# Average and Local Structure of Apatite-Type Germanates and Implications for Oxide Ion Conductivity

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**ABSTRACT:** Materials with the apatite structure have a range of important applications in which their function is influenced by details of their local structure. Here we describe an average and local structural study to probe the origins of high temperature oxide ion mobility in La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> and La<sub>8</sub>Bi<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> oxygen-excess materials, using the low-conductivity interstitial oxide-free La<sub>8</sub>Sr<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>2</sub> as a benchmark. For R = La and Bi we locate the interstitial oxygen, O<sub>int</sub>, responsible for conductivity by Rietveld refinement and relate the  $P6_3/m$  to  $P\overline{1}$  phase transitions these phases undergo on cooling to oxygen ordering. Local structural studies using neutron total scattering reveal that well-ordered GeO<sub>5</sub> square pyramidal groups form in the structure at low temperature, but that O<sub>int</sub> become significantly more disordered in the high-conductivity for R = Bi to the presence of several O<sub>int</sub> sites of similar energy in the structure, which correlates with its less-distorted low temperature average structure.

# Introduction

Apatites are materials with the general formula  $A_{10}(MO_4)_6X_{2\pm x}$ , where A = alkaline or rare earth metal, M = Ge, Si or P and X = halides, O<sup>2-</sup> or [OH]<sup>-</sup>. Compounds adopting this structure type are ubiquitous both as natural biomaterials and as synthetic functional materials with a range of technological applications. They are therefore of significant research interest in areas as diverse as materials chemistry, physics, engineering, medicine, archaeology and conservation science.

In all its roles, the fine details of the apatite structure, and in particular short-range local structure, have a critical impact on its physical or structural properties. For example, symmetry-lowering deviations from the ideal apatite structure and differing arrangements of ions in its channels are important to the function of natural bone and tooth enamel (carbonated calcium phosphate hydroxyapatite), and the use of apatite-type synthetic materials in bone grafts and implants.<sup>1-5</sup> In particular, the details of the local coordination environment of Ca<sup>2+</sup> ions are necessary for understanding bone formation and diseases, but this information is hard to obtain given the experimental difficulty of techniques such as <sup>43</sup>Ca solid state NMR.<sup>6-8</sup> Similar research questions regarding apatite-type materials are important in archaeological science and anthropology.<sup>9-12</sup>

In the context of modern technological applications, departures from the ideal centrosymmetric crystal structures make some apatite-type materials second harmonic generation (SHG) active and potentially suitable for non-linear optical (NLO) applications.<sup>13-14</sup> Apatite-type oxides are also promising hosts for the development of phosphors for solid-state lighting. In this application, the local environment and symmetry of the crystallographic sites occupied by the dopant activator ions significantly influence luminescence probabilities and energy transfer processes, and hence the emission properties of the phosphors.<sup>15-18</sup> Certain apatite-type lanthanum germanates and silicates also exhibit high oxide ion conductivities, making them applicable in oxygen sensors and pumps,<sup>19</sup> separation membranes,<sup>20-21</sup> and solid oxide fuel cells (SOFCs).<sup>22-24</sup> These applications are the focus of this paper.

The ideal apatite structure (x = 0 in A<sub>10</sub>(MO<sub>4</sub>)<sub>6</sub>X<sub>2+x</sub>) can be described as a framework in which columns of face-sharing AO<sub>6</sub> trigonal prisms share corners with TO<sub>4</sub> tetrahedra (dark green and grey polyhedra, respectively, in Figure 1a) to form channels running down the crystallographic c-axis; these channels are filled by  $A_6X_2$  constituents (represented by light green and pink spheres in Figure 1a). The space group is usually hexagonal  $P6_3/m$ , although lower-symmetry variants are also known.<sup>25</sup> The AO<sub>6</sub> trigonal prism twist angle  $\phi$  has been used as a quantitative parameter indicative of the geometry of the framework and symmetry of the material.<sup>26</sup> If the size of the channels is large relative to the units occupying it, the A cations in the channels tend to be under-bonded and the tetrahedral T cations over-bonded; to remove this unfavourable situation, the framework trigonal prisms twist. A recent density functional theory (DFT) computational study <sup>27</sup> of a series of hexagonal and triclinic apatites with A cations of very similar sizes, A<sub>10</sub>(PO<sub>4</sub>)<sub>6</sub>F<sub>2</sub> (A=Ca, Sr, Pb, Cd, Hg), found the symmetry lowering to be an electronic effect associated with mixing of the occupied  $(n-1)d^{10}$  and the empty  $ns^0$  states for A = Cd and Hg. In these cases the authors propose a particular displacive mode, associated with cooperative tilts of the TO<sub>4</sub> tetrahedra belonging to the  $\Gamma_4^+\Gamma_6^+$  irreducible representation (irrep), as driving the  $P6_3/m$  to  $P\overline{1}$  phase transition.



**Figure 1.** (a) The hexagonal  $P6_3/m$  apatite structure, exemplified by the high-temperature average structure of La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> viewed down the *c*-axis; (b) Labels of different potential O<sub>int</sub> sites (orange spheres, A-F) in the  $P\overline{1}$  apatite structure. The grey tetrahedra show GeO<sub>4</sub> groups and are included to highlight their proximity to the O<sub>int</sub>. Symmetry-unique GeO<sub>4</sub> tetrahedra are numbered.

Oxygen-excess (x = 1) La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> and La<sub>8</sub>Bi<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> are both excellent oxide ion conductors, with  $\sigma \approx 7.8 \times 10^{-3}$ and  $1.3 \times 10^{-2}$  S/cm at 775 °C, respectively.<sup>28</sup> The room temperature structure of  $La_{10}(GeO_4)_6O_3$  has been described as triclinic from low-resolution powder neutron diffraction data,<sup>29</sup> while the high temperature structure has not been reported. The structure of La<sub>8</sub>Bi<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> has been studied by highresolution synchrotron X-ray and neutron diffraction data and found to be triclinic at room temperature and hexagonal at 800 °C.<sup>30</sup> There is a general consensus in the literature, both from diffraction-based investigations of the average structure and from computational studies, that the most likely location for the extra oxygen (O<sub>int</sub>) atoms in apatite-type germanates is between the opposite faces of adjacent GeO4 tetrahedra (locations represented by orange spheres in Figure 1). In space group  $P6_3/m$ , these locations correspond to a single unique crystallographic site (Wyckoff site 6g). When the symmetry is lowered to triclinic  $P\overline{1}$ , however, they become 6 independent crystallographic sites). The potential locations of interstitial Oint sites in the unit cell are shown in Figure 1b and labelled A-F. Pramana et al.<sup>29</sup> located the O<sub>int</sub> atoms on the B-sites of La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub>, while Tate et al.<sup>30</sup> found the O<sub>int</sub> atoms to be distributed over several site of La<sub>8</sub>Bi<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub>. The experimental limitations in these works were the reliance on difference Fourier maps generated from relatively low-resolution powder neutron diffraction data in the former case, and the use of the HAADF-STEM method which is an inherently twodimensional probe, in the latter.

The aim of this work was to experimentally probe the local structure of two apatite-type oxide ion conductors,  $La_{10}(GeO_4)_6O_3$  and  $La_8Bi_2(GeO_4)_6O_3$ , as a function of temperature and correlate the local structure with the measured properties and with density functional theory (DFT) simulations of their oxide ion conductivity. In addition to these triclinic examples, we also studied hexagonal  $La_8Sr_2(GeO_4)_6O_2$  (x = 0, no interstitial oxygen) to provide a reference point for the interpretation of structure-property relationships in the two triclinic

oxygen-excess materials. We used variable temperature neutron total scattering and Reverse Monte Carlo (RMC) modelling to derive local structural models which account for the higher conductivity found for  $La_8Bi_2(GeO_4)_6O_3$ .

#### Experimental

#### Synthesis

Two 5 g samples of  $La_8R_2(GeO_4)_6O_{2+x}$  for each of R = Sr, La and Bi were prepared by hand-grinding stoichiometric amounts of  $La_2O_3$  (Acros, 99.9%, heated at 950 °C for 12 h prior to synthesis), SrCO<sub>3</sub> (Fisher, 99.9%), Bi<sub>2</sub>O<sub>3</sub> (Fisher, 99.9%) and GeO<sub>2</sub> (Acros, 99.999%). The samples were heated at 1100 °C for 16-36 h followed by a further heating at 1300 °C and 1100 °C for R = Sr, La and R = Bi respectively, with intermittent grinding. The two 5 g batches were mixed and then heated at 1300, 1200 and 1100 °C for R = Sr, La and Bi respectively. All products were pale cream in colour. Samples were also heated at 1000 °C for 12 h immediately prior to neutron/synchrotron powder diffraction studies on HRPD and I11. For  $La_8Bi_2(GeO_4)_6O_3$  this treatment appeared to give rise to small impurity phases, and a fresh 5 g sample was prepared for later neutron total scattering data collections on POLARIS. All heating and cooling rates were set to 10 °C min<sup>-1</sup>. Phase purity was assessed using laboratory X-ray powder diffraction data collected on a D8 Bruker Advance with Cu K $\alpha_{1/2}$  radiation, a Ni filter and Lynxeye detector. Samples were mounted on Si zero-background slides and typically scanned over a  $10^{\circ} \le 2\theta$  $\leq 80^{\circ}$  range in 0.02° steps for 1 second per step.

#### Synchrotron Powder Diffraction

Synchrotron X-ray powder diffraction data were collected using a wavelength of  $\lambda = 0.8259211$  Å, determined from a Si standard, on samples loaded into 0.3 mm diameter fused silica capillaries at the Diamond Light Source beamline I11. A hot air blower was used to control the temperature for variable temperature data collections. The samples were heated from 30–800 °C in 25 °C steps with a scan time of 1 hour at 30 °C and 800 °C and 5 minutes for the intervening scans. The temperature was calibrated with an Al<sub>2</sub>O<sub>3</sub> standard. Room temperature experiments on La<sub>8</sub>Sr<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>2</sub> to investigate unusual peak shapes used 30 minute data collections with  $\lambda = 0.824681$  Å.

#### **Neutron Scattering**

High resolution neutron powder diffraction data were collected for La<sub>8</sub>R<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>2+x</sub> in 11 mm vanadium cans (sample heights of ~2.5 cm and masses of ~3.0 g) on the High Resolution Powder Diffraction (HRPD) instrument at the ISIS Neutron and Muon Source. A 6 hour scan was collected at room temperature for all samples. For La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> 6 minute measurements were recorded from 100-840 °C in 20°C increments to follow cell parameter evolution, followed by six 1 hour measurements at 850 °C. Equivalent data were recorded for La<sub>8</sub>Bi<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> but with a maximum temperature of 800 °C. For La<sub>8</sub>Sr<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>2</sub>, 6 minute measurements were recorded from 90-850 °C. Data of suitable quality for PDF analysis were recorded using the ISIS POLARIS diffractometer on ~2.7 g samples of La<sub>8</sub>R<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>2+x</sub> in 8 (R = La, Sr) or 6 (R = Bi) mm vanadium cans with sample heights of ~4.7 cm.  $6 \times$ 1 h measurements were recorded for each sample at room temperature.  $6 \times 1$  h measurements were performed at 850 °C for R = La and 800 °C for R = Bi.

Data were processed for Rietveld refinement using routines within Mantid software and an absorption correction applied.<sup>31</sup> PDF data were processed using GudrunN<sup>32</sup> version 5 to produce S(Q) data and the STOG<sup>33</sup> software package was used to create G(r) and F(Q) files. G(r) data were produced using  $Q_{\text{max}}$ = 35 Å<sup>-1</sup> and a Lorch function<sup>34</sup> was applied to remove Fourier ripples due to the limited  $Q_{\text{max}}$ . Low-*r* data were Fourierfiltered as follows: R = Sr,  $r \le 1.6$  Å (room temperature); R = La:  $r \le 1.52$  Å (room temperature) and  $r \le 1.55$  Å (850 °C); R = Bi:  $r \le 1.55$  Å (room temperature) and  $r \le 1.57$  Å (800 °C).

#### **Rietveld Refinement**

Rietveld refinements were performed using TOPAS v6.35-36 Average structural models were derived from starting models published by Pramana *et al.* (R = La,<sup>29</sup> Sr<sup>37</sup>) and Tate *et al.* (R= Bi<sup>30</sup>). For combined refinements, HRPD banks 1 ( $2\theta$  = 168.567°), bank 2 ( $2\theta = 90.2481^\circ$ ) and synchrotron data were simultaneously fitted. Instrument calibration constants, wavelengths and instrumental contributions to peak shapes were determined using Si or LaB<sub>6</sub> standards. Backgrounds were described using a 12<sup>th</sup>-order Chebyshev polynomial, and an additional broad pseudo-Voigt peak was used to describe scattering from the capillary. Peak shapes were described using conventional functions for these diffractometers, with a spherical harmonic function applied to describe hkl-dependent sample strain broadening.<sup>38</sup> When fitting multiple data sets the same spherical harmonic function was used for each, with appropriate scaling onto each x-axis. An absorption correction was refined for X-ray data, and the value was consistent with that expected based on the capillary diameter and packing density. For analysis of rapid variable-temperature data sets used to extract unit cell parameters, the peak shapes, absorption corrections and atomic coordinates were fixed at values derived from long data collections. Details of the refined structural parameters and agreement factors obtained are included in tables in the Electronic Supplementary Information (ESI).

## **PDF** Analysis

Total scattering analysis was performed using TOPAS v635-<sup>36</sup> for small-box analysis by converting the G(r) data produced by STOG into D(r) normalised to the sum of the scattering of all atomic pairs according to standard formalisms.<sup>39</sup> RMCProfile<sup>40</sup> was used for big-box analysis. The G(r) and F(Q) produced by STOG and the Bragg scattering data from POLARIS bank 3 ( $2\theta = 52.2461^{\circ}$ ) were used in the RMCProfile refinement. The G(r) data were fitted as D(r).  $5 \times 5 \times 4$  (~50×50×30 Å;  $\alpha \approx \beta \approx 90^\circ$ ,  $\gamma \approx 120^\circ$ ) supercells were generated from the Rietveld models containing a total of 4,200 sites for R = Sr. For R = La, Bi, an extra 600 or 1200 sites were used to describe interstitial oxygen (Oint) positions A - F (Figure 1b), and equivalent vacancy positions. Site swapping procedures were used during the RMCProfile modelling to test data sensitivity to different Oint distributions. Minimum distance restraints were applied to prevent unreasonably short atomic approaches, and weak bond valence restraints were applied to maintain chemically plausible structures. Convergence and a good fit to experimental data was typically achieved after approximately  $5 \times 10^6$  moves (approximately 15 hours on a standard desktop PC). A minimum of 8 different configurations were used for each set of data fitted and the resulting configurations averaged during data analysis. "Cloud plot" figures were drawn using the VESTA software package.<sup>41</sup>

#### **Results and Discussion**

#### **Average Structure and Phase Transitions**

Powder diffraction showed that high purity samples of La<sub>8</sub>R<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>2+x</sub> (R = Sr, La, Bi) could be prepared. In the large samples prepared for neutron scattering experiments, the R = Sr sample contained an  $\lesssim 1$  weight % Sr<sub>2</sub>GeO<sub>4</sub> impurity, the R = La sample a  $\lesssim 1$  % La<sub>2</sub>GeO<sub>5</sub> impurity and the R = Bi sample a  $\lesssim 1$  % La<sub>2</sub>Ge<sub>2</sub>O<sub>7</sub> impurity. Rietveld plots of combined synchrotron X-ray and HRPD neutron refinements are given in the supplementary Figures S1 to S6 for room temperature (R = Sr, La, Bi) and 850/800 °C (R = La, Bi) data.

There has been some debate in the literature as to the true symmetry of the R = Sr (x = 0) phase; the presence of apparent peak splittings and broadening in the reported room temperature powder diffraction patterns prompted some authors to propose monoclinic symmetry at room temperature.<sup>26</sup> We also observed unusual peak splittings in samples synthesised at 1300 °C, but found that they could be explained by assuming that a mixture of hexagonal phases with slightly different cell parameters was present, suggesting compositional inhomogeneity. The fits obtained to high-resolution synchrotron X-ray diffraction data using a multi-phase model in which cell parameters vary linearly with composition are shown in Figure S1. Heating to higher temperature (e.g. 1350 °C for 16 h) produced a phase with sharp peaks consistent with hexagonal  $P6_3/m$  symmetry, as shown by the fits in Figure S2. We therefore conclude this is the correct space group for roomtemperature R = Sr. Consistent with earlier studies, Rietveld refinement using both low- and high-temperature data suggested that Sr is located primarily (95%) on the A1 trigonal prismatic 4f site (dark green polyhedra in Figure 1).

Table 1. Important structural parameters. Notes: <sup>*a*</sup> room temperature values; <sup>*b*</sup> Unit cell distortions as % difference of *a* and *b* from their average and sum of angle differences from 90/120°. <sup>*c*</sup> Volume coefficient of thermal expansion quoted from 100 to 700 °C; <sup>*d*</sup> room temperature average Ge–O bond length from Rietveld refinements; <sup>*e*</sup> room temperature range of tetrahedral bond angles with standard deviation of values in () parentheses; <sup>*f*</sup> *fwhm* extracted by fitting a Gaussian curve to the range of tetrahedral bond angles in RMCProfile configurations.

Cell Param. <sup>a</sup>	a (Å)	b (Å)	<i>c</i> (Å)	α (°)	β(°)	γ (°)	$V(Å^3)$
Sr	9.90260(2)	9.90260(2)	7.32501(3)	90	90	120	622.067(3)
La	9.9412(1)	9.9190(1)	7.30337(6)	90.914(1)	88.1994(16)	120.8202(9)	618.15(1)
Bi	9.9042(1)	9.89656(1)	7.34419(7)	90.568(1)	89.174(7)	120.172(1)	622.27(1)
Site Occ.	А	В	С	D	Е	F	
Sr	0	0	0	0	0	0	
La	0.08(1)	0.46(1)	0	0	0	0	
Bi	0.22(1)	0.23(1)	0.06(1)	0	0	0	
Quantity	$\delta(a,b) \ (\%)^b$	$\delta(\text{ang})(^\circ)^b$	$\alpha_{\rm V} \; (\times 10^{-6} \; {\rm K}^{-1})^c$	Twist (φ,°)	Ge–O (Å) $^d$	Tet angles <sup>e</sup>	RMC tet <i>fwhm<sup>f</sup></i>
Sr	0	0	26.5(2)	23	1.741	103.8-114.9 (3.7)	20.3
La	0.11	3.54	29.1(6)	19–27=23(5)	1.777	91.8-136.6 (10.4)	23.2
Bi	0.04	1.56	29.0(5)	18-27=22(3)	1.765	90.8-128.9 (9.3)	30.2

Room temperature powder patterns for R = La and Bi are significantly more complex, in line with the  $P\overline{1}$  symmetry previously reported.<sup>29-30</sup> Samples showed a significant *hkl*-dependence to their peak shape, which is consistent with the strain broadening expected from the symmetry-lowering phase transition (discussed below) that occurs as they are cooled from high temperature. Peak shapes could be described by convoluting a *hkl*-dependent spherical harmonic function describing strain broadening onto the instrumental and isotropic sample contributions to the peak shape. We note that this approach is not possible in the RMCProfile modelling discussed below. Bi was found to occupy solely the A2 site within uncertainty.

One of the aims of this study is to identify the positions of the additional oxygen in for R = La and Bi (x = 1 in  $La_8R_2(GeO_4)_6O_{2+x}$ ) compared to Sr (x = 0); we call this 27<sup>th</sup> oxygen per formula unit Oint. The distribution of the excess oxygen atoms over likely sites A-F (Figure 1b) was determined by combined Rietveld fitting of synchrotron X-ray and HRPD high resolution neutron powder data. After refining a model with no Oint, oxygens were introduced on 2i Wyckoff sites close to positions A-F with occupancy 1/12 (giving one  $O_{int}$  per formula unit or x = 1). Their occupancies were refined with a soft restraint applied to maintain the overall O<sub>27</sub> composition, a constrained minimum value of zero and a soft restraint was applied to keep them close to the midpoint of two tetrahedra (an inversion centre in  $P\overline{1}$ ). Repeated cycles of full Rietveld refinement and randomisation of occupancies and coordinates consistently showed low or zero occupancy of sites D, E and F. Final O<sub>int</sub> occupancies at sites near A/B/C were 0.08(1)/0.46(1)/0.00(1) corresponding to a total O content of 27.08(4) for R = La, and 0.22(1)/0.23(1)/0.06(1) total = 27.00(6) for R = Bi. Removing restraints on the O<sub>int</sub> occupancy gave a total oxygen content per cell of 27.19(5) and 27.02(6) for R = La and Bi respectively. Since the change in fit quality was minimal in these free refinements (e.g. R<sub>wp</sub> 4.146 to 4.141%.), and given the correlation between oxygen occupancies and other parameters, we choose to report models

soft-restrained to an overall O<sub>27</sub> composition. Performing equivalent analysis using lower resolution but higher signalto-noise Polaris data for R = La gave essentially identical occupancies of 0.06(1)/0.43(1)/0.00(1) for A/B/C sites. For R = Bi, Polaris data were recorded on a physically different sample and gave A/B/C/D/E/F occupancies of 0.13(1)/0.21(1)/0.13(1)/0.00(1)/0.02(1)/0.01(1). These differences could reflect the difficulties in determining oxygen content by Rietveld refinement, as discussed below, or minor changes in oxygen distribution due to different sample thermal histories.

Even using neutrons, the contribution of the single O<sub>int</sub> per unit cell to the total scattering is relatively small (2.5 %), such that the determination of O<sub>int</sub> positions is challenging from powder diffraction data. It is also worth noting that the symmetry reduction from  $P6_3/m$  to  $P\overline{1}$  could give rise to six different domains in a single crystal based on the different basis vector choices relative to the parent structure. For a polycrystalline sample this means there are six equivalent cell parameter choices with similar but subtly different values. These are tabulated for R = La in supplementary Table S6. Choosing a different set of cell parameters in the triclinic fit is equivalent to selecting a different domain, and will change the labelling of the various  $O_{int}$  sites (ESI Table S6). Care must therefore be taken when comparing different refined models. Despite these caveats, our data suggest the B site has significant occupancy in both the La and Bi phase, with some density on site A. Ge sites Ge2 and Ge3, which lie on the BD and AC Oint chains respectively, show the most significant departure from tetrahedral bond angles, consistent with this occupation pattern. In contrast, tetrahedral bond angles in Ge1, which lie along the EF chains, remain relatively undistorted with standard deviations of the 6 tetrahedral angles similar to those in the hexagonal R = Sr case. Average Ge1O<sub>4</sub> bond angles and standard deviations of 109.4/4.6; 109.2/5.4 and 109.6/3.7° were obtained for R = La, Bi and Sr respectively. Values for all tetrahedral are given with other important structural parameters for the three phases in Table 1.



**Figure 2.** Unit cell edges and angles derived from variable temperature synchrotron powder diffraction studies of La<sub>8</sub>R<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>2+x</sub>. (a) R = Sr; (b) volume of all phases; (c)–(d) R = La; (e)–(f) R = Bi. For R= La, Bi a triclinic model was used at all temperatures. Note that  $\gamma$  ( $\approx 120^{\circ}$ ) is plotted on the right hand scale of (d) and (f) to emphasise the cell angle-convergence at the phase transition to *P*6<sub>3</sub>/*m*.

Figure 2 shows the unit cell parameter evolution for each sample between room temperature and ~850 °C extracted by Rietveld refinement using synchrotron X-ray data. Values obtained from high resolution neutron powder diffraction experiments are essentially identical and are shown in ESI Figure S7 Each material shows positive thermal expansion with  $\alpha_{Vol} = 26.5(2), 29.1(6) \text{ and } 29.0(5) \times 10^{-6} \text{ K}^{-1} \text{ between } 100 \text{ }^{\circ}\text{C}$ and 700 °C for R = Sr, La and Bi respectively. For R = La and Bi we see a clear indication of a phase transition to hexagonal symmetry at T = 780 and 740 °C with a and b cell edges converging, angles  $\alpha/\beta$  approaching 90° and angle  $\gamma$  approaching 120°. Whilst a symmetry analysis using ISODISTORT<sup>42</sup> shows that a  $P6_3/m$  to  $P\overline{1}$  transformation could occur via a number of intermediate subgroups (e.g.  $P\overline{3}$ ,  $P2_1/m$ ), it can be described using a single  $\Gamma_4^+\Gamma_6^+$  irrep describing O<sub>int</sub> ordering, and is allowed to be continuous by Landau theory. We see no evidence in the diffraction data for simple intermediate phases, which would all require at least one 90° cell angle. We do see a subtle change in thermal expansion for R = Bi in the X-ray and neutron data around 250 °C, which could suggest the onset of partial O mobility. One notable feature in Figure 2 and Table 1 is the smaller metric distortion from hexagonal for R =Bi than R = La, manifested in the smaller differences between the unit cell parameters a and b, and the smaller deviations of the unit cell angles from 90 and  $120^{\circ}$  for R = Bi; we return to this point below.

As discussed in the Introduction, previous workers have described the P6<sub>3</sub>/m to P-1 phase transition in Cd and Hg fluorophosphate  $O_{int}$ -free apatites in terms of a soft mode  $\Gamma_4^+\Gamma_6^+$  irrep distortion which causes a cooperative tilting of tetrahedra when frozen into the structure. Instead of using conventional crystallographic fractional coordinates, we can describe our room temperature structures of the P-1 La phase in terms of the high temperature  $P6_3/m$  parent structure plus the amplitude of 63 symmetry-adapted-distortion-modes using the language of Campbell, Stokes and co-workers.<sup>42</sup> We find that 16 of the 63 allowed modes have irrep  $\Gamma_4^+\Gamma_6^+$ . These  $\Gamma_4^+\Gamma_6^+$  modes appear to dominate the structural distortion, with 6 of the 9 largest amplitude modes belonging to this irrep. These modes alone describe the essence of the GeO<sub>4</sub> displacements and distortions. We see similar behaviour for R = Bi, and histograms of mode amplitudes are included for both phases in ESI Figure S8. Comparing mode amplitudes between R = La and R = Bi we see, as expected, that the two structures display very similar distortions. On average, mode amplitudes (which give the root-summed-squared displacement of atoms in the child cell) for R = Bi are around 2/3 of their values for R = La, another representation of its lower distortion. The most significant difference between the two structures is in the O4 (O') zcoordinate. O4 is coordinated exclusively by A2 cations and its position could be influenced by the lone pair cations on these sites, as indicated in previous theoretical studies.<sup>43</sup>



**Figure 3.** Representative RMCProfile fits at room temperature of  $La_8R_2(GeO_4)_6O_{2+x}$ : (a) R = Sr Bragg; (b) R = Sr PDF; (c) R = La Bragg; (d) R = La PDF; (e) R = Bi Bragg; (f) R = Bi PDF.

#### Local Structure

Further insight into the coordination environments of  $O_{int}$ , their influence on the surrounding coordination polyhedra and the implication of their temperature evolution on conductivity requires a local rather than average structural probe. We have used big-box modelling using 5×5×4 supercells within the RCMProfile package to simultaneously fit Bragg, D(r) and F(q) data from each sample.

For R = Sr excellent agreement with experimental data was obtained from each of 8 different RMCProfile starting configurations. Representative fits to Bragg and PDF data are shown in Figure 3a/b (F(Q) fits shown in ESI Figure S9). As expected for an ordered structure, the "cloud plots" of atomic density produced by folding the supercell configurations back into a single unit cell show good agreement with the average structural model (Figure 4a) and with the size of the atomic displacement parameters. Bond lengths and bond angle distributions around GeO<sub>4</sub> were as expected for regular tetrahedra. Fitting the observed distributions (ESI Figure S10a/b) in all 8 configurations to Gaussian functions gave a mean Ge–O bond length of 1.7584(7) Å and O–Ge–O angles of 108.91(3)°; the full width at half maximum (fwhm) of the distributions were 0.176(1) Å and 20.33(7)° respectively. The corresponding bond length determined from Rietveld refinement was 1.741(8) Å and angles ranged from 103.9 to 114.9°. The difference between the Rietveld- and RMCProfile-derived bond lengths is presumably caused by the well-known shortening of refined bond lengths when harmonic displacement parameters are used for groups undergoing correlated motion.<sup>44</sup>



**Figure 4.** Cloud plots showing volumes enclosing oxygen sites in (a)  $La_8Sr_2(GeO_4)_6O_2$  and (b-d)  $La_{10}(GeO_4)_6O_3$ . (a) shows clouds for all atoms. (b) Shows only GeO<sub>5</sub> groups where O<sub>int</sub> is part of Ge coordination sphere at room temperature. (c) Shows only GeO<sub>5</sub> groups at 800 °C. (d) Shows only GeO<sub>4</sub> groups at 800 °C. Small orange spheres show ideal O<sub>int</sub> sites and green spheres A sites. Tetrahedra show GeO<sub>4</sub> groups in the high temperature  $P6_3/m$  structure. In (b) the displacement of clouds from tetrahedral positions shows the local distortion of GeO<sub>5</sub> groups. Yellow surface drawn at 2% of maximum grid value in each plot; contours blue to red where cloud surfaces interact the cell edges show the probability of population of different regions.

For R = La and Bi, four models were tested to describe the Oint distribution and how it influences conclusions about local structure. These involved starting Oint atoms on sites to mimic the distribution indicated by Rietveld refinement, or statistically placing them across all available sites. During RMCProfile modelling O<sub>int</sub> was then either limited to swapping between just the Rietveld sites or allowed to visit all sites A-F. After convergence we find that the O<sub>int</sub> distribution broadly follows that derived by Rietveld refinement (ESI Table S7). This occurs despite RMC methods favouring random models and is caused, at least in part, by the GeO<sub>4</sub> distortions present in the Rietveld-derived average model, meaning that certain O<sub>int</sub> sites provide more favourable local environments in terms of an absence of short contacts with neighbouring oxygens. Importantly, analysis of different models showed that the local structural information derived and discussed below was independent of the Oint model used. The results are therefore based on the average of 8 starting configurations in which Oint was allowed to populate all possible sites starting from a statistically distribution.

Figure 3 shows typical fits to the room temperature Bragg and D(r) data for R = La (Figure 3c, 3d) and R = Bi (Figure 3e, 3f); Figure 5 shows equivalent fits above the  $P\overline{1}$  to  $P6_3/m$ transition. F(Q) plots are included in ESI Figure S9. We see good agreement between observed and calculated data in all cases. The largest discrepancies are the reflections for R = La at ~13500 µs ( $d \approx 4$  Å) which derive from the parent (2, -1, 1): whilst the overall intensity of this reflection is captured correctly, we are not able to describe the *hkl*-dependent peak shape during RMCProfile modelling. We note that this peak is fitted much better in the Rietveld plot of Figure S3d. Since the overall intensity is modelled correctly, this will not influence our conclusions on local structure.

The most important information from PDF modelling concerns the local coordination geometry around Oint and how it changes with temperature. By analysing each Ge site in the big-box configuration we can probe individual GeO<sub>n</sub> groups and how they vary depending on the location of Oint. Histograms of bond distances and angles around Ge groups (ESI Figure S10) show that the  $GeO_n$  polyhedra fall into two distinct categories: GeO4 tetrahedra (84% of total) which are remote from O<sub>int</sub> sites and GeO<sub>5</sub> polyhedra (16%) where O<sub>int</sub> enters the coordination sphere. This is consistent with  $^{17}\mathrm{O}$  NMR studies on La\_8Y\_2Ge\_6O\_{27}.^{45} Whilst it would also be possible to produce GeO<sub>6</sub> groups if two nearby O<sub>int</sub> sites were simultaneously occupied and close to the same Ge (e.g. B and D close to Ge2 in Figure 1b), the number of such groups is extremely small in our configurations. We can produce cloud plots showing the local geometries near Oint by folding all the GeO<sub>4</sub>O<sub>int</sub> groups from all configurations back into a single crystallographic cell as shown in Figures 4b and 4c for R = La. Since the plots of Figure 4 use a single unit cell, they average the big-box configuration in a fashion similar to the average crystallographic model meaning that the "double clouds" or "peanut" Oint shapes represent sites that are not occupied simultaneously. We can see, however, that O<sub>int</sub> sites are relatively localised at low temperature (Figure 4b) but become significantly more diffuse at high temperature (Figure 4c).



**Figure 5.** Representative RMCProfile fits of La<sub>8</sub>R<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> at 850 °C (R = La) and 800 °C (R = Bi); (a) R = La Bragg; (b) R = La PDF; (c) R = Bi Bragg; (d) R = Bi PDF.

The cloud plots of Figure 4 help understand the origin of the high oxide ion conductivity in oxygen excess apatite-type germanates. Recent density functional theory (DFT) studies of the La<sub>10-y</sub>Bi<sub>y</sub>Ge<sub>6</sub>O<sub>27</sub> (y=0, 2, 4) series, supported by neutron scattering experiments, have calculated the oxide ion migration pathways in these materials and provided new insight into the lone pair effects associated with the presence of Bi(III).<sup>43, 46</sup> The ab initio simulations identified four oxide ion exchange mechanisms contributing to the ionic conductivity, with the main one based on oxide ion transport in the *c*-direction via the ...GeO<sub>4</sub> – GeO<sub>5</sub> – GeO<sub>4</sub>... network. This mechanism, prevalent at all simulation temperatures for all compositions in the series, gives rise to ionic displacement clouds that are very similar in appearance to the atomic density map shown in Figure 4c.

Focussing on the geometries of individual  $GeO_n$  polyhedra, we find that GeO<sub>4</sub> tetrahedra are only slightly more distorted than in the R = Sr case (fwhm of angle distribution 20.3°,  $23.2^{\circ}$  and  $30.2^{\circ}$  for R = Sr, La, Bi respectively), whereas the GeO<sub>5</sub> groups display a much broader distribution of bond angles with maxima at ~90, 140 and 160°. Bond angle histograms are given in ESI Figure S10. We can gain more insight into the geometry of the GeO<sub>5</sub> polyhedra from the histograms of Figures 6 and 7. Figure 6 shows a histogram of the difference-in-distance between Oint and its two neighbouring Ge (i.e., the Ge–O·····Ge local asymmetry). For both R = La and R = Bi we observe a strong preference for each  $O_{int}$  to have one short and one long Ge-O distance at room temperature. This is consistent with local bond valence considerations. If we take a typical Ge–Ge separation of 4 Å, an O placed midway between the two would have a bond valence sum of ~1.0. Displacing  $O_{int}$  locally to give bond lengths of 1.6 and 2.4 ( $\Delta r$ 

= 0.8 Å as observed experimentally, Figure 6a) gives a more reasonable bond valence sum of ~1.7.



**Figure 6.** Histograms of  $\Delta$ Ge–O<sub>int</sub> distances in (a) La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> at room-temperature (purple line) and 850 °C (gold bars) and; (b) La<sub>8</sub>Bi<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> at room-temperature (blue) and 800 °C (red). Distances  $\Delta$ *r* defined in inset to (b).



**Figure 7.** Histogram of  $\tau$  values found in (a) La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> at room-temperature (purple line) and 850 °C (gold bars); (b) La<sub>8</sub>Bi<sub>2</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> at room-temperature (blue line) and 800 °C (red bars).

On heating above the  $T_c \approx 750$  °C phase transition we see a marked change in the  $\Delta r$  histograms. The sharp peaks at  $\Delta r = \pm 0.8$  Å move to slightly smaller values and we see a significant increase in the number of O<sub>int</sub> located symmetrically between the two GeO<sub>4</sub> tetrahedra ( $\Delta r \approx 0$ ). These can be viewed as Ge<sub>2</sub>O<sub>9</sub> units as opposed to GeO<sub>5</sub> + GeO<sub>4</sub>. Using  $|\Delta r| \le 0.25$  Å as a cut off (based on the shape of the Figure 6 histograms), we can quantify the % of O<sub>int</sub> in Ge<sub>2</sub>O<sub>9</sub> groups at room/high temperature as 6(2)/20(2) % and 15(3)/29(5) % for R = La and Bi respectively.

We can also probe the local coordination geometry of each GeO<sub>5</sub> group using the  $\tau_5$  parameter discussed by Addison and co-workers and defined as  $\frac{\beta-\alpha}{60^\circ}$ , where  $\beta$  and  $\alpha$  are the largest and second largest bond angles respectively.<sup>47</sup>  $\tau_5$  varies from 0 for a square pyramidal site to 1 for a trigonal bipyramid. Figures 7a and 7b superimpose low and high temperature  $\tau_5$  histograms for R = La and R = Bi. In the low temperature  $P\overline{1}$  structure we see that the GeO<sub>5</sub> groups are predominantly square planar in nature, whereas there is a shift in the distribution to predominantly trigonal bipyramidal at high temperature.

#### Conclusions

Our studies have provided important new insight into the structural properties of  $La_8R_2(GeO_4)_6O_{2+x}$  phases and how these influence conductivity. For the low-conductivity, x = 0,  $O_{int}$ -free R = Sr phase we find that it adopts the simple  $P6_3/m$  structure at all temperatures. Apparent peak splittings in the powder diffraction patterns of samples prepared at 1300 °C or lower reflect compositional inhomogeneities. RMCProfile modelling of its local structure provides a benchmark for the other  $O_{int}$ -containing higher conductivity materials.

 $La_{10}(GeO_4)_6O_3$  and  $La_8Bi_2(GeO_4)_6O_3$  both adopt triclinic  $P\overline{1}$ symmetry at room temperature and hexagonal  $P6_3/m$  symmetry at high temperature, as previously reported. Rietveld refinement shows that the single O<sub>int</sub> atom per formula unit lies between GeO<sub>4</sub> tetrahedra. Multiple sites can be occupied: the B site is strongly-favoured in La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> and the A, B and C sites are occupied in  $La_8Bi_2(GeO_4)_6O_3$ . The preferential occupation of certain sites drives the phase transition to  $P\overline{1}$ . The less-complete site ordering for R = Bi presumably occurs due to the distribution of La and Bi atoms on the A2 site (2:1 on average, but there will be local deviations from this) leading to a range of energies for nearby Oint sites. Previous computational studies have also shown that the Bi lone pair gives rise to significant local structural distortions. Due to the similar neutron scattering lengths of La and Bi, our local studies can't probe this aspect of the structure. The wider Oint distribution for R = Bi is presumably the reason why it shows a smaller distortion from hexagonal symmetry at room temperature than R = La. The presence of similar-energy  $O_{int}$  sites can then explain the lower phase transition temperature to  $P6_3/m$  and underpin the higher  $O^{2-}$  conductivity of R = Bi. Less stable Oint sites may also redistribute at low temperature, leading to the small discontinuity in the cell parameters observed at around 250 °C.

Local structural analysis shows that the  $O_{int}$  atoms are predominantly present in GeO<sub>5</sub> square planar units at room temperature, rather than in more symmetrical Ge<sub>2</sub>O<sub>9</sub> units. At high temperature, the GeO<sub>5</sub> polyhedra adopt a more trigonal bipyramid character, and there is an increase in the number of units that can be classified as Ge<sub>2</sub>O<sub>9</sub>. This shows that, whilst the high temperature structure retains significant aspects of the low temperature structure, the phase transition leads to increased disorder of the O<sub>int</sub> oxygens. The less distorted R = Bi material also appears to have a higher proportion of Ge<sub>2</sub>O<sub>9</sub>like groups at low temperature than R = La. This is again correlated to the higher O<sub>int</sub> disorder caused by a range of local environments.

Previous computational studies have suggested that the most common conduction pathway in La<sub>10</sub>(GeO<sub>4</sub>)<sub>6</sub>O<sub>3</sub> occurs through an interstitialcy mechanism that involves movement of O<sup>2–</sup> along the *c*-axis between GeO<sub>4</sub> units *via* normal-site O<sup>2–</sup> and O<sub>int</sub>. Our experimental local structure analysis results are consistent with this hypothesis and suggest a distribution of sites is present between the GeO<sub>4</sub> tetrahedra at high temperature. Our static local structure snapshots hence sample the dynamic distribution of atoms that are present in the highlyconducting phases.

In summary, total scattering studies have provided important insight into the local structure rearrangements that occur to accommodate the oxygen interstitial in  $La_{10}(GeO_4)_6O_3$  and  $La_8Bi_2(GeO_4)_6O_3$ , and how these structures re-arrange and become more disordered in the oxygen conducting high temperature phase.

# ASSOCIATED CONTENT

**Supporting Information**. Supporting information contains: all Rietveld plots discussed in the text; plots of unit cell parameters against temperature for all phases derived from neutron diffraction data; histograms of mode amplitudes and resulting atomic displacements; fits to neutron F(Q) data; histograms of bond lengths and angles at low and high temperature from RMCProfile

models; Rietveld-derived fractional coordinates;  $O_{int}$  distributions from RMCProfile models. This material is available free of charge via the Internet at http://pubs.acs.org.

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