\text{\AA}^2)$	Occ.
Ca1	4a	0.7360(6)	0.2654(8)	0	[4.2, -9.1, -]	0.91(5)	1.016(3)
Ca2	8b	0.7578(4)	0.2657(5)	0.1861(1)	[9.0, -6.7, 0.8]	0.79(5)	1.014(2)
Mn	8b	0.2500(5)	0.2500(20)	0.0978(1)	[-0.8, 1.6, 0.0]	0.52(4)	1.019(3)
O1	4a	0.2513(10)	0.2947(11)	0	[2.5, 2.8, -]	2.2(1)	0.95(3)
O2	8b	0.0258(10)	0.5297(9)	0.1045(1)	[0.6, 2.1, -0.1]	1.4(1)	0.91(1)
O3	8b	0.9649(10)	0.0335(9)	0.0889(1)	[-2.0, 2.9, -1.6]	1.5(1)	0.90(1)
O4	8b	0.2455(7)	0.2118(9)	0.1964(1)	[0.8, 3.9, -2.4]	2.0(1)	0.96(1)

Table 1. Refined RT A2₁am structural parameters with a best fit of R=3.7%. Column δs gives the difference in picometres between this refinement and the synchrotron X-ray + neutron study of Lobanov et al.. Previous work (Hubert et al., 2019) indicates that D-LACBED refinements overestimate atomic displacement parameters B and that the apparent occupancy is influenced by charge transfer between atoms: here all cations appear to have an occupancy slightly greater than one, and oxygen anions less than one.

Second, D-LACBED data is also sensitive to thermal atomic displacement factors B and apparent occupancy. The influence of B on D-LACBED patterns is typically ~ 50 times less than that of atomic coordinates. Optimising the fit to experiment improved R by at most 0.5% and had no noticeable effect on atomic coordinate refinement. The lower sensitivity is reflected in the lower precision of B-factors in Table 1. Our previous study (Hubert $et\ al.$, 2019) found that thermal atomic displacement parameters in GaAs were significantly overestimated in D-LACBED refinements. This also is the case here; Guiblin $et\ al.$ give $B_{Ca}=0.58$, $B_{Mn}=0.30$ and $B_O\approx 0.65$ at RT. Since the IUCr macros version 2.1.11: 2020/03/03

main effect of thermal atomic vibrations on LACBED patterns is to change absorption, the discrepancy may indicate that an improvement in the subtraction of diffuse background intensity in the original CBED patterns is required. Nevertheless, as shown in Section 3.5, B-factors determined from D-LACBED data increase with temperature, as expected. An additional improvement in R of at most 0.5% is obtained by allowing occupancy to vary, shown in the final column of Table 1. Oxygen stoichiometry may be affected by, for example, loss of oxygen under the electron beam and/or during heating in the vacuum of the microscope. However the best fit for oxygen occupancy at RT, between 0.9 and 0.95, suggests a very large oxygen loss that does not seem reasonable. This discrepancy is almost certainly due to the neutral independent atom model (IAM) used here for structure factors; a lower measurement of occupancy corresponds to a shallower atomic potential and a positively charged atom, consistent with the O^{2+} anions in this material. This explanation is supported by occupancies very slightly in excess of unity for both Ca and Mn cations. We observe similar effects consistent with charge transfer between cations and anions in all other oxides studied by D-LACBED using a neutral IAM.

Finally, D-LACBED data is also sensitive to lattice parameter, but unlike the other structural refinements this changes the position of the Bragg condition in the pattern as well as changing intensities from place to place. Our cross-correlation Eq. (1) requires both the angular range of the D-LACBED pattern to be known, and the position of the Bragg condition within it to be aligned to that of the simulated LACBED pattern with sub-pixel precision. We find that the presence of distortions such as intermediate lens astigmatism prevents a precise and absolute calibration of angular range to better than 0.5%, which is vastly inferior to X-ray measurements and not sufficient to allow reliable refinement of lattice parameter using D-LACBED. We thus use values from other studies (Senn et al., 2015).

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3.4. Sensitivity of D-LACBED data

The structure given in Table 1 clearly shows that structural refinement from D-LACBED data is viable even for quite complex inorganic materials. We now consider the sensitivity of LACBED patterns to the different structural parameters; we find that they depend strongly upon which atom, and which coordinate, is being refined. The effect of a structural parameter can be quantified by taking the difference between zero-mean normalised LACBED patterns for a reference structure S and an altered one S', in which a small change in the i^{th} structural parameter is made, in the current case a small change in atomic coordinate δs_i . The result is an image that has positive and negative values corresponding to increases or decreases in intensity produced by the small change in structural parameter, which we call a δ image. Dividing by the magnitude of the displacement allows a direct comparison between different coordinates,

$$\delta_{ij} = \left(\frac{I_j' - \bar{I}_j'}{\sigma_j'} - \frac{I_j - \bar{I}_j}{\sigma_j}\right) \frac{1}{\delta s_i}.$$
 (2)

The δ images are shown for all 54 patterns j and all 19 allowed atomic displacements i for $A2_1am$ Ca₃Mn₂O₇ in the Appendix. We may assess the relative effect of each structural parameter by taking the root mean square (RMS) of the δ image, i.e.

$$\Delta_{ij} = \left(\delta_{ij}^{\bar{2}}\right)^{1/2},\tag{3}$$

plotted in Fig. 6. A larger Δ_{ij} corresponds to a larger change in the intensities of LACBED pattern j. Thus, in Fig. 6(a), each point corresponds to an individual Δ_{ij} for the refined RT structure, ordered by the structural parameter i. A similar plot can be used to assess the goodness of fit between simulation and experiment, as shown in Fig. ??, which shows the general improvement in fit for all patterns as the structure is refined from I4/mmm to $A2_1am$, but also the improvement in the fit for specific patterns such as $1\ \bar{1}\ 11$ and $1\ \bar{1}\ 15$. The different atomic coordinates have differing

effects on the LACBED patterns, and it is instructive to examine this in more detail. An obvious dependence on atomic number is present, with the largest changes produced by the z coordinate of Mn (Z=25) followed by Ca (Z=20) and oxygen (Z=8). The ratio of maximum intensity changes produced by equal movements of the three elements, $I_{max}(Mn): I_{max}(Ca): I_{max}(O)$ is 7.0:4.3:1, which indicates a dependence of roughly $Z^{1.6}$. A second clear trend is the dependence of Δ_{ij} on the ordinate being refined; z coordinates have roughly 4 to 10 times larger influence on the patterns than y and x coordinates. This can be partly understood by noting that the z coordinate is perpendicular to the incident beam direction; movements along z lie completely in the [110] projection, whereas movements along both x and y are close to 45° to the beam direction and so produce displacements in this projection that are smaller by a factor of $\cos(45^\circ)$. However, the y coordinate of Mn produces the smallest intensity changes of all, which seems in contradiction with the dependence upon atomic number noted above.

800

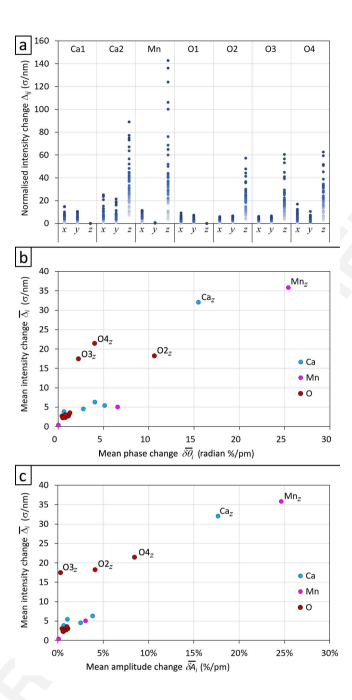


Fig. 6. a) Normalised RMS changes in intensity Δ_{ij} of the 54 unique [110] LACBED patterns for the 19 allowed atom displacements within the constraints of $A2_1am$ (N.B. Ca1 z and O1 z are fixed at zero). The change in intensity is given in units of standard deviation of the image per nm movement. b) The correlation between the average of all LACBED intensity changes $\bar{\Delta}_i$ and the average change in structure factor phase $\delta \bar{\theta}_i$ for the 19 allowed atom displacements. c) The same correlation for the mean change in structure factor amplitude $\delta \bar{A}_i$.

Further insight may be gained into the sensitivity of LACBED patterns to changes in atomic coordinates by considering the corresponding structure factors. The change in the structure factor of the j^{th} reflection F_j due to a change in the i^{th} structural parameter is

$$\delta F_{ij} = \sum_{cell} f_n \exp\left(2\pi \iota \mathbf{g}_j \cdot \delta \mathbf{r}_{ni}\right),\tag{4}$$

where the sum is performed over all n atoms in the unit cell, f_n is the scattering factor of the n^{th} atom and $\iota = \sqrt{-1}$. We find that plotting δF_{ij} against Δ_{ij} yields no correlation (plots showing δF_{ij} for all 19 coordinate parameters in $A2_1am$ Ca₃Mn₂O₇ are shown in the Appendix). This lack of correlation is due to the transfer of intensity between different **g**-vectors due to dynamical scattering; notably, large intensity changes occur even for LACBED patterns that have precisely zero change in amplitude or phase. However, for a given structural change i there is a positive correlation between the *average* change in structure factor of all reflections $\delta \bar{F}_i$ and the *average* of all LACBED intensity changes $\bar{\Delta}_i$, shown in Figs. 6b and c in terms of mean change in structure factor amplitude $\delta \bar{A}_i$ and phase $\delta \bar{\theta}_i$.

In summary, structural parameters that have large effects on structure factors do produce large changes in the intensity of LACBED patterns, but these intensity changes are redistributed through the set of patterns by dynamical scattering. Equally, changes that have little effect on structure factors have a lesser effect on LACBED patterns. Although the magnitude of a change in intensity in any individual pattern may effectively be uncorrelated to that of the underlying cause, the complete set of LACBED patterns shows a similar overall sensitivity to that expected from kinematical scattering. This explains the relative lack of sensitivity of the D-LACBED data to the Mn x coordinate, which produces only very small changes in the structure factors of the LACBED patterns at the [110] zone axis (the point close to the origin in Figs. 6b and c).

We now understand that the average magnitudes of intensity changes $\bar{\Delta}_i$ are correlated with the average change in structure factor, but have not considered why these should be so different for the particular case of Ca₃Mn₂O₇. Certainly the dependence on scattering factor implicit in the structure factor equation ((4)) explains the effect of atomic number, but why is there such a difference between different ordinates? We may understand this more clearly by examining the allowed displacements within the $A2_1am$ space group, as shown for the Ca2 atom in Fig. 7. The Ca2 atom lies in the rocksalt layer of the Ruddlesden-Popper structure with a multiplicity of eight in the unit cell. Displacing a Ca2 atom along x results in a movement of all Ca2 atoms in the same direction when viewed in the [110] projection as shown in Fig. 7a. Conversely, a displacement along y moves atoms at different depths in opposite directions, and what appears as a single atom column in I4/mmm splits into a pair of overlapping atoms at different depths. Finally displacement along z moves the Ca2 atoms further towards the centre of the rocksalt layers. Movements of the other atoms are similar as illustrated in video V1. The very small Δ_{ij} values for y movements of the Mn atom may be caused by two factors: first, the the centroid of the atom column remains unchanged and a y displacement therefore does not change the phase of low order structure factors; and second the close and symmetrically arranged O2 and O3 atoms overlap the Mn atom completely. The latter probably indicates that D-LACBED refinement loses some precision when atoms overlap in projection and may be a limiting factor in the technique.

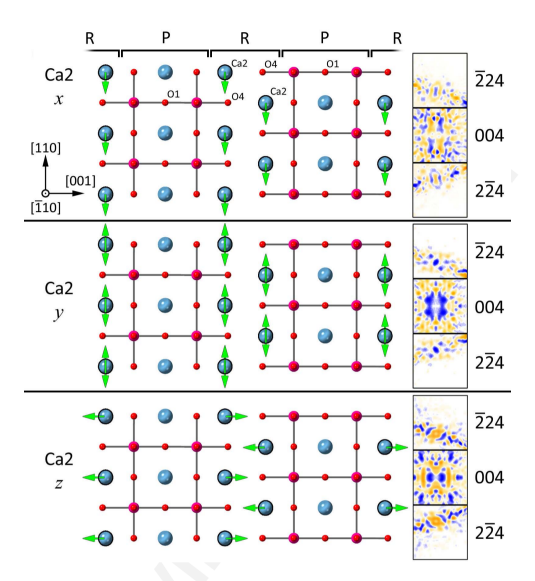


Fig. 7. Left: the movements of atom Ca2 from the prototype structure within space group $A2_1am$ when displaced along the x, y and z direction respectively as seen in the $[1\bar{1}0]$ projection. Colour coding is Ca blue, Mn magenta and oxygen red; perovskite/rocksalt layers are labelled P and R respectively. There are eight symmetrically equivalent Ca2 atoms in the unit cell, which lie in the rocksalt layers, while Ca1 atoms lie in the perovskite layers. Right: three δ_{ij} patterns ($\bar{2}24$, 004, $2\bar{2}4$) showing the resulting changes in intensity (yellow = positive, blue = negative). Displacement of Ca2 along x breaks mirror symmetry and produces antisymmetric δ_{ij} patterns. Displacement of Ca2 along y moves atoms at different depths in opposite directions, maintaining mirror symmetry in the projection and δ_{ij} patterns, while displacement along z moves Ca2 away from the perovskite layers, also maintaining mirror symmetry. See also video V1 and figure ?? in the appendix.

In Fig. 7 the atoms are displaced from the tetragonal I4/mmm structure. The effect on the symmetry of the whole LACBED dataset is illustrated using a triplet of δ images (the 004 pattern and the Bijvoet pair $\bar{2}24$, $2\bar{2}4$) on the right. Displacements along y and z do not break mirror symmetry in the [110] projection and thus the δ patterns maintain this symmetry. Conversely, a shift along x does break mirror symmetry and the δ patterns are antisymmetric, with opposite changes in intensity for Bijvoet pairs. The presence of 2-fold symmetry in all LACBED patterns produces lines of zero intensity change in the systematic 001 row in the latter case, a similar effect to the well-known Gjonnes-Moodie bars but in this case in the differential of the LACBED pattern δ_i . More complicated changes can be produced by displacement of the oxygen atoms, for example movements of atoms O2 and O3 have δ images without mirror or antimirror symmetry. However, mirror symmetry can be maintained by coupled movement of the oxygen atoms, for example when O2z and O3z change by equal amounts in the same direction, or when O2x and O3x change by equal amounts in opposite directions (see Fig. ??).

In summary, while it is operationally straightforward to refine structural parameters using a simple fit parameter and a downhill gradient optimisation, the sensitivity of the technique depends on the details of the structure being examined. The symmetry visible in D-LACBED patterns provides extra information, which can be used to constrain the permissible structural distortions in a structure. We make use of this additional constraint in the following section.

3.5. Refinements at elevated temperature

Measurements were taken and refinements performed for nominal sample temperatures up to 400° C, with the starting point for each refinement the best fit determined for the temperature below. Refinements are constrained to the $A2_1am$ space group

for all temperatures. (While (Senn et~al., 2015) proposed the space group Acaa at temperatures where the anti-phase X_3^- distortion is absent, this gave identical fits since in-phase and anti-phase octahedral twists about [001] appear exactly the same in the [110] projection of these structures. For simplicity here we ignore the possibility of an anti-phase X_1^- distortion and the Acaa phase on the understanding that the results are also compatible with it.) Selected D-LACBED patterns ($\bar{2}22$, $2\bar{2}2$ and $\bar{1}17$, $1\bar{1}7$) are shown in Fig. 8 on the left, with best-fit simulations on the right. A good fit is obtained for all cases and the change in symmetry is readily apparent, with the development of (110) mirror symmetry essentially complete by 200°C, indicating that the X_3^- distortion has dropped to zero.

008

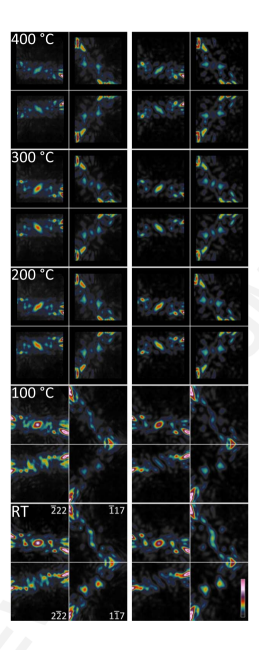


Fig. 8. Comparison of experimental D-LACBED patterns (left) and best-fit simulated LACBED patterns (right) at a range of temperatures. The $\bar{2}22$ and $\bar{1}17$ Bijvoet pairs are strongly asymmetric at RT due to the presence of both X_2^+ and X_3^- distortions (see Fig. 1). Mirror symmetry is present at 200°C and higher.

In refinements at 200°C or higher, mirror symmetry was preserved in the simulations by coupling the movement of oxygen atoms in octahedral rotations about the long c-axis corresponding to the X_2^+ distortion mode, as noted above. Structures are given in the Appendix, Tables ?? to ??. Fig. 9(a) shows the magnitude of the distortions IUCr macros version 2.1.11: 2020/03/03

as a function of temperature. Although previous investigations found that tetragonal I4/mmm symmetry is obtained at temperatures 400°C and higher, the octahedral rotations about c do not drop to zero at 400°C in our measurement and in fact increase between 300°C and 400°C. This may be because the temperature of the TEM specimen at the electron probe is rather lower than the nominal value, obtained from a thermocouple in the specimen holder. The thermal atomic displacement parameters (B-factors) extracted from D-LACBED data are shown in Figs. 9(b) and (c). The Mn and Ca atoms show a monotonic increase in B-factor with temperature, as expected. Interestingly, oxygen atoms O1 and O4, whose displacement from the sites shown in Fig. 7 correspond to the X_3^- distortion, show a strong decrease in B-factor up to 300°C, i.e. even when the X_3^- distortion is zero. This may show the presence of strong anharmonicity or rapid hardening of a soft phonon mode above 200°C.

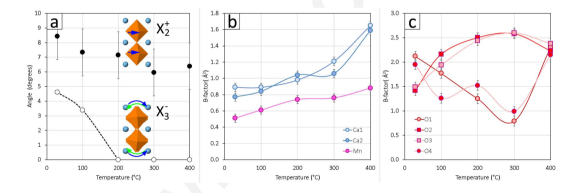


Fig. 9. (a) The magnitude of oxygen octahedral rotations corresponding to the distortion modes X_3^- and X_2^+ (or X_1^-). (b) Thermal atomic displacement parameters (*B*-factors) for Ca and Mn as a function of temperature. (c) *B*-factors for oxygen atoms showing an anomalous decrease for O1 and O4 up to 300°C.

4. Discussion

We begin by addressing the question posed at the end of the introduction - the information that is produced and the ease of use of the technique in comparison with other methods. The structural measurements made here are quite routine for single crystal IUCr macros version 2.1.11: 2020/03/03

neutron diffraction or synchrotron X-ray diffraction, but have to date not been possible using electron diffraction. While a large amount of data and simulated LACBED patterns are presented here (see Appendix), this is primarily for the sake of completeness and to explore the possibilities offered by the D-LACBED method and should not be taken as the level of detail or data processing necessary to obtain a result. In fact, data acquisition is straightforward and takes only a short amount of time, and the level of expertise required is little more than that needed to collect a conventional CBED pattern. Data processing into a form suitable to match simulated patterns typically takes less than an hour. Refinement by matching to Bloch wave simulations, although robust due to the presence of a global minimum, currently requires access to good computing resources (preferably a cluster with 200 cores or more), but these are widely available and the computation requirement compares favorably with other techniques such as molecular dynamics or density functional theory. Improvements in both experimental and computational methods could easily reduce time and effort by some orders of magnitude, while the relationship between intensities in dynamical diffraction patterns and structural parameters provide an extra dimension to analysis which is available if required. Additionally, refinement of individual structure factors, as typically practised in CBED studies, (Zuo, 1998; Zuo & Spence, 2017) is manifestly possible using LACBED data. As illustrated here, the D-LACBED technique brings the advantages of transmission electron microscopy, i.e. the ability to collect data from nanoscale regions and the opportunity to directly investigate the same region using diffraction contrast or high resolution imaging, as well as structural determination with sub-nanometre precision. Perhaps the biggest limitation of the technique, at least in its current form where HOLZ information is essentially absent, is the lack of information on atomic displacements parallel to the electron beam. In the case of [110] Ca₃Mn₂O₇ the displacement of one atom parallel to the beam must be accompanied

by the displacement of another in a different direction that is related by symmetry, thus ensuring that all atomic coordinates can be refined. This is not the case in general. It may also be the case that atoms that are overlapping in the particular point of view chosen cannot be separated. To get the most from the technique, there may have to be some trade off between the choice of an incident beam direction that is sensitive to all atomic displacements while maintaining enough dynamical diffraction to be sensitive to the relative phases of the diffracted beams. Alternatively, refinements may be made using data collected from different zone axes.

In the current refinement these limitations prevent the two models of structure at elevated temperature, Acam and Acaa, from being distinguished, although a repeat experiment at a different zone axis could do so. However, the decay in antiphase oxygen octahedral tilt (X_3^-) with increasing temperature (Fig. 9) is quite clear, as is the relative lack of change in octahedral twist about the c-axis (X_2^+) . Our results are in good agreement with more sophisticated studies such as that of (Ye et al., 2018).

5. Summary and conclusions

We have demonstrated that D-LACBED data can be used to determine the relatively subtle structural distortions in $Ca_3Mn_2O_7$ with a single, global minimum in fit parameter that is straightforward to obtain using gradient descent optimisation. Refinement of the structural model for RT data in the $A2_1am$ space group gives an excellent fit to simulation. The result agrees with previous X-ray and neutron diffraction measurements to a mean accuracy better than 2 pm. Measurements at elevated temperature confirm the decay in anti-phase oxygen octahedral tilting perpendicular to the c axis, producing an Acam or Acaa structure at $200^{\circ}C$ and above. The octahedral twists about c are robust and persist to a nominal $400^{\circ}C$, in agreement with previous work by others. The high sensitivity of the technique to atomic displacements can be un-

derstood by examining the changes in structure factor they produce. While dynamical diffraction effects ensure that changes in intensity for an individual pattern are generally unrelated to the change in its structure factor, taken as a whole an ensemble of D-LACBED patterns demonstrates an average sensitivity that correlates well to the average change of all structure factors. The additional information provided by the symmetry of the D-LACBED patterns allows additional constraints to be applied to a structural model. The main weakness of the technique in its current form is the inability to distinguish between structures that appear the same in projection for the chosen orientation of the crystal. Thermal atomic displacement parameters can also be obtained but are much larger than those obtained by other methods, possibly indicating that improvements in the absorption model or data processing are required. Apparent atomic occupancies may be influenced by charge transfer between atoms, and show the limitations of the neutral independent atom model used here.

Finally, we note that the method can in principle be used on any modern computercontrolled transmission electron microscope, giving an additional quantitative diffraction method that can be applied at the nanoscale, complementing the already considerable strengths of TEM in imaging and spectroscopy.

Acknowledgments

We would like to acknowledge the University of Warwick Research Technology Platform on Scientific Computing for assistance in the research described in this work. P. V. and S.K. were supported by the Czech Science Foundation (Project No. 18-09265S) and by MŠMT Project No. SOLID21 - CZ.02.1.01/0.0/0.0/16_019/0000760.

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